

Dr. Fauci's Long Criminal History

Part 1



(Fauci black eye signifies [membership in Satanic cult that rules the world](#)) Fauci is also a Jesuit!

"The coronavirus tests do not at all prove presence of a deadly virus in any patient. ...it is perhaps the greatest criminal fraud in medical history."

Anthony Fauci has more influence over Covid-19 policy than the President. A closer look at Anthony Fauci's career paints a very alarming picture. The most important thing you can learn about Dr. Fauci is that he is a Jesuit-trained educator, a Jesuit for life! The history of the Jesuits is plain and simple – Evil! Pope Francis is a Jesuit. They serve a different god.

Tony Fauci has held the top post at the NIAID in Washington for an astonishing 36 years. Today he is well past retirement age at 79, and holds the funds to determine which drug companies or university researchers will get precious government funds or not from NIAID's annual \$5 billion budget.

[The Talented Dr. Fauci](#) by F. William Engdahl

THE BLACK EYE CLUB

Who is punching them?



According to [John Bolton](#)'s nephew, Greg T Dixon, a Masonic High School friend and informant deeply connected with Freemasonry, the elite have produced themselves a designer drug – which causes *black eye* when taken... “*the darker the eye the better the high.*”

COINCIDENCE?



Whenever elite are seen sporting this high, other elite instantly are envious (provided they have taken the ‘black eye’ vacation at least once before themselves).

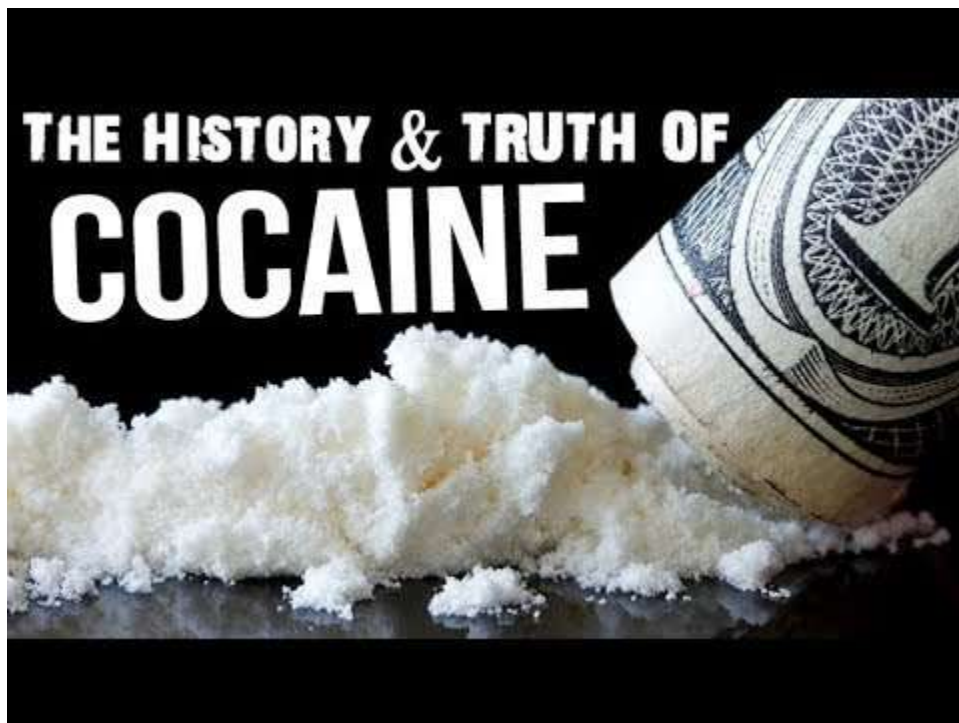
However, there is a caveat to this rule, which is taking the black eye high too many times, particularly in a row, is detrimental to one’s health.

Therefore, must be administered with caution, care and the utmost consideration. Which further leads to elite envy – flaunted before us by proxy of the U.S. media. In fact, its

recommended the drug only be administered 1-2 times – years apart – with three being the limit in a lifetime.

It's not clear the exact composition of the drug that produces the *black eye high*, but is believed to be a derivative of [lithium](#), which also is well-known to be unhealthy to take for extended periods of time. Jay Myers Documentaries.

This most likely is the reason fellow elite *love* to share photos of themselves sporting black eye, thus demonstrating they are part of the club.



(Cocaine also is another *favorite* drug among the club, where if the police knew the caliber of elite snorting it, they would no longer bother arresting people for using it, based on this reality alone.)

