EXPERIENCE WITH PREPARATION OF AN INACTIVATED VACCINE AGAINST AUJESZKY'S DISEASE

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Abstract


Cell lines IBRS-2, PK-Č or primary porcine kidney cells are the most suitable ones for Aujeszky's disease virus propagation for preparation of an inactivated vaccine. Glutaraldehyde-inactivated virus of the Aujeszky's disease proved to be safe in mice and rabbits. Effectiveness of the vaccine was tested in rabbits. Their vaccination followed by revaccination generated immunity for 6 months. Lyophilized inactivated vaccine can be stored for long periods without loss of effectiveness. The vaccine is reconstituted immediately before use by the supplied diluting fluid containing a lipoid adjuvant.

Aujeszky's disease, preparation, inactivated vaccine, cell cultures, glutaraldehyde, safety and potency.

Pork meat production is largely dependent upon sufficient numbers of healthy feedlot piglets. Their successful rearing, however, may be threatened, especially under large-scale systems, by Aujeszky's disease.

In Czechoslovakia, live vaccines have been used to date for immunoprophylaxis of Aujeszky's disease. They were prepared from avirulent Aujeszky's disease viral strains and at the time of their introduction they yielded better results than aluminium hydroxide-inactivated vaccines. The avirulent vaccines decreased considerably animal loss in pig populations.

Recent knowledge of new inactivation substances and the use of adjuvants for potentiation of the antibody response when administered with various antigens enabled the preparation of a new, potent inactivated vaccine against Aujeszky's disease.

Both live avirulent and inactivated vaccines against Aujeszky's disease must contain sufficient amounts of virus. So, for instance, only a sufficient viral content (10^4 TCID_{50}) of the avirulent strain BUK-TK-300/9,2 can induce a good antibody response in sheep (Zuffa 1972). For preparation of inactivated vaccines viral suspensions with substantially higher viral titers are employed (from 10^6.5 to 10^8/cm³). For preparation of an inactivated vaccine the Aujeszky's disease virus propagated in hamster kidney cell line BHK-21 was used by Wittmann and Jakubik (1977). They obtained a viral titer of 10^{5.5}—10^{6} TCID_{50}/cm³.

For virus inactivation, various agents have been used. Ethylenimine was used by Wittmann and Jakubik (1977), acetylatedimine by Gutenkunst (1978), betapropiolactone by Frescura et al. (1977), glutaraldehyde or formaldehyde by Toma et al. (1975), formaldehyde by Zuffa and Neurath (1962), alcohol and saponine by Zuffa et al. (1978). Inactivation by gamma-irradiation at a dose of 0.5 Mrad was used by Dilovski (1973). Possibilities of Aujeszky's disease virus inactivation by ultraviolet light were explored by Lai and Jong (1980).

To increase the antibody response to inactivated vaccines against Aujeszky's disease, Toma et al. (1975) and Delagneau et al. (1975) employed vaseline oil, mannit and emulgin, Wittmann and Jakubik (1977) used DEAE-D, Frescura et al. (1977) used Marcol 52 and Arlacel 80, Lukert et al. (1978) used Tween 80, mineral oil and Arlacel. Gutenkunst (1978) followed the immune response after a vaccine supplemented with lauric acid and aluminium hydroxide.
Žuffa et al. (1978) employed Marcol, Arlacel and Tween 80. All above-mentioned vaccines supplemented with agents potentiating the immune response were liquid.

The present paper deals with preparation of a lyophilized inactivated vaccine against Aujeszky’s disease and its testing in experimental animals — rabbits.

**Materials and Methods**

**Aujeszky’s disease virus**

Cell cultures were infected with Aujeszky’s disease (AD) virus isolated from cattle and propagated in cell line PK-Č.

Potency of the prepared inactivated vaccine was tested by challenging the experimental rabbits infected with a virulent AD virus — lyophilized strain B/200 (Institute for State Control of Veterinary Biologicals and Drugs, Brno).

**Cell cultures**

Cell lines PK-Č and primary cell cultures from pig kidney cultivated in Earle’s solution with LAH were tested for suitability for AD virus propagation. We further tested a cell line from porcine kidney cortex IBRS-2 cultivated in Hanks’ medium, a rabbit kidney cell line RK13, and a monkey kidney cell line VERO cultivated in MEM (Eagle) medium.

**Virus inactivation**

A suitable concentration of the inactivation agent was determined using the viral titer of $10^{8}$ TCID$_{50}$/1 cm$^2$. From inactivation agents glutaraldehyde was tested at final concentrations of 0.025%, 0.05%, 0.1% and 0.2% added to the viral samples. The mixture of the virus and glutaraldehyde was incubated for 2 hours at 34°C in a water bath in tightly closed vials. The inactivated substrate at each of the above-mentioned dilutions was injected intracerebrally into mice weighing 10 g at doses of 0.03 cm$^2$. The animals were observed for 6 days after inoculation.

