Acaciaside-B-enriched fraction of *Acacia auriculiformis* is a prospective spermicide with no mutagenic property

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Abstract

As a part of our continued venture to develop a safe and effective spermicide, we have identified a triterpene glycoside (Acaciaside-B (Ac-B))-enriched fraction (Ac-B-en) isolated from the seeds of *Acacia auriculiformis* and evaluated its spermicidal potential in vitro. Sperm motility was completely inhibited within 20 s at a minimum effective concentration (MEC) of 120 µg/ml. Tests for sperm viability by dual fluoroprobe staining showed the effect to be spermicidal with an EC₅₀ of 35.20 µg/ml. A series of investigations including tests for hypo-osmotic swelling, membrane lipid peroxidation, and electron microscopy document that the spermicidal effect of the fraction involves loss of sperm plasma membrane integrity and dissolution of the acrosomal vesicle – the two most important structural components that play diverse roles in physiological functions of sperm including fertilization. The fraction at 10 × MEC exerted no detrimental effects on in vitro growth of *Lactobacillus acidophilus*, which is considered the major constituent of vaginal microflora that maintains vaginal health. Ames tests performed with different strains of *Salmonella typhimurium* including TA 97a, 98, 100, and 102, which detect mutagens causing bp substitution or frameshifting at G-C or A-T bp, demonstrate no mutagenic potential of the fraction. Significant spermicidal potential with no possible mutagenic effect and adverse impacts on lactobacilli growth attests to the credential of Ac-B-en as a prospective future spermicide for the development of a safe and effective vaginal contraceptive formulation.


Introduction

The population explosion has been a major global concern. Globally, over 200 million pregnancies occur every year and, of these, ~50% are unwanted. A number of contraceptive options are available on the market, but successful prevention of unplanned pregnancies relies not only on access to available products, but also on the product’s acceptability and couple’s willingness and ability to use them effectively. Thus, in making contraceptive choices, couples balance their sexual lives, their reproductive goals, and each partner’s health and safety. The search for a choice that satisfies all three objectives presents significant challenges for women and men throughout the world. Furthermore, what constitutes an ideal or suitable contraceptive method differs not only among individuals, but also as individuals enter different life phases. Therefore, it is important for women and men to have access to a wide variety of contraceptive options, which will allow them to select the suitable ones. Another major problem that has further complicated the situation is the increasing incidence of sexually transmitted infections (STIs) that mainly occurs through heterosexual contact. Women face the greatest risk of acquiring STIs because of substantial mucosal exposure to seminal fluids and the high prevalence of nonconsensual and unprotected sex (Smits et al. 1999, Clara et al. 2004). No cure exists for many STIs of viral etiology; particularly infections caused by human immunodeficiency virus (HIV) and herpes simplex virus. Consequently, prevention is the strategy of choice for controlling the spread of infection. This scenario has given rise to growing interest in combining contraception with STI prevention. The barrier methods such as condoms may prevent pregnancy and block HIV transmission when used appropriately; however, personal preferences and cultural practices have limited the use of condoms. There is a need for female-controlled methods to prevent unplanned pregnancy as well as infection. Development of vaginal microbicides with discerning spermicidal property, especially for formulations that may be available over the counter, constitutes one of the cornerstones of the prevention science agenda for curbing the rising HIV epidemic and unplanned pregnancy. The proposed chemical barriers can be used alone or in combination with a mechanical barrier to
provide increased protection against pregnancy or STIs, or back-up in case of mechanical barrier failure. An ideal product would be female controlled, with minimal systemic exposure and adverse effects, and would have the ability to coat the vagina, cover the cervix and be retained for an extended period of time. The development of such products is of urgent need and has been the focus of a great deal of research activity for the last decade. Nonoxynol-9 (N-9), which was originally thought to have spermicidal as well as virucidal potential, was so far the most widely used effective molecule for spermicidal formulations. But N-9 has been subsequently shown to cause lesion of vaginal epithelium and damages the vaginal microflora to render the user vulnerable to STIs (Richardson et al. 1998). So the development of a suitable alternative is an urgent global need. Many novel compounds that have antifertility and possible antimi-
properties (Setty et al. 1976). Saponins, isolated from Indian medicinal plants, have been reported to have potential spermicidal activity (Farnsworth & Waller 1982). Medicinal plants have long been known as a source of diverse kinds of saponins and terpenoids that represent the effective component of majority of the currently available spermicidal preparations (Farnsworth & Waller 1982). Saponins, isolated from Indian medicinal plants, have been reported to have potential spermicidal properties (Setty et al. 1976). Acacia auriculiformis is a common Indian medicinal plant, the seeds of which are a rich source of triterpinoid saponins including Acaciaside-A (Ac-A) and Acaciaside-B (Ac-B). These acaciasides have been known for anti-helminthic activity, and can induce membrane damage by generating superoxide anions and initiating lipid peroxidation (Sinha Babu et al. 1997). Earlier investigations from this laboratory have demonstrated that an isolate of the extracts of A. auriculiformis seeds comprising of a mixture of Ac-A and Ac-B possesses spermicidal activity (Pakrashi et al. 1991). Subsequent studies have shown that Ac-A and Ac-B individually possess spermicidal properties; but Ac-A is a mutagen, while Ac-B is spermicidal at significantly lower concentrations, and also demonstrates anti-HIV potential in vitro but having no mutagenicity (D Pal, unpublished observations). However, from the practical point of view, the major limitation is its poor yield through extraction, and synthetic preparation of Ac-B is also not possible. During the extraction and isolation procedure, a crude fraction was obtained with high yield that is mostly comprised of Ac-B (> 38% by weight) with no trace of Ac-A, henceforth referred to as Ac-B-enriched (Ac-B-en) fraction. We took an interest in investigating the spermicidal potential and mutagenicity of Ac-B-en fraction and explored whether it can substitute pure Ac-B without compromising the efficacy.

