Vaccines and Production of Negative Genetic Changes in Humans
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Vaccination and Genetic Change: Mobility of Genetic Material Between Life Forms:

One of the indications that vaccinations may in fact be changing the genetic structure of humans became evident in September of 1971, when scientists at the University of Geneva made the discovery that biological substances entering directly into the bloodstream could become part of human genetic structure. Originally, Japanese bacteriologists discovered that bacteria of one species transferred their own specific antibiotic resistance to bacteria of an entirely different species. Dr. Maurice Stroun and Dr. Philip Anker in the Department of Plant Physiology at the University of Geneva, began to accumulate evidence that the transfer of genetic information is not confined to bacteria, but can also occur between bacteria and higher plants and animals. According to an article in World Medicine on September 22, 1971, "Geneva scientists are convinced that normal animal and plant cells shed DNA, and that this DNA is taken up by other cells in the organism."

In one experiment, scientists in Geneva extracted the auricles of frog hearts and dipped them for several hours in a suspension of bacteria. Afterward, they found a high percentage of RNA-DNA hybridization between bacterial DNA extracted from bacteria of the same species as that used in the experiment and titrated DNA extracted from the auricles which had been dipped in the bacterial suspension. Bacterial DNA had been absorbed by the animal cells. This phenomenon has been dubbed transcession. There is evidence that this kind of phenomenon is happening all the time within the human body. It is conceivable, for example, that heart damage following rheumatic fever could be the result of the immune system reacting to its own cells producing a foreign RNA complex after absorption of foreign DNA.

In Science magazine, November 10, 1972, bacterial RNA was demonstrated in frog brain cells after a bacterial peritoneal infection. In the April 1973 issue of the Journal of Bacteriology, transcription of spontaneously released bacterial DNA was found to be incorporated into cellular nuclei of frog auricles. Studies by Phillipe Anker and Maurice Stroun have indicated spontaneous release of DNA material from mammalian cells, spontaneous transfer of DNA from bacteria to higher organisms, spontaneous transfer of DNA between cells of higher organisms, release of RNA by mammalian cells, and biological activity of released complexes containing RNA.

Malignant Cellular Transformations Caused By Foreign DNA:

There is evidence that freely circulating foreign DNA can cause malignancy. In a 1977 issue of International Review of Cytology, Volume 51, Anker and Stroun discuss the possible effects of foreign DNA causing malignant cell transformations. When foreign DNA is transcribed into a cell of a different organism, "this general biological event is related to the uptake by cells of spontaneously released bacterial DNA, thus suggesting the existence of circulating DNA. In view of the malignant transformations obtained with DNA, the oncogenic (cancer-causing) role of circulating DNA is postulated." The discovery in 1975 that viruses causing cancer in animals had a special enzyme called reverse transcriptase makes the problem even more interesting. These kind of viruses are called RNA viruses. When an RNA virus has the reverse transcriptase enzyme within its structure, it allows the virus to actually form strands of DNA which easily integrate with the DNA of the host cell which it infects. Studies by Dr. Robert Simpson of Rutgers
University indicate that RNA viruses which do not cause cancer can also form DNA, even without the presence of reverse transcriptase. DNA formed in this way from an RNA virus is called a provirus. It is known that some non-cancerous viruses have a tendency to exist as proviruses for long periods of time in cells without causing any apparent disease. In other words, they remain latent. Some examples of common RNA viruses that do not cause cancer, per se, but have the capacity to form proviruses are influenza, measles, mumps and polio viruses. In the October 22, 1967 British Medical Journal, it was brought out by German scientists that multiple sclerosis seemed to be provoked by vaccinations against smallpox, typhoid, tetanus, polio, tuberculosis and diptheria. Even earlier, in 1965, Zintchenko reported 12 cases in which MS became evident after a course of anti-rabies vaccinations.

Remember that millions of people between 1950 and 1970 were injected with polio vaccines containing simian virus 40 (SV-40) transferred from contaminated monkey kidney cells used to culture the vaccine. It is impossible to remove animal viruses from vaccine cultures. You are reminded that SV-40, the 40th virus to be discovered in simian tissue, is a cancer-causing virus. Immunization programs against influenza, measles, mumps and polio are in fact seeding humans with RNA and forming proviruses which become latent for long periods in throughout the body, only to re-awaken later on. Post-polio syndrome is a good example of this problem. Other examples may include the so-called mesenchymal and collagen diseases, such as rheumatoid arthritis, multiple sclerosis and lupus erythmatosis, where antibodies are formed by the immune system against the person’s own tissues - tissues which have been impregnated with foreign genetic material. According to a special issue of Postgraduate Medicine in May 1962, "although the body generally will not make antibodies against its own tissues, it appears that slight modification of the antigenic character of tissues may cause it to appear foreign to the immune system and thus a fair target for antibody production." Two years later in 1964, studies were conducted on the polyoma virus, a tumor-producing DNA virus. It was discovered that the persistent genetic DNA material in the polyoma virus brought about malignant transformations in hamster embryo cell cultures. This was reported in the November 23, 1964 issue of the Journal of the American Medical Association.

Even common non-tumor viruses, including those in smallpox vaccine and polio virus 2, can act as carcinogens. It was reported in Science on December 15, 1961 that these common viruses acted as catalysts in producing cancer when given to mice in combination with known organic carcinogens in amounts too small to induce tumors themselves. This means that some vaccinations will induce cancer, when combined with the growing problem of environmental pollution from toxic by-products of agriculture (pesticides on and in food) and industry. Of course, this information is hidden from the public, which is why the FDA, EPA and the agricultural industries can get away with "sanctioning" small amounts of pollutants in food, water and air. The connection has not been made public, much to the joy of the chemical industry, the National Cancer Institute and the growing cancer industry, which continues to fraudulently solicit public donations to justify its own existence. As an aside, it has already been admitted that polio vaccinations have caused 100% of all polio in the United States since 1980 and the predominant cases of all paralytic polio since 1972 (Science, April 4, 1977). It is suspected that the Salk and Sabin vaccines, made of monkey tissue culture, have also been responsible for the major increase in leukemia in the United States.

The use of viruses, bacteria and animal tissue cultures in mass immunization campaigns, considering that this information has been known for 20 years, constitutes an intentionally created hazard to humans. The global impact on the wide range of genotypes relative to human beings is
difficult to assess, but the outcome is definitely negative, and permitting the seeding of latent proviruses in humans, knowingly, can have no other rationale other than future medical profiteering, and constitutes a criminal conspiracy of vast proportions which is tantamount to a genocidal policy against the population, further constituting crimes against humanity, which is internationally punishable by death. But, of course, especially in the United States, this fact is ignored and suppressed from public knowledge, despite a 1984 plea by some U.S. physicians to the United Nations in a report. The fact that this goes on with the full knowledge of the world medical community makes this an international conspiracy where the population has no recourse, given that vaccinations are becoming mandatory and a prerequisite for many social programs.

Persistence of long-term viruses and foreign proteins and their relationship to chronic and degenerative disease was also pointed out by Dr. Robert Simpson of Rutgers University in 1976, when he addressed science writers at an American Cancer Society seminar, saying "these proviruses could be molecules in search of a disease." Dr. Wendell Winters, a virologist at the University of California noted, "immunizations may cause changes in slow viruses and changes in the DNA mechanism." Although host cells containing latent viral particles operate more or less normally, they begin to synthesize viral proteins under the guidance of the viral DNA, eventually creating the circumstances for various autoimmune diseases, including diseases of the central nervous system, which unfortunately add to the growing load of aberrant social behavior patterns.