



World Record Holder Has Heart Damaged by Vaccine

Analysis by [Dr. Joseph Mercola](#)

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STORY AT-A-GLANCE

- › Florian Dagoury is a world record holder in static breath-hold freediving; he held his breath for an astonishing 10 minutes and 30 seconds
- › After receiving his second dose of Pfizer's COVID-19 injection, he experienced increased heart rate and a reduction in his breath-holding capacity
- › A cardiologist diagnosed him with myocarditis and pericarditis, both recognized adverse effects linked to the shots
- › In another instance, 34-year-old Jeremy Chardy, a professional tennis player ranked 73rd in the world, suspended his season due to a severe adverse reaction to the COVID-19 shot
- › Veteran triathlete Antoine Méchin, 32, is also facing the potential end to his career after receiving Moderna COVID-19 injections and developing a pulmonary embolism
- › While health officials remain silent about COVID-19 injection reactions, the growing number of reports of adverse reactions cannot be silenced forever

Another elite athlete has experienced devastating injuries after receiving COVID-19 shots. Florian Dagoury is a world record holder in static breath-hold freediving. The freediver – from France and now based in Thailand – held his breath for an astonishing 10 minutes and 30 seconds.

After receiving his second dose of Pfizer's COVID-19 injection, he experienced increased heart rate and a reduction in his breath-holding capacity. A cardiologist diagnosed him with myocarditis, or inflammation of the heart muscle, and pericarditis, which is inflammation of the outer lining of the heart.¹ Both are recognized adverse effects linked to the shots.

While the U.S. Centers for Disease Control and Prevention continues to claim that these effects are "rare" after COVID-19 injections, as of November 10, 2021, 1,793 reports of myocarditis or pericarditis among people ages 12 to 29 years who received COVID-19 injections have been reported to the Vaccine Adverse Event Reporting System (VAERS).²

World Record Holder May See End of His Career

As a result of the shots, Dagoury's career may be over. He shared his experience with Pfizer's COVID-19 shots in his own words on Instagram:³

"Myocarditis, Pericarditis and Trivial Mitral regurgitation! Thank you Pfizer. Just want to share my annoying experience after vaccination and perhaps have some testimonials and similar stories around Freedivers. Did you get better?"

After my 2nd dose I noticed that my heart rate was way higher than normal and my breath hold capacities went down significantly. During sleep, I'm at 65-70bpm instead of 37-45bpm. During the day I'm now always over 100bpm instead of 65bpm, even when I sit down and relax.

I once even reached 177bpm while having dinner with friends !!!! 10 days after my 2nd jab, I went to see a cardiologist and he told me it's a common side effect of Pfizer vaccin, nothing to worry about, just rest it will pass.

40days after 2nd jab, I had no progress so I went to see another cardiologist and got diagnosed with Myocarditis, Pericarditis and Trivial Mitral

regurgitation! Which is basically an inflammation of the heart muscles caused by the immune system and some tiny leaks of blood from the valves that no longer close properly.

I'm now struggling to reach 8min breath hold, 150m dyn and I even have a strong urge to breathe doing 40m dives. 30% decrease on my diving performance roughly.

My first thought and recommendation to Freedivers around the world is to choose a vaccine which is done the old fashion way like Sputnik, Sinovac, Sinopharm etc...instead of those new mRNA vaccines."

Professional Tennis Player Ends Season After Shots

In another instance, 34-year-old Jeremy Chardy, a professional tennis player ranked 73rd in the world, suspended his season due to a severe adverse reaction to the COVID-19 shot, which left him unable to engage in intense activity. Speaking with The COVID World, Chardy said:⁴

"Since I had my vaccine (between the Olympics and the US Open), I have a problem. I am struggling. I can't train, I can't play. In my head, it's difficult because I don't know how long it will last. For now, my season is over and I don't know when it will resume.

It's frustrating because I started the year really well. I was playing very good and then I went to the Olympic Games where I also felt very good. It's frustrating, especially that I don't have ten years left to play. I regret having the vaccine, but I could not have known that this would happen.

I'll be 35 in February so right now I might be a bit negative but this is the first time I have ever thought that this season might be my last. I don't want to think about it ... it's difficult because I was having fun and I want to play longer."

Triathlete's Career May Be Over Due to COVID-19 Injections

Veteran triathlete Antoine Méchin, 32, is also facing the potential end to his career after receiving Moderna COVID-19 injections. After his second dose, he began to experience shortness of breath and low-back pain, which turned out to be a pulmonary embolism.

