The 1950s race to be the first with a polio vaccine was led by Jonas Salk and Albert Sabin. Both designed polio vaccines intended to make people immune by exposing them to millions of polio virus. Both would be administered in multiple doses to several hundred million children.

Making so much vaccine required a vast amount of polio virus. There was a fierce debate over what kind of cell to grow this virus in. Some advocated breeding it in fertilized chicken eggs, others in human placental cells grown in laboratory vessels, and others in dishes containing the cells of wild-caught monkeys. Salk and Sabin decided to use monkeys, since they could provide large organs on which the virus would grow readily, and would be a few pence cheaper than the alternatives.

Salk and Sabin must have known that monkey viruses were a serious danger. Sabin had lost a colleague to a monkey virus (Simian Virus B) during vaccine-related research in 1932. And, Dr Herald R Cox, the Principal Bacteriologist of the United States Public Health Service, had forbidden his scientists from using monkeys to make a polio vaccine because of the danger monkey virus represented. Nonetheless Salk and Sabin pressed on with monkeys. They both selected the rhesus monkey found in the temples of northern India. They used their kidneys, since these are large and easy to remove and their testicles, since these are even easier to extract. They calculated they could grow enough viruses on a single kidney to make around 6,000 doses of the vaccine – enough for 2,000 children at 3 doses each. In 1955 this meant they required the kidneys of some 47,710 monkeys for the US – and some 8,000 for the UK vaccine.1

The monkeys were flown via London to the US. On average, half of the monkeys died on route or were rejected as too infected or ill to use on arrival. But some two million wild-caught monkeys arrived in good enough shape to be killed in the West for polio vaccine production and testing over the next decades.

In 1955, the UK adopted the Salk vaccine against the recommendation of its local manufacturer, Wellcome, which wanted instead to use a vaccine it thought safer as it was not grown in monkeys but in fertilized chicken eggs. Sweden and Canada would also refuse to use monkey cells – instead they grew their vaccines’ polio virus on human cells multiplied in laboratories.

In 1954, the scientist in charge of the US government’s safety testing laboratory, Bernice Eddy, made a shocking discovery. Her monkeys, after being dosed with the monkey kidney preparation, had collapsed and died. This should have been the end for the Salk vaccine – but astonishingly it wasn’t. Instead, Eddy was silenced by her employer, the federal National Institutes of Health.

Eddy continued to worry. In 1959 she took matters into her own hands. She went back unauthorised to put the Salk polio vaccine through more tests. She was horrified to find that, when she injected its growth medium into 23 hamsters, 20 of them grew large cancer tumours. She investigated further and found the Salk preparation had infected the hamsters with a monkey virus. This would be named Simian Virus 40 (SV40) as it was the 40th monkey virus discovered. Again her boss would react with fury, and ordered her to remain silent. This time she didn’t. In 1960, at a meeting of the New York Cancer Society, she told them what happened when she had tested the Salk vaccine. She was immediately demoted by the National Institutes of Health. They took her laboratory from her and delayed publication of her research. Meanwhile the Salk vaccine was

**Poisonous vaccines**

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Making the vaccine

To mass vaccinate, the vaccine scientists had to produce a stable 'seed-stock' of poliovirus from which they could breed the huge amounts of virus needed for the vaccine. The process they used was called 'virus propagation'.

They made a suspension in water of diseased spinal tissue from polio victims, and injected this into the living brains of monkeys. They repeated this process countless times, which created millions of monkey viruses. One of these researchers, Leonard Hayflick, wrote in 1958: 'Monkey viruses and DNA fragments, all circulating among great numbers of DNA fragments and much cellular debris, all potentially highly dangerous. This was inevitable, given Salk and Sabin's choice of product methods and the technology available to them.'

proving ineffective. Children vaccinated with it were still coming down in hundreds with polio. The Journal of the American Medical Association would carry an article admitting, 'It is now generally recognized that the vaccine introduced in the USA has been useless for the eradication of poliomyelitis.' By 1959, preparations had begun to replace it with its main rival, the Sabin oral vaccine.

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behind the stories, news of Eddy's unknown research had reached Merck, Sharpe and Dohme, who were then manufacturing both the Salk and Sabin polio vaccines. They put two scientists, Ben Sweet and Maurice Hilleman, on to checking to see if her research on the Salk vaccine also applied to the Sabin. They found it did. In a 1960 paper they reported the 'Sabin live polio virus vaccine was contaminated' and 'SV40 has oncogenic [cancer-causing] properties in hamsters.' They added that this 'raises the important question of the existence of other' monkey viruses.

Asked many years later why they had not warned the public, Hilleman replied; 'Because you could start a panic. They wanted to alarm the public. It would take two years before all the contaminated stocks of vaccine were exhausted. The authorities didn't want to alarm the public. It would take two years before all the contaminated stocks of Salk vaccine were exhausted.'