It would appear that there are two main categories within the “Black Eye” Club. Those seated firmly into positions of government/politics and those in the entertainment/Hollywood. Are there others in the fields of science, and military? It remains to be seen. Maybe they are merely sampling the fountain of youth referred to as adrenochrome The Secret Immortality Serum all the Illuminati use. Hunter S. Thompson's book *'Fear and Loathing in Las Vegas'* claims that adrenochrome is obtained from the adrenal gland of living humans. It is often falsely reported that it would say in the book that it is obtained from the human adrenal gland, but this is only reproduced in the film version. There is more to this evil practice than the public knows.

Steve Jackson Games published its intriguing game "Illuminati" in 1982. The Illuminati Card Game is a playing card game by Steve Jackson Games. The object of the game is to control the world via competing secret societies by sinister means - sort of like an evil form of Monopoly. The interesting part is that the game contains cards depicting a terrorist attracts on the Twin Towers and Pentagon eerily reminiscent of the 9/11 attacks. A few years ago a second edition was published.





The history of the Illuminati dates to July 4, 1776 with Jesuit Adam Weishaupt. It is a code word for Jesuits and means "Enlighten Ones". It initially was founded to bring the world's brightest and smartest people in the world of Europe. It went underground to avoid the powers threatened by the Illuminati. Adam Weishaupt gained the financial support of the House of Rothschild to advance its agenda to control the world. Today Pope Francis above is the "white" pope, and his counter-part referred to as the "black" pope are both Jesuits! Arturo Marcelino Sosa Abascal SJ is the thirty-first and present Superior General of the Society of Jesus.

**CHILDREN DON'T
JUST DISAPPEAR**

**ADRENOCHROME AND
CHILD TRAFFICKING**

IG: @TheShadowWatchman

Famous Satanist, Aleister Crowley, taught the scientific fact that the blood of children is an anti aging device in itself but is 10x more efficient when it's adrenalized children's blood. Child Trafficking is a massive and rich industry especially in Hollywood and the alphabet agencies but it's always been exposed as satanic ritual abuse where there is ritual sacrifice, rape, and blood drinking. Not only do they drink the children's blood but they torture and rape them so their blood gets adrenalized so it will work more effectively on these wicked demons. This is why the higher up Jewish Zionist elites continue aging and never seem to die. They are secretly afraid of the punishment they will get for their wickedness.

Let's go back to 1984 when Fauci was named head of NIAID during the Reagan era.

That year an AIDS researcher, Robert Gallo, working under Dr. Anthony Fauci, held a press conference to announce that he had "discovered" the AIDS virus. He said it was HIV- human immuno deficiency virus.

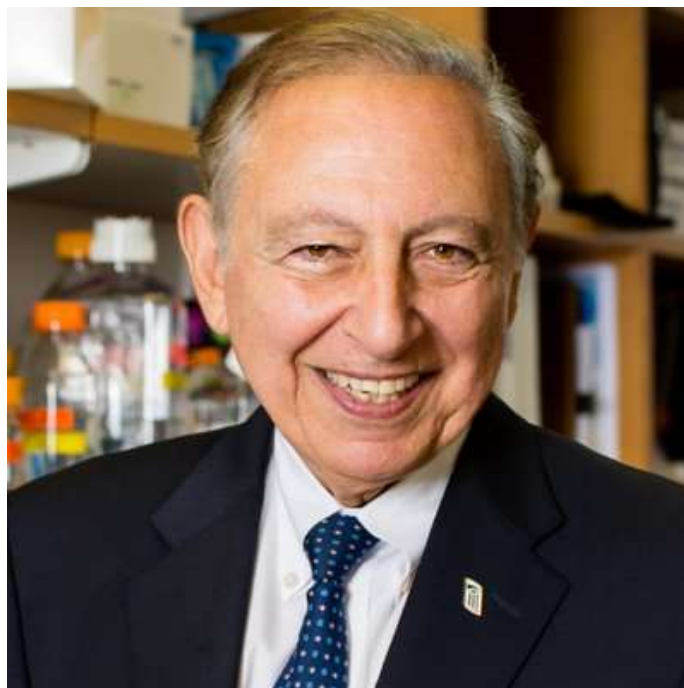
The shocking announcement which went around the world was in complete disregard of scientific procedures of prior peer-reviewed published scientific evidence, including the required electron microscope analyses.

