A Caulobacter crescentus Microbicide Protects from Vaginal Infection with HIV-1JR-CSF in Humanized Bone Marrow-Liver-Thymus Mice

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ABSTRACT Over 2 million people are infected with HIV-1 annually. Approximately half of these new infections occur in women residing in low-income countries, where their access to and control over HIV-1 preventative measures are often limited, indicating that female-controlled prevention options for HIV-1 are urgently needed. Microbicides that can be topically applied to the vaginal tract in advance of sexual activity represent a promising female-controlled prevention option for HIV-1. We have previously described the development of an HIV-1-specific microbicide using the surface or S-layer recombinant protein display capabilities of the nonpathogenic, freshwater bacterium Caulobacter crescentus. Recombinant C. crescentus bacteria were created that displayed proteins that interfere with the HIV-1 attachment and entry process and that were able to provide significant protection of TZM-bl cells from infection with HIV-1 pseudovirus. These studies have been expanded to investigate if these recombinant C. crescentus bacteria are able to maintain efficacy with replication-competent HIV-1 and both TZM-bl cells and human peripheral blood mononuclear cells (PBMCs). In addition, we utilized the humanized bone marrow-liver-thymus (BLT) mouse model to determine if vaginal application of recombinant C. crescentus at the time of HIV-1JR-CSF infection could provide protection from HIV-1 infection. Recombinant C. crescentus bacteria expressing Griffithsin, GB virus C E2 protein, elafin, α-1-antitrypsin, indolicidin, and the fusion inhibitor T-1249 were able to protect 40 to 75% of the BLT mice from vaginal infection with HIV-1JR-CSF, with C. crescentus bacteria expressing Griffithsin being the most effective. Taken together, these data suggest that a C. crescentus-based microbicide could be a safe and effective method for HIV-1 prevention.

IMPORTANCE Human immunodeficiency virus (HIV) disproportionally infects young women in sub-Saharan Africa. Current HIV-1 prevention options have had limited success among women, suggesting that alternative, female-controlled prevention options need to be developed. Microbicides that can be applied to the vaginal tract are a promising prevention option. In this study, we describe the testing of 15 potential candidates for inhibition of HIV-1 infection in a humanized mouse model of HIV-1 infection. Four of these candidates were able to provide significant protection from vaginal infection with HIV-1, with the most successful candidate protecting 75% of the mice from infection. This study describes the preclinical testing of a new strategy that could be a safe and effective option for HIV-1 prevention in women.

KEYWORDS Caulobacter crescentus, HIV, microbicide, humanized mice

HIV-1 prevention for women is a global health priority. Approximately 1 million women are infected with HIV-1 each year (1). In sub-Saharan Africa, 75% of new HIV-1 infections in 15- to 19-year-olds are among women, making them twice as likely to acquire HIV-1 infection as men (1, 2). Existing HIV-1 prevention options, like con-
doms, male circumcision, and viral suppression, are not always feasible for women, as these rely on their partners for use. Thus, prevention options that women can use and control are urgently needed. With the difficulties in developing an effective vaccine and the inconsistent results of preexposure prophylaxis (PrEP) in women (1, 3–5), alternative female-controlled HIV-1 prevention options are urgently needed.

Recent microbicide clinical trials have focused on either a 1% tenofovir gel or a dapivirine vaginal ring for HIV-1 prevention (2, 6–8). Tenofovir gel demonstrated 39% efficacy in the CAPRISA 004 clinical trial (2). However, in both the FACTS 001 and VOICE trials, it had no protective effect on HIV-1 acquisition (2–4, 9). While adherence was a concern, follow-up studies have indicated that vaginal microbiota, particularly *Gardnerella* and *Prevotella*, can interfere with the effectiveness of tenofovir-based microbicides (10). In the ASPIRE and The Ring Study trials, the dapivirine vaginal ring reduced HIV-1 acquisition by 27% and 31%, respectively (7, 8). However, among women under the age of 21 years, the ring was 15% effective in The Ring Study and had no efficacy in the ASPIRE trial (7, 8). These clinical trial results indicate that the development of alternative microbicide strategies is urgently needed.

