Development of a lipopolysaccharide (LPS)-supplemented adjuvant and its effects on cell-mediated and humoral immune responses in male rats immunized against sperm

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Abstract. Supplementation with lipopolysaccharide (LPS) from non-pathogenic Escherichia coli was found to enhance the adjuvant effects of a veterinary vaccine adjuvant (ISA 71VG®). Sperm immunization using 71VG as an adjuvant in the immature period induced infertility in 25% of male rats, whereas this increased to 62.5% after immunization with 71VG + LPS or Freund’s complete adjuvant (FCA). Mean testicular weight of non-sterile males in the 71VG + LPS group was significantly lower than that in the 71VG or FCA group. Histological examination of testicular tissue from sterile males demonstrated severe impairment of spermatogenesis due to experimental autoimmune orchitis, a cell-mediated autoimmune condition. The serum anti-sperm titer was elevated in the three sperm-immunized groups relative to male rats treated with adjuvant alone, but the titer was higher in the 71VG + LPS and FCA groups than in the 71VG group. We consider that this LPS-supplemented adjuvant stimulates both humoral and cell-mediated immune responses to an extent comparable to FCA.

Key words: Adjuvant, Autoimmune orchitis, Lipopolysaccharide (LPS), Rat, Sperm immunization

Fertility control is a non-lethal approach for reducing the population of overabundant wildlife. Gonadotropin releasing hormone and porcine zona pellucida based immunooncontraceptive vaccines are already in practical use in the United States of America [1]. Successful immunoocontraception has been achieved with these two antigens through combination with a veterinary adjuvant, AdjuVac®. This powerful adjuvant was developed by modifying a mycoplasma vaccine, Mycopero®, and was proven not to cause inflammation at the injection site [2]. However, its use in Japan is probably impermissible, since AdjuVac contains killed bacteria (Mycoplasma spp.) thus contravening the domestic Policy for Control of Infectious Diseases in Domestic Animals. Freund’s complete adjuvant (FCA) is a strong adjuvant containing Mycoplasma spp., but it was proven to cause inflammation at the injection site [2], thus making it suitable only for experimental use in terms of animal welfare. With the long-term aim of achieving effective immunooncontraception for density control of wildlife in Japan (e.g., sika deer [3]), the present study was undertaken to develop an alternative adjuvant that would overcome the two problems described above, allowing its registration as a vaccine adjuvant for field use. To achieve efficacy and gain public acceptance, we investigated the effects of adding non-pathogenic Escherichia coli lipopolysaccharide (LPS) to montanide ISA 71VG®, a mineral oil-based water-in-oil-type veterinary vaccine adjuvant. The source and amount of LPS were based on data in previous reports [4, 5]. LPS is a structural component of the outer membrane of Gram-negative bacteria. It consists of three major domains: O-specific chain, core, and lipid A. Lipid A binds to Toll-like receptor 4 on immune cells to activate both innate and adaptive immune responses [6]. A derivative of LPS, mono-phosphoryl lipid A, has been approved as a human vaccine adjuvant [7].

A historic study by Freund et al. in 1953 showed that administration of testis-derived antigens or sperm emulsified in FCA was able to induce experimental autoimmune orchitis (EAO) in guinea pigs [8]. Later, methods to induce EAO in mice [9, 10] and rats [11, 12] were developed with some modification of the immunization procedure, notably intravenous (i.v.) injection of killed Bordetella pertussis to enhance the adjuvant effect of FCA. Subsequent studies involving transfer of T cells from immunized males to syngeneic recipients revealed that the EAO was a result of cell-mediated immunity [13–16], CD4+ T cells in particular playing a leading role [17]. Our preliminary experiments showed that it was possible to induce EAO in rats by immunization in the immature period with sperm emulsified in FCA, without subsequent injection of Bordetella pertussis. After several trials with different doses of sperm, we found that two inoculations of 2 × 10⁷ epidydimal sperm emulsified in FCA induced EAO in over 50% of recipients with a concomitant increase in the serum level of anti-sperm IgG. Therefore, we considered it would be feasible to evaluate the adjuvant’s effectiveness in terms of both cell-mediated and humoral immune responses by replacing FCA in our immunization program.
In order to evaluate the effects of adjuvants on the induction of EAO, we examined the fertility of immunized males at 10 and 20 weeks of age by mating them with intact female rats. The results are shown in Table 1. All of the males examined showed normal libido and copulated with pro-oestrous females. Next morning vaginal plugs were detected in all of the mated female rats. There was no significant difference in the number of implantations among 3 control groups (non-treated, treated with FCA, or treated with LPS-supplemented ISA 71VG), indicating that these adjuvants do not intrinsically interfere with male fertility. One out of 8 male rats immunized with sperm emulsified in FCA (referred to hereafter as sperm+FCA) was infertile at 10 weeks of age. This male and 4 additional males in the group were sterile at 20 weeks of age, while the other 3 males remained fertile with the number of implantations being comparable to those in the 3 control groups. In the group treated with sperm and ISA 71VG supplemented with LPS (referred to hereafter as sperm + 71VG + LPS), 2 out of 8 rats were sterile at 10 weeks of age, and these 2 and an additional 3 males exhibited sterility at 20 weeks of age. The mean number of implantations for non-sterile males indicated in parentheses.