**Results**

**Spermicidal potential**

The effect of test compounds on sperm motility has been presented in Fig. 1. With an increase in the concentration of Ac-B, Ac-B-en, and N-9 (not shown in the figure), there were dose-dependent increases in the immobilization of sperm. The minimum effective concentration (MEC) of Ac-B and Ac-B-en that induced 100% immobilization of sperm in 20 s was found to be 60 and 120 μg/ml respectively, as against 550 μg/ml for N-9. In neither of the treated groups, any revival of motility was recorded following washing and incubation in Baker's buffer.

**Sperm viability assay**

Proportionate distribution of green- (live) and red- (dead) stained spermatozoa was recorded using dual emission filter for SYBR-14 and propidium iodide (PI). In the control set (without Ac-B-en treatment), ~ 95% were viable (green stained). With an increase in the concentration of Ac-B-en, there was a proportionate increase in the population of red-stained dead spermatozoa (Fig. 2). The presence of no viable spermatozoa was recorded following exposure to Ac-B-en at the concentration of 120 μg/ml. A linearized dose–response curve was plotted (Fig. 3). The EC$_{50}$ of Ac-B-en was calculated to be 35.20 μg/ml.

![Figure 1 Dose-dependent sperm immobilizing activity of Ac-B and Ac-B-en](image)

Figure 1 Dose-dependent sperm immobilizing activity of Ac-B and Ac-B-en. The percentage of motile human spermatozoa was determined after 20 s following exposure to the test compounds at different concentrations. All data were adjusted to a normal control motility of 95%. Each point of Ac-B represents the mean ± s.d. values of five independent experiments; while there were 21 determinations against each concentration of Ac-B-en.
Effect on sperm membrane integrity

Over 92% of the control spermatozoa responded to hypo-osmotic solution by swelling and curling of the tail, while 100% of the sperm treated with Ac-B-en at MEC exhibited no response to hypo-osmotic exposure (Fig. 4).

Electron microscopic photographs of human sperm are presented in Fig. 5. As compared with intact plasma membrane and acrosomal vesicles of 92% untreated sperm, all (100%) Ac-B-en-treated sperm exhibited disintegrated plasma membrane with damaged acrosomal cap of various degrees ranging from perforations and vesiculation to complete disintegration.

Transmission electron microscopy

Ultrastructural microphotograph showed considerable membrane damage in 100% of the Ac-B-en-exposed sperm. As compared with the intact plasma membrane surrounding the head of 84% of the control spermatozoa (Fig. 6A), all treated sperm exhibited dissolution of the acrosomal cap, expansion and separation of the plasma membrane from the nucleus (Fig. 6B).

Effect on sperm lipid peroxidation

The spectrophotometric readings demonstrated a dose-dependent increase in the concentration of malondialdehyde (MDA; nmol/10^8 sperm) in concert with an increase in Ac-B-en concentration (Fig. 7). As analyzed by ANOVA, the MDA production by sperm treated with MEC of Ac-B-en (24.99 ± 0.242 nmol/10^8 sperm) was significantly higher (P<0.01) than that of control sperm (20.66 ± 0.266 nmol/10^8 sperm).

In vitro effect on Lactobacillus acidophilus

Ac-B-en at 1×MEC (120 μg/ml) and 10×MEC (1.2 mg/ml) did not significantly affect the growth of Lactobacillus colonies during the 36 h culture period. On the contrary, N-9 at 1×MEC (550 μg/ml) significantly inhibited the growth of Lactobacillus continuing throughout the 36 h span of the culture (P<0.001; Fig. 8).

Test for mutagenicity

Table 1 represents the number of revertant colonies in different treatment groups. Positive control plates, as expected, exhibited higher number of revertants over the spontaneous revertant rate for all tested strains.
In the Ac-B-en-treated groups, by contrast, the revertant colony sizes of neither of the bacterial strains differed significantly from those of the corresponding negative control, regardless of the presence or absence of S9 mix.