**Vaccine preparation**

The virus for vaccine preparation was propagated in 3-to-4-day old cell line IBRS-2, infected with 0.05 TCID$_{50}$ of virus per cell. The infected cell culture was cultivated in the maintenance medium without serum at 37°C for 36 to 48 hours. Within this time a marked cytopathic effect occurred. The harvested viral suspension was homogenized, stored at +4°C and then tested for sterility. The virus titer was determined. Preparation of the vaccine required $10^7$ to $10^8$ of viral TCID$_{50}$ per 1 cm$^2$. Before the vaccine preparation proper the cell debris was eliminated by centrifugation or filtration. The virus was inactivated by glutaraldehyde at 0.15% concentration for 2 hours at 34°C. From the inactivated product a sample was taken for inactivation control in mice. The product was mixed with the lyophilic medium and lyophilized.

The lyophilized vaccine was reconstituted in a solvent serving also as a lipoid adjuvant. The solvent was composed of distilled water, paraffin oil, pharmaceutical lanolin and Tween 80.

**Safety and potency tests**

were performed in groups of rabbits weighing 2—2.5 kg (3 animals per group). The rabbits were inoculated i. m. (thigh muscles) with various amounts of the vaccination dose for pigs (see Table 1) amounting 5 cm$^2$ (Jefábek and Dedek 1981). Eleven days later the rabbits were revaccinated with the same dose and in the same way. Ten days later (i.e. 21 days after the vaccination) all rabbits, including the control animals, were challenged by the virulent AD virus at a dose of 1 cm$^2$ containing $10^6$LD$_{50}$ for rabbits. The viral suspension was administered into the thigh musculature of the intact hind limb. An observation period of 14 days followed.

**Evaluation of the safety test**

None in the vaccinated rabbits may show symptoms of AD during the 11-day observation period after the first vaccination and they may not exhibit overall or local postvaccination reactions.

**Evaluation of the potency test**

The vaccine must protect 100% of the experimental animals against a challenge by AD virus in groups 1, 2, 3 (see Table 1). Animals of the groups 4 and 5 may die. All animals of the control group should die within the 14-day observation period after the challenge (Delagneau et al. 1975).
Observation of immunity development in vaccinated rabbits

For the experiment 13 rabbits weighing 2.5 kg each were employed. They were vaccinated intramuscularly with 1 cm³ of the vaccine and revaccinated with the same dose 21 days later. Virus-neutralizing antibodies were assayed prior to vaccination (with negative results) and 21, 35, 97 and 153 days thereafter. For the neutralization tests a micromethod was employed using volumes of 0.05 cm³ and working viral dilutions of 100 to 500 TCID₅₀.

For challenges AD viral doses of 10 000 LD₅₀ per rabbit were used and the animals were challenged 1, 3, 5, 6, 9, 10 and 11 months after vaccination.

Expiration of the vaccine

was determined in lyophilized vaccine samples stored at +4 °C and inoculated into rabbits at various time intervals from 6 to 31 months.

Results

Suitability of cell cultures for AD virus propagation

is shown in Table 2. Highest viral titers were obtained in the IBRS-2 cell line, in primary porcine kidney cells and in the PK-Č cell line. Markedly lower titers were obtained in RK₁₃ and VERO cell lines. In testing the most sensitive cell system the cell lines PK-Č and RK₁₃ yielded best results. By one order less sensitive were the line IBRS-2 and primary porcine kidney cells.

<table>
<thead>
<tr>
<th>Titration cell system</th>
<th>PK-Č</th>
<th>IBRS-2</th>
<th>Primary pig kidney</th>
<th>RK₁₃</th>
<th>VERO</th>
<th>Mean</th>
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</thead>
<tbody>
<tr>
<td>PK-Č</td>
<td>8.3</td>
<td>8.5</td>
<td>8.5</td>
<td>6.5</td>
<td>6.4</td>
<td>7.6</td>
</tr>
<tr>
<td>IBRS-2</td>
<td>7.7</td>
<td>7.7</td>
<td>-</td>
<td>5.3</td>
<td>5.0</td>
<td>5.4</td>
</tr>
<tr>
<td>Primary pig kidney</td>
<td>8.3</td>
<td></td>
<td>7.8</td>
<td>4.8</td>
<td>4.8</td>
<td>6.4</td>
</tr>
<tr>
<td>RK₁₃</td>
<td>8.0</td>
<td>8.2</td>
<td>8.0</td>
<td>5.9</td>
<td>5.8</td>
<td>6.4</td>
</tr>
<tr>
<td>Mean</td>
<td>8.1</td>
<td>8.1</td>
<td>8.1</td>
<td>5.6</td>
<td>5.5</td>
<td>7.2</td>
</tr>
</tbody>
</table>