It was a case of "science by press conference" as a critical scientist Prof. Peter H. Duesberg described it. Duesberg was an award-winning researcher at Berkeley who isolated the first cancer gene through his work on retroviruses in 1970, and mapped the genetic structure of these viruses.

For Gallo and Fauci, that was unimportant as millions in research funds flowed into NIAID to research the new virus, HIV. Fauci and Gallo claimed that AIDS was highly contagious, also by sexual transmission, especially among homosexual men.

Notably, before the Gallo claim to have found the HIV AIDS virus, NIAID had been doing research on the role of drugs, poppers or nitrites, proven immune-suppressants, in the deaths of the earliest AIDS patients. Dr. Gallo is pictured below.

That was quickly dropped in favor of researching a "cure" for AIDS. Media was told that AIDS was the "public health threat of the Century."



Gallo went on to make millions on his patented blood test for HIV, despite the fact that the test was often giving false positives and did not test directly for the alleged virus but for active antibodies, something immunology practice said was not valid, as antibodies merely suggested a past infection response and not necessarily presence of HIV. At this time in the 1980'S Fauci was responsible for AIDS research at NIAID, a post he still holds....

Yet this fraud has shaped the career of Dr. Tony Fauci for more than 35 years. Fauci as head of NIAID has taken millions from the Bill & Melinda Gates Foundation as well as the Clinton Foundation along with tens of billions from U.S. taxpayers for this bogus research. Suspiciously, the 2006 article by Giraldo and de Harven was suddenly retracted by the journal in 2019 just before the coronavirus Wuhan outbreak.

Despite the fact that he knew the established rules of virology, Fauci, as head of NIAID, recommended the Burroughs Wellcome chemotherapy drug, AZT as a "preventive drug" for HIV diagnosed patients even without symptoms!

Burroughs Wellcome gave NIAID the study that was deliberately biased for AZT. Dr. Fauci even backed AZT for pregnant women despite the grave risk to the fetus. One mark of pregnancy in all women is a higher level of antigens as the natural immune system fights any infection to protect the fetus. AZT or Retrovir, a failed leukemia drug, has been proven to be a highly toxic drug. It was approved for AIDS testing in a record 5 days by Fauci and the US Government in 1987. Today despite more than thirty years funded research and billions of dollars, no effective vaccine for HIV/AIDS exists.... [Editor note: Burroughs Wellcome is a UK pharma operation, closely aligned with the Rothschild Pirbright Institute.]

FAUCI AND COVID 19

In October, 2019 Dr. Fauci and his NIAID got \$100 million from the Gates Foundation to develop "gene-based" therapies for HIV and sickle cell disease. That means at the time of the first claims of novel coronavirus in Wuhan China, Fauci was still promoting a 35-year fraud around HIV. Fauci is also part of the Gates Foundation cabal. In 2012 Fauci was named one of the five Leadership Council of the Gates Foundation-created Global Vaccine Action Plan.

This is highly relevant to his role today as the Trump Administration coronavirus "pope." Has his NIAID or any other laboratory in the world rigorously, with electron microscopy, isolated and purified samples of patients tested SARS-CoV-2 positive for Covid-19? Or are the virus proofs as faulty as Fauci and the AIDS clique have made for HIV?

In addition NIAID is working with Gilead to conduct Phase II human trials on Gilead's drug, remdesivir, as a potential treatment for hospitalized adult patients diagnosed with COVID-19.

A Coincidence?

Relevant also is the fact that all top scientific advisers to the U.S. President's Task Force on COVID-19 are tied since decades to the bogus and destructive HIV/AIDS research and propagation of false theories.



Alongside Tony Fauci of NIAID are Deborah L. Birx, M.D., Obama appointee as US Global AIDS Coordinator who worked under Tony Fauci at NIAID from 1983-1986. Far right is Robert Redfield, current Director of the Centers for Disease Control and Prevention, center of the recent coronavirus testing scandal. Redfield cofounded with the discredited Robert Gallo, the Institute of Human Virology based at University of Maryland. Redfield and Birx also coauthored numerous scientific articles on purported HIV vaccines, none of which have been effective.

Fauci, Birx and Redfield, all incestuously complicit in the HIV/AIDS frauds and malpractice, today hold the future of not only American public health, but also of the entire world economy in their hands. Not a good situation. As their work on the proved HIV=IDS fraud shows, the coronavirus tests do not at all prove presence of a deadly virus in any patient. If this is so, it is perhaps the greatest criminal fraud in medical history.

