The Inhibitory Effects by Combined Doses of DDS and several Immunostimulants on the Growth of Leprosy Bacilli Inoculated into Footpads of Hybrid Nude Mice, Jcl:AF-\textit{nu}

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The rapid eradication of leprosy bacilli (LB) with a combination of chemotherapy (CT) and immunostimulation (IS) is an urgent problem in the treatment of leprosy. For examining this synergic effect, a favorable animal model is indispensable whose T cell immunity is enough to be augmentative by IS whereas LB can enough proliferate. Another indispensability is the use of a bacteriostatic drug which can firstly become bactericidal when combined with IS.

A strain of nude mice, Jcl:AF-\textit{nu} established by the hybridization of male FNS/N (\textit{nu}/\textit{nu}) mice with female IAI (+/+) mice in Japan Clea Co. was known to be resistible against infections more than athymic BALB/c mice. On the other hand, the mechanism of action of DDS was well-known to be bacteriostatic.

We examined the inhibitory effects by the combinations of DDS and few immunostimulants on the growth of LB inoculated into footpads (fps) of female Jcl:AF-\textit{nu} mice.

Materials and Methods

Examined agents: DDS, an immunopotentiative $\beta$-1,3-glucan, ATSO and a water-soluble arthrogenous lipoidal amine, 4-aminomethyl-(2,3-(di-n-decyloxy)-n-propyl)-4-phenylpiperidine hydrochloride (CP-46665) were kindly supplied by Daiichi pharm. Co., One Pharm. Ind. Co. and Dr. K. E. Jensen of Central Research, Pfizer Inc., U.S.A., respectively. N-acetyl-muramyl-L-alanyl-D-isoglutamine (MDP) was purchased from Sigma Chemical Co. and its synthetic derivative, N2-MDP-N6-stearoyl-L-lysine (MDP-Lys(L18), muroctasin) was kindly supplied by Daiichi Pharm. Co.

Dosing of agents: DDS and ATSO were given by mixing with a heat-stable pellet form chow, MB-6E (Funabashi Nojyo Co.) in the content ratios of 0.01 or 0.002\% and 0.04\%, respectively. When both DDS and ATSO were dosed, equal quantity of DDS-and ATSO-chows were well mixed. CP-46665 and MDP were dissolved in physiological saline (PS) and filtered through Millex-GS (pore size 0.22 \textmu m, Japan Millipore Ltd.). They were intraperitoneally injected to each mouse once weekly in doses of 12 \textmu g /0.2 ml and 100 or 75 \textmu g /0.2 or 0.15 ml, respectively. Muroctasin aseptically measured into a sterilized glass homogenizer was irradiated for 1 hr by an UV-lamp (National GL-15) at about 30 cm from the lamp. It was homogenized with PS containing a very small amount of tween 80. The homogenate was further diluted to 500 \textmu g /ml with PS. A portion (75 \textmu g /0.15 ml) was intraperitoneally injected to each mouse once per
Experimental animals: Female Jcl:AF (nu/nu) mice aging 4 or 5 weeks were purchased from Japan Clea Co. All of them were fed with MB-6E chow in inside of vinyl isolators (Sanki Kagaku Kogyo Co.). The sterilized water was freely given.

LB inoculated: Strain of LB was Thai-53 received from Dr. K. Kohsaka of this Institute, which had been passaged 5 or 6 times through fps of athymic BALB/c female mice. The 6th passage was used in experiment 1 and the 7th in experiment 2.

Counting of acid-fast bacilli (AFB): Four or six fps of 2 or 3 animals were well minced, triturated and kneaded with 4-or 6-ml PS, respectively. After centrifuged at 330xg for 3 min, 10 μl of the supernatant or its diluted bacterial suspensions were dried with 10 μl each of formol-milk and fixed by exposing to formalin gas for 3 min followed by heating on vapour for 2 min. The staining method was: in 1% carbol fuchsin solution for 28 min; in 1% HCl/70% EtOH for 8-9 sec; in a Löffler solution (methylene blue solution (30 ml) diluted with 100 ml of 0.01% KOH) for 28 sec, at room temperature.

