

## **TO WHOM IT MAY CONCERN**

My name is Sucharit Bhakdi. I am an M.D. who has spent his life practicing, teaching and researching Medical Microbiology and Infectious Diseases. I chaired the Institute of Medical Microbiology and Hygiene at the Johannes Gutenberg University of Mainz, Germany, from 1990 until my retirement in 2012. I have published over 300 research articles in the fields of immunology, bacteriology, virology and parasitology, and served from 1990 to 2012 as Editor-in-Chief of Medical Microbiology and Immunology, one of the first scientific journals of this field that was founded by Robert Koch in 1887.

My name is Arne Burkhardt. I am a Pathologist. I have been teaching at the Universities of Hamburg, Bern and Tübingen. I was invited for visiting professorships/study visits in Japan (Nihon University), the United States (Brookhaven National Institute), Korea, Sweden, Malaysia and Turkey. I headed the Institute of Pathology in Reutlingen for 18 Years, after which I worked as a practicing Pathologist with consulting contracts with Laboratories in the US. I published more than 150 scientific articles in German and International Journals as well as contributions to handbooks in German, English and Japanese. For many years I certified Institutes of Pathology in Germany.

We herewith present scientific evidence that calls for an immediate stop of the use of gene-based COVID-19 vaccines. We first lay out why the agents cannot protect against viral infection. While no positive effects can be expected, we show that the vaccines can trigger self-destructive processes that lead to debilitating illness and death.

### **Why the vaccines cannot protect against infection**

A fundamental mistake underlying the development of the vaccines was to neglect the functional distinction between the two major categories of antibodies which the body produces in order to protect itself from pathogenic microbes:

- The first category (secretory IgA) is produced by immune cells (lymphocytes) which are located directly underneath the mucous membranes that line the respiratory and intestinal tract. The antibodies

produced by these lymphocytes are secreted through and to the surface of the mucous membranes. These antibodies are thus on site to meet air-borne viruses, and they may be able to prevent viral binding and infection of the cells.

- The second category of antibodies (IgG and circulating IgA) occur in the bloodstream. These antibodies protect the internal organs of the body from infectious agents that try to spread via the bloodstream.

Vaccines that are injected into the muscle – i.e., the interior of the body – will only induce IgG and circulating IgA, not secretory IgA. Such antibodies cannot and will not effectively protect the mucous membranes from infection by SARS-CoV-2. Thus, the observed “breakthrough infections” among vaccinated individuals merely confirm the fundamental design flaws of the vaccines. Measurements of antibodies in the blood can never yield any information on the true status of immunity against infection of the respiratory tract.

The inability of vaccine-induced antibodies to prevent coronavirus infections has been reported in recent scientific publications.

### **The vaccines can trigger self-destruction**

A natural infection with SARS-CoV-2 (coronavirus) will in most individuals remain localized to the respiratory tract. In contrast, the vaccines cause cells deep inside our body to express the viral spike protein, which they were never meant to do by nature. Any cell which expresses this foreign antigen will come under attack by the immune system, which will involve both IgG antibodies and cytotoxic T-lymphocytes. This may occur in any organ. We are seeing now that the heart is affected in many young people, leading to myocarditis or even sudden cardiac arrest and death. How and why such tragedies might causally be linked to vaccination has remained a matter of conjecture because scientific evidence has been lacking.

This situation has now been rectified. Histological analyses have been performed on organs of 15 persons who died after vaccination. The age, gender, vaccination schedule, and time of death after injection of each patient are listed in the table below.

Case #	Gender	Age (years)	Vaccine (injections)	Time of death after last injection
1	female	82	Moderna (1. and 2.)	37 days
2	male	72	Pfizer (1.)	31 days
3	female	95	Moderna (1. and 2.)	68 days
4	female	73	Pfizer (1.)	unknown
5	male	54	Janssen (1.)	65 days
6	female	55	Pfizer (1. and 2.)	11 days
7	male	56	Pfizer (1. and 2.)	8 days
8	male	80	Pfizer (1. and 2.)	37 days
9	female	89	Unknown (1. and 2.)	6 months
10	female	81	Unknown (1. and 2.)	unknown
11	male	64	AstraZeneca (1. and 2.)	7 days
12	female	71	Pfizer (1. and 2.)	20 days
13	male	28	AstraZeneca (1.), Pfizer (2.)	4 weeks
14	male	78	Pfizer (1. and 2.)	65 days
15	female	60	Pfizer (1.)	23 days

The following points are of utmost importance.

1. Only 4 patients had been treated in the ICU for more than 2 days. The majority were never hospitalized and died at home (5), on the street (1), at work (1), in the car (1), or in home-care facilities (1). Therefore, in most cases, therapeutic intervention is unlikely to have significantly influenced the post-mortem findings.
2. Not a single death was brought into any possible association with the vaccination by the coroner or the public prosecutor; this association was only established by our autopsy findings.
3. Conventional post-mortems also uncovered no obvious hints to a possible role of vaccination, since the macroscopic appearance of the organs was overall unremarkable. In most cases, “rhythmogenic heart failure” was postulated as the cause of death.

But histological analyses then brought about a complete turnaround. A summary of the fundamental findings follows.

Histopathological findings of similar nature were detected in organs of 14 of the 15 deceased. Most frequently afflicted were the heart (14 of 15 cases) and the lung (13 of 15 cases). Pathologic alterations were furthermore observed in the liver (2 cases), thyroid gland (Hashimoto`s Thyreoiditis, 2 cases), salivary glands (Sjögren`s Syndrome; 2 cases) and brain (2 cases).

A number of salient aspects dominated in all affected tissues of all cases:

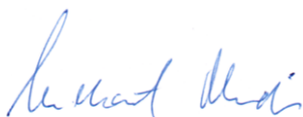
1. inflammatory events in small blood vessels (endothelitis), characterized by an abundance of T-lymphocytes and sequestered, dead endothelial cells within the vessel lumen;
2. the extensive perivascular accumulation of T-lymphocytes;
3. a massive lymphocytic infiltration of surrounding non-lymphatic organs or tissue with T-lymphocytes,

Lymphocytic infiltration was occasionally with signs of intense lymphocytic activation and follicle formation. If present, this was regularly accompanied by tissue destruction (9 cases).

This combination of multifocal, T-lymphocyte dominated pathology that clearly reflects the process of immunological self-attack is without precedent. Because vaccination was the single common denominator between all cases, there can be no doubt that it was the trigger of self-destruction in these deceased individuals.

That myriad adverse events deriving from such auto-attack processes must be expected to very frequently occur in all individuals particularly following booster injections is self-evident.

Beyond any doubt, injection of gene-based COVID-19 vaccines places lives under threat of illness and death. We also note that both mRNA and vector-based vaccines are represented among these cases, as are all four major manufacturers.



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