We have previously demonstrated up to 72% protection from HIV-1 infection *in vitro* using the surface or S-layer recombinant display capabilities of the nonpathogenic, freshwater bacterium *Caulobacter crescentus* (11–13). In these studies, 15 unique recombinant *C. crescentus* bacteria with the ability to prevent the attachment or entry of HIV-1 into a target cell were created (11, 12). The recombinant bacteria expressed a wide variety of anti-HIV proteins, including the carbohydrate binding agents cyanovirin-N (14), microvirin (15), and griffithsin (16, 17), ligands (macrophage inflammatory protein 1α [MIP-1α]) (18), decoy receptors (CD4, mimetic CD4M33F23) (18, 19), fusion inhibitors (Fuzeon [20], T-1249 [21], CS2 variant [22]), and the antimicrobial peptides BmKnn2 (23), α-1-antitrypsin (A1AT) (24), indolicidin (25), and elafin (26). The success of these recombinants for HIV-1 prevention in initial studies indicated that further studies using more physiologically relevant models are warranted.

While *C. crescentus* is a nonpathogenic bacterium, it is a Gram-negative bacterium that could stimulate an immune response *in vivo*. Previous work in our lab (13) and by collaborators (27) has demonstrated that *C. crescentus* appears to be safe for topical application to the vaginal tract. Importantly, there was no significant production of inflammatory cytokines, immune cell recruitment, or antibody production after vaginal application of *C. crescentus* in an immunocompetent mouse model (13). Furthermore, *C. crescentus* cannot be cultured from the peritoneal cavity of immunocompetent mice within 10 days following intraperitoneal injection (27). These data suggest that *C. crescentus* will likely be safe for use as a topical mucosal agent.

Herein, both *in vitro* and *in vivo* studies were undertaken to further test the ability of recombinant *C. crescentus* to prevent HIV-1 infection. *In vitro* studies using replication-competent HIV-1 isolates from several strains indicated that both TZM-bl cells and human peripheral blood mononuclear cells (PBMCs) were protected from HIV-1 infection. In addition, the recombinant *C. crescentus* was applied to the vaginal tract of humanized bone marrow-liver-thymus (BLT) mice (28–30), and HIV-1 infection was measured. We found that 40 to 75% of mice were protected from vaginal infection with HIV-1 using 6 different recombinants, with the *C. crescentus* recombinant expressing griffithsin (Cc-griffithsin) being the most effective at preventing HIV-1 acquisition. Taken together, these data suggest that a *C. crescentus*-based microbicide could be a worthwhile option for developing a safe and effective method for HIV-1 prevention.

**RESULTS**

Many recombinant *C. crescentus* bacteria provide protection from replication-competent HIV-1 infection. We have previously demonstrated a successful proof of concept for a recombinant *C. crescentus*-based microbicide expressing antiviral lectins, fusion inhibitors, decoys, and antiviral peptides (11–13), demonstrating inhibition of infection using HIV-1 pseudoviruses representing the two most common viral clades, B and C. These studies were repeated using replication-competent HIV-1 and both
TZM-bl cells and human PBMCs. Previous studies have indicated that $1 \times 10^8$ recombinant bacteria provide the best protection from HIV-1 infection while minimizing nonspecific inhibition by control (no insert) C. crescentus bacteria (Cc-control) (13). HIV-1 infection rates were significantly decreased with all recombinant bacteria across various HIV-1 strains in both cell types. While the most effective recombinant varied depending on the virus/cell combination, several recombinants were able to provide a >90% decrease in HIV-1 infection in both TZM-bl cells and PBMCs with specific viral strains (Table 1; Fig. 1). With some of the recombinants, there was a wide range of effectiveness depending on the viral strain used, which is why we anticipate combining recombinants to improve efficacy in a final microbicide product. Notably, HIV-1 was suppressed, on average, by 50% with each recombinant, results that were consistent with those of our previous studies.