The symptoms, which included breathing problems and arm pain, started after the first dose, but doctors brushed off his shortness of breath as related to stress and fatigue. About a month after his second dose, shortness of breath and body pain returned. Only after testing at a sports clinic was the pulmonary embolism revealed. Méchin said:⁵

"I am now getting treatment and I hope to regain my lung capacity (in 3-6-9-12 months?) Until then: rest and low intensity for several months. Damaging healthy people to preserve the health of the weakest, a choice of backward logic. I would not get vaccinated again if it had to be done again."

Professional Mountain Biker Sidelined by Shots

Kyle Warner is another example of healthy people in peak physical condition being harmed by COVID-19 injections. Warner, a 29-year-old professional mountain bike racer, got his second dose of Pfizer's COVID-19 shot in June 2021. He suffered a reaction so severe that, as of October, he was still spending days in bed, easily overwhelmed by too much mental or physical exertion.⁶

Within seconds of the second dose, Warner experienced a metallic saline taste in his mouth, which can be an indication that the shot went into a vessel instead of the muscle.⁷ About two weeks later he started having strange reactions in his heart. Throughout the day, he started experiencing periods of accelerated heart rate.

As is often the case, an ER doctor completely brushed off his symptoms, telling him that he's not having a reaction to the shot but instead was having an anxiety attack.

Days after being sent home from the ER – and still experiencing heart problems, including cramping and burning – Warner went to a different hospital, where he was diagnosed with pericarditis along with postural orthostatic tachycardia syndrome (POTS) and reactive arthritis.

POTS is a blood circulation disorder that affects the autonomic nervous system and can be triggered by injections, including mRNA COVID-19 shots.⁸ One of the key symptoms of POTS is a significant increase in heart rate when a person stands up, and the elevated heart rate remains elevated for a longer than normal period. Fatigue, nausea, dizziness, heart palpitations and exercise intolerance can also occur.

While his symptoms of pericarditis have cleared, he's still struggling with the symptoms of reactive arthritis and POTS, which can last for 12 to 18 months or more. And Warner, being very fit and accustomed to listening to his body, caught the problem early – many others may not.

“I believe where there is risk, there needs to be choice,” he said.⁹ But right now, people are being misled. “People are being coerced into making a decision based on lack of information versus being convinced of a decision based on total information transparency.”¹⁰

Pfizer COVID-19 Shot Increases Myocarditis Risk Threefold

Myocarditis and pericarditis cause symptoms such as chest pain, shortness of breath and a fluttering or pounding heart. Cases have occurred most often after mRNA COVID-19 injections (Pfizer-BioNTech or Moderna), particularly in male adolescents and young adults, according to the CDC. Further, myocarditis occurs more often after the second injection, usually within a week.¹¹

A large study from Israel¹² revealed that the Pfizer COVID-19 mRNA jab is associated with a threefold increased risk of myocarditis,¹³ leading to the condition at a rate of one to five events per 100,000 persons.¹⁴ Other elevated risks were also identified

following the COVID jab, including lymphadenopathy (swollen lymph nodes), appendicitis and herpes zoster infection.¹⁵

The real-world case-control study from Israel included a mean of 884,828 people, aged 16 years and older, in each of two groups: one vaccinated and one control.¹⁶ The increased risk of myocarditis was clear, with researchers noting:¹⁷

“The risk appears to be highest among young men. We found that the risk of myocarditis increased by a factor of three after vaccination, which translated to approximately 3 excess events per 100,000 persons; the 95% confidence interval indicated that values between 1 and 5 excess events per 100,000 persons were compatible with our data.

Among the 21 persons with myocarditis in the vaccinated group, the median age was 25 years (interquartile range, 20 to 34), and 90.9% were male.”

When myocarditis occurs, it reduces your heart’s ability to pump and can cause rapid or abnormal heart rhythms that can be deadly. In severe cases, myocarditis can cause permanent damage to the heart muscle and lead to heart failure, heart attack, stroke and sudden cardiac death.¹⁸

In August 2021, New Zealand reported the death of a woman following Pfizer’s COVID-19 jab, which they believe was due to vaccine-induced myocarditis.¹⁹ Another devastating report in VAERS states that a 15-year-old boy from Colorado, with no preexisting conditions or allergies, died from cardiac failure two days after receiving Pfizer’s COVID-19 vaccine.²⁰

Unlike in the U.S., where the CDC has recommended Pfizer’s COVID-19 shot for children 5 years and up, the U.K. is taking a more cautious and sensible approach. Due to myocarditis risks in youth, Britain’s Joint Committee on Vaccination and Immunization (JCVI) recommended against COVID-9 injections for healthy 12- to 15-year-olds.