Discussion

With an overall objective to search for a spermicidal agent that may effectively serve as the active constituent of vaginal contraceptives, the present investigation evaluated Ac-B-en for its sperm-immobilizing efficacy in a series of in vitro experiments. The results document that Ac-B-en, like Ac-B, exerted a dose-dependent irreversible sperm-immobilizing effect. The MEC of Ac-B-en was found to be 120 μg/ml, which was twice that of pure Ac-B, and the effect was adjudged spermicidal because no revival of sperm motility was demonstrated after the Ac-B-en-exposed sperm were incubated in Baker’s buffer.

The plasma membrane plays a vital role in the process of sperm migration and fertilization (de Lamirande et al. 1997). A number of spermicidal agents are known to execute their effects by structural and functional modulation of the plasma membrane. We therefore took an interest in examining whether the spermicidal effect of Ac-B-en was mediated by adverse modulation of sperm membrane. Two commonly employed techniques were adopted: hypo-osmotic swelling (HOS) test; and dual staining with SYBR-14 and PI.

HOS is a response that reflects the functional integrity of sperm membrane (Jayendran et al. 1984). Following exposure to hypo-osmotic environment, the intact sperm membrane permits free passage of fluids into the cell to reach osmotic equilibrium. As a result, the sperm volume increases and plasma membrane bulges. Since the plasma membrane around the sperm tail fiber is more loosely attached than that around other parts, the sperm tail is particularly susceptible to hypo-osmotic exposure and responds by coiling. This characteristic feature was exhibited in >92% of the sperm in the control set, while the Ac-B-en-exposed sperm showed no such morphological distortion. This observation suggests that the functional integrity of the sperm membrane was lost following exposure to Ac-B-en. This suggestion was additionally supported by the differential reaction of the normal and Ac-B-en-exposed sperm to a living cell nucleic acid stain SYBR-14 and a membrane-impermeable dye PI. A morphologically intact membrane in live sperm offers selective permeability and, therefore, debar entry of fluorescent dye like PI. But as the sperm die, they lose their ability to resist the influx of PI, which upon entering the sperm replaces or quenches the SYBR-14 staining and turns the sperm red. This forms the basis of the dual staining technique to differentiate between live and dead sperm (Garner & Johnson 1995). We observed that following dual fluorescent staining, almost 5% of the total population of control spermatozoa showed PI staining, while the remainders appeared entirely green due to SYBR-14 staining only. By contrast, exposure to gradually increased concentration of Ac-B-en produced a gradual and proportional decrease in SYBR-14-stained green sperm population with parallel increase in PI-stained red sperm. At 120 μg/ml concentration of Ac-B-en, all sperm were positively stained with PI.

Figure 4 Response of control (A) and Ac-B-en-treated (B) human sperm population following exposure to hypo-osmotic solution and evaluated under a phase contrast microscope. Over 92% of control-untreated sperm exhibited HOS response typically characterized by tail coiling, whereas sperm exposed to Ac-B-en at MEC showed no response. Each bar represents the mean ± S.D. of five observations. P<0.0001. Bar = 50 μm.

In the Ac-B-en-treated groups, by contrast, the revertant colony sizes of neither of the bacterial strains differed significantly from those of the corresponding negative control, regardless of the presence or absence of S9 mix.

Figure 5 High resolution scanning electron micrographs (X18,000) of human sperm treated without and with Ac-B-en at MEC. (A) Control sperm shows intact acrosomal cap and plasma membrane around the head and neck regions, while (B) Ac-B-en-treated sperm demonstrates dissolution of the acrosomal cap; (C) histogram shows increased percentage (P<0.0001) of sperm with distorted acrosome in the Ac-B-en-treated groups. The values are based on examining 50 sperm in each group. Bar = 1 μm.
together, these observations left no doubt that Ac-B-en led to the loss of functional and morphological integrity of the sperm membrane. Consistent with these results, the electron microscopy findings also demonstrated that the effect of Ac-B-en on sperm membrane involved the loss of plasma membrane architecture with dissolution of the outer acrosomal membrane.

Nonionic surfactants are amphipathic molecules consisting of a hydrophobic (alkylated phenol derivatives, fatty acids, long-chain linear alcohols, etc.) and a hydrophilic part (generally ethylene oxide chains of various length). Due to this favorable physicochemical property, nonionic surfactants interact not only with proteins but also with membrane phospholipids to modify their structure and permeability. Mammalian sperm cells present highly specific lipid composition comprising a high proportion of polyunsaturated fatty acids, plasmalogens, and sphingomyelins. This unusual structure of the sperm membrane renders it highly susceptible to damage by nonionic surfactants. It was suggested that the double bonds present in nonionic surfactants form allelic radicals that may react with the molecular oxygen present in the membrane and cause lipid peroxidation (Nandi et al. 2004). This mechanism perhaps explains the increased lipid peroxidation of the sperm membrane preparation and loss of membrane integrity under the exposure to Ac-B-en.