**GB virus C E2 protein blocks HIV-1 infection in vitro.** GB virus C (GBVC; human pegivirus, formerly hepatitis G virus) causes a persistent viral infection that does not cause any known disease pathology and appears to improve the survival of HIV-positive individuals and delay progression to AIDS (31–33). While many factors have been linked to this favorable survival (34, 35), the E2 protein of GB virus C (GBVCE2) is a putative fusion peptide that can interfere with HIV binding or fusion (36, 37). The E2 protein of GB virus C was expressed in the S layer of C. crescentus (Fig. 2a) and used in in vitro viral blocking assays with both TZM-bl cells and PBMCs (Fig. 2b and c). The E2 protein was successfully expressed in the S layer of C. crescentus, and recombinant C. crescentus bacteria expressing GBVCE2 (Cc-GBVCE2) were able to provide 57% protection from HIV-1 in TZM-bl cells and 54% protection in PBMCs (Fig. 2). Furthermore, when Cc-GBVCE2 was tested for the ability to prevent HIV-1 infection with additional viral strains, it provided an average 65% reduction in HIV-1 infection in the TZM-bl cell line and 76% in PBMCs. These results suggest that Cc-GBVCE2 is an excellent candidate for additional microbicide testing.

**Cc-griffithsin, Cc-GBVCE2, Cc-elafin, and Cc-A1AT provide significant protection from HIV-1 infection in BLT mice.** These in vitro studies suggested that recombinant C. crescentus could be an excellent option for an HIV-1-specific microbicide and that further preclinical testing was warranted. Therefore, humanized BLT mice (28–30) were used to test the ability of the recombinant C. crescentus bacteria to provide protection from vaginal infection with HIV-1. Human immune cell reconstitution of BLT mice was confirmed by flow cytometry (Fig. 3a; Table 2). Vaginal infection of the mice with HIV-1JR-CSF in the presence of $10^8$ Cc-control bacteria indicated that C. crescentus did not impact susceptibility to HIV-1 infection (Fig. 3b and c). We have previously demonstrated by titration that $10^8$ recombinant C. crescentus bacteria provide significant protection from infection in vitro without causing nonspecific inhibition of infection and have demonstrated that this dose can provide significant protection from vaginal infection with herpes simplex virus 2 (HSV-2) in C57BL/6 mice (13).

While human immune cell reconstitution was variable across different cohorts of BLT mice, each cohort contained mice infected with HIV-1 only and mice infected with HIV-1 and treated with Cc-control, both groups of which were measured to be HIV-1 positive following infection, suggesting that each cohort was susceptible to vaginal infection with HIV-1. Furthermore, each recombinant C. crescentus bacterium was tested in multiple cohorts of BLT mice to verify that protection from HIV-1 infection was observed across more than one cohort.

Cc-griffithsin was the most effective recombinant bacterium, protecting 6 of 8 mice (75%, $P = 0.003$) from vaginal infection with HIV-1, confirmed by both a p24 enzyme-linked immunosorbent assay (ELISA) and reverse transcription-quantitative PCR (RT-qPCR) (Fig. 4a and 5a). To ensure that all HIV-1-positive mice were detected, HIV-1 infection was measured by both p24 ELISA and RT-qPCR for each mouse, and a mouse was considered HIV-1 negative only if both assays had no virus detected. Interestingly, one mouse was HIV-1 negative by the p24 ELISA but had detectable HIV-1 RNA by RT-qPCR. Although the p24 levels in the blood of HIV-1-negative mice that received
### TABLE 1 Recombinant C. crescentus

