Abstract

Background: Evaluation of microbicides for prevention of HIV-1 infection in macaque models for vaginal infection has indicated that the concentrations of active compounds needed for protection by far exceed levels sufficient for complete inhibition of infection in vitro. These experiments were done in the absence of seminal plasma (SP), a vehicle for sexual transmission of the virus. To gain insight into the possible effect of SP on the performance of selected microbicides, their anti-HIV-1 activity in the presence, and absence of SP, was determined.

Methods: The inhibitory activity of compounds against the X4 virus, HIV-1 IIIB, and the R5 virus, HIV-1 BaL was determined using TZM-bl indicator cells and quantitated by measuring β-galactosidase induced by infection. The virucidal properties of cellulose acetate 1,2-benzenedicarboxylate (CAP), the only microbicide provided in water insoluble, micronized form, in the presence of SP was measured.

Results: The HIV-1 inhibitory activity of the polymeric microbicides, poly(naphthalene sulfonate), cellulose sulfate, carrageenan, CAP (in soluble form) and polystyrene sulfonate, respectively, was considerably (range = 4 to = 73-fold) diminished in the presence of SP (33.3%). Formulations of micronized CAP, providing an acidic buffering system even in the presence of an SP volume excess, effectively inactivated HIV-1 infectivity.

Conclusion: The data presented here suggest that the in vivo efficacy of polymeric microbicides, acting as HIV-1 entry inhibitors, might become at least partly compromised by the inevitable presence of SP. These possible disadvantages could be overcome by combining the respective polymers with acidic pH buffering systems (built-in for formulations of micronized CAP) or with other anti-HIV-1 compounds, the activity of which is not affected by SP, e.g. reverse transcriptase and zinc finger inhibitors.

Background

Sexual virus transmission plays the major role in the worldwide HIV-1 epidemic [1]. In the absence of effective anti-HIV-1 vaccines, great emphasis has been put on the development of topical microbicides to be applied vaginally in the form of gels, creams or solid dosage formulations expected to inactivate HIV-1 infectivity or to interfere with steps in the virus life cycle, preferably blocking virus entry into susceptible cells. The model of choice for evaluating candidate anti-HIV-1 microbicides in vivo...
are female rhesus macaques to whom anti-HIV-1 products and either simian immunodeficiency virus (SIV) or HIV-1/SIV hybrid viruses (SHIVs) are consecutively applied in the vagina [2-7]. Results obtained in this animal model have indicated that the concentrations of anti-HIV-1 compounds in microbicide formulations adequate to prevent vaginal infection exceed by several orders of magnitude concentrations sufficient for complete inhibition of infection in in vitro systems [8-10]. The macaque model overlooks the role of human seminal plasma (SP), a common source of male to female sexual transmission of HIV-1, in infection and the ultimate effectiveness of microbicides. Because of impediments for including SP into the macaque model studies, the effect of this “natural diluent for HIV-1” on virus inhibitory activity of several candidate microbicides was investigated. They included the polymers: carrageenans, poly(naphthalene sulfonate) (PRO 2000), cellulose sulfate, cellulose acetate 1,2-benzenedicarboxy late (CAP) and polystyrene sulfonate, some of which are being evaluated in phase III clinical trials for efficacy [10-12]. Antiretroviral drugs specifically targeted to HIV-1 reverse transcriptase, UC781 [12,13] and TMC 120 [14,15], respectively, and to the zinc fingers of the HIV-1 nucleocapsid protein NCp7 [16-18] were included in control experiments.

**Methods**

**Reagents**

Aquateric (the micronized form of CAP containing ≈34% Poloxamer and distilled acetylated monoglycerides) was obtained from the FMC Biopolymer Corporation, Philadelphia, PA. The following polymers were obtained from commercial sources different from proprietary products being developed as microbicides: carrageenans κ and λ (Sigma, St. Louis, MO; mixed at a 1:1 (w/w) ratio in all experiments); cellulose sulfate (Across Organics, Piscataway, NJ); poly(naphthalene sulfonate) (BASF, Parsippany, NJ); and polystyrene-4-sulfonate (Polysciences, Inc., Warrington, PA).