Wei Shen Lim, COVID-19 chair for JCVI, stated, “The margin of benefit is considered

too small to support universal Covid-19 vaccination for this age group at this time.”²¹

COVID-19 Injection Victims Need To Be Heard

While health officials remain silent about COVID-19 injection reactions, the growing number of reports of adverse reactions cannot be silenced forever. Websites like C19 Vax Reactions,²² started by former Green Bay Packers offensive lineman Ken Ruetters, whose wife Sheryl suffered a severe neurological reaction to Moderna's COVID-19 shot, exist online for people to share their stories.

There you can read over 500 real testimonies of adverse reactions to the shots and view dozens of videos detailing individuals' reactions. The Real, Not Rare website has also collected dozens of stories from people who have been injured by COVID-19 shots,²³ and there's a good chance you know someone personally who's been injured by the injections as well.

It's important that these voices are heard, so if you or a loved one has been injured by a COVID-19 injection, please share your story with us and encourage others you know who have a story to share theirs as well.

Sources and References

- ^{1, 3} [The COVID World November 5, 2021](#)
- ² [U.S. CDC November 16, 2021](#)
- ⁴ [The COVID World September 24, 2021](#)
- ⁵ [The COVID World September 29, 2021](#)
- ^{6, 9} [YouTube, Dr. John Campbell, Kyle's Vaccine Complication October 21, 2021, 1:01](#)
- ⁷ [YouTube, Dr. John Campbell, Kyle's Vaccine Complication October 21, 2021, 21:59](#)
- ⁸ [Cureus. 2021 May; 13\(5\): e14837](#)
- ¹⁰ [YouTube, Dr. John Campbell, Kyle's Vaccine Complication October 21, 2021, 41:51](#)
- ¹¹ [U.S. CDC November 12, 2021](#)
- ^{12, 14, 15, 16, 17} [The New England Journal of Medicine August 25, 2021](#)
- ¹³ [MedPage Today August 25, 2021](#)
- ¹⁸ [Mayo Clinic, Myocarditis](#)

- ¹⁹ [New Zealand Ministry of Health August 30, 2021](#)
- ²⁰ [Yahoo May 6, 2021](#)
- ²¹ [BBC News September 3, 2021](#)
- ²² [C19 Vax Reactions](#)
- ²³ [Real Not Rare, Real Stories](#)



How Did Carcinogenic Generic Pill Get Past the FDA?

Analysis by [Dr. Joseph Mercola](#)

✓ Fact Checked

STORY AT-A-GLANCE

- › Since 2018, the carcinogenic compound NDMA has been found in several different drugs, including three blood pressure medications (valsartan, losartan and irbesartan), two heartburn medications (Zantac and Axid) and the diabetes drug metformin
- › In the case of valsartan, three companies whose drugs were recalled in 2018 had all purchased the active ingredient from a Chinese manufacturer called Zhejiang Huahai Pharmaceutical Co.
- › The U.S. Food and Drug Administration checks less than 1% of imported drugs for impurities or potency, and in five years sent warning letters to only 25% of companies suspected of faking quality data
- › While generics are a boon to patients in that they're far less expensive while still providing the same benefits, there's more room for error as they also receive far less scrutiny by regulators, and manufacturers are trusted to regulate themselves
- › An estimated 80% of all active drug ingredients are manufactured in China and India, and overseas plants are rarely inspected by U.S. authorities

This article was previously published September 30, 2020, and has been updated with new information.

Previously, I reported that carcinogenic N-nitrosodimethylamine (NDMA) had been found in certain blood pressure, heartburn and diabetes medications. As of February 2020, drugs recalled due to contamination with this poison included:¹

- Valsartan, losartan and irbesartan (high blood pressure medications)
- Zantac² and Axid (heartburn medications)
- Metformin (diabetes medication)

In the case of valsartan, the three companies whose drugs were recalled in 2018 had all purchased the active ingredient from a Chinese company called Zhejiang Huahai

Pharmaceutical Co. It's one of China's largest manufacturers of generics.³

Since 2018, the recall has been expanded dozens of times to also include losartan and irbesartan, made by more than 10 different companies with distribution in some 30 countries.⁴

As reported⁵ by Bloomberg in December 2019, the U.S. Food and Drug Administration checks less than 1% of imported drugs for impurities (or potency for that matter). Clearly, the regulatory system, which is meant to safeguard patients, is broken, and trust in drug manufacturers is often misplaced.

Disturbingly, Bloomberg's report⁶ suggests the NDMA contamination at Huahai may have been intentional, at least in the sense that profitability was prioritized over thorough quality testing and perfecting of novel manufacturing methods.