Salk then sent his patented vaccine 'seed' to various manufacturers where it would be mixed with vast quantities of minced monkey kidney on which the virus would multiply a million-fold. Sabin thus injected the diseased tissue into the brains of about one hundred chimpanzees. A leading scientist, Leonard Hayflick, wrote in 1958: 'Monkey kidney and human viruses, all circulating among great numbers of DNA fragments and much cellular debris, all potentially highly dangerous. This was inevitable, given Salk and Sabin's choice of product methods and the technology available to them.'

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Hence, the poliovirus was weakened by mutation, brought about through rapidly multiplying monkey and human viruses, all circulating among great numbers of DNA fragments and much cellular debris, all potentially highly dangerous. This was inevitable, given Salk and Sabin's choice of product methods and the technology available to them. But again none of this was reported in the newspapers for fear of public panic. The US health authorities did not want to alarm the public. It would take two years before all the contaminated stocks of Salk vaccine were exhausted.

Sabin thus injected the diseased tissue into the brains of 14 monkeys. After the eighth injection, he noticed some of the monkeys were rapidly and repeatedly moving from one lot of monkey cells to another. It was also weakened by having to adjust to growing in different cell types. Both Indian rhesus and African green monkeys cells were employed - thus giving the vaccine 'seed' every opportunity to become contaminated with incompatible viruses from two continents before being bottled as the patented 'Sabin Original Merck' polio virus seed lot. This was 'safety tested' by being injected into the brains of about one hundred chimpanzees.

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In the self-defence the US health authorities have since repeatedly claimed that the measures they took in March 1961 ensured that the polio vaccine was totally clear of SV40 from then on. But this would be exposed as a lie when the post-vaccine correspondence between government and vaccine manufacturers became public in the course of the Salk scandal investigation in 1961. In 1961, the government's man in charge of vaccine safety, a Dr Murray, secretly authorised Lederle Inc. (the major Sabin polio vaccine manufacturer in the US) to use SV40 contaminated vaccine.

On top of this, the same internal memo revealed that the company was not only using the SV40-free African Green Monkeys to make the vaccine but was 'harvesting kidneys' from a monkey species from the Philippines, the cynomolgus, that did carry SV40. And another memo found out into the open revealed that Lederle had totally ignored the FDA regulation that bound manufacturers to ensure 'each seed virus used for manufacturing shall be demonstrated to be free of extraneous microbial agents'. Lederle had even bothered to check to see if they had made sure the Russians would be irresponsibly continued to endorse the polio vaccine.

In 1973, with the withdrawal of Salk, Lederle became the only manufacturer of the Sabin vaccine in the US, and that same year, researchers at the US Bureau of Biologics found its polio vaccine contaminated between 1,000 and 10,000 simian viruses per millilitre of vaccine. In 1978, John Martin, Director of the Virus Oncology Laboratory at the US government's Bureau of Biologics, inspected the samples of polio vaccine held at his lab. He reported: 'There was a lot of extraneous DNA in the vaccine.' But he was told to do nothing about it, since a protest might cause Lederle to stop production and 'vaccine manufacturing was an essential component of industry; this country's protection against potential biological warfare'. John Martin would later discover in damaged human brain cells another monkey virus, SVCMV. He found this was from the African Green Monkey, the same species that are currently used to make the polio vaccine. Thus monkey viruses and DNA fragments continued to be administered to hundreds of millions of children under the guise of the polio vaccine.

The consequences are now coming out in scores of scientific papers. The first human cancers containing SV40 were discovered around 1970. One of these was that of Mark Moreno. He had a large brain tumour removed in 1970, and has since had several operations. His tumour was riddled with SV40. (He is currently suing for compensation.) Many similar cases have since been found.

Yet in 1984 the US Health Minister would assure Parliament that, although the polio vaccine was once contaminated with SV40, American research had showed SV40 to be harmless.

Is the Current Polio Vaccine Safe?

Michael Stewar, Professor of Immunology at the London School of Hygiene and Tropical Medicine, headed a team working on new vaccines, so I asked him about children who fell severely ill after receiving polio vaccines based on living viruses. One of my questions was: ‘Could their parents possibly be right in suspecting the vaccine?’ His reply was: ‘What else would you expect?’ I expressed surprise. He continued, ‘We all know the current living viral vaccines are dangerous – that is why I am heading a team to try to develop safer vaccines.’

Quite simply we still do not have the technology available to completely purify these vaccines; at least at a price the manufacturers are willing to pay. WHO instead recommended a level for maximum vaccine contamination. It was recommended in the mid 1990s that ‘the amount of extraneous DNA [contaminating] biological products should be limited to 100 picograms [100 billionths of a gram] per dose’. This level however seemingly proved ‘unrealistically low’. As the recommended maximum was increased ten billion fold (100 billionths of a gram) in 1993, and the level has been increased again to 11 nanograms (ten billionths of a gram). However, a safety-supervising scientist admitted in 1999 that ‘for live viral vaccines, … it may not be possible to limit the total amount of DNA to ten nanograms’. In case this level of contamination seems inconsequential, it has been believed that ten nanograms is greater than the approximate weight of 250 million poliovirions or 200 million SV40. The setiveness of this level of contamination is still undermined, but it has been noted that the presence of a single SV40 virus, or a piece of DNA, in a cell, may suffice for that cell to be damaged, and possibly made cancerous.