F. William Engdahl is strategic risk consultant and lecturer, he holds a degree in politics from Princeton University and is a best-selling author on oil and geopolitics, exclusively for the online magazine "New Eastern Outlook."

F. William Engdahl only scratched the surface on the Jesuit Dr. Tony Fauci. Dr. Fauci stole the academic research paper of Dr. Judy Mikovits who worked at and for the NIH and allowed Dr. Robert Gallo to write a similar paper under his name copying the research of Dr. Mikovits. Dr. Fauci destroyed Dr. Judy Mikovits career twice, as she shares her story on the Internet. She was falsely sent to prison, discredited, and her career destroyed.

Dr. Fauci's History in Spending Billions of Government Funds on Vaccine Research with Little to Show for it

Dr. Fauci and COVID-19 Priorities: Therapeutics Now or Vaccines Later?

by Lyn Redwood, RN, MSN, President; Mary Holland, Children's Health Defense General Counsel & Vice Chair; and the Children's Health Defense Team

The rapidity with which normal life has ground to a halt as a result of coronavirus-related edicts has stunned citizens around the world, generating massive social and economic upheaval.

Meanwhile, media coverage of COVID-19 has whipped up unprecedented levels of public anxiety and fear, laying the psychological groundwork for people to eagerly embrace "magic bullet" medical solutions, no matter how experimental.

In the U.S., the World Health Organization (WHO) is now compounding the domestic panic, warning that America could become the new coronavirus "epicenter."

Across the country, a debate is raging about the nation's medical response and how best to apportion available resources. Many argue, quite reasonably, for the importance of identifying safe, effective and affordable therapies that can provide immediate help to those who are sick. On March 22, *The New York Times* reported that there are at least 69 existing drugs or compounds that might be effective in treating the coronavirus.

In China, researchers are studying intravenous vitamin C as a potential nontoxic treatment, while a paper published by French researchers on March 20 described promising COVID-19 results from the off-label use of hydroxychloroquine (an antimalarial) and azithromycin (an antibiotic).

The head of the French team, Didier Raoult, MD, PhD, is one of the world's top infectious disease and virology experts, with roughly 2,000 peer-reviewed publications and multiple awards to his name.

Raoult and coauthors point out that a major advantage of “repositioning” older drugs for this coronavirus is that their safety profile, side effects, dosing and drug interactions are already well documented.

However, Ian Lipkin, MD, of Columbia University recently told MSNBC, with a grin, that investments tend to go toward treatments that are “sexy and new and patentable” rather than to “tried-and-true, classical sort of methods repurposing drugs and strategies that have already been shown to work.”

Fauci’s tired rhetoric

For biopharma companies that are poised to profit from COVID-19-related misfortune, older drugs that have outlived their patent terms are not terribly helpful for the bottom line.

Could this be why leading White House coronavirus advisor Dr. Anthony Fauci, MD, long-time head of the National Institute of Allergy and Infectious Diseases (NIAID), recently pooh-pooed the published chloroquine evidence as merely “anecdotal”?

Fauci is a stalwart enthusiast of “patentable” vaccines, skilled in attracting massive government funding for vaccines that either never materialize or are spectacularly ineffective or unsafe.

For example, Fauci once shilled for the fast-tracked H1N1 influenza (“swine flu”) vaccine on YouTube, reassuring viewers in 2009 that serious adverse events were “very, very, very rare.”

Shortly thereafter, the vaccine went on to wreak havoc in multiple countries, increasing miscarriage risks in pregnant women in the U.S., provoking a spike in adolescent narcolepsy in Scandinavia and causing febrile convulsions in one in every 110 vaccinated children in Australia—prompting the latter to suspend its influenza vaccination program in under-fives.

In 2010, then-Senator and physician Tom Coburn, MD, called out Dr. Fauci for misleadingly touting “significant progress in HIV vaccine research.” Coburn stated ,

“The study [Fauci] referred to was a clinical trial in Thailand finding a vaccine to be 31% effective at preventing HIV infection. Unfortunately, the results of this study have been found to be statistically insignificant and the findings of the study have received much skepticism.

[. . .]

Most scientists involved in AIDS research believe that an HIV vaccine is further away than ever . . . and may never be possible. . . .”

Senator Coburn also noted that Dr. Fauci’s agency had spent over \$5.2 million over a four-year period on lavish “HIV vaccine awareness” events. Without the least hint of embarrassment, however, Dr. Fauci reappeared on YouTube in 2016 to once again push his HIV vaccine agenda, even citing the unimpressive Thailand trial.

