Making of Vaccines against Human Chorionic Gonadotrophin for Control of Fertility of Women without Impairment of Ovulation and Menstrual Regularity

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Abstract

Reviewed briefly is the development of a unique vaccine against hCG, which prevents unwanted pregnancy without blocking ovulation and disturbing menstrual regularity. A recombinant version of this vaccine, amenable to large-scale production by industry, in both DNA and protein form has been made. Priming with DNA followed by protein improves further its immunogenicity. Mycobacterium indicus pranii (MIP), a cultivable non-pathogenic mycobacteria has been used in the recombinant vaccine as a potent adjuvant. MIP was developed as an immunotherapeutic vaccine against leprosy. It is approved by the Drugs Controller General of India and USFDA, and has many additional applications in treatment of tuberculosis, curing of ugly ano-genital warts and in prevention and treatment of some cancers.

Keywords: Recombinant vaccine; DNA priming; Potent adjuvant; Mycobacterium indicus pranii (MIP)

Introduction

Reproduction is made possible by suppression of the immune response which would normally be raised to the foetus carrying foreign genes of father, besides that of the mother. The objective of this article is not to discuss the considerable work that has been done on this issue, but to project how the immune system can be mobilized to prevent unwanted pregnancy without impairment of ovulation and menstrual regularity. The target of the potential fertility control vaccine is the human chorionic gonadotrophin (hCG). hCG is not secreted by any organ and is not normally present in circulation of the non-pregnant healthy female, the rationale on which diagnosis of pregnancy is based by detection of hCG in blood or urine for occurrence of pregnancy. hCG emerges soon after fertilization of the egg as a product of the early embryo. This was reported initially by Bob (Robert) Edwards, who eventually was honored by a Nobel prize [1].

The fact that hCG plays a crucial role in the implantation of the embryo onto the endometrium was demonstrated by John Hearn, who reported that marmoset embryos exposed to anti-hCG immunoglobins fail to implant, whereas the same embryos exposed to normal immunoglobins implant perfectly [2]. Thus our choice of hCG as a target for the eventual birth control vaccine in 1970s is justified by the above quoted work reported in the literature many years after we published our first paper on an anti-hCG vaccine developed by us [3].

hCG is a hormone composed of two subunits: alpha and beta. The alpha subunit of hCG is common to three other pituitary hormones, TSH, FSH and LH, the beta subunit in each case imparts the individual hormonal properties. It was thus logical for us to employ the beta subunit of hCG for the vaccine. The beta subunit of hCG is however not immunogenic in women. She makes enormous quantities of hCG during pregnancy and her immune system is fully tolerant to it. To make it immunogenic, we linked it with a carrier tetanus toxoid (TT), available at cheap rates in unlimited amounts from industrial sources.

An additional benefit of using TT as a carrier was that in case antibodies were also generated to the carrier, these will accord protection to the woman against tetanus. In those years a large number of deaths due to tetanus used to take place in India following delivery occurring in the field or at home, under aseptic conditions.

Ability of the antibodies generated by the hCGβ-TT vaccine to neutralize hCG bioactivity was tested by administration of a load dose of 5000 IU of hCG to a woman immunized by the vaccine. The vaccine generated in addition antibodies protective against tetanus [3] (Figure 1).

Enhancement of Immunogenicity

Though hCG β-TT generated anti-hCG antibodies in all four women immunized with it, the antibody titres were not sufficiently high to counteract against the high amounts of hCG encountered in early pregnancy. An improved vaccine was made by associating non-covalently hCG beta with alpha subunit of ovine LH to create a hetero-species dimer (HSD) which was
linked to TT. HSD-TT induced higher antibody response than hCG β-TT vaccine as shown in Table 1.

Figure 1: The antibody response against hCG and tetanus in a woman immunized by hCG β-TT vaccine. The antibodies bound with hCG administered to the woman intravenously bringing down the anti-hCG antibody titres, which recovered to the original level in course of time. Antibodies were also raised against tetanus by this vaccine which was additional and independent of those against hCG. Thus, the strategy of immunization with beta subunit of hCG linked to TT as a carrier molecule fulfilled the requirement of a vaccine against hCG, in addition to antibodies against tetanus toxoid.

