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Journal of Microbiology, Ipidemiology and Immunobiology, No. 10, 1955; 53-56.

V. S. Silchenks

On the History of Vaccination against Tularemia.

Seventy years ago H. A. Gaiskii was born. A noted Soviet Scienist who began his work early. He took an early part in anti-plague work, adding to it throughout his life.

One particularly great service of his was the preparation of an anti-tularemia vaccine(live), appreciatively high in anti-epidemic properties content.

Work on the anti-tularemia vaccine was began in the USSR in 1931 by Khatenever and Levchenke and Sinai. They prepared a glycerine vaccine on killed tularemia bacteria. Later Khatenever tested the heated, formalinized vaccines and others. Tests were conducted on guines pigs, According to Khatenever, the Quinosel vaccine was the most effective.

In 1931 Khatenever was the first im the workte vaccinate humans against tularemia. Forty-one people were vaccinated with a glycerine vaccine. Unfortunately, the vaccinated wire under observation for only three weeks.

In 1934 Miller and Grahebina occupied themselves with the study of tularemia vaccines. They vaccinated rabbits, gophers and white mice with live and killed (agar and glycerine) vaccines. Upon a subsequent infection of these animals a majority of them died.

In 1935 Sinai tested the protective properties of heated glycerine vaccine on white mice, but single, or quadruple immunisations did not protect them from death. In 1936 Khatenever and Levchenko prepared a vaccine from live, weakly virulent cultures and a polivalent vaccine from killed cultures of B. Talarense. The live, weakly virulent strains were poorly effective and allowed death in a majority of the animals upon infection. The polivalent vaccine, according to Khatenever, gave better results.

In 1937 Miller and Grahebina reported on the local immunisation against bularemia by the skin method, for which they used a partially lysed culture, soda and water antigens. The authors concluded that this method increased the resistance of the organism.

Rhatenever, and then Burgheev under his supervision, immunized rabbits with a thermo-extract. According to the author a just majority of the rabbits survived.

Tosts were made on the use of serums from immunised animals, but according to Miller and Grahebina the serums proved weak in a prophylastic sense.

Along with the study of the killed tularemia vaccines there were vaccinations of a small group of humans. Grahebina vaccinated 46 rats with a formalinized vaccine, but 22 of them quickly died. Gerbunov used a polivalist vaccine on 9 workers of the lab. The results were insignificant, 4 of them quickly became ill. Only in one case did Thatemover note good results during the use of a killed tularemia vaccine. In 1944, 595 people of the Tyumensk region were vaccinated,. after 4-5 months none of them became ill with tularemia.

In foreign countries work along this same line was being conducted. Francis (USA) tested formaline and phenol vaccines. He used sub-lethal deces of tularemia cultures and vaccines of avirulent cultures. The tests gave poor results. Filtrates of virulent cultures gave no peritive results.

Aaki, Kondo and Taxawa (Japan, 1927-1928) immunised rabbits and guines pigs with a suspension of brain from dead animals. The suspension was first boated to 600 for 15 minutes. According to them the results were goed.

M. Endo (Japan 1930 and 1934) used a heated phonol and formalize vaccine and noted the survival of white mice and guines pigs upon infection with B. tularense. He prepared the vaccine from aviralent strains.

In 1932 Downs propared a vaccine with the addition of 0425 formaline to a suspension of microbes in a physiological solution. The vaccinations were given 6-Stimes. The animals lived, but were ill for periods up to 90 days.

Os and Talai Vasfi (Turkey, 1940) immunised animals with a safe endetoxin.

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Gotshlich, Galem Sand Bilal and Tansin Berkin (Turkey 1940) used vaccines of live, weaky strains of tularemia. More than 1/3 of the immunized pigs died from the action of the endetoxin, Almost half of the mice died from the tularemia process, the remaining animals survived a subsequent infection.

Foshey, Hesselbrock, Wittenberg and Hodenberg (USA 1942) prepared a vaccine from virulent strains of B. tularense, worked with a water solution of sodium mitrate and acetic acid. Although this was comsidered to be a most effective vaccine in the USA, its results were not long lambing or very definate, it did not fully pretect the vaccinated person.