Fauci's mobilization of billions for a never-completed Zika vaccine followed a similar playbook. And now, Fauci is predictably shining a spotlight on risky and uncertain coronavirus vaccines that may not be available for two years, rather than prioritizing the short-term therapies that patients need right now.

First off the block

According to the WHO, up to 35 COVID-19 vaccines are in the offing, including experimental messenger RNA (mRNA) vaccines and formulations that attach coronavirus to genetically modified measles vaccines. He has not mentioned the patented vaccine for COVID-19, which the CDC already owns, or the patented vaccine by the Pirbright Institute.

As biopharma companies position themselves to reap blockbuster profits, the first off the block is a vaccine thrown together at record speed by Dr. Fauci's NIAID in collaboration with Massachusetts-based biotech firm Moderna.

NIAID and Moderna began developing the vaccine before a single COVID-19 case had appeared in the U.S., completing the first batch of vaccine "within 42 days of the company obtaining genetic information on the coronavirus."

The vaccine, which bears the ho-hum name of mRNA-1273, uses an unproven mRNA technology platform. mRNA vaccines appeal to industry because of the potential for "rapid, inexpensive and scalable manufacturing," but researchers at Harvard and other medical sciences institutions have issued warnings about the vaccines' propensity to produce higher rates of side effects, including local and systemic inflammation and worrisome autoimmune responses.

Noting that the "non-native modified nucleotides" used in mRNA vaccines and components of mRNA vaccines' delivery systems have potentially toxic effects, these researchers recommend taking precautions during preclinical studies and clinical trials.

Dr. Fauci, on the other hand, praises the "new era of vaccinology"—of which the mRNA-1273 effort is a part—celebrating its use of "atomic level structural information for vaccine design, gene-based vaccine platforms, modern protein engineering and potent adjuvants."

On March 16, NIAID launched a Phase 1 trial of mRNA-1273 in 45 healthy adults after making the decision—deemed "morally questionable" by some—to sidestep the standard process for vaccine development.

That process ordinarily requires "that a manufacturer show a product is safe [in animal models] before it goes into people." Leading virologist Shibo Jiang, MD, PhD, recently condemned this "quick-fix" approach, arguing that "safety always comes first" and that it is important not to "cut corners" by skipping animal studies. Moderna's chief medical officer disagrees, saying,

“I don’t think proving this in an animal model is on the critical path to getting this to a clinical trial.”

For his part, Dr. Fauci has expressed willingness to expedite the vaccine’s approval process as soon as NIAID deems the Phase 1 trial successful.

Vaccine-hesitant experts

Ordinarily, vaccine scientists line up in lockstep to pledge their allegiance to the Faucian worldview that vaccination is the “mainstay” of prevention and offers the primary solution for challenges such as the coronavirus situation.

In an interesting turn of events, however, some of the pharma-funded media’s favorite vaccine spokesmen—slick, high-level medical professionals that manufacturers ordinarily can count on to endorse any vaccine—are urging caution.

Peter Hotez, MD, PhD, Dean of the National School of Tropical Medicine at Baylor College of Medicine, is no stranger to coronaviruses, having developed a vaccine for an earlier coronavirus in 2016 that stopped just short of commercial development.

Despite having “tried like heck” to obtain funding to move his vaccine into clinical trials, Hotez just told a U.S. Congressional Committee (on March 5) that coronavirus vaccines are scientifically challenging and have a “unique potential safety problem,” namely a “kind of paradoxical immune enhancement phenomenon.”

When Hotez observed this immune pathology in his coronavirus laboratory animals, he thought, “Oh my God, this is going to be problematic.”

Paul Offit, MD, of the Children’s Hospital of Philadelphia, another media darling who has profited handsomely from his insider status, stated in a March 10th YouTube interview that influenza deaths are “*far worse*” than COVID-19 deaths, “*yet we don’t quarantine for influenza, we don’t shut down schools for influenza, we don’t cancel meetings for influenza, we don’t cancel schools and churches and synagogues.*”

Also Offit expressed worry about the push to “rush [a vaccine] through,” particularly in the absence of “any history of making a coronavirus vaccine.” Offit concluded:

And certainly the FDA has got to regulate this product because right now everybody in the United States will probably take it in a second, even if it wasn’t tested.