One major shortcoming of the presently marketed spermicides is their adverse effects on vaginal ecology. The vagina and cervix constitute a complex ecosystem that includes the vaginal and cervical epithelia, microbial flora, vaginal fluid, and cervical mucus. The normal vaginal flora of healthy women of childbearing age is dominated by lactobacilli. Lactobacillus produces a number of compounds including lactic acid, hydrogen peroxide, lactacin, and acidolin that maintain a low, acidic pH (3.5–5.0; Hawes et al. 1996), and thereby protect against the pathogens that cause STIs including HIV (Kempf et al. 1991, Klebanoff & Coombs 1991). In order to predict any possible adverse effects of Ac-B-en on vaginal microflora, we investigated the impacts of the test fraction on the growth of L. acidophilus, a major species that represents the vaginal lactobacilli population. The present study demonstrates that as compared with control, Ac-B-en up to a tenfold concentration of MEC did not inhibit the growth of L. acidophilus colonies during the period of 36 h of culture. It may therefore be envisioned that Ac-B-en, unlike the existing spermicides, possibly would not disturb the vaginal ecology to invite opportunistic infections. However, the L. acidophilus complex is highly heterogeneous and varies between geographically separated locations (Vásquez et al. 2002). Ac-B-en may be effective against a particular species, but its action against other strains or subspecies of Lactobacillus may be investigated to give more power to the prediction of safety against commensal microorganisms of the human vagina covering different geographical locations.

Because of the intimate contact of a spermicidal agent with human gametes and the possibility that components of the spermicide may be absorbed systemically raise concerns about the possible mutagenic effect of the test substances (Lambert et al. 2004). We evaluated the possibility of mutation via the bacterial reverse mutation test in which histidine-requiring strains of Salmonella typhimurium were used to detect point mutations involving substitution, addition, or deletion of one or few DNA bp. Each of the four strains of Salmonella

Figure 6 Transmission electron micrographs (×100 000) of human sperm samples incubated in the absence or presence of Ac-B-en (at MEC). (A) Control spermatozoa show proper acrosomal cap with intact plasma membrane, while (B) Ac-B-en-treated spermatozoa exhibit dissolution of the acrosomal cap; (C) histogram shows increased percentage (P<0.0001) of distorted sperm in the Ac-B-en-treated group. The values are based on examining 50 sperm in each group.

Figure 7 Generation of malondialdehyde (MDA) as a function of lipid peroxidation by human sperm treated with or without Ac-B-en at varying concentrations. Values on Y-axis represent mean±s.d. value of five determinations. The graph shows dose-dependent increase in the MDA generation following exposure of motile spermatozoa to Ac-B-en (P<0.01).
contains a different type of mutation in the histidine operon. *Salmonella typhimurium* TA100 detects mutagens that cause bp substitutions at G-C pairs; TA97a and TA98 detect frameshift mutagens that damage the correct reading frame of histidine synthesis at G-C bp (Halder *et al.* 2005). Since Ac-B-en increases lipid peroxidation and these strains may not detect the oxidizing mutagens, we also employed the strain TA102 (Windebank *et al.* 2006), which has A-T bp in the primary reversion site. The test was performed with or without metabolic activities by cofactor-supplemented post-mitochondrial fraction (S9). For all test strains, regardless of the presence or absence of S9 mix, the number of revertant colonies in the Ac-B-en-treated groups did not differ significantly from that in the corresponding negative control. The reproducibility of the test results was confirmed. The positive controls used in the assays in the presence or absence of S9 mix showed clear positive responses by the respective test strains, as evidenced by the number of revertant colonies being greater than twofold of the respective negative control value. From the results described above, it may be concluded that Ac-B-en is expected to be safe in terms of mutagenicity.

We further emphasize that although Ac-B-en, like the most popularly used spermicide N-9, is a nonionic surfactant; it differs from N-9 in various respects. Ac-B is a natural compound having molecular weight 1876, which is about threefold that of the synthetic molecule N-9 (molecular weight 617), which indicates a lesser chance of its absorption through vaginal epithelia. Also, the MEC is about fourfold lower than that of N-9 (550 µg/ml). Moreover, N-9-containing preparations...

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**Figure 8** Optical density as the measure of turbidity denoting growth of bacteria colonies during 36 h of culture in the absence (control) and presence of different test compounds. There was a gradual increase in the growth of colonies that reached a plateau after 24 h of culture. Irrespective of the dose of Ac-B-en (1 × MEC and 10 × MEC), the growth of bacterial colony was comparable with that of the control. N-9, however, exerted a constant inhibition over bacterial growth throughout the entire culture period. Each points represent the mean ± S.D. value of five independent cultures.