Volumetry of fp swelling: The instrument used was a digital volumeter, MK-550 (muromachi Kikai Co.). The volume of liquid in its cell was adjusted by a sterilized water containing a small amount of tween 80. The volumetry was performed in inside of a clean bench, Hitachi PCW-13038 SG3. In experiment 1, it was once examined just before the last sacrifice of animals. In experiment 2, it was 3 times examined immediately after sacrifice of animals by their neck bone puncture.

Results

The chemical structures of examined immunostimulants are shown in Figure 1.

![Chemical Structures of Examined Immunostimulants](image-url)

Fig. 1 Chemical Structures of Examined Immunostimulants
Experiment 1: Groups of 10 female Jcl:AF-nu mice aging 4 weeks were inoculated with $3 \times 10^7$ LB into both hind fps. Two groups were fed with 0.005%-DDS chow or 0.005%-DDS plus 0.02%-ATSO chow during the 9th-21st weeks after inoculation. The latter group was further given ATSO through 0.02%-ATSO chow during the 22nd-32nd weeks. The other two groups were fed with 0.005%-DDS chow during the same period, but further dose with CP-46665 or MDP during the 9th-32nd weeks. The result is shown in Figure 2. The growth of LB in untreated female Jcl:AF-nu mice was comparable to that in athymic BALB/c female mice (1).

However, when it was compared with a result found in the latter strain of nude mice (1), the growth inhibitory effect of DDS alone was more clearly exhibited in this strain of hybrid nude mice than that in athymic BALB/c mice. No synergic effect was found by the combination of DDS and ATSO. On the other hand, the combined doses of DDS and CP-46665 or MDP completely inhibited the growth of LB until 49 weeks after inoculation. In these 2 groups, a number of AFB were found to be beaded and short rods.

Fig. 2 Inhibitory Effects by Combined Doses of DDS and several Immunostimulants on the Growth of Leprosy Bacilli Inoculated into Footpads of Hybrid Nude Mice, Jcl:AF-nu (Female)

Inoculation: $3 \times 10^7$ leprosy bacilli (Thai-53, the 6th passage) into each of both hind footpads
Periods of doses: DDS, during the 9th-21st weeks after inoculation; the others, during the 9th-32nd weeks
Dosages: ATSO, given through 0.02%-ATSO chow; MDP, 100μg / mouse or CP-46665, 12μg / mouse, intraperitoneally once per week
The result in the volumetry of fp swelling showed no significant difference among treated groups. The difference was found only between treated groups and the control. The volumetry of all the fps was thought to be useful for selecting those animals whose swelling of fps was medium throughout the group.

![Fig. 3 Result in the Volumetry of Footpads in Experiment 1 Examined Just Before the Last Sacrifice of Animals](image)

The small marks on the right side of deviation symbols are the volumes of right hind footpads and those on the left side are the volumes of left hind footpads.

Experiment 2: Groups of 9 female Jcl:AF-nu mice aging 5 weeks were inoculated with $6 \times 10^6$ LB (the 7th passage) into both hind fps. One group was intraperitoneally dosed with muroctasin once weekly covering the 9th-32nd weeks. Another group was further given DDS during the 9th-21st weeks through 0.001%-DDS chow. A group given DDS alone was not examined due to a clear result in experiment 1. The counting of AFB and the volumetry of fp swelling were started from the 217th day after inoculation. The results are shown in Figures 4 and 5.