A profitable crisis

In early March, Congress passed an emergency coronavirus spending bill, much of which will “directly benefit the drug industry.” Legislators who tried—and failed—to include meaningful affordability provisions in the spending bill worry that *“A danger remains that the federal government will simply write a blank check signed to big pharma as a result of this crisis.”*

Clues that pharma is embracing the opportunities furnished by the COVID-19 crisis come from the financial markets. Market reports indicate that the health care industry has been able to “withstand” the wider stock market plunge due to big gains by pharmaceutical, biotech and medical diagnostic companies involved in developing coronavirus-related products.

After Moderna announced, in late February, that it had shipped off its mRNA-1273 to NIAID for the Phase 1 trial, the company instantly became “one of the hottest biotech stocks on the market.” (Moderna’s NASDAQ ticker symbol is, handily enough, MRNA.)

Investment advisors have pointed out that the likely promotion of mRNA-1273 as prevention is a “big deal” that stands to make Moderna a fortune, because “millions” of uninfected people will want to “pre-emptively protect themselves.”

Coronavirus drug and vaccine manufacturers are also sitting pretty because of liability immunity conferred under the 2005 PREP Act (Public Readiness and Emergency Preparedness Act) and a follow-up Department of Health and Human Services (HHS) Declaration specific to COVID-19 published in the Federal Register on February 4, 2020.

The PREP Act protects the manufacturers of medical “countermeasures”—including vaccines, medications, medical devices and other products—from the risk of damages in the event of a declared public health emergency such as the currently declared coronavirus pandemic.

The tort immunity described under the COVID-19 Declaration pertains to “any claim of loss caused by, arising out of, relating to, or resulting from the manufacture, distribution, administration, or use of medical countermeasures . . . except for claims involving ‘willful misconduct’ as defined in the PREP Act.” The authors of a legal blog point out that while the new HHS declaration “is couched in a lot of administrative word salad,” the “prime takeaway is that the scope of tort immunity being conferred . . . is quite broad.”

In fact, the PREP Act became law over significant consumer and congressional opposition. Senator Ted Kennedy and 20 colleagues in Congress wrote a letter to the Speaker of the House and Majority Leader to repeal the Act, characterizing it as “a travesty of the legislative process” and stating that it could “be used to allow manufacturers of virtually any drug or vaccine to escape responsibility for gross negligence or even criminal acts.”

The dissenting lawmakers also accused the Act’s sponsors of creating “an empty shell of a compensation program for injured patients with none of the funding needed to make compensation a reality.”

Safety first

A look at the response to past influenza and coronavirus epidemics provides little reassurance that safety considerations will come first. In addition to the already mentioned adverse events associated with the 2009 swine flu vaccine, we have witnessed treatments that “may have been harmful” and whistleblower lawsuits against false claims of efficacy.

For the moment, our government is prioritizing vaccine development (with the enticing promise of lucrative patents) over existing therapeutics (such as vitamin C and already-FDA-approved drugs) that do not offer comparable financial windfalls.

In light of the immunity from liability guaranteed by the PREP Act during declared emergencies, fast-tracked vaccines are a sweetheart deal for both biopharma and government.

A safer and common-sense approach would direct resources toward examining the merits of existing therapeutics that can be put to immediate use.

The government must not allow Big Pharma and biotech companies to cash in on this catastrophe with speculative, patentable vaccines at the expense of the therapeutics needed to save lives now.

Read the full article at ChildrensHealthDefense.org.

Comment on this article at HealthImpactNews.com.

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Dr. Fauci Lied – Hydroxychloroquine rated ‘most effective therapy’ by doctors for coronavirus in global survey of 6,000 doctors in 30 countries By Valerie Richardson - The Washington Times. *Drug known for treating malaria used by U.S. doctors mostly for high-risk COVID-19 patients.*

Published on: **Apr 10, 2020**

FRN presents Dr. Fauci’s own statements as part of the evidence that he has lied to the public in this ABC news clip below.

By Valerie Richardson – The Washington Times – An international poll of more than 6,000 doctors released Thursday found that the antimalarial drug hydroxychloroquine was the most highly rated treatment for the novel coronavirus.

[The survey](#) conducted by Sermo, a global health care polling company, of 6,227 physicians in 30 countries found that 37% of those treating COVID-19 patients rated hydroxychloroquine as the “most effective therapy” from a list of 15 options.

Of the physicians surveyed, 3,308 said they had either ordered a COVID-19 test or been involved in caring for a coronavirus patient, and 2,171 of those responded to the question asking which medications were most effective.

The U.S. Food and Drug Administration gave chloroquine and its next-generation derivative, hydroxychloroquine, emergency-use authorization Monday for treating the novel coronavirus, although the drug was already being used off-label by some doctors and hospitals for COVID-19 patients.