What Is NDMA?

NDMA is a water-soluble chemical known to cause cancer in animals. In humans, it's classified⁷ as a probable carcinogen and causes serious liver damage and liver failure.⁸

According to the Environmental Protection Agency's technical fact sheet,⁹ NDMA, which can form in both industrial and natural chemical processes, is a member of N-ni-trosamines, a family of potent carcinogens.

"Potential industrial sources include byproducts from tanneries, pesticide manufacturing plants, rubber and tire manufacturers, alkylamine manufacture and use sites, fish processing facilities, foundries and dye manufacturers," the EPA notes. However, we now know the chemical can also be produced during the manufacturing of drugs.

Historically, there are several cases¹⁰ in which NDMA was used as a poison. In 1978, a German teacher's wife died after he put NDMA in her jam and a Nebraska man was

sentenced to death that same year for spiking lemonade with it, killing two people.

In 2013, a Chinese medical student died as a result of an April Fool's prank when NDMA was put into the water cooler, and in 2018, a Canadian graduate student poisoned a post-doctoral fellow by injecting it into an apple pie. Meanwhile, hundreds of millions of patients around the world have been taking drugs contaminated with this poison, oftentimes daily, for years on end.

Can FDA Ensure Drug Safety?

Bloomberg's report¹¹ reviews the history of how carcinogens like NDMA have crept into the generic drug supply, and raises serious questions about the FDA's ability to ensure drug safety.

The article features the story of Karen Brackman, who after taking generic valsartan for two years suddenly found herself with a diagnosis of a rare and aggressive liver cancer, despite having no family history of cancer, and no specific risk factors for it.

As reported by Bloomberg,¹² some of the contaminated valsartan pills contained as much as 17 micrograms of NDMA per pill, an amount estimated by European health regulators to give 1 in 3,390 people cancer. Brackman suspects she's one of the unlucky ones.

While generics are a boon to patients in that they're far less expensive while still providing the same benefits, there's more room for error as they also receive far less scrutiny by regulators, and manufacturers are trusted to regulate themselves.

Most Active Ingredients Are Manufactured in China and India

An estimated 80% of all active drug ingredients are manufactured in China and India, and overseas plants are rarely inspected by U.S. authorities. At present, the U.S. has just one FDA inspector's office in China. In the case of valsartan, even when a plant is

inspected and found wanting, it can take years before problems are addressed – if ever.

"Huahai, the first manufacturer found to have NDMA in its valsartan, is also the one whose product had the highest concentration," Bloomberg reports.¹³

"When an FDA inspector visited in May 2017, he was alarmed by what he saw: aging, rusty machinery; customer complaints dismissed without reason; testing anomalies that were never looked into.

He reported that the company was ignoring signs its products were contaminated. Senior FDA officials didn't reprimand Huahai; they expected the company to resolve the problem on its own. Huahai didn't ...

It wasn't until a year later that another company ... found an impurity in Huahai's valsartan and identified it as NDMA. That was when the FDA demanded drugmakers begin looking for NDMA in their valsartan. They found it again and again."

As David Gortler, a drug safety consultant and former FDA medical officer, told Bloomberg, "Valsartan is just the one we caught. Who knows how many more [tainted drugs] are out there?" Well, we now know the NDMA contamination affects many other drugs as well, including metformin, used by more than 78.6 million Americans as of 2017.¹⁴

Huahai's Mistake

Bloomberg goes on to recount some of the historical details of Huahei, from its inception in 1989 to its current status as one of the largest generics companies in China, and the first Chinese company to gain FDA approval to export finished drugs to the U.S. – a generic HIV medication.

When Novartis' patent on Diovan (the brand name for its valsartan drug) expired in

2011, Huahai became one of the companies to manufacture valsartan for generic drug companies. Valsartan, being a simple compound to make and used daily by millions, looked like it could be just what Huahai needed to grow and improve its bottom line.

Now, as explained by Bloomberg, if a company like Huahai wants to create its own version of a generic drug and then export it to the U.S., they must first get FDA approval. However, if they're just manufacturing and supplying the active ingredient to a U.S. company that then produces the finished product, then FDA approval is not required. All they have to do is inform the FDA if there are any changes to the manufacturing process.

In the case of Huahai's valsartan, the company did make a change to its manufacturing process, but downplayed its significance. In November 2011, Huahai stopped using the solvent used by Novartis in the manufacturing of the brand name drug, and started using another called dimethylformamide (DMF).