| Protein | Name          | Sequence                                                                 | Reference in which construct was first described | Mean (range) % HIV-1 inhibition Live virus-infected TZM-bl cells | Mean (range) % HIV-1 inhibition Live virus-infected PBMCs |
|---------|---------------|--------------------------------------------------------------------------|--------------------------------------------------|---------------------------------------------------------------|----------------------------------------------------------|
| Blank control | Cc-control   | NA                                                                      | 11                                               | 110.1 (80.8–137.7)                                            | 100 (99.9–100)                                           |
| Griffithsin | Cc-griffithsin | SLTHRKF GGSGGS PSGLSIAVRSGYLDAAIIDGVHGGSGGNLSPTFTFSGGEYISNMTRT            | 12                                               | 61 (19.6–85.2)                                                | 56.7 (25.4–100)                                          |
| Microvirin | Cc-microvirin | MPNSHTCSSINYPDSTSILSASCQARDGEWLPTELRLSDHIGMDGELGQDGNQFOET               | 12                                               | 61.8 (11.9–82.4)                                              | 56 (–7.9–100)                                            |
| Cyanovirin | Cc-cyanovirin | LQKFSQTCYNSAIQGSLTVSTCERTNGGYNTSIDLSWENVGSLKMWGQS                      | 12                                               | 51.6 (21.6–87.6)                                              | 71.6 (15.9–100)                                          |
| Fuzeon, T-20 | Cc-Fuzeon   | NMYTLSHSLIEQSNODAENQEEQEEQELLEDKWA SLLNWMFM                            | 12                                               | 58.8 (29.3–78.9)                                              | 62 (8.3–100)                                              |
| T-1249      | Cc-T1249     | WQWSWEQATLLEQAQIQQEKNEYELQKLDKWASLLWWEF                              | 12                                               | 40 (4.9–54.2)                                                 | 27.6 (30.9–100)                                          |
| C52         | Cc-C52       | NHTTWMERDREINNTYSLHSLEIERSQNEQQNEQEEQELLEDKWA SLLNWMFN                 | 12                                               | 50.7 (12.2–81.3)                                              | 86.8 (46.5–100)                                          |
| MIP-1α      | Cc-MIP1α     | APLAADPTACCSYTSRQPQNFIAFDYETSSQC SLPVFILTKRGRQVCAD PSEEWQKYSVDSLELSA     | 11                                               | 60.3 (23–83)                                                   | 69.2 (11.2–97.2)                                         |
| CD4         | Cc-CD4       | GDTVELTCTASQKKSIQFHWKNSNQKLGNQGSLTKEGPSKLNDRADSRRSL WDQGNFPLILKNNIELDSITYCEVEQ   | 11                                               | 55.5 (10.6–72.5)                                              | 74.6 (18–97.5)                                            |
| CD4 mimicetic | Cc-CD4M33F23 | NLHFCQRLKSSSLGLGKCSGFCACV                                               | 12                                               | 36.2 (16–74.3)                                                | 26.9 (36.5–93.2)                                         |
| BmKn2       | Cc-BmKn2     | FIGIAARLLSRF                                                              | --                                               | --                                                             | --                                                       |
| α-1-antitrypsin | Cc-A1AT   | LEAIPCSPPEFLPGKFVFLMIEQNTKSLEMG                                       | --                                               | --                                                             | --                                                       |
| Elafin      | Cc-elafin    | AOEVPKVPSGTPSQPIIURCAMLNPNRRKLDTDCPGI1KCCCEGSCGMAFCVPQ                  | --                                               | --                                                             | --                                                       |
| Indolicidin | Cc-indolicidin | ILPKWKWPWPWPWR                                                       | --                                               | --                                                             | --                                                       |
| GB Virus C E2 protein | Cc-GBVCE2 | WDRGNTLLCDDPNPWWPPPAPFCQAVG                                             | Herein                                           | 64.9 (19.5–86.4)                                              | 75.7 (27.2–87.9)                                         |

*a* The mean and range presented are a summary of data from HIV-1 strains 89.6, BaL, JR-FL, and pybK-CSF.

*b* The mean and range presented are a summary of data from HIV-1 strains 89.6, JR-FL, and SF162.

*c* NA, not applicable.

*d* Data represent the mean (range) percent infection.