Table 1. Effects of sperm inoculation on fertility in male rats

| Treatment* | n | Fertility** (10 weeks of age) | Fertility ** (20 weeks of age) | Testis weight*** (g) |
|------------|---|------------------------------|-------------------------------|---------------------|
|            |   | /all / fertile / (n)          | /all / fertile / sterile (%)  | /all / sterile (n)  |
| none       | 4 | 16.0 ± 0.50 / 16.0 ± 0.50 (4) | 14.1 ± 0.34 / 14.1 ± 0.34 (4) | 0.4 / 0.4 (0)     |
| + sperm    | 4 | 16.4 ± 1.38 / 16.4 ± 1.38 (4) | 14.8 ± 0.63 / 14.8 ± 0.63 (4) | 0.4 / 0.4 (0)     |
| ISA 71VG + LPS | 4 | 14.4 ± 0.80 / 14.4 ± 0.80 (4) | 15.0 ± 0.84 / 15.0 ± 0.84 (4) | 0.4 / 0.4 (0)     |
| + sperm    | 8 | 13.3 ± 1.94 / 15.2 ± 0.42 (7) | 4.8 ± 2.50 / 14.2 ± 1.42 (3)  | 5/8 (62.5)         |
| ISA 71VG + LPS | 10 | 10.1 ± 3.38 / 13.5 ± 0.00 (3) | 11.0 ± 3.67 / 14.7 ± 0.33 (3) | 1/4 (25.0)         |
| + sperm    | 8 | 11.0 ± 2.43 / 14.7 ± 0.56 (6) | 3.1 ± 1.61 / 8.3 ± 1.59 (3)   | 5/8 (62.5)         |

* Each male was administered adjuvant with/without sperm (2 × 10⁷) at 2 and 4 weeks of age. ** Fertility was examined by mating tests. Each male was placed with a pro-oestrous female and the female was examined for the number of fetuses at 12–14 days after mating. *** Testicular weights were measured at 21–22 weeks of age, and a few at 30 weeks of age. † Means ± SEM were calculated from numbers of males in column “n”. †† Means ± SEM were calculated from the numbers of fertile males indicated in parentheses. a–c Values with different superscripts are significantly different (P < 0.05).

In the immature period had no influence on spermatogenesis after maturation. Mean testicular weight was significantly lower in the sperm + FCA and sperm + 71VG + LPS groups than in the control sperm + 71VG groups. However, it was noteworthy that the testicular weight of fertile males in the 3 sperm-immunized groups was relatively lower than that in the control groups, the sperm + 71VG + LPS group in particular showing a significantly lower value than in the non-treated group. In these smaller testes, some of the seminiferous tubules showed loss or partial depletion of germ cells (Fig. 1B). The testes of sterile males in the 3 immunized groups showed bilateral atrophy, the weight being 0.44–0.67 g, i.e., less than half of that in the 3 control groups. Histological examination of these testes demonstrated EAO with severe impairment of spermatogenesis, resulting in complete loss of differentiated germ cells (Fig. 1C and D). It was noteworthy that marked morphological changes were not evident in the interstitial tissues of the testis, despite the marked damage to seminiferous tubules. Previous studies of EAO had shown that immunization with testicular antigen induced infiltration of considerable amounts of immune cells into the testicular interstitial tissue of mice [10] and rats [11, 12]. These immune cells were shown to produce pro-inflammatory cytokines, such as TNF-α and interferon-γ, which are thought to damage germ cells directly or through modification of Sertoli cell function [17, 18]. The impaired spermatogenesis we observed here, in the absence of marked inflammatory cell infiltration, does not seem to match this type of pathogenesis. In the affected testis of 3 immunized groups, small number of TNF-α-positive cells attaching to the seminiferous tubules were detected (data not shown). These cells might influence Sertoli cell functions, resulting in impaired spermatogenesis. Further investigations are ongoing to clarify the mechanism responsible for spermatogenesis impairment without accumulation of immune cells.

