This Is How Graphene Oxide in All Covid ‘Vaccines’ is Slowly Killing the Vaccinated

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How Long Do the Vaccinated Have to Live?

By Steven Fishman

I deferred this question to a friend of mine, Dr. Mylo Canderian, Ph.D. [born Milos Iskanderianos, Corfu, Greece, 1938], who developed the patent for Graphene Oxide for use as a Hematological Bioweapon in 2015.

In full transparency, Dr. Canderian is what I would call a “Genocidal Globalist,” who follows Precept Ten of the Georgia Guidestones, which is very seldom discussed, stating “Be not a Cancer upon the Earth; Leave Room for Nature.”

Dr. Canderian is a Medical Contributor to the World Health Organization and is also very supportive of Klaus Schwab and the “Great Reset,” ushering in one world digital currency which is a secondary goal of the WHO for 2022.

Dr. Canderian is of the opinion that 95% of the world’s population are “Useless Eaters” who need to be euthanized as quickly as possible.

“Look at downtown Chicago, Baltimore, or Los Angeles,” he has stated, “and you will clearly see why the Useless Eaters must be put down like rabid dogs.”

He has expressed his disdain for “Infectious Educators” who promote Critical Race Theory, and is confident that the “vaccine” will put an end to “Human Cancer Upon the Earth.”

Dr. Canderian is an ardent supporter of Freemasonry’s Duty and Obligation to rid the world of the “Plague of Humanity.”

Yet on a personal level, he and I share a passion for the same exotic dish served at L’emince de Veau in Geneva: Cream of Hummingbird Soup followed by Elk Tongue.

We both are fans of Chef Gaston Sere de Rivieres, who is a culinary genius.

So, I asked Mylo, “How can the “vaccinated” know with certainty how long they have to live once they have been jabbed?”

He presented me with the information, called the “End of Cycle Formula.”
He explained how easy it is to calculate.

“The Power of Simplicity,” he said. “There is a maximum cycle of ten years from injection to End of Cycle,” [or death], he elaborated. “And it is extremely easy to determine.”

He said any hematologist can see it within seconds under a microscope, and even more readily under an electron microscope. “The percentage of blood affected [or contaminated] by or with Graphene Oxide is the reciprocity of the End of Cycle calculation,” he divulged.

In other words, an “inoculatee” [as he calls anyone jabbed with the Experimental Use Authorization Eugenics Depopulation Lethal Injection Bioweapon] having 20% Graphene Oxide deterioration in their blood will, barring any other input criteria, live for 8 years. [10 years less 20%].

Someone with 70% Graphene Oxide deterioration will not live more than 3 years. [10 years less 70%].

Dr. Jane Ruby recently was interviewed by Stew Peters on his podcast and showed examples of what the deteriorated blood looks like when exposed to Graphene Oxide.

Graphene Oxide, for those who are unaware, is the component of Messenger RNA spike proteins and prions, which is at war with the heart, lungs, brain and blood for oxygen.

Graphene Oxide is an oxygen sponge which deprives the body of necessary oxygen and causes many complications, including but not limited to anaphylactic shock, toxic blood clotting, fatal lung paralysis, mitochondrial cancer, and endothelial cancer.”

Dr. Mylo Canderian’s viewpoint is much the same as Klaus Schwab, Bill Gates, and the Big Pharma CEO’s: LET THEM ALL DIE!

I asked Mylo what the effect of second and third shots and boosters do and how that changes the End of Cycle table.

Mylo replied: “It is all measurable through hematological testing. The more shots and boosters the imbeciles get, the worse their blood will look under a microscope, and the quicker they will turn to fertilizer.”

Finally, I asked him how the plot to kill so many billions of people could be kept so secret by such a group of elites.

His answer was: “You don’t know much about Freemasonry, do you, Steve?”
And there you have it.

http://stopthecrime.net/wp/2021/08/08/long-do-the-vaccinated-have-to-live/

Pfizer ‘Vaccine’ Contains 99% Graphene Oxide After Electron Microscope Analysis

Posted on June 30, 2021 by State of the Nation

No mRNA (Gene Therapy) in Pfizer “Vaccine”

(above, Pfizer vaccine; r. Graphene)

Spanish researchers put the Pfizer vaccine under an electron microscope and found it contains 99% graphene oxide and hardly anything else.