Unexpectedly, muroctasin itself was found to decrease the count of detected AFB and the fp swelling. The combined dose of muroctasin with the chow containing low concentration of DDS (0.001%) further decreased the count of AFB, but could not achieve the complete inhibition. If each nude mouse weighing 20 g takes daily 3 g of chow, the averaged daily dose of DDS is calculated to be 1.5 mg/kg. In contrast to the complete inhibition after administration of MDP and 0.005%-DDS chow, the inhibitory effect at this dose of DDS was supposed to be insufficient even though combined with muroctasin.
Fig. 4 Inhibitory Effects by Muroctasin Alone or Combined Dose of DDS and Muroctasion on the Growth of Leprosy Bacilli Inoculated into Footpads of Female Jcl:AF-\textit{nu} Mice

Inoculation: \(6 \times 10^6\) LB (the 7th passage) into each of both hind footpads
Periods of Doses: DDS, during the 9th-21st weeks; muroctasin, during the 9th-32nd weeks
Dosages: DDS, given through 0.001%-DDS chow; muroctasin, 75\(\mu\)g / mouse, intraperitoneally once per week

Fig. 5 Result in the Volumetry of Footpads in Experiment 2, Examined Just After Sacrificed Animals
Discussions

It is a well-known fact that both the adaptability of a tumor cell line to a strain of nude mice and the inhibitory effect of an antitumor agent on the tumor cells proliferating in the nude mice are varied with the strain of nude mice. This fact suggests at the same time that an immunological level of nude mice varies with strains.

Macdonald, et al. (2) and Lawetzky, et al. (3) reported the presence of T cell antigen receptor expression in nude mice if the nude mice were aged more than 4-5 months.

Gillis, et al. (4) reported that the nude mouse spleen cells could become responding to alloantigen sensitization in mixed lymphocyte cultures by the presence of a T cell growth factor. Wagner, et al. (5) also reported that the injection of allogenic stimulator cells and IL-2 could differentiate in vivo the nude mouse lymphocytes into alloreactive cytotoxic T lymphocytes. Both reports indicated the presence of T cell immunity in nude mice if they had received an IS pre-treatment.

On the other hand, Kawashima, et al. had reported that a cell population of prethymic T cell precursor was mainly found in the spleen of athymic BALB/c mice whereas it was mainly found in the peripheral lymph nodes in athymic Swiss mice (6). They also suggested that the prethymic cells in both strains of nude mice were derived from the degenerating thymus glands in the nude mouse fetuses. The reason why we dosed the immunostimulants covering 24 weeks or till the end of the 8th month after birth of nude mice was based on these preceding findings in which the possible differentiation of cell subpopulations in aged nude mice was suggested to be responsible for the competence of their T cell immunity.

Practically, Pimm, et al. (7) had found the regression of 2 immunogenic tumor cell lines, a rat sarcoma Mc7 and a rat hepatoma D 23 in their transmitted nude mice after vaccinated them with BCG.

In experiment 1, an antitumor β-1,3-glucan, ATSO with a mild immunopotentiative activity was the substance purified from a crude fraction (corresponding to krestin) of Basidiomycete (Coriolus versicolor) by eliminating its protein component (8). The growth of LB was completely inhibited by a combined dose of DDS and MDP or CP-46665, but could not be inhibited by a combination of DDS and ATSO.

The arthrogenous lipoidal amines have been oncologically used as immunostimulants. CP-20961 (avridine), a strictly lipophilic lipoidal amine, was once examined by us. However, the emulsion dissolving avridine seemed to be injurious and the nude mice intraperitoneally injected the emulsion (an ethanolic solution of avridine-tween 80-soy bean oil (3:1:45.9)) very gradually died. Since avridine was presumed to be unsuitable for such a long term treatment, CP-46665, a freely water soluble lipoidal amine, was examined, whose arthritogenetic potential was found to be comparable to that of avridine (9).