Next, we examined serum titers of anti-sperm antibody to evaluate the effects of adjuvant on humoral immunity. The non-treated group showed a very slight increase in the serum anti-sperm titer as the rats aged (Fig. 2A), the highest value (3.2 × 10³ cps) being detected at 20 weeks of age. The titers in the 71VG + LPS group were indistinguishable from those in the non-treated group. In the FCA-treated group, however, the mean titer was higher than in the non-treated and 71VG + LPS groups after 10 weeks of age, and the difference between this and the other two control groups was significant at 15 weeks of age. Relatively higher titers in 2 out of 4 rats in the FCA-treated group (23.7 and 27.7 × 10³ cps) reflected the mean titer at 20 weeks of age (14.3 ± 6.7 × 10³ cps). The reason for this increase was unclear. Establishment of the humoral immune reaction against sperm was reflected as elevated titers in all of the 3 sperm-immunized groups. In the sperm + FCA group, individual titers ranged between 30.4 and 329.0 × 10³ cps at 8 weeks of age, being at least 20-fold higher than those in the FCA-treated group (Fig. 2C), where the average count was 1.5 ± 0.4 × 10³ cps at the
same age. Individual peak titers in the sperm + FCA group ranged between 117.9 and 418.0 × 10³ cps at 10 or 15 weeks of age, and a declining tendency was detected at 20 weeks of age. In the sperm + 71VG group, individual titers ranged between 69.5 and 204.1 × 10³ cps at 8 weeks of age, increasing slightly thereafter with individual peaks of between 121.3 and 236.3 × 10³ cps (Fig. 2B). These results indicate that 71VG activated the humoral immune response moderately in comparison with FCA. Enhancement of the immune response by LPS supplementation was clearly evident in the sperm + 71VG + LPS group. The titers at 8 weeks of age ranged from 133.0 to 390.1 × 10³ cps, and at this point the titer reached individual peaks of 2 out of 8 rats; in the other 6 rats the titers continued to increase until 15 weeks of age, reaching individual peaks of between 182.9 and 477.4 × 10³ cps (Fig. 2D). Endogenous antigens produced in parallel with spermatogenesis might have stimulated the memory cells generated by immunization during the immature period, the activated immune cells then producing the IgG we detected in 3 sperm-immunized groups. Although the blood–testis barrier in the seminiferous epithelium as well as the tight junctions formed in the epithelium of the efferent and epididymal duct separate sperm antigens from the immune system, it has been reported that there are some specific regions where the barrier is not complete [17]. Discordance of individual reproductive abilities and levels of serum anti-sperm antibody titer in the immunized groups suggests that cell-mediated immunity was fundamentally involved in this disorder [13–16]. The results might also suggest different autoantigenicity of sperm antigens in the humoral and cell-mediated immune response. It is possible that autoimmune orchitis is induced by the immune cells activated with orchitogenic autoantigens among sperm antigens; therefore, we started experiments for identification of sperm antigens.

Fig. 1. Testicular morphology of sperm-immunized rats. Control males (A) administered 71VG+LPS showed normal spermatogenesis. The prevalence of autoimmune orchitis was observed in males immunized with sperm emulsified in 71VG + LPS. In cases of moderate orchitis (B), seminiferous tubules with partial (☆) or almost complete loss of germ cells (★) were observed adjacent to normal tubules. In cases of severe orchitis (C and D), almost all of the tubules had lost germ cells. Inflammatory cells were observed in the blood vessels (arrowheads), but not in the interstitial tissue. The box in C is magnified (D). Scale bar = 100 µm.
causative of the orchitis.