There is practically NO evidence that this “vaccine” is gene therapy. There is ZERO genetic material: mRNA or DNA or spike protein. It means this “vaccine” has nothing to do with a so-called virus. This product wasn’t developed to avoid an infection caused by a virus. The true purpose of this product remains hidden. No official statement from governments or health institutions has been made. The toxicity of graphene oxide is reason enough to stop the global vaccination program.
Makow – If this is true, it would discredit a lot of MDs who are saying it is gene altering. Could that be the purpose?

Jim Stone says this graphene story is nonsense. Says graphene is black. The author refutes him below article.

**Related**– Covid Vaccine Delivers Nanoparticles for Mind Control

——— Vaccines Deliver Graphene Oxide Nanotubes for 5G Mind Control

by Des Marxants

(henrymakow.com)

The whole world is following the story released on the Spanish web site program la Quinta Columna.

A June 28th report from the University of Almeria (in Spain) supports the findings of the two researchers hosting the daily broadcast.

Surprising conclusions on the first results of the microscopy on one covid-19 mRNA Comirnaty (Pfizer) sample.

Suspicion of the two researchers turned out to be true. There is almost NO biological matter in the Pfizer covid-19 vial. What the lab in the University of Almeria found has nothing to do with mRNA.

The main matter found was a substance more alike to nanoparticles of graphene or a very similar nanomaterial. Extraction and quantification of mRNA in the sample, identified that 99% of the whole substance inside the vial was highly probably graphene oxide or something very similar, just very little genetic material was found. Until now, no lipid nanocapsules were identified.

The study mentions that there is a reduced form of graphene oxide (GO) When graphene oxide is mixed with hydrogen it becomes MAGNETIC (MGO). It is also known as reduced graphene oxide (this happens when it is injected blood too.)

Graphene oxide layers from scientific literature and from a commercial graphene oxide sample compared against the microscopy performed on the Pfizer sample show identical patterns in both electron and optical microscopy, leaving doubt we’re talking about the same material.
The study offers solid evidence of possible graphene derivatives because the structure seen in the microscopy is characteristic. Academics who previously worked with graphene oxide were consulted and agreed there is no doubt about the presence of graphene oxide in the vial.

The study required by Biostatistician Ricardo Delgado contrasts a graphene oxide sample bought in the market against covid-19 Pfizer mRNA *Comirnaty* [commercial name of the Pfizer/BioNTech product “vaccine.”]

This preliminary study includes optical and electron microscopy conducted by Prof. Dr. Pablo Campra team director in charge. Digital signature here.

It has been only one brand jab but more tests are needed to be done with other vials. More studies are to come with some other different technics. La Quinta Columna makes a call to other institutes and universities to join this research crusade for the truth.

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Related— Jim Stone says this Graphene story is a hoax. If vaccine was mostly graphene, it would be jet black…also can’t pass through a needle.

Des Marxants replies: “In coating technology graphene oxide can be used also. Multilayer graphene oxide films are optically transparent and impermeable under dry conditions. Exposed to water (or water vapor), they allow passage of molecules smaller than a certain size. Glassware or copper plates covered with such a graphene “paint” can be used as containers for corrosive acids. Graphene-coated plastic films could be used in medical packaging to improve shelf life”… wish him luck with his black nose when the nasal version of the covid-19 vaccine get released.

Why is Graphene Transparent? and Stone has made a fool of himself

— A French company is already ready to market 5 G / Covid vaccine packages for “augmented human beings” having received nanoparticles allowing Human-Machine fusion.


among the partners of this society – at the bottom of the page – there is the Thinktank “Next Humanity” which has already defined what will be our future as augmented beings.
Is this why Graphene Oxide was put into the Covid vaccines for connecting to 5G Grid Magnetofection?

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Neurophilosophy

Genetically engineered ‘Magneto’ protein remotely controls brain and behaviour

“Badass” new method uses a magnetised protein to activate brain cells rapidly, reversibly, and non-invasively

Mo Costandi
The Guardian
Researchers in the United States have developed a new method for controlling the brain circuits associated with complex animal behaviours, using genetic engineering to create a magnetised protein that activates specific groups of nerve cells from a distance.

Understanding how the brain generates behaviour is one of the ultimate goals of neuroscience – and one of its most difficult questions. In recent years, researchers have developed a number of methods that enable them to remotely control specified groups of neurons and to probe the workings of neuronal circuits.