Table 1: hCG neutralization potency of anti-βhCG and anti-HSD antisera generated in rats and bonnet monkeys, a sub-human primate species.*Rats received three injections of 10 µg gonadotropin equivalent adsorbed on alum at monthly intervals. SPLPS (200 µg) as adjuvant was included in the first injection only. Bleeds were tested 1 week after the last immunization. Bonnet monkeys (Macaca radiata) were given three injections of 50 µg gonadotropin equivalent adsorbed on alum at monthly intervals. SPLPS (1 mg) was included as adjuvant in the first injection only. Bleeds were tested 2 weeks after the last immunization [4].

<table>
<thead>
<tr>
<th>Animal immunized</th>
<th>Immunogen*</th>
<th>HCG binding capacity Mean ± S.E.M. (pg) (l)</th>
<th>HCG neutralization potency Mean ± S.E.M. (pg) (B)</th>
<th>B/I x100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rats (n=6)</td>
<td>β hCG-TT/CHB</td>
<td>27.1 ± 1.7</td>
<td>17.1 ± 1.2</td>
<td>63 ± 1.5</td>
</tr>
<tr>
<td></td>
<td>HSD-TT/CHB</td>
<td>32.5 ± 1.4</td>
<td>26.1 ± 0.8</td>
<td>80 ± 2.3</td>
</tr>
<tr>
<td>Bonnet monkeys (n=5)</td>
<td>β hCG-TT/CHB</td>
<td>22.2 ± 2.3</td>
<td>10.1 ± 1.8</td>
<td>44 ± 3.7</td>
</tr>
</tbody>
</table>

Does Anti-hCG Vaccine Prevent Pregnancy?

After due Regulatory and Ethics committees’ approval, Phase II efficacy trials were conducted in 148 sexually active women of proven fertility. A putative threshold of 50 ng/ml bioeffective anti-hCG titres was set for testing whether at these titres; it prevents pregnancy (Figure 2).

Figure 2: Anti-hCG response to the HSD vaccine in 4 sexually active women of proven fertility. MRG 30 year old and TRW 23 year old had 2 children each; HJN 32 year & SVN 29 year old had 2 children each and 1 elective termination of pregnancy. All of them remained protected from becoming pregnant over 26-32 cycles. The dots (....) at top edge represent the menstrual events which remained regular, solid lines denote the period over which they were exposed to pregnancy. Booster injections were given to keep antibody titres above 50 ngm/ml [5].

Although every woman immunized with HSD-TT vaccine made antibodies against hCG, 110 women had titres above 50 ng/ml for duration of three months or longer. IUD was removed in these women to expose them to pregnancy. Only 1 pregnancy occurred in 1224 of observation cycles indicating a high efficacy of anti-hCG antibodies to prevent pregnancy [5]. All women kept ovulating normally and continued to have regular menstrual cycles. The block of pregnancy was reversible. Fertility was regained on antibodies declining below 30 ngm/ml (Figure 3).
In Figure 3, a 30-year-old subject with two children and one termination of pregnancy remained protected from becoming pregnant over 12 cycles shown by a solid line at the top. The menstrual cycles, which remained regular after immunization, were indicated by (...) in the text. She conceived in the cycle when titles were below 20 ng/ml [5].

The antibody response induced by the vaccine was reversible. The children born to previously immunized women were normal in their developmental landmarks and cognitive abilities as compared to their siblings [6].

**Development of a Recombinant Anti-hCG Vaccine Amenable to Large Scale Industrial Production**

A recombinant hCG-β-LTB vaccine was made, which adsorbed on alum, along with *Mycobacterium indicus pranii* (MIP) used as adjuvant induced high anti-hCG titles in 100% of Balb c mice [7]. The vaccine was also immunogenic in every mouse of three other genetic strains of mice [8]. Figure 4a and 4b indicate the potent adjuvanticity of MIP.

Figure 3: Regain of fertility on decline of antibodies. STS 30-year old with 2 children and 1 termination of pregnancy remained protected from becoming pregnant over 12 cycles shown by solid line at the top. (...) Indicate the menstrual cycles, which remained regular after immunization. She conceived in the cycle when titles were below 20 ng/ml [5].