Virus inactivation

is shown in Table 3, indicating a perfect inactivation of the AD virus by glutaraldehyde at 0.1 % and higher concentrations.
Safety and potency tests

All 14 batches of the vaccine were perfectly safe. Potency of the vaccine was tested in 13 batches. Two of them did not give satisfactory results although their AD viral titer had been sufficiently high prior to inactivation and lyophilization. The reason of this failure remains unknown. The remaining 11 batches of the inactivated vaccine gave satisfactory results.

Post-vaccination immunity in rabbits

In rabbits vaccinated with 1 cm³ of the vaccine and revaccinated 21 days later the highest antibody titer of 51 was found on day 35. On day 97 after vaccination an average titer of 10 was found (Fig. 1).

The control of vaccination effectiveness by challenges of the vaccinated rabbits had shown 100 % of the animals to be protected 1, 3, 5 and 6 months after vaccination. Fifty per cent of the animals were found to be protected 9 and 10 months after vaccination. The control, non-vaccinated rabbits died on days 5 and 6 after challenge. The results are shown in Table 4.

Expiration time of the vaccine

Table 5 shows good effectiveness of all vaccine samples tested in the period of 6 to 31 months.
**Table 5**

Results of potency tests carried out with vaccines stored at +4 °C for determination of expiration time

<table>
<thead>
<tr>
<th>Batch No.</th>
<th>Potency of vaccine stored at +4 °C</th>
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<tbody>
<tr>
<td></td>
<td>Months after production</td>
</tr>
<tr>
<td></td>
<td>6</td>
</tr>
<tr>
<td>V - 01 04 76</td>
<td></td>
</tr>
<tr>
<td>V - 02 06 76</td>
<td></td>
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<tr>
<td>V - 05 03 76</td>
<td></td>
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<tr>
<td>V - 06 04 77</td>
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<td>V - 07 04 77</td>
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<td>V - 11 09 77</td>
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<tr>
<td>V - 12 09 77</td>
<td></td>
</tr>
</tbody>
</table>

**Discussion**

Vaccines containing a live, avirulent AD virus generally confer excellent immunity but the persisting residual virulent virus may limit their use (Delagneau et al. 1975; Lai and Jong 1980). Therefore in several countries including Czechoslovakia possibilities have been explored to produce inactivated vaccines against AD suitable for immunoprophylaxis in swine.

This paper presents preliminary results obtained at preparation of the inactivated vaccine against AD. Among various cell substrates for virus propagation the most suitable were the cell line IBRS-2, PK-Ç and primary porcine kidney cells. The viral titer yields in these cell line cultures are comparable with data of other authors (Delagneau et al. 1975; Frescura et al. 1977). Glutaraldehyde-inactivated AD virus was entirely safe for mice and rabbits. Our results of AD virus inactivation by glutaraldehyde are comparable to those published by Toma et al. (1975).

The two-component form of the vaccine (i.e. lyophilized inactivated viral antigen and the lipoid adjuvant) proved advantageous since in this type of vaccines the decrease in effectiveness is smaller than in those adjuvanted during the manufacturing process. Effectiveness of this vaccine proved to be high in rabbits as shown by good immunogenicity of the glutaraldehyde-inactivated AD virus. Vaccination and revaccination of rabbits with 1 cm³ of the vaccine generated a 100 per cent immunity 6 months thereafter. A protection period of 7 months in rabbits experimentally infected by AD virus reported Toma et al. (1975). Another study of our laboratory presents the results reporting on the use of the inactivated vaccine in immunoprophylaxis of the Aujeszky's disease in pigs (Jerábek and Dedek 1981).
Zkušenosti s přípravou inaktivní vakcíny proti Aujeszského chorobě


Опыт изготовления инактивированной вакцины против болезни Ауски

Для размножения вируса болезни Ауски с целью изготовления инактивированной вакцины наиболее удобно использовать клеточную линию IBRS-2, линию PK-Č или первичные свиные почечные клетки. Глютаральдегидом инактивированный вирус болезни Ауски является совершенно безвредным для мышей и кроликов. Эффективность инактивированной вакцины была испытана на кроликах. Вакцинацией и ревакцинацией был у всех кроликов обеспечен иммунитет продолжительностью 6 месяцев. Лиофилизированная инактивированная вакцина может сохраняться долговременно без потери ее эффективности. Разбавитель, с помощью которого вакцина разбавляется непосредственно перед применением, содержит липидное вспомогательное средство.

References