Saiki, et al. (10) and Parant, et al. (11) examined some lipophilic derivatives of MDP such as benzoquinonyl MDP and 1-O-MDP-glycerol-3-mycolate, respectively. The development of MDP derivatives having lipophilic radicals at its muramyl ring such as 6-O-mycoloyl-MDP (12) or those substituted some lipophilic acyl-NH radicals for NH2 in its peptide side residue had been endeavored. Amongst these examinations, Parant, et al. alone found a nonspecific IS activity in a desmuramyl compound, 1-O-(L-alanyl-D-isoglutamine-L-alanyl) glycerol-3-mycolate. Though an
emulsion of MDP and trehalose-6,6'-dimycolate regressed the proliferation of some tumor cell lines and eliminated lymph node metastasis (13), MDP itself and its derivatives have played a main part in these experimental IS and finally reached to muroctasin(14).

It was reported that MDP was analgesic(15) in spite of its inflammatory actions(16). In addition, muroctasin was found to have an inhibitory effect on edema formation(14) in contrast to the induction of edema by MDP(17). These pharmacodynamic characteristics of muroctasin are presumed to be desirable for treatment of leprosy. Its IS effect superior to MDP was reported in details(18).

The complete growth inhibition of LB after administration of 0.005%-DDS chow and MDP in addition to the partial growth inhibition by muroctasin alone indicated the enhancement of antibacterial effect due to an IS effect by these 2 immunostimulants.

The results in experiments 1 and 2 may suggest the usefulness of this strain of hybrid nude mice to be a favorable animal model for examining the inhibitory effects of CT-IS combinations on the growth of LB.

**Summary**

The inhibitory effects by the combined doses of bacteriostatic DDS and several immunostimulants on the growth of leprosy bacilli (LB) inoculated into the footpads of nude mice were examined, using a strain of hybrid nude mice named Jcl:AF-nu, which is resistible to infections more than strain BALB/c-ncu.

The results found were:

1. LB could proliferate up to 10⁹ level in the footpads of this strain of hybrid nude mice.
2. The combined dose of muramyl dipeptide (MDP) or a water-soluble lipoidal amine, CP-46665 and DDS mixed with chow in the content ratio of 0.005% completely inhibited the growth of LB. Whereas, the growth inhibition by DDS alone was only partial.
3. A derivative of MDP, muroctasin could partially inhibit the growth of LB withou combined dose of DDS.
4. The combined dose of an antitumor β-1,3-glucan named ATSO could not enhance the partial growth inhibition due to the dose of DDS through 0.005%-DDS chow.

Based on these results, the possibility of Jcl:AF-nu mice as a favorable animal model for examining the synergic inhibitory effect on the growth of LB due to the combined dose of an antileprous chemotherapeutic and an immunostimulant was discussed.

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**References**


交雑ヌードマウス Jcl:AF-nu 足蹊に移植したらい菌の増殖に対する DDS と二、三の免疫促進物質の併用投与による阻止効果

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キーワード：交雑ヌードマウス、免疫促進、リボイダルアミン、ムラミルジペプチド、ムクロタシン

BALB/c ヌードマウスよりも感染抵抗性の強い交雑ヌードマウス Jcl:AF-nu を用いて、足蹊接種らい菌増殖に対する静菌作用しか持たない DDS と二、三の免疫促進物質の併用投与による阻止効果を調べたところ、次のような結果を得た。

1. らい菌は当交雑ヌードマウス足蹊で、10²レベルまで増殖した。

2. ムラミルジペプチド (MDP) または水溶性リボイダルアミン CP-46665を0.005% DDS 混合飼料で飼育の当交雑ヌードマウスに併用投与するとらい菌増殖を完全に阻止し、一方 DDS のみでは不完全な阻止しか示さなかった。

3. MDP の誘導体であるムクロタシンは、DDS を併用投与しなくても不完全阻止ながららい菌増殖を抑制した。

4. ATSO と呼ばれる抗がん性 β-1, 3-グルカンは、0.005% DDS 混合飼料で飼育のヌードマウスに併用投与しても DDS の阻止効果を増強しなかった。

以上の結果に基づき、Jcl:AF-nu マウスは抗らい化学療法薬と免疫促進物質の併用投与による相乗的らい菌阻止効果を調べるのに好適な動物モデルになりそうな可能性につき討議した。