The HIV-1 non-nucleoside reverse transcriptase inhibitors, UC781 and TMC120 were obtained by custom synthesis from Albany Molecular Research, Inc., Albany, NY. Zinc finger inhibitors #89 and #247 were a gift from Dr. Ettore Appella and Dr. Marco Schito (National Cancer Institute, Bethesda, MD). Stabilite SD60 Polglycitol, hydroxypropyl methylcellulose E4M and Avicel PH 105, respectively, were from SPI Polysols, New Castle, DE, Dow Chemical Co., Midland, MI and the FMC Biopolymer Corporation, Philadelphia, PA. SP was purchased from Vital Products, Inc., Boynton Beach, FL. HIV-1 IIIB and BaL were from Advanced Biotechnologies, Inc., Columbia, MD. Hela-CD4-LTR-β-gal, MAGI-CCR5 and TZM-bl cells were obtained from the AIDS Reagent and Reference Reagent Program (operated by McKesson BioServices, Rockville, MD) and contributed by Drs. M. Emerman, J. Overbaugh and J. C. Kappes and X. Wu (Tranzyme, Inc.), respectively. Dulbecco’s modified Eagle medium (DMEM) was from GIBCO Invitrogen Corporation, Carlsbad, CA. The Galacto-Light Plus chemiluminescence reporter assay for β-galactosidase was from Applied Biosystems, Foster City, CA.

**Inhibition of infection by anti-HIV-1 compounds in the presence or absence of seminal plasma**

Seventy μl of serially two-fold diluted compounds in DMEM medium (final concentrations after dilution: 1.25 to 10,000 μg/ml) were mixed with 70 μl of HIV-1 IIIB and BaL, respectively, and 70 μl of either SP or DMEM medium. The mixtures were added to TZM-bl indicator cells (in 96-well plates) which can be infected by both X4 and R5 HIV-1, enabling quantitative analysis of HIV-1 infection using either β-galactosidase (β-gal) or luciferase as a reporter [19]. After 90 min at 37°C, the virus (and SP) containing supernatants were removed from the cells. The cells were washed once with DMEM medium, supplemented with the same medium, and incubated for 48 hr at 37°C. The removal of SP after 90 min was necessary to avoid problems caused by the cytotoxicity of SP evident after prolonged incubation [20-22]. Finally, the culture supernatants were removed and the cells were washed once with phosphate buffered saline, pH 7.2 (PBS). Subsequently, 50 μl of lysis buffer from the Galacto-Light Plus kit were added to the wells for 1 hr at 20°C. Aliquots (20 μl) of the cell lysates were transferred into wells of 96-well microplates and β-gal was quantitated using the Galacto-Light Plus System chemiluminescence reporter assay in a MicroLight ML 2250 luminometer (Dynamitex Laboratories, Inc. Chantilly, VA). The concentration of viruses was selected so as to provide a readout of ≤70 in the absence of SP and drug, respectively. The percentage of inhibition was calculated using the Microsoft Excel computer program. ED₅₀ values were calculated using an online computer program [23]. The inhibitory activity of the compounds in the absence of SP was measured also at a lower (1/6) virus dose (readout ≤20) to determine whether or not this would result in higher ED₅₀ values. This was not the case, confirming that the presence of SP (causing partial virus inactivation or inhibition) truly diminishes the inhibitory activity of the polymers under investigation.

**Measurements of HIV-1 infectivity**

The following two formulations were tested for virucidal activity against HIV-1 IIIB and BaL, respectively, in the presence of SP: (1) Aquateric, 18%; glycerol, 58%; Stabilite SD60, 20%; Hydroxypropyl methylcellulose E4M 1%; Avicel PH 105, 3%; and (2) 18% Aquateric in a universal placebo gel [24].
Two-fold serial dilutions of HIV-1 IIIB treated with Aqua-
teric formulations, and separated from these polymers by 
bear precipitation with 3% PEG or by centrifugation at 14,000 
rpm for 1 h, and control virus (100 µl), respectively, were 
added to HeLa-CD4-LTR-β-gal cells which had been 
plated a day before infection in 96-well plates at 1 × 10^4 
cells/well in 100 µl of DMEM medium containing 10% 
fetal bovine serum (FBS). After incubation at 37°C for 48 
h, the culture supernatant fluids were removed and the 
cells washed once with PBS. β-gal in the cells was mea-
ured as described above. The infectivity of treated and 
control HIV-1 BaL was measured by the same method 
except that MAGI-CCR5 cells were used.