**What is MIP?**

*Mycobacterium indicus pranii* is a non-pathogenic mycobacteria originally developed by us as an immunotherapeutic and immuno-prophylatic vaccine against leprosy [9]. It was initially coded as M.w. Its ancestry and gene sequence has been determined. As no mycobacterium of this constitution existed in the World Data Bank, it has been named as *Mycobacterium indicus pranii* [10,11]. MIP shares antigens with both M. leprae and M. tuberculosis. Figure 5 is an electron micrograph of this bacillus.

Figure 4: Enhancement of antibody response to hCG-β-LTB vaccine in BalbC mice by MIP. Mice were immunized intramuscularly with 2 μg of the vaccine adsorbed on alum with or without MIP. Primary immunization consisted of 3 injections given at fortnightly intervals followed by a booster on day 60 or 120. The symbols represent the titles in a given mouse. Bars give the geometrical Means [7].

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Besides its properties of invigorating strongly cellular and humoral immune response, MIP as adjunt to drugs, expedites bacterial clearance in both multibacillary leprosy patients and category II, difficult to treat tuberculosis patients [12]. It brings about a dramatic recovery of ugly ano-genital warts [13]. *Mycobacterium indicus pranii* has received the approval of the Drugs Controller General of India and also of USFDA. It is licensed to Cadilla Pharma, and is available to public.

**Further Enhancement of Antibody response to the Recombinant hCG-β-LTB Vaccine**

Recombinant vaccines can be made in both DNA and Protein versions. We employed eukaryotic plasmid VR1020 (DJ) for making the DNA version of the hCG-β-LTB vaccine. Plasmid VR1020 (DJ) is modified version of VR1020 (Vical Inc., San Diego, CA, USA) in which human CpG motifs are incorporated in the plasmid backbone for better adjuvanticity. Our immunization protocol demands initially 3 primary immunizations with the vaccine. While priming thrice with proteinic version of the vaccine achieved fairly high titles employing MIP as adjuvant, priming twice with the DNA version of the vaccine at fortnightly interval followed by the 3rd primary injection of the protein version, generated higher antibody response (Figure 6) than priming thrice with only the proteinic vaccine [14]. It may be stated that DNA vaccine is cheaper to make and is thermostable at room temperature thus requiring no cold chain.

Figure 5: Electron micrograph of autoclaved *Mycobacterium indicus pranii* (MiP).
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Figure 6: (A) Anti hCG antibodies generated in Balb/c mice after 3 injections of protein form of the vaccine administered intra-muscularly. Symbols denote the titres in each mouse and bars represent the Geometric Mean. (B) Anti hCG antibodies titres in Balb/c mice immunized twice with DNA form of the recombinant hCGβ-LTB vaccine for the two primary injections followed by the 3rd injection with protein form of the vaccine. Symbols give the titres in each mouse and bars represent the Geometric Mean. On day 60 the antibody titres in 35 out of 40 mice were much higher than the scale [14].

Present Status of the Anti-hCG Vaccine

The recombinant hCGβ-LTB vaccine has received the approval of RCGM, the National Review Committee on Genetic Manipulation. It has undergone extensive toxicity studies in 2 species of rodents and in marmosets, a sub-human primate species. The vaccine is fully safe and devoid of any side effects and toxicity. Technology has been transferred to a Company, which will make available the vaccine made under GMP conditions for clinical trials to be conducted under the auspices of the Indian Council of Medical Research after due Ethical and Regulatory approvals.

Summary and Concluding Comments

A lot of research is in progress to develop vaccines for control of fertility. Vaccines against antigens on Spermatozoa and Zona pellucida of egg are being made. Amongst the hormones, vaccines against LHRH are effective for control of estrus of dogs and fertility of wild animals [15,16]. Against hCG, the only vaccine which has reached the stage of Phase II efficacy trials, is the one devised by us [5]. A recombinant vaccine amenable to industrial production has received the approval of RCGM and completed toxicity studies. It is expected to go back for clinical trials in the near future. An alternate vaccine based on using the C-terminal peptide of hCGβ demanded the use of a strong oily adjuvant [17]. Besides side-effects, it generated antibodies of lower affinity than the Km of hCG for binding to its receptors. Trials on the vaccine were abandoned in Sweden due to adverse reactions.

References