*e* —, data are presented elsewhere (13).
FIG 1 Viral blocking assays. A total of 10,000 TZM-bl cells or PHA-stimulated PBMCs were incubated for 48 to 72 h with 200 TCID₅₀ of HIV-1₈₉₆ or HIV-1₉₉/K-RCSF and 10⁸ recombinant *C. crescentus* bacteria. HIV-1 infection was measured using a β-galactosidase assay (TZM-bl cells) or p24 ELISA (PBMCs). To minimize assay-to-assay variability, the wells containing cells and HIV-1 were set as 100% infection, and the results for the other wells were normalized to the 100% infection. Each experiment was performed in quadruplicate and repeated three times. An unpaired two-tailed t test (for the Cc-control group and 1 group receiving recombinant *C. crescentus* bacteria) or ANOVA with Bonferroni's correction for multiple comparisons (for the Cc-control group and 2 groups receiving recombinant *C. crescentus* bacteria, with each group being compared to the Cc-control group and each other) were used to evaluate the significance of differences between groups as appropriate. Results are reported as the mean ± SEM. (a) Antiviral lectins. (b) Fusion inhibitors. (c) MIP-1α. (d) CD4-based inhibitors. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001; ns, not significant (P > 0.05).
HIV-1 plus Cc-griffithsin (0 pg/ml p24) were not statistically significantly different from those of HIV-1-negative mice that received HIV-1 plus Cc-control, there was a significant reduction in HIV-1 RNA in the blood of mice that received HIV-1 plus Cc-griffithsin and that were HIV-1 negative compared to that in the blood of mice that received HIV-1 plus Cc-control. While the other antiviral lectins provided protection levels similar to those provided by Cc-griffithsin in vitro, recombinant C. crescentus bacteria expressing microvirin (Cc-microvirin) and cyanovirin (Cc-cyanovirin) were not effective at preventing infection in BLT mice. Cc-microvirin protected 37.5% (3 of 8, \( P = 0.05 \)) of the BLT mice from HIV-1 infection (Fig. 6a). In a pilot experiment, Cc-cyanovirin protected 1 of 3 BLT mice from vaginal infection with HIV-1 (Fig. 6b). As this protection level was not statistically significant and at some times postinfection the HIV-1 levels were higher in mice that received HIV-1 plus Cc-cyanovirin, additional experiments were not undertaken.

Three additional recombinants were able to provide significant protection from vaginal infection in BLT mice: Cc-GBVCE2, recombinant C. crescentus bacteria expressing elafin (Cc-elafin), and recombinant C. crescentus bacteria expressing \( \alpha-1 \)-antitrypsin (Cc-A1AT). There were undetectable viral loads in 6 of 10 mice (60%, \( P = 0.0124 \)) receiving Cc-GBVCE2 by both p24 ELISA and RT-qPCR following vaginal infection with HIV-1 (Fig. 4b and 5b). Cc-elafin provided 56% protection (\( P = 0.0036 \)) from vaginal infection with HIV-1, with five of nine mice remaining HIV-1 negative, as determined by a lack of detection of p24 or viral RNA (Fig. 4c and 5c). Three mice that received Cc-elafin and that became HIV-1 positive had a delay in detectable levels of p24 in the blood until day 42 postinfection, although at this time p24 levels were significantly higher than the peak levels reached with HIV-1 alone. Cc-A1AT was able to protect 5 of 10 mice (50%, \( P = 0.0487 \)) from vaginal infection with HIV-1, based on a lack of detection of p24 and viral RNA (Fig. 4d and 5d). Interestingly, those mice that received...
HIV-1 plus Cc-A1AT and that became HIV-1 positive appeared to have higher levels of virus in the blood, based on p24 levels.

Eight of the recombinant C. crescentus bacteria (those expressing Fuzeon [Cc-Fuzeon], T1249 [Cc-T1249], C52 [Cc-C52], CD4 [Cc-CD4], CD4M33F23 [Cc-CD4M33F23],...
MIP-1α (Cc-MIP1α), indolicidin (Cc-indolicidin), and BmKn2 (Cc-BmKn2) provided some level of protection from vaginal infection with HIV-1, based on both p24 ELISA and RT-qPCR results, although this protection was not statistically significant (Fig. 6c to j).

**DISCUSSION**

In this preclinical microbicide study, recombinant bacteria expressing the antiviral lectin griffithsin (Cc-griffithsin) were able to protect 75% of BLT mice from vaginal infection with HIV-1. Based on these results and the promising safety profile of griffithsin (16, 38), Cc-griffithsin represents an excellent option for additional testing as a potential HIV-1 microbicide.

Cc-GBVCE2 is another promising microbicide candidate, as it was able to protect 60% of BLT mice from vaginal infection with HIV-1. While these studies focused on the use of Cc-GBVCE2 as a topical agent to prevent HIV-1 transmission, it is possible that Cc-GBVCE2 may also have utility as a therapeutic option for HIV-1 infection. It has been reported that people coinfected with GB virus C and HIV-1 have a better prognosis and slower progression to AIDS, suggesting that GB virus C has beneficial effects on HIV-1 disease through multiple mechanisms (34, 35, 39–42), including the E2 portion of GB virus C being a putative fusion peptide that interferes with HIV binding or fusion and that prevents cell-to-cell spread (42–44). As such, Cc-GBVCE2 may be able to limit the spread of HIV-1 in infected individuals, as well as lower viral shedding to prevent transmission.