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**Table 1** Mutagenicity test outcome (number of revertant colonies) in *Salmonella typhimurium* strains TA97a, TA98, TA100, and TA102 on exposure to Ac-B-en with and without S9 activation.

<table>
<thead>
<tr>
<th>Salmonella strains</th>
<th>Without S-9 activation</th>
<th>With S-9 activation</th>
</tr>
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<tbody>
<tr>
<td>TA 97a</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (DMSO)</td>
<td>54.33 ± 6.51*</td>
<td>58.5 ± 5.97*</td>
</tr>
<tr>
<td>Positive control (4-NPD/B(a)P)</td>
<td>295.70 ± 8.96†</td>
<td>146.5 ± 20.01‡</td>
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<tr>
<td>Ac-B-en (0.12 mg/ml)</td>
<td>49.0 ± 6.08*</td>
<td>54.5 ± 6.35*</td>
</tr>
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<td>Ac-B-en (1.20 mg/ml)</td>
<td>51.33 ± 4.04*</td>
<td>56.0 ± 3.92*</td>
</tr>
<tr>
<td>Ac-B-en (12 mg/ml)</td>
<td>52.0 ± 6.25*</td>
<td>58.0 ± 4.32*</td>
</tr>
<tr>
<td>TA 98</td>
<td></td>
<td></td>
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<tr>
<td>Control (DMSO)</td>
<td>21.75 ± 1.71*</td>
<td>19.5 ± 6.56*</td>
</tr>
<tr>
<td>Positive control (4-NPD/B(a)P)</td>
<td>230.30 ± 12.92‡</td>
<td>95.5 ± 9.57†</td>
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<tr>
<td>Ac-B-en (0.12 mg/ml)</td>
<td>19.75 ± 6.65*</td>
<td>19.1 ± 5.88*</td>
</tr>
<tr>
<td>Ac-B-en (1.20 mg/ml)</td>
<td>20.5 ± 3.42*</td>
<td>18.6 ± 3.42*</td>
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<tr>
<td>Ac-B-en (12 mg/ml)</td>
<td>15.25 ± 4.57*</td>
<td>18.0 ± 2.94*</td>
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<tr>
<td>TA 100</td>
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<td></td>
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<tr>
<td>Control (DMSO)</td>
<td>66.0 ± 12.53*</td>
<td>65.67 ± 6.11*</td>
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<tr>
<td>Positive control (NaN₃/B(a)P)</td>
<td>387.7 ± 12.58‡</td>
<td>494.3 ± 14.57‡</td>
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<td>Ac-B-en (0.12 mg/ml)</td>
<td>68.33 ± 9.02*</td>
<td>66.3 ± 6.11*</td>
</tr>
<tr>
<td>Ac-B-en (1.20 mg/ml)</td>
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<td>66.67 ± 4.73*</td>
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<tr>
<td>Ac-B-en (12 mg/ml)</td>
<td>67.33 ± 3.51*</td>
<td>64.1 ± 11.00*</td>
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<tr>
<td>TA 102</td>
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<tr>
<td>Control (DMSO)</td>
<td>118.0 ± 7.94*</td>
<td>121.1 ± 6.31*</td>
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<tr>
<td>Positive control (Daunomycin/B(a)P)</td>
<td>269.0 ± 12.12‡</td>
<td>453.6 ± 15.20‡</td>
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<tr>
<td>Ac-B-en (0.12 mg/ml)</td>
<td>116.0 ± 5.00*</td>
<td>115 ± 2.38*</td>
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<tr>
<td>Ac-B-en (1.20 mg/ml)</td>
<td>110.7 ± 15.53*</td>
<td>119 ± 8.70*</td>
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<tr>
<td>Ac-B-en (12 mg/ml)</td>
<td>117.3 ± 4.16*</td>
<td>117 ± 5.24*</td>
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*Values with different superscripts against each strain under identical culture condition differ significantly (*P* < 0.001). 4-NPD, 4-nitrophenylene diamine; NaN₃, sodium azide; B(a)P, benzo(a)pyrene.

*Represents mean ± S.D. counts of five representative plates.
disturb the vaginal microflora by inhibiting the growth of lactobacilli, leading to an increased risk of developing consequential infections in the genitourinary tract (Richardson et al. 1998); and several studies have documented that N-9-containing vaginal spermicide formulations are mutagenic both with and without liver activation (Lambert et al. 2004). Thus, Ac-B-en appears superior to N-9 with respect to effective concentration, mutagenicity, and vaginal health.