The most powerful of these is a method called **optogenetics**, which enables researchers to switch populations of related neurons on or off on a millisecond-by-millisecond timescale with pulses of laser light. Another recently developed method, called **chemogenetics**, uses engineered proteins that are activated by designer drugs and can be targeted to specific cell types.

Although powerful, both of these methods have drawbacks. Optogenetics is invasive, requiring insertion of optical fibres that deliver the light pulses into the brain and, furthermore, the extent to which the light penetrates the dense brain tissue is severely limited. Chemogenetic approaches overcome both of these limitations, but typically induce biochemical reactions that take several seconds to activate nerve cells.

**Remote control of brain activity with heated nanoparticles**

The new technique, developed in Ali Güler’s lab at the University of Virginia in Charlottesville, and described in an advance online publication in the journal *Nature Neuroscience*, is not only non-invasive, but can also activate neurons rapidly and reversibly.

Several earlier studies have shown that nerve cell proteins which are activated by heat and mechanical pressure can be genetically engineered so that they become sensitive to radio waves and magnetic fields, by attaching them to an iron-storing protein called ferritin, or to inorganic paramagnetic particles. These methods represent an important advance – they have, for example, already been used to regulate blood glucose levels in mice – but involve multiple components which have to be introduced separately.

The new technique builds on this earlier work, and is based on a protein called TRPV4, which is sensitive to both temperature and stretching forces. These stimuli open its central pore, allowing electrical current to flow through the cell membrane; this evokes nervous impulses that travel into the spinal cord and then up to the brain.
Güler and his colleagues reasoned that magnetic torque (or rotating) forces might activate TRPV4 by tugging open its central pore, and so they used genetic engineering to fuse the protein to the paramagnetic region of ferritin, together with short DNA sequences that signal cells to transport proteins to the nerve cell membrane and insert them into it.

Try watching this video on www.youtube.com


When they introduced this genetic construct into human embryonic kidney cells growing in Petri dishes, the cells synthesized the ‘Magneto’ protein and inserted it into their membrane. Application of a magnetic field activated the engineered TRPV1 protein, as evidenced by transient increases in calcium ion concentration within the cells, which were detected with a fluorescence microscope.

Next, the researchers inserted the Magneto DNA sequence into the genome of a virus, together with the gene encoding green fluorescent protein, and regulatory DNA sequences that cause the construct to be expressed only in specified types of neurons. They then injected the virus into the brains of mice, targeting the entorhinal cortex, and dissected the animals’ brains to identify the cells that emitted green fluorescence. Using microelectrodes, they then showed that applying a magnetic field to the brain slices activated Magneto so that the cells produce nervous impulses.

To determine whether Magneto can be used to manipulate neuronal activity in live animals, they injected Magneto into zebrafish larvae, targeting neurons in the trunk and tail that normally control an escape response. They then placed the zebrafish larvae into a specially-built magnetised aquarium, and found that exposure to a magnetic field induced coiling manoeuvres similar to those that occur during the escape response. (This experiment involved a total of nine zebrafish larvae, and subsequent analyses revealed that each larva contained about 5 neurons expressing Magneto.)

Researchers read and write brain activity with light

In one final experiment, the researchers injected Magneto into the striatum of freely behaving mice, a deep brain structure containing dopamine-producing neurons that are involved in reward and motivation, and then placed the animals into an apparatus split into magnetised a non-magnetised sections. Mice expressing
Magneto spent far more time in the magnetised areas than mice that did not, because activation of the protein caused the striatal neurons expressing it to release dopamine, so that the mice found being in those areas rewarding. This shows that Magneto can remotely control the firing of neurons deep within the brain, and also control complex behaviours.

Neuroscientist Steve Ramirez of Harvard University, who uses optogenetics to manipulate memories in the brains of mice, says the study is “badass”.

“Previous attempts [using magnets to control neuronal activity] needed multiple components for the system to work – injecting magnetic particles, injecting a virus that expresses a heat-sensitive channel, [or] head-fixing the animal so that a coil could induce changes in magnetism,” he explains. “The problem with having a multi-component system is that there’s so much room for each individual piece to break down.”

“This system is a single, elegant virus that can be injected anywhere in the brain, which makes it technically easier and less likely for moving bells and whistles to break down,” he adds, “and their behavioral equipment was cleverly designed to contain magnets where appropriate so that the animals could be freely moving around.”

‘Magnetogenetics’ is therefore an important addition to neuroscientists’ tool box, which will undoubtedly be developed further, and provide researchers with new ways of studying brain development and function.

Reference