Interestingly, some mice appeared to be protected from HIV-1 infection initially but were measured to be HIV-1 positive at later time points. Further studies are necessary to investigate the potential mechanism responsible for this. Furthermore, some mice that became HIV-1 positive after receiving a recombinant C. crescentus bacterium that provided significant protection from HIV-1 infection in other mice appeared to have higher levels of HIV-1 infection than control mice, which may be a result of the variability of immune reconstitution inherent in this model.

Three of the microbicide candidates that were able to provide significant protection from vaginal infection with HIV-1 also target HSV-2: Cc-griffithsin, Cc-elafin, and Cc-A1AT (13, 45, 46). HSV-2 infection is a major risk factor for HIV-1 acquisition, increasing the risk of HIV acquisition by 2- to 4-fold (47–49). As HSV-2 can increase HIV-1 acquisition, this suggests that these recombinant C. crescentus bacteria could have a significant impact on HIV-1 infection rates, not only directly by preventing HIV-1 infection but also by preventing HSV-2 acquisition. In particular, Cc-A1AT provided 50% protection from HIV-1 infection in BLT mice and provided an 86% increase in survival.
following vaginal infection in C56BL/6 mice, whereas Cc-griffithsin provided 75% protection from HIV-1 infection and 57% protection from HSV-2 disease (13). While these studies investigated using the recombinant C. crescentus individually, it is likely that combining the successful recombinant C. crescentus candidates will

FIG 4 Recombinant C. crescentus bacteria protect from vaginal infection with HIV-1_{JR-CSF} in BLT mice. Humanized BLT mice were infected intravaginally with 10,000 TCID<sub>50</sub> of HIV-1_{JR-CSF} in the presence or absence of 10<sup>8</sup> recombinant C. crescentus bacteria. Blood was collected from the mice biweekly for 6 weeks and analyzed by p24 ELISA and RT-qPCR. Data are presented as a modified survival curve indicating when mice seroconverted, and statistics were performed as a log-rank test comparing HIV-1-infected mice receiving Cc-control and HIV-1-infected mice receiving a recombinant C. crescentus bacterium. p24 ELISA and RT-qPCR data are presented as the mean ± SEM, and the presented statistics were performed by the Kruskal-Wallis test and represent the comparison between HIV-1-infected mice receiving Cc-control and HIV-1-infected mice receiving a recombinant C. crescentus bacterium. *, P < 0.05; **, P < 0.01. (a) Cc-griffithsin (n = 8); (b) Cc-GBVCE2 (n = 10); (c) Cc-elafin (n = 9); (d) Cc-A1AT (n = 10).
increase HIV-1 protection beyond those levels observed with each one individually. In addition, as all recombinant *C. crescentus* bacteria provided at least some low-level protection from HIV-1 infection, it is possible that combining multiple candidates that target different aspects of the HIV-1 attachment and entry process, including those that did not significantly reduce HIV-1 acquisition in BLT mice, may generate the most effective microbicide cocktail. In support of this, we have previously demonstrated that combining Cc-CD4 and Cc-MIP1α in vitro increases HIV-1 protection from 50 to 75% to 97% (11).