Measurement of HIV-1 infectivity in seminal fluid is 
impeded by the cytotoxicity of SP [20,21] contributed to 
by spermine [22]. Therefore, HIV-1 was separated from SP 
ingredients by precipitation with 3% PEG or centrifuga-
tion (14,000 rpm for 1 h) and the infectivity of the resus-
pended pellets was measured. Such cytotoxic effects were 
iminized by using for titrations of virus infectivity pel-
lets after precipitation with 3% PEG or after centrifugation 
(14,000 rpm for 1 h).

**Micromethod for CAP quantitation**

CAP in the form of a complex with ruthenium red was 
quantitated spectrophotometrically [25].

**Results**

**The HIV-1 inhibitory activity of several polymeric 
candidate microbicides is diminished in the presence of 
seminal plasma**

Inhibition of HIV-1 IIIB (a virus utilizing CXCR4 as cellu-
lar coreceptor = X4 virus) and HIV-1 BaL (a virus utilizing 
CCR 5 as cellular receptor = R5 virus) [8,26] infection, 
respectively, of TZM-bl cells [19] by polymeric candidate 
microbicides in the absence or presence of SP (final con-
centration 33.3%) was investigated. The polymers 
included: carrageenan, poly(naphthalene sulfonate), cel-
lulose sulfate, cellulose acetate 1,2-benzene dicarboxylate 
(CAP) and polystyrene sulfonate. Dose response curves 
for inhibition of HIV-1 IIIB infection (Fig. 1) show the 
suppressive effect of SP on the virus inhibitory activities 
of the polymers. Similar results were obtained for HIV-1 BaL 
(data not shown). The results are summarized in Table 1 
showing values for 50% inhibition of infection (ED_{50}) 
corresponding to the distinct polymers. The ED_{50} values 
are consistently higher for HIV-1 BaL than for HIV-1 IIIB. 
This is consistent with lower positive charges on V3 loops 
of R5 gp120 envelope glycoproteins as compared with X4 
gp120, leading to diminished binding of negatively 
charged polymers [27,28]. The presence of SP led to a = 4 
to = 73-fold increase of ED_{50}, resulting in ED_{50} values for 
HIV-1 BaL (a representative of more frequently sexually 
transmitted R5 viruses [8,26]) > 100 µg/ml. This raises the 
question whether or not sufficient concentrations of the 
respective inhibitors can be reached in the vaginal envi-
ronment at which cells playing a major role in the virus 
replicative cycle (dendritic cells, macrophages and CD4+ 
T cells) occur [11].

In contrast with the polymeric candidate microbicides, 
the anti-HIV-1 activities of reverse transcriptase inhibitors 
UC781 and TMC120, and of zinc finger inhibitors #89 
and #247, respectively, were not affected by the presence 
of SP (data not shown). This will support confidence for 
results of efficacy tests for these antiretrovirals drugs in ani-
mal model systems for male to female virus transmission. 
The presence of SP alone (33.3 to 80%) resulted in 4- to 
22-fold, and 2- to 10-fold inhibition of HIV-1 IIIB and BaL 
infection, respectively, in different experiments.