Eachll, Beames, Ceriell and Feshay (USA 1950) used phonol and acetone extracts of D. tularense. This vaccine was rated 2/3 effective.

Our survey of all Soviet and foreign literature indicates that the vaccines prepared from killed cultures of B. tularense are ineffective. They required upto 5 applications, and this did not insure a stable, long lasting protection.

The same difficulty was at first experienced with live cultures, this was because the strains were not of a weak virulence and did not pessess high immunegenic qualities.

Specific prophylactics against tularemia were more recently developed by H. A. Gaiskii, who, since 1935, together with B. Y. Elbert, cendnoted studies on tularemia immunity. They found an abd strain of B. tularense with a weak virulence, but with high immunogenic properties (Moscow strain). This strain was tested on 10 volunteers and proven harmless. At the same time it built up antibodies in the orgemism. This strain was lost, and only in 1941 did Gaiskii find a substitute. One of the weak strains, virulent for white mice and avirulent for guines pigs, was named 'Bulennii He, 15, the second---Ondatra IV dry. '..

Further work on the characteristics of strains of B. Tularense was done by Faibich, Maiskii, Ruelyanov and others.

Gaiskii used his weak strains for the preparation of live tularemia vaccines also. In 1942 the first liquid live tularemia vaccine for subsutaneous injection was prepared (called Virusvaccine).

This vaccine was tested on 50 humans in 1942, 6 people acted as control. All were working in the anti-plague lab and volunteers.

These tests confirmed the effectiveness and harmlessness of the vaccine. In 1942 1300 people were vaccinated, in 1943-4214 people. This was the first attempt at mass vaccination against tularemia.

The Gaiskii vaccine had good anti-epidemic properties, but had one deficiency, it quickly deteriated at room temperature. This was, bad for shipping and storage.

In 1944 Gaiskii propared a dry tularemia vaccine. It survived 5 months at 0-2 C. He never finished his work.

Faibich continued Gaiskii's work on the dry vaccine. Using a high vacuum to dry a frozen suspension in a special medium.

Faibich and Tamarin prepared a live dry tularemia vaccine for subcutanteous and cutaneous use. This vaccine survived for two years at a coel temperature. This vaccine is considered as one of the best.

In 1945 Elbert continued a study of the cutaneous method of vaccination. He prepared, together with Tinker, Puchkov and others, a live liquid tularemia vaccine, which allowed for the quick vaccination of large numbers of people.

The en-skin method of vaccination also allowed for the quick detection of immunity. The average skin reaction takes place in 10-15 days. Some reactions are in as little as 2 days, and some 20 days.

Kosmachevski noted other ractions during the introduction of the live vaccine (rise in temperature in 50%, enlarged lymphatic nodes in 30%, etc.). Gaiskii and Khishinskaya noted appearances in 20-40% of all those vaccinated.

The vaccination against tularemia causes the formation of a reaction which can be used to detect the degree of immunity, in many cases 5 years later.

Agglutining in the blood form after 2-4 weeks, and can be detected for 3-5 years. A shorter length of time than the allergic reactions.

Mibert observed that no infections with tularemin took place among these vaccinated, among the non vaccinated the rate was 4,35

Since 1946 the live tularemia vaccime is used extensively for the provention of epidemics. It has proven very effective if the vaccinations can be started with in 3-5 days after the initial start of infections. Thus, both the liquid and dry vaccines can be used, both are very effective.

Observations of health workers (those in contact with tularemia) who had been vaccinated indicate that not one single case of infection feemlted over a 6 year period.

Infection with tulremin of newly vaccinated personnel is rare after the first week. Nost of the cases of infection after vaccination (77.3%) are noted in the first week to twolve days, theothers are from 12-16 days, after which there are very few. This, evidently, is because in the first week the antibodies have not yet been built up to a point of resistance.

The works of various authors list the period of immunity to be from 6 months to 10 years. We believe the average time is 4 yrs., (absence of illness, immunological reaction, no reaction to revacaination.).

Masch credit must be given to Gaiskii, Elbert and Faibich for their work in the study of the talaremia vaccines.