**FIG 5** Individual viral loads in HIV-1-positive BLT mice. BLT mice were infected intravaginally with HIV-1m.CSF in the presence of 10⁸ recombinant *C. crescentus* bacteria, and HIV-1 infection levels were measured biweekly by p24 ELISA and RT-qPCR. Results are shown for each individual mouse that was measured to be HIV-1 positive. Mouse identifiers for p24 ELISA and RT-qPCR results are matched for each recombinant *C. crescentus* bacterium when a mouse was measured to be HIV-1 positive by both techniques. (a) Cc-griffithsin; (b) Cc-GBVCE2; (c) Cc-elafin; (d) Cc-A1AT.
Our studies have focused on investigating a dosing strategy that would be coitally dependent, similar to a vaginal gel, as the first phase of testing. As a coitally dependent option may not be feasible in a real-world situation, we have begun to undertake additional studies to improve the utility of a \textit{C. crescentus}-based microbicide. Preliminary results from recombinant \textit{C. crescentus} in BLT mice are shown in FIG 6. BLT mice were infected intravaginally with 10,000 TCID\textsubscript{50} of HIV-1\textsubscript{JR-CSF} in the presence or absence of \(10^8\) recombinant \textit{C. crescentus} bacteria. Data are presented as a modified survival curve indicating when the mice were measured to be HIV positive, and statistics were performed as a log-rank test comparing HIV-1-infected mice receiving \textit{Cc-control} and HIV-1-infected mice receiving a recombinant \textit{C. crescentus} bacterium. (a) \textit{Cc}-microvirin (\(n = 8\)); (b) \textit{Cc}-cyanovirin (\(n = 3\)); (c) \textit{Cc}-Fuzeon (\(n = 8\)); (d) \textit{Cc}-T1249 (\(n = 10\)); (e) \textit{Cc}-CS2 (\(n = 8\)); (f) \textit{Cc}-CD4 (\(n = 7\)); (g) \textit{Cc}-CD4M33F23 (\(n = 7\)); (h) \textit{Cc}-MIP1\(\alpha\) (\(n = 7\)); (i) \textit{Cc}-indolicidin (\(n = 7\)); (j) \textit{Cc}-BmKn2 (\(n = 3\)).
nary data indicate that *C. crescentus* is able to prevent infection up to 8 h after application, and additional studies are ongoing to determine if this can be extended further. In addition, preliminary studies in an HSV-2 model suggest that *C. crescentus* may have some ability to prevent infection when applied after exposure. We have not investigated the use of recombinant *C. crescentus* as a long-term prevention option, such as on a vaginal ring, but it should be possible to assemble the recombinant S-layer proteins on a vaginal ring to provide long-term HIV-1 prevention. While *C. crescentus* is a nonpathogenic bacterium found in freshwater sources, including drinking water in the United States (50), the possibility does exist for immune responses to develop after prolonged usage. Our previous studies support a lack of immune response after the short-term use of *C. crescentus* in mouse models (13, 27, 51). In particular, the lipopolysaccharide (LPS) of *C. crescentus* has been found to produce greater than 100-fold less tumor necrosis factor than LPS isolated from *Escherichia coli* (51). To minimize the risk of an immune response, creating an abiotic delivery system containing just the recombinant S-layer proteins would greatly reduce the chance of an immune response.

HIV-1 remains a major global health priority, particularly for young women. A recombinant *C. crescentus*-based microbicide represents a safe and cost-effective option that should undergo further investigation for female-controlled prevention of HIV-1 as well as HSV-2. If a *C. crescentus* microbicide is developed, it could be combined with other prevention strategies for sexually transmitted infections and pregnancy as part of a comprehensive reproductive health package. In addition, adding a successful microbicide to the current HIV prevention tool kit will fulfill the need for a discreet, easy-to-use product that is targeted for women and that does not require male consent or continuous prophylactic antiretroviral use. Further preclinical studies are warranted to continue the investigation of a recombinant *C. crescentus*-based microbicide in a nonhuman primate model or clinical setting.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** Recombinant *C. crescentus* strain JS4038 was grown in PYE medium (0.2% peptone, 0.1% yeast extract, 0.01% CaCl₂, 0.02% MgSO₄) with 2 μg/ml chloramphenicol at 30°C. Gene segments were synthesized by Integrated DNA Technologies, Inc. (Coralville, IA), with codon usage being adapted for *C. crescentus*. *C. crescentus* bacteria displaying chimeric S-layer proteins have been previously described (11–13). The amino acid sequence for Cc-GBVCE2 is WDRGNVTLLCDCP. The synthesized DNA segment also specified BglII and SpeI restriction sites on the side and an NheI site on the 3’ end to facilitate directional cloning into p4BRsaA(723)/GSCC digested with Bgll and Nhel (52, 53).