This turns out to have been a massive mistake, as side reactions ended up producing NDMA, which could not be removed from the drug. "The chemists at Huahai either didn't realize that or didn't consider it a potential hazard," Bloomberg writes, adding that, in 2018, after the recall began, vice chairman of Huahai, Jun Du, told an FDA inspector that "The purpose of the change was to save money," thus increasing their profits.

The cost-savings were so substantial, it allowed Huahai to dominate the global market share for valsartan. Making matters worse, since Huahai's patent was public, other generic companies copied the new, toxic, process. According to Bloomberg,¹⁵ this is "one reason so much of the world's valsartan supply is now contaminated."

Incompetence or Intentional Poisoning?

It's hard to justify a defense of ignorance, though, seeing how the 2017 FDA

inspector's report noted multiple problems at the plant, including suspicious contaminants showing up in quality tests.

Du claimed the tests showed "ghost peaks ... from time to time for undetermined reasons." In another instance, he referred to the residual spike showing in testing as "noise." Huahai never investigated to determine what the contaminants might be, or how they got there. Instead, they simply omitted the incriminating tests from official reports.

The FDA inspector recommended the agency issue a warning letter, which would have meant Huahai would have to pass another inspection before continuing its manufacturing. But the FDA didn't send a warning letter. Instead, they urged Huahai to resolve the issues on their own – which they didn't.

Disturbingly, a lax FDA approach to inspections that reveal faked quality testing is not unusual. Bloomberg spoke to Michael de la Torre, who runs a database of FDA inspections. According to Torre, in the five years up to 2019, the FDA issued warning letters in response to faked data just 25% of the time.

“ The only element who cares in this whole global supply chain is patients. ~ David Light, CEO Valisure LLC ”

Bloomberg also recounts a number of quality problems discovered at Indian drug manufacturing plants. Clearly, FDA is failing in its mission to regulate the generics industry overseas.

The industry is expected to regulate itself, and profit wins over quality concerns most of the time when no one is around to hold the companies accountable. A company is only as ethical and conscientious as the people running it.

Quality problems are really not uncommon. The New Haven, Connecticut-based

online pharmacy Valisure LLC tests every drug it orders, and reports rejecting more than 10% of all batches it receives – in some cases due to inaccurate amounts of active ingredient, in others due to contaminants or other inconsistencies in quality.¹⁶

Kevin Schug, analytical chemistry professor at the University of Texas, told Bloomberg¹⁷ Huahai "certainly should have caught" the NMDA contamination, and "should have modified the procedure to correct it." Former FDA medical officer Gortler agreed, saying, "Any well-trained analytical chemist would know to check. If it's not intentional, it's incompetence. At some point, those are the same."

Valisure CEO David Light told Bloomberg that while people in the industry are well aware of the problems, the overwhelming consensus is that it's not "their" problem. "There's no liability at any one point," he said. "The only element who cares in this whole global supply chain is patients."

The FDA didn't send a warning letter¹⁸ to Huahai until November 2018, stating the obvious: The company should have anticipated the possibility that changing the process to use DMF solvent might cause problems, and when testing revealed anomalies, they should have identified the impurity.

Brackman filed a lawsuit against Huahai in April 2019. About 140 others have also sued Huahai and other drugmakers involved in the valsartan recall, and lawyers are reviewing several hundred additional cases, Bloomberg reports.

Bottom Line

This devastating and pervasive toxic exposure results largely from people's reliance on using drugs as symptomatic bandages that in no way, shape or form treat the cause of the disease. They trust their physicians to help them but sadly they have been captured by the drug industry and are nearly universally clueless on how to identify and address the underlying cause of most diseases.

That is why it is crucial to understand that YOU are responsible for your own health

and need to use physicians as your consultants, and not implicitly trust them. If you provide your body with what it needs, it typically tends to self-correct and get better so you can avoid these dangerous medications which, rarely, if ever, resolve the foundational cause.

Fortunately, this COVID-19 crisis has shown us the two most important physical strategies to optimize your health: vitamin D and metabolic flexibility. The ability to eliminate insulin resistance is a strategy that addresses the majority of illnesses that you will ever encounter in your lifetime.

This is why time-restricted eating, eliminating industrially processed seed oils like soy, corn and canola oils, eating a cyclical ketogenic diet, exercising and sleeping well can improve, if not eliminate, most conditions that you would need to take medications for. As you can see, drugs can harm you just because they were made with shortcuts to increase company profits.

When you follow these health principles you will decrease, if not eliminate, your need for these dangerous medications. You will also enjoy a high degree of health and freedom from the pain, disability and suffering associated with these conditions.