Kabir et al. (2008) have demonstrated that Ac-B inhibits transmission of HIV in vitro at concentrations lower than that of its cytotoxic doses. HIV is known to require intact lipid rafts (highly specialized subregions in cell membranes) for entry into cells and budding of fully infectious particles (Campbell et al. 2001). The lipid-dispersing effect of Ac-B that may disrupt the lipid rafts is postulated to attribute to the anti-HIV property of Ac-B (Kabir et al. 2008). It is also important to note that Ac-B at doses 100-folds of hemolytic index did not show any toxicity on the vaginal epithelium (HN Ray, unpublished observations). Since Ac-B represents the major constituent of Ac-B-en, it seems probable that Ac-B-en would be safe for vaginal epithelia; and its possible anti-HIV potential seems worthy to be explored.

Taking all observations into consideration, we conclude that significant spermicidal activity with apparently no possible mutagenic or adverse effects on vaginal ecology highlights the credentials of Ac-B-en as a prospective alternative to pure Ac-B for the development of spermicidal formulation. However, nonclinical pharmacology/toxicology issues and regulatory considerations that are pertinent to the development of topical spermicides need to be addressed.

Materials and Methods

Isolation of Ac-B-enriched fraction

The air-dried and powdered seeds (1.5 kg) of A. auriculiformis were extracted with methanol (41×3) by percolation. Methanol extract was evaporated to dryness under reduced pressure. The residue (73 g) was suspended in water and extracted successively with ethyl acetate and petroleum ether. The residue (73 g) was suspended in water and extracted successively with ethyl acetate and petroleum ether. The residue (73 g) was suspended in water and extracted successively with ethyl acetate and petroleum ether. The residue (73 g) was poured into a large amount of n-butanol to form a precipitate. The precipitate was filtered and dried in a vacuum desiccator. This process was repeated three times to get Ac-B-en (18 g).

Chemicals

Unless otherwise stated, chemicals used for the preparation of reagents were of analytic grade and purchased from Sigma–Aldrich. Purified water (Milli-Q Biocell System, Millipore Corporation, Bedford, MA, USA) was used throughout the investigations. Disposable plastic wares were purchased from Tarsons Products Pvt Ltd (Kolkata, India). The media for culture of bacteria were purchased from Hi-Media Laboratories Pvt Ltd, Mumbai, India. Dual fluorescent live/dead staining kit was purchased from Invitrogen. The mutagenicity test was performed by using Oxoid nutrient broth No. 2 (Remel Inc., Lenexa, KS, USA). All common chemicals including sodium chloride (NaCl), potassium chloride (KCl), calcium chloride (CaCl2), potassium dihydrogen phosphate (KH2PO4), disodium hydrogen phosphate (Na2HPO4), magnesium sulphate (MgSO4), sodium bicarbonate (NaHCO3), fructose, sodium citrate, thiobarbituric acid (TBA), and trichloroacetic acid (TCA) were purchased from Merck Specialities Pvt Ltd (Mumbai, India).

Working media

Working spermatozoa suspensions were made in Biggers, Whitten, and Whittingham’s medium (BWW; 94 mM NaCl, 4.7 mM KCl, 1.7 mM CaCl2, 1.2 mM KH2PO4, 1.2 mM MgSO4, 7 H2O, 25 mM NaHCO3, 0.5 mM sodium pyruvate, 19 mM sodium lactate, 5 mM glucose, 0.4% BSA, and 0.1% antibiotic (penicillin/streptomycin) solution, pH 7.2).

Semen collection and processing

Human semen samples, collected by masturbation after 3–4 days of sexual abstinence, were obtained from Institute of Reproductive Medicine, Kolkata, with due approval of the ethics committee. Donors gave informed, written consent. Samples having >60×106/ml sperm count with >85% motility and normal morphology were used for the study. The highly motile spermatozoa with forward motility were washed with BWW medium, separated from immotile or sluggishly motile cells by the ‘swim-up’ technique (WHO 1999), and finally resuspended in pre-equilibrated BWW medium to obtain working spermatozoa suspension having concentration of 25–30×106 cells/ml.

Determination of spermicidal activity

The spermicidal effect of Ac-B and Ac-B-en was evaluated by a modified version of Sander and Cramer test (Sander & Cramer 1941) followed by test for reversal of motility. Solutions of pure Ac-B and Ac-B-en having concentrations ranging between 10 and 200 μg/ml were prepared by serial dilutions in BWW.
medium. An aliquot of 20 µl of sperm suspension was mixed with 100 µl of BWW medium (control), or each concentration of Ac-B and Ac-B-en solution at a ratio 1:5 and gently vortexed for 1–2 s. A drop was immediately placed on a glass slide, covered with a cover slip and five different fields were quickly examined under a phase contrast microscope (×100). The results were observed for 20 s and counted for motile sperm. The sperm that lost complete motility within 20 s following exposure to drugs were subsequently tested for motility revival.