In conclusion, it is evident that the effectiveness of montanide ISA 71VG, a commercial veterinary vaccine adjuvant, for activating the immune response can be enhanced by LPS supplementation. With regard to safety, we did not note any abnormal signs during the experiments, for example injection site inflammation, weight loss or other external features. However, further verification of the safety of this adjuvant will be needed.

Methods

Immunization and sample collection

All the rats used (Iar: Wistar-Imamichi strain) were purchased from the Institute of Animal Reproduction (Kasumigaura, Japan) and maintained in the Institute of Agrobiological Sciences, NARO. All of the experimental procedures were approved by the Committee on the Care and Use of Experimental Animals at the Institute of Agrobiological Sciences, NARO.

Epididymal sperm were collected from the cauda epididymis of adult males, divided into small aliquots, and stored at –80°C until use. For immunization, the thawed sperm were suspended in saline. LPS (from E.coli O127:87, Sigma, St. Louis, USA) was added if necessary at 0.1 mg/kg BW. The suspension was emulsified in an equal volume of FCA (Wako, Osaka, Japan) or montanide ISA 71VG (a gift from Seppic, Paris, France). Immature rats at 12–14 days of age were divided into 6 groups: non-treated, treated with adjuvant alone (FCA or 71VG + LPS), and 3 sperm-immunized groups with FCA, 71VG or 71VG + LPS. Rats were injected with 100 μl of the emulsion including 2 × 10^7 sperm subcutaneously in the back under light ether anesthesia. The second immunization with 200 μl of the
emulsion including $2 \times 10^7$ sperm was performed 2 weeks later. Controls were administered an emulsion of saline and adjuvant. Blood samples were collected from the jugular vein under ether anesthesia at 8, 10, 15 and 20 weeks of age. Fertility of the treated males was examined at 10–11 and 20–22 weeks of age by mating tests; each male was mated with an adult female rat at pro-oestrus overnight. Successful mating was confirmed by the presence of vaginal plugs the following morning. Females were examined for implantation between 12 and 14 days after mating. For the mating test, each male was tested at least twice with an interval of 3–4 days. Testes were collected from the males at 21–22 weeks of age, and at 30 weeks in some cases. After being weighed, the testes were fixed in Bouin's fixative. Paraffin sections were prepared and stained with hematoxylin and eosin for morphological examination.

**Determination of anti-sperm antibody titer**

The anti-sperm antibody titer in serum was determined as follows. Antigens extracted from the sperm were adsorbed onto the wells of a 96-well plate (FluoroNunc, Thermo Fisher Scientific, Waltham, MA, USA) at a concentration of 5 µg/ml (total protein concentration determined by Bradford assay) with coating buffer [19] at 25°C. The antigens used were prepared as follows. Epididymal sperm were suspended in 0.5% CHAPS (Dojindo, Kumamoto, Japan) PBS at a concentration of $10^8$ sperm per ml. The suspension was frozen and thawed three times, centrifuged at 17,000 $\times$ g for 30 min at 4°C, and the supernatant was collected. Rabbit anti-rat IgG (Merck Millipore, Temecula, CA, USA) was labeled with an Eu-chelate of N1-(p-isothiocyanatomethyl)diethylenetriamine-N1,N2,N3,N3-tetraacetic acid (Eu-Labeling reagent; Perkin Elmer, Waltham, MA, USA) by isothiocyanatobenzyl-diethylenetriamine-N1,N2,N3,N3-tetraacetic acid (Eu-Labeling reagent; Perkin Elmer) was added to each well, and the fluorescence in each well had been rinsed, 100 µl of enhancement solution (Perkin Elmer) was added to each well (2 × 10$^6$ counts per second [cps] per 100 µl), and the wells were incubated for 3 h. After the wells had been rinsed, 100 µl of enhancement solution (Perkin Elmer) was added to each well, and the fluorescence in each well was measured with a fluorometer (Arvo, Perkin Elmer). Anti-sperm antibody titer was expressed as cps for each well.

**Statistical analysis**

Statistical analysis was performed using SAS add-in 6.1 for Microsoft® Office. When significance was detected by one-way ANOVA, the significance of differences in means among the groups was determined by Tukey’s test ($P < 0.05$).

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