**Direct HIV-1 inactivation in the presence of seminal plasma**

Anionic polymeric candidate microbicides are thought to 
interfere with HIV-1-cell interactions involving cellular 
receptors either directly in productive infection (CD4 and 
CXCR4/CCR5) or in cell to cell transfer of virus (DC-SIGN 
etc.) [11]. However, some polymers rapidly (≤ 5 min) 
inactivate HIV-1. Under these circumstances, there is no 
need for the antiviral compounds to reach cells involved 
in virus dissemination or infection, the virus being ren-
dered non-infectious before encountering cellular/tissue 
surfaces. Among the aforementioned polymers, cellulose 
sulfate, polystyrene sulfonate, poly(naphthalene sul-
fonate) (at a concentration of 10 mg/ml; at concentra-
tions ≤ 1 mg/ml the compounds may not be virucidal 
[29]), CAP (micronized), as well as the acidifying formu-
lation, BufferGel, inactivate not only X4 viruses but also 
the R5 virus, HIV-1 BaL [30]. Poly(naphthalene sulfonate) 
and CAP, respectively, rapidly elicit in the virus envelope 
the formation of “dead-end” gp41 six-helix bundles ren-
dering the virus irreversibly incapable to fuse with suscep-
tible target cells, a prerequisite for infection [30]. CAP 
being the only candidate microbicide provided in a water 
insoluble micronized form, and therefore, very unlikely to 
penetrate into tissues or cells [31], was selected for further 
studies regarding the effect of SP on its virus inactivating 
properties.

CAP formulations (expected to contain about 240 to 480 
mg CAP per dose for human use) have a good acidic buff-
ering capacity, are non-toxic to vaginal and cervical 
explants or reconstituted tissues [32], and provide a pH ≤ 
5.5 if the volume of SP is ≤ 1.8 ml per 100 mg CAP (Fig. 
2). CAP remains water insoluble under these conditions 
(Fig. 3) and will remain in the form of particles about 1 
micron in diameter. Changes in pH and CAP solubility 
during the course of addition of SP to formulated CAP (= 
120 mg/ml) are shown in Fig. 4. It is evident that even at
an excess of SP over the volume of the formulation used (3:1), the pH remains < 5.5 and the maximum concentration of CAP becoming soluble is ≤ 7 μg/ml. This concentration is sufficient for partial inhibition of infection by the laboratory strain HIV-1 IIIB [33] but insufficient for inhibition of most other HIV-1 strains [34]. Thus, the predominant contribution to anti-HIV-1 activity of formulated CAP is expected to be attributable to the insoluble micronized particles even at an excess of SP. These particles adsorb both X4 and R5 viruses, elicit 6-helix bundle formation in their envelope glycoproteins and render the viruses non-infectious [30]. Both HIV-1 IIIB and BaL are inactivated within 5 min at 37°C by CAP in micronized form even in the presence of a volume excess of SP (Table 1).

Table 1: Decreased anti-HIV-1 activity of polymeric candidate microbicides in the presence of human seminal plasma

| Polymer                          | HIV-1 IIIB | HIV-1 BaL |
|---------------------------------|------------|-----------|
|                                 | ED50 μg/ml | ED50SP μg/ml | ED50sp ED50 ratio | ED50 μg/ml | ED50SP μg/ml | ED50sp ED50 ratio |
| Poly (naphthalene sulfonate)     | 1.1        | 17.3       | 15.7          | 37.5       | 141         | 3.8           |
| Cellulose sulfate                | 1.6        | 64.7       | 40.4          | 7.1        | 401         | 56.5          |
| Carrageenan                      | 1.4        | 63.4       | 45.6          | 6.3        | 149         | 23.7          |
| Cellulose acetate 1,2-benzenedi-carboxylate | 1.0        | 72.7       | 72.7          | 9.8        | 141         | 14.4          |
| Polystyrene sulfonate            | 2.2        | 36.1       | 16.4          | 12.8       | 389         | 30.4          |