The spermatozoa treated with Ac-B and Ac-B-en at 60 and 120 µg/ml respectively, which induced immobilization of 100% sperm, were washed twice in pre-equilibrated BWW medium, resuspended in fresh 250 µl of Baker’s buffer (glucose 3%, Na₂HPO₄·2 H₂O 0.31%, NaCl 0.2%, and KH₂PO₄ 0.01%), and incubated at 37 °C for 60 min. At the end of the incubation, a wet preparation of the treated spermatozoa from each set was made on a glass slide. The preparation was examined under a phase contrast microscope (at 100×) to observe any recovery of motility. Even if a single sperm in the ten fields that were examined showed any sign of jerking or viability, the dose was not recorded as ‘effective dose’. The MEC of Ac-B-en that caused 100% immobilization within 20 s with no subsequent revival of motility was considered to be the MEC.

**Sperm viability assay**

Effect of Ac-B-en on viability of human spermatozoa was evaluated using a dual fluorescent live/dead staining kit consisting of SYBR-14 and PI (Flajshans et al. 2004). Briefly, a 50-fold dilution of the SYBR-14 stock solution was prepared in anhydrous DMSO. Sperm suspensions were treated with 100 µl solution of Ac-B-en of varying concentrations (0–120 µg/ml) or diluted 1:5 with BWW medium (control), identically to that during determination of spermicidal activity by Sander–Cramer method. Immediately after 20 s, 2 ml of BWW medium were added, and sperm were washed and centrifuged twice. The sperm pellet was resuspended in 1 ml BWW medium to which 5 µl of diluted SYBR-14 dye were added. After 5 min of incubation, 5 µl of PI were added and finally incubated for 10 min at 37 °C. The fluorescent staining of sperm was monitored and photographed with a Zeiss Axioshot epifluorescent microscope (Carl Zeiss Inc., Thornwood, NY, USA) equipped with a FITC filter set (Zeiss #487909). The proportionate number of live sperm over total number of cells (%) which was graphically plotted against each concentration of the test solution, was linealized and EC₅₀ (the concentration that induced death of 50% spermatozoa) was assessed. Each point on Y-axis represents mean ± s.d. (%) of five independent observations with respective concentrations of Ac-B-en.

**HOS test**

The physiological integrity of the sperm plasma membrane was evaluated by HOS test (Jayendran et al. 1984) using the kit developed by National Institute of Health & Family Welfare, New Delhi, India, for sperm function tests. Sperm were treated without (control) or with Ac-B-en at MEC for 20 s and exposed to 500 µl of hypo-osmotic solution. This was mixed gently and incubated at room temperature for 5 min followed by the addition of 50 µl of color stop solution. After thorough mixing, a tiny drop of the mixture was placed on a glass slide and covered with cover slip. The percentage of spermatozoa exhibiting characteristic swelling or tail coiling was counted in a total population of 1000 sperm per specimen under a phase contrast microscope (×400). The percentage of HOS response was calculated by counting the number of sperm showing a characteristic morphologic changes divided by total number of sperm and multiplied by 100. The data were collected from five independent sets of experiment and presented as mean±s.d.

**Electron microscopy**

Topographical imaging of membrane domains over the sperm head (Wilborn et al. 1983) was done by high resolution scanning electron microscopy (SEM). Control (untreated) human spermatozoa and Ac-B-en-exposed ones at MEC were fixed in 1% paraformaldehyde and 1% glutaraldehyde in 0.05 M phosphate buffer (PB; pH 7.2) for 3 h followed by alternate spinning (5 min at 900 g) and washing for three times. After the final washing, suspensions were placed on 0.1% poly-L-lysine-coated glass chips and allowed to adhere for 1 h at room temperature. Samples were post-fixed in 1% osmium tetroxide in 0.5 M PB for 1 h at room temperature and subsequently dehydrated through an ascending series of ethanol, critical point dried and coated with Au–Pd (80:20) using a sputter coater (Emitech K-575X). All samples were examined using a Field Emission SEM (Philips FEI) at an accelerating voltage of 30 kV. Spermatozoa were observed initially under low magnification (×2000–5000) and representative spermatozoa were photographed under intermediate magnification (×18 000–20 000). For each group, 50 spermatozoa were scanned for the intactness of the acrosomal region. The difference between the treated and untreated groups with respect to intactness of the acrosomal vesicle was done by χ²-test.

**Transmission electron microscopy**

Transmission electron microscopy was performed according to the method as described by Souada et al. (2007). Briefly, control (untreated) and Ac-B-en-treated spermatozoa were fixed in 2.5% glutaraldehyde in 0.2 M PB (pH 7.4) for 2 h at 4 °C, and then post-fixed overnight in 1% osmium tetroxide. The sample was dehydrated in alcohol, embedded in spurr resins, and polymerized for 48 h. Ultrathin sections were cut and stained with uranyl acetate and lead citrate and observed under the transmission electron microscope (Philips TECHNEI) at low (<25 000) and high (<100 000) magnification. Fifty sperm from the untreated as well as treated groups were evaluated for the intactness of sperm head membranes and photograph of one representative sperm from each group has been presented. The difference between the treated and untreated groups with respect to intactness of the sperm head membranes was done by χ²-test.