And we still do not know what effect this concentration of DNA, and of SV40 DNA, debris, bacteria and toxins, and the possible resultant re-combinations and mutations of viruses, has had on the same four billion children
In 1988, a review of a study conducted between 1959 and 1965 on 58,807 pregnant women discovered that the risk of brain tumours among offspring of mothers who had received the Salk vaccine was 13 times higher than the risk among offspring of mothers who had not. The conclusion was that the cancers were probably caused by a still-unidentified infection originating in the polio vaccine, which (according to the reviewers) was known to have been contaminated with numerous simian viruses.

Also in 1988, Michele Carbone, a researcher in Chicago, found SV40 in around 85 per cent of the cancers associated with asbestos. It appeared to make this toxin more dangerous. He found it switches off a key human gene, the p53, which helps to protect us from cancers.

In 1997 I attended a National Institutes of Health emergency workshop in Washington called, because laboratories worldwide had found SV40 in over 33 per cent of all the human bone cancers tested and in over 85 per cent of the childhood brain tumours. The FDA that same year also reported: 'The discovery in 1960 that a DNA tumour [carcinogenic] virus, designated simian virus 40 (SV40), was an inadvertent contaminant of rhesus monkey cells...it confronted the scientific and regulatory community with the very problem that they had sought to avoid in vaccine development...'

In 1998 SV40 was found for the first time in English cancers. At that time no laboratory in England was equipped for such a search. It was only found because I went looking for it with colleagues while working on a documentary for Channel 4’s Dispatches. Our team used a laboratory in Italy to test about 20 cancer samples from English patients. We found SV40 present in a bone cancer and in a terminal case of mesothelioma.

Two very recent studies, from Finland and Turkey, found no SV40 in domestic mesothelioma (cancer caused by asbestos) samples but did find it in American and Italian samples. Neither Turkey nor Finland used SV40-contaminated vaccines, while Italy and the US did. Today Finland has one of the lowest rates of mesothelioma in the Western world.

In the last few years SV40 has been linked to more and more cancers, such as Non-Hodkin’s lymphoma, the fifth most common cancer in the US and one that has been rapidly increasing since the contaminated polio vaccine was released.

A recent German study found that if one put SV40 into lactating female rats they all got breast cancer, (as did 70 per cent of the non-lactating) but the SV40 did not stay in the tumours it helped create.

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It seems from all the research that SV40 is dangerous because it is badly adjusted to living in us, perhaps because it only recently infected humans and has not yet adapted to us. It attaches to our cells in such a way that it disables two key immune system defences. It also damages our chromosomes by adding or deleting whole sections. Once inside a cell, Joseph Testa reported, ‘it looks as if somebody set off a bomb inside the cell’s nucleus.’

to which the contaminated polio vaccine has been given in repeated doses through their most vulnerable years.

The evidence seems to lead to the inescapable conclusion that the polio vaccine has been an unmitigated disaster. It was made to stop epidemics of infantile paralysis but they are still happening, and mistakenly tried to do so by targeting a virus that, given the evidence, is most likely never to have been the principal cause of this disease. Instead it has spread monkey viruses and other contaminants around the world, perhaps causing far more serious illness than the poliovirus ever did.

At the root of this disaster as always, lies money. The drug companies made the choice for the UK and much of the rest of the world. They chose to continue to use monkey kidneys instead of safer cells since it was for them a few pennies cheaper a dose, despite knowing that these kidneys carried monkey viruses into the vaccine, despite knowing from early on that at least one of these was linked to cancers. They have thus knowingly and dangerously contaminated our children – and, tragically, are still doing so.

Resources

1. To follow up the footnotes in this report please visit The Ecologist website: www.theecologist.org
2. The author’s website at www.vaccines.plus.com contains many of the documents to which she refers in this special report. These resources can also be obtained on a CD via this website.
3. Other valuable resources:
   • Jim West’s Images of Polioeventitis Website – a new definition of polio
     www.gnc-cties.com/harpub/model.htm
   • A very useful collection of research on vaccines
     www.whatisbo
   • Testimony to Congress on SV40 in 2003
     www.cbsnews.com/4800-10756380-100-1828117900
   • Research on polio vaccine, SIV and HIV
     venus.soci.niu.edu/~sociclass/bmartin/dissent/dissent/SIVandAIDS/
   • A parents’ website – note the great 2003 letter to the US congress www.curealzheimer.org/
   • The No Spray Coalition – trying to stop organophosphate spraying against West Nile Virus – www.nospay.org/
   • Virusesmyth.com – a website that very extensively documents research by top academics on how chemical pollutants could also have led to AIDS.