SP = seminal plasma added

Figure 1
Inhibition of HIV-1 IIIB infection by negatively charged polymers in the presence (33.3%; v/v), and absence of seminal plasma, respectively. TZM-bl indicator cells and β-galactosidase readout were used for quantitative analysis of HIV-1 infection.
Similar results were obtained with an Aquateric formulation in a “universal placebo” gel [24] (Fig. 5). Virucidal activity against HIV-1 BaL was still detectable at relatively high dilutions of the formulation in SP (resulting in pH ≥ 7.0), not expected to occur in vivo. The virucidal activity of CAP was similar to that observed in the absence of SP (see Fig. 2 in our earlier publication) [35]. In contrast, acidic pH alone is much less virucidal than CAP in micronized form, the half-life for virus infectivity being 6, 10 and >120 min at pH 3.5, 4.0 and 4.5, respectively [36] and fails to elicit “dead-end” six-helix bundle formation in HIV-1 gp41 (data not shown). Thus, combination of a low pH buffering system with a compound targeted to functionally important sites on HIV-1 envelope glycoproteins (e.g. CD4 and/or CXCR4/CCR5 coreceptor binding sites, and gp41 regions involved in 6-helix bundle formation) provides an attractive approach for development of virucidal microbicides. In the case of CAP, the low-pH buffering capacity is an inherent property of the polymer.

**Discussion**

Results presented here indicate that SP caused a decrease in the HIV-1 inhibitory activity of all polymers being considered as candidate microbicides while inhibition of virus infection caused by reverse transcriptase and zinc finger inhibitors remained unaffected. Semen contains the polyamines spermine, spermidine and putrescine [37,38] which are positively charged at neutral pH while the polymeric anti-HIV-1 inhibitors all have negative charges. Thus, complex formation between the polyamines and the anti-HIV-1 polymers seems likely. This is supported by the observed complex formation between seminal plasma and Aquateric.
between polyamines and sulfonic and carboxylic poly-
anions [39,40]. In agreement with this, SP interfered with
the inhibitory effect of a selected candidate microbicide,
cellulose sulfate, on binding of monoclonal antibodies to
the positively charged V3 loop of HIV-1 IIIB gp120 [41]
(data not shown). This interference was completely abro-
gated by addition of sodium maleate with a negative
charge in the maleate moiety, forming complexes with
polyamines. However, maleate failed to restore the inhib-
itory activity of the polymeric candidate microbicides
against HIV-1 infection to levels observed in the absence
of SP (data not shown). Thus, electrostatic interactions
between SP polyamines and the polyanions tested seem
to provide an incomplete explanation for the suppression
of anti-HIV-1 activities of polymeric candidate microbi-
cides by SP.

The decreased anti-HIV-1 inhibitory activity in vitro of
polymeric candidate microbicides in the presence of SP sug-
gest that these compounds may become less effective in vivo in animal models and human clinical trials than
would be expected from macaque efficacy studies using a
semen-free virus challenge [5,6]. This possibility confirms
the need for development of combination microbicides in
which candidate polymers would be supplemented with
anti-HIV-1 compounds remaining fully effective in the
presence of SP, and preferably having a mechanism of
action distinct from that of the polymeric microbicides.
The simplest approach towards this goal, which may also
bypass potential regulatory hurdles resulting from two
active ingredients in a single formulation, is to formulate
the anti-HIV-1 polymers in buffer systems maintaining an
acidic pH (similar to that of a normal vaginal environ-
ment).

**Figure 3**

**Dissolution rate of CAP as a function of pH.** Data are derived from J. Spitael and R. Klinget: Solubility and dissolution rate
of enteric polymers, *Acta Pharmaceutica technologica* 1979; S7:163–168 [56].
ment) even in the presence of semen, and causing inactivation of HIV-1 and other sexually transmitted disease (STD) pathogens. Such buffering bioadhesive formulations include ACIDFORM [42,43] based on small mole-

Table 2: Inactivation of HIV-1 IIIB and BaL by formulated Aquateric (18%) (5 min, 37°C) in the presence of seminal plasma

| Seminal Plasma: Formula Ratio | % Virus Inactivation |
|-----------------------------|---------------------|
|                             | HIV-1 IIIB | HIV-1 BaL |
| Ratio pH Aquateric Pellet Supernatant Aquateric Pellet Supernatant |
| 0.33 3.9                  | ≥ 99.8       | ≥ 99.8       | ≥ 99.6       | ≥ 99.6       |
| 1.0 4.8                   | ≥ 99.8       | ≥ 99.8       | ≥ 99.6       | ≥ 99.6       |
| 3.0 5.43                  | ≥ 99.8       | ≥ 99.8       | ≥ 99.6       | ≥ 99.6       |