**Determination of lipid peroxidation**

The magnitude of lipid peroxidation of sperm was measured by determining MDA production using TBA as per the method of Buege & Aust (1978) subsequently modified by Suleiman et al. (1996). Ac-B-en was serially diluted in BWW medium to make solutions of final concentrations ranging between 20 and 120 µg/ml and mixed with sperm suspension at 5:1 ratio. Briefly, 1 ml suspension of spermatooza (25–30 × 10⁶ cells/ml) treated with or without different concentrations of Ac-B-en was mixed with 2 ml of TBA–TCA reagent (15%, w/v TCA; 0.375%, w/v TBA; and 0.25 M HCl). The mixture was boiled in a water bath for 30 min. After cooling, the suspension was centrifuged at 1500 g for 10 min. The supernatant was then separated, and absorbance was measured at 535 nm. The results were expressed as a simple concentration of MDA (nmol/10⁸ sperm) as determined by the specific absorbance coefficient (1.56 ± 10⁸/mol per cm²).

\[
\text{MDA produced (µmol/ml)} = \frac{\text{OD} \times 10^6 \times \text{total volume (3 ml)}}{1.56 \times 10^5 \times \text{test volume (1 ml)}} 
\]

\[
= \frac{\text{OD} \times 30}{1.56}.
\]

**Effect on L. acidophilus in vitro**

The effect of Ac-B-en on the growth of *L. acidophilus* was evaluated in vitro. Lympholized *L. acidophilus* obtained from Lactobacil, Organon India Ltd, Gujarat, India, and was cultured in sterile Lactobacilli MRS broth medium maintained at 37 °C and 5% CO₂ for 24 h before each experiment. Briefly, culture tubes were prepared with 2 ml of autoclaved MRS broth to which was added 100 µl of vehicle as control or Ac-B-en solution to a final tube concentration of 120 µg/ml and 1.2 mg/ml, which were representative of 1× MEC and 10× MEC, respectively. N-9 was used at 550 µg/ml as a positive control. The tubes were inoculated with spores of ~ 10⁻⁵ *L. acidophilus* population. The culture tubes were maintained at 37 °C in an atmosphere containing 5% CO₂ and 95% air for a total period of 36 h with intermittent measurement of optical density (turbidity) at 600 nm at 6 h intervals. This experiment was performed five times in duplicates. Results are presented as mean ± S.D.

**Mutagenicity test**

Mutagenicity of the test substance was assessed in a bacterial reverse mutation assay by using a preincubation method (Maron & Ames 1983) that detects frameshift mutagens (*Salmonella* strains TA 97a and TA 98), bp mutagen (TA 100), or both (TA 102). The test was conducted following preincubation in the presence or absence of S9 mix without or with Ac-B-en at doses of 84, 840, and 8400 µg/plate that represented final tube concentration of 1 ×, 10 ×, and 100 × of MEC. Ac-B-en was diluted in DMSO, which also served as the negative control. In experiments without external metabolic activation, 4-nitrophenylene diamine (20 µg/plate) served as the positive control for TA 97a and TA 98, while sodium azide (NaN₃; 1.5 µg/plate) and daunomycin (6 µg/plate) were used for TA100 and TA102 respectively. For experiments with metabolizing enzyme system, however, benzo(a)pyrene (1 µg/plate) served the purpose for all strains. For each treatment, 0.1 ml of the test substance, negative control, or positive control solution were mixed with 0.5 ml of 0.1 mol/l sodium PB (pH 7.4) in a sterilized tube, to which 0.1 ml of the respective bacterial suspension (~10⁸ bacteria/ml) was subsequently added. Assays with metabolic activation involved substitution of 0.1 mol/l sodium PB with 0.5 ml of 59 mix. The mixture was incubated with gentle shaking for 20 min at 37 °C (pre-incubation). After this incubation period, 2 ml top agar (0.5% agar dissolved in 0.5% sodium chloride in double-distilled water supplemented with 0.05 mM D-biotin/l-histidine solution) was added and transferred to a petridish (85 mm in diameter) with ‘minimal agar’ (1.5% agar with 5% glucose and 4% Vogel–Bonner Medium-E in double-distilled water). After the overlaid agar had solidified, the plates were incubated for 48 h at 37 °C and the numbers of revertant colonies were subsequently evaluated manually. The level of microbial toxicity was confirmed for background lawn with a stereoscopic microscope. All samples were tested five times in two independent experiments. Results are presented as mean number of revertants ± S.D., and the statistical analysis was done by one-way ANOVA.

**Statistical analysis**

The results were analyzed by paired two tailed t-test, χ²-test, and one-way ANOVA, as applicable, using the GraphPad Prism 3.0 software (GraphPad Software, Inc., San Diego, CA, USA). *P* < 0.05 was considered significantly different.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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