The survey also found that the most commonly prescribed treatments are analgesics (56%), azithromycin (41%) and hydroxychloroquine (33%).

Azithromycin, known by the brand name Zithromax or Z-Pak, was rated the second-most effective therapy at 32%, followed by “nothing,” analgesics (including acetaminophen), anti-HIV drugs and cough medicine.

Hydroxychloroquine, which is sold under the brand name Plaquenil, was prescribed mainly in the United States for the most severe cases, but not so in other countries.

“Outside the U.S., hydroxychloroquine was equally used for diagnosed patients with mild to severe symptoms whereas in the U.S. it was most commonly used for high risk diagnosed patients,” the survey found.

The 30 nations surveyed included those in Europe, Asia, North America and South America, as well as Australia. No incentives were provided to participate in the poll, conducted March 25-27, according to Sermo.

Hydroxychloroquine usage was most widespread in Spain, where 72% of physicians surveyed said they had prescribed it, followed by Italy at 49%, and least popular in Japan, where 7% had used it to treat COVID-19.

The poll found 23% of U.S. medical professionals had prescribed the drug, which has been FDA-approved for malaria, lupus and rheumatoid arthritis.

Debate about hydroxychloroquine has raged in the United States since President Trump touted it two weeks ago as a potential “game-changer” in the fight against the deadly pandemic, prompting critics to accuse him of peddling unproven remedies, or “snake oil,” as USA Today put it.

Sermo CEO Peter Kirk called the polling results a “treasure trove of global insights for policy makers.”

“Physicians should have more of a voice in how we deal with this pandemic and be able to quickly share information with one another and the world,” he said. “With censorship of the media and the medical community in some countries, along with biased and poorly designed studies, solutions to the pandemic are being delayed.”

The survey also found that 63% of U.S. physicians believe restrictions should be lifted in six weeks or more, and that the epidemic’s peak is at least 3-4 weeks away.

The survey also found that 83% of global physicians anticipate a second global outbreak, including 90% of U.S. doctors but only 50% of physicians in China.

On average, U.S. coronavirus testing takes 4-5 days, while 10% of cases take longer than seven days. In China, 73% of doctors reported getting test results back in 24 hours.

In cases of ventilator shortages, all countries but China said the top criteria should be patients with the best chance of recovery (47%), followed by patients with the highest risk of death (21%), and then first responders (15%).

In China, the survey said doctors prioritized patients at greatest risk of death.

Data suggests that 6% of the U.S. population – 20 million Americans are harboring a retrovirus in their bodies that can develop into an acquired immune deficiency- AIDS. This is not the well-known AIDS caused by HIV, but Acquired Immune Deficiency Syndrome (AIDS) associated with other retroviruses. These non-HIV retroviruses were unintentionally introduced into humans over the past 75 years. It began with trials of polio vaccines and yellow fever vaccines given in the early 1930s. This is when the first recorded cases of Chronic Fatigue Syndrome and autism appeared. It involved the use of laboratory mice to prepare vaccines for human use. [1]

Even though 20 million Americans are likely to be infected, not everyone will develop serious illness. **Retroviruses in the human body are like sleeping giants, very clever and very stealthy. They can infect a person and stay with them for their entire lifetime. Sometimes they shorten life substantially. But, they are just as likely to bring a person to total disability and deny people the opportunity for a normal life. They are quiet until they are activated in immune deficient people.**