Virus concentrates of HIV-1 IIIB and BaL, respectively, were diluted 200-fold in seminal plasma and then mixed with Aquateric formulation 1 (see Methods section) at volume ratios indicated in the Table. After incubation for 5 min at 37°C, the samples were cooled in ice and centrifuged at 10,000 rpm for 10 min. The supernatant fluids were neutralized by addition of 1 M Na2HPO4. The pellets containing Aquateric with adsorbed virus were resuspended and brought to pH 7.0 by addition of 1 M Na2HPO4. Both the supernatants and the redissolved pellets were mixed with a solution of PEG 8000 (final concentration 3%). The resulting pellets were resuspended in DMEM medium and tested for infectivity as described in the Methods Section. Virus preparations diluted in seminal fluid but not treated with the Aquateric formulation represented controls.

Figure 4
CAP solubility in seminal plasma-Aquateric formulation mixtures. Increasing volumes of seminal plasma were added to one ml of formulation 1 (see Methods section). The pH was measured; the mixtures were centrifuged to pellet most of Aquateric and CAP in the supernatant fluids was quantitated [25].
cule components, and preferably polymers with built-in acidic buffering capacity: Carbomer 974P (BufferGel) [44] and CAP (Fig. 2, 3, 4, 5; Table 1).

To fully appreciate the role of SP ingredients on sexual transmission of HIV-1, one should consider also the observations that they appear to cause a decrease of infectious virus titer (see Results; [36,45]). Major contributors to this phenomenon are probably products resulting from oxidation of SP polyamines by diamine oxidase [46] which is also present in this biological fluid [47,48]. The oxidation is augmented by peroxidase [46]. Peroxidase is present in a healthy vaginal environment [49,50]. This suggests that a combination of SP ingredients and a “normal” vaginal environment might provide a natural defense system against infection by HIV-1 and other STD pathogens and might contribute to the low incidence of HIV-1 transmission per unprotected coital act [8]. In fact, several viruses and parasites were shown to become inactivated by polyamine oxidation products [51-55]. Efforts to augment this natural defense mechanism might potentially become part of the microbicide development strategy.

**Conclusion**

Candidate anti-HIV-1 topical microbicides have been evaluated for virus inhibitory activities *in vitro* and efficacy in animal models mostly without considering the possible effects semen/seminal plasma may have on the ultimate performance of these compositions. Studies in macaques challenged vaginally by SIV or SHIV, after application of anti-HIV-1 compounds, indicated that drug doses required for protection against infection exceeded by several orders of magnitude concentrations sufficient for virus inactivation.
for blocking virus replication in vitro. Since semen/seminal plasma is a vehicle for male to female sexual transmission of the virus, it is of crucial interest to determine its potential impact on HIV-1 inhibitory activities of candidate products. Such studies are difficult, if not impossible, in animal models. Therefore, at least a relevant assessment in in vitro systems is called for. Results presented here indicate that the anti-HIV-1 activity of synthetic polymers, acting as HIV-1 entry inhibitors (carrageenan, cellulose sulfate, poly(N-phthalene sulfonate), polysulfonate and CAP in water soluble form) is greatly diminished in the presence of SP. The impact of this finding on performance of these candidate microbicides in vivo remains unknown. On the other hand, HIV-1 inhibition by reverse transcriptase inhibitors, UC781 and TMC120 and by zinc finger inhibitors #89 and #247, respectively, was not affected by SP. Combination of polymeric HIV-1 entry inhibitors with active compounds having a distinct mechanism of action, and reliance on polymers having direct virucidal activity and built-in low pH buffering capacity (contributing to virus inactivation) (CAP in micronized form; BufferGel) are expected to overcome potential problems resulting from interference of semen components with the performance of some microbicides being considered for, or already in clinical trials.

Competing interests
The author(s) declare that they have no competing interests.

Authors' contributions
Author 1 ARN developed the concepts representing the basis of the manuscript and designed most experiments; author 2 NS carried out most experiments and contributed to the development of experimental techniques; author 3 YYL did all the tissue culture work and infectivity assays. All authors have read and approved the final manuscript.

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