**Preparation of *C. crescentus* cells.** *C. crescentus* S-layer display constructs were grown in PYE medium to an optical density at 600 nm of approximately 1 (3.1 × 10⁹ cells/ml). Cells were centrifuged and suspended in sterile water. This was repeated one time for *in vitro* experiments and three times for *in vivo* experiments.

**Cell lines.** 293T cells were a gift from Ninan Abraham (University of British Columbia) and were maintained in Dulbecco’s modified Eagle medium (DMEM) with 7.5% fetal bovine serum (FBS), 100 U/ml penicillin, and 0.1 mg/ml streptomycin. TZM-bl cells were obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH, and maintained in DMEM with 7.5% FBS (Gibco), 100 U/ml penicillin, and 0.1 mg/ml streptomycin (Gibco) as previously described (13). 174xCEM cells were obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH, and maintained in RPMI 1640 medium supplemented with 10% FBS as previously described (13).

**Primary cells.** Whole blood was collected and processed using Lymphoprep medium (Stemcell Technologies Inc.). PBMCs were grown in RPMI 1640 supplemented with 20% FBS.

**HIV-1.** The following reagents were obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: HIV-1BaL was from Jay Levy (54); pYK-JRCSF (catalog number 2708) was from Irvin S. Y. Chen and Yoshio Koyanagi (55–57) and was a gift from Zabrina Brumme (Simon Fraser University); HIV-1Sx1 was from Jay Levy (58); HIV-1296 was from Suzanne Gartner, Mikulas Popovic, and Robert Gallo (59, 60); and HIV-1m1 was from Irvin S. Y. Chen (56, 61, 62).

**HIV-1 propagation.** HIV-1296 was propagated in 174xCEM cells with 7.5 μg/ml DEAE-dextran (Sigma) and was harvested at the peak cytopathic effect. HIV-1Pac-B_C12 was propagated in 293T cells using the Lipofectamine 2000 reagent (Invitrogen) according to the manufacturer’s instructions. HIV-1m1 and HIV-1Sx1 were propagated in 5 × 10⁹ phytohemagglutinin (PHA)-stimulated PBMCs using 10 μg/ml hexamethylene bis(ethyleneimine) (Polybrene; Sigma). HIV-1296 was propagated in PHA-stimulated PBMCs with 7.5 μg/ml DEAE-dextran.

**Virus titration.** Serial dilutions of virus were prepared in 96-well plates using medium containing 7.5 μg/ml DEAE-dextran. TZM-bl cells (10,000) were added. The plates were maintained at 37°C in 5% CO₂ for 48 h. Infection of cells was measured indirectly using a mammalian β-galactosidase assay kit (Pierce),
followed by reading of the absorbance at 415 nm. An absorbance of greater than 0.2 was considered a positive infection. The 50% tissue culture infectious dose (TCID<sub>50</sub>) per milliliter was determined for each viral stock by identifying the dilution of virus in which 50% of the TZM-bl cells were infected.

**Virus blocking experiments.** The *C. crescentus* constructs were grown and prepared as described above. Experiments were carried out in quadruplicate wells of 96-well plates. The volume of virus added was determined by calculating the 200 TCID<sub>50</sub> value, and 1 × 10<sup>6</sup> *C. crescentus* cells were added to each well. The virus and *C. crescentus* constructs were incubated for 1 h at 37°C before adding 10,000 TZM-bl cells or PBMCs and 7.5 μg/ml DEAE-dextran to each well. Cc-MIP1α was incubated with the TZM-bl cells or PBMCs for 1 h before virus addition. The level of infection was determined after 48 to 72 h by use of a β-galactosidase assay kit (TZM-bl cells) or a p24 ELISA (ZeoMetrix Corporation and ProSci Incorporated) according to the manufacturer’s instructions (PBMCs). For the β-galactosidase assays, the data are presented and determined as a percentage of infection of the wells with Cc-bl plus HIV-1 with the background from uninfected TZM-bl cells subtracted. p24 ELISA data were normalized to infection of the untreated control wells with PBMCs and HIV-1, set as 100%.

**Preparation of humanized bone marrow-liver-thymus (BLT) mice.** NSG mice were obtained from The Jackson Laboratory and maintained in the Modified Barrier Facility at the University of