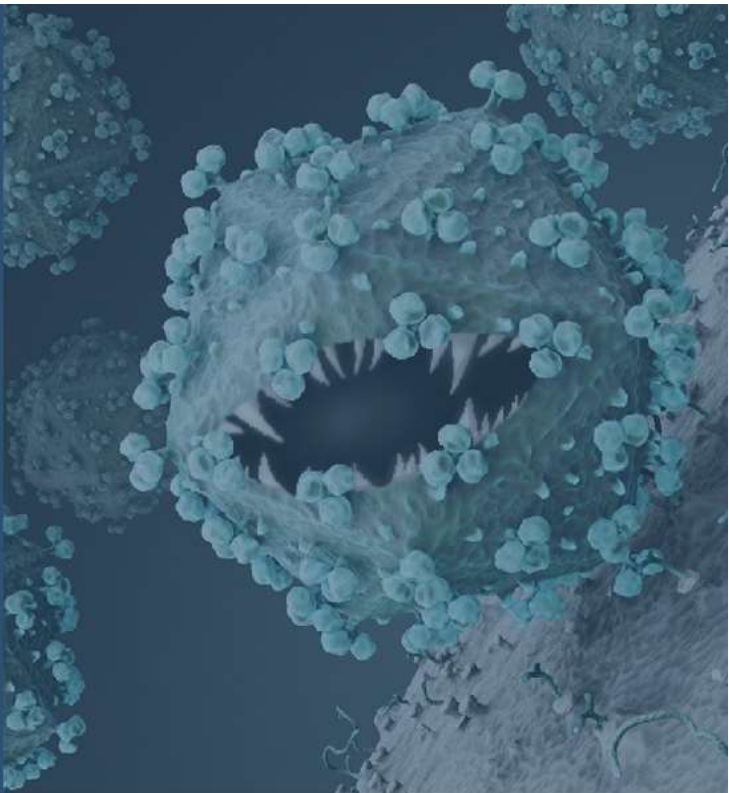
These viruses are associated with many diseases. Once activated, they create diseases such as Myalgic Encephalomyelitis, also called Chronic Fatigue Syndrome (ME/CFS), Chronic Lyme disease, Chronic Lymphocytic Leukemia, autism spectrum disorder (ASD), numerous cancers, and a wide range of other autoimmune, neuroimmune, and central nervous system diseases. Please see reference number 2 at the end of this article for a comprehensive list of diseases. [2]

Meet Dr. Mikovits Ph.D. – Fired, Jailed and Broke For Her Horrifying Discovery of Retrovirus Contamination



Dr. Judy Mikovits discovered vaccines are contaminated with retroviruses infecting 20 million Americans.

Like sleeping giants, retroviruses are quiet until they are activated in immune deficient people. These gammaretroviruses have been associated with cancers, chronic liver disease, AIDS, ALS, ME/CFS, and Autism.



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BY HEALTH FREEDOM IDAHO

**Dr. Judy Mikovits presents
Drugs, Disease and Deception:
Silenced No More**

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Tickets: HealthFreedomIdaho.org/NHS

Dr. Mikovits is one of the leading scientists in the area of retrovirus related illness. Her research showed how retroviruses are linked to the plague of modern illnesses that are bankrupting the U.S. healthcare system. With a well-established history of working for the National Cancer Institute as a cancer research, Dr. Mikovits worked with human retroviruses like HIV. Her work focused on immunotherapy research and involved HIV. In 2009, she was working on autism and related neurological diseases. She found that many of the study subjects has cancer, motor-neuron disorders and chronic fatigue Syndrome (CFS). She believed a virus may have been responsible for these symptoms, and through her research, she isolated the viruses that turned out to come from mice.

She soon realized that these protein and viral contaminants were being introduced into the human population via contaminated vaccines. “Twenty-five million Americans are infected with the viruses that came out of the lab... into the humans via contaminated blood and vaccines.” In response to this discovery, she was fired from her job, indicted, prosecuted, jailed and ordered to retract her research and publicly claim she “made it all up.” She refused to cover up the scientific evidence and was targeted and punished by the “vaccine deep state” establishment.

The result of her unwavering determination to stick to the truth of her research and to stand up against those who want to keep the truth hidden, resulted in her being taken to criminal court and civil court. She was gagged for four years by fabricated criminal charges in Nevada, and could not speak openly about retrovirus science or the government cover-up without risking further persecution/prosecution.

Dr Mikovits is silenced no more.
She will join us as the featured speaker at the
[Natural Health Symposium in Boise July 21, 2018.](#)

In one of the most shocking science videos you’ll see this year, reveals the disturbing true story of how she was thrown in prison for blowing the whistle on deadly viral contamination of human vaccines.

[Dr. Mikovitz’ book, ‘Plaque:’ One Scientist’s Intrepid Search for the Truth about Human Retroviruses and Chronic Fatigue Syndrome \(ME/CFS\), Autism, and Other Diseases.](#)

I first heard of Dr. Judy Mikovitz and her story about Dr. Fauci three years ago on a health sight that I regularly subscribe to and was stunned by her employer, not once, but actually twice. Dr. Anthony Fauci is the Bernie Madoff of Science using the HIV Virus treatment as a Ponzi scheme to make millions. A number of web sites have posted her story of how Dr. Anthony Fauci stole her work only to be published by one of Dr. Fauci’s close friends.

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[alt="Toxic PFAS Can Cross the Placenta Exposing Unborn Children">](#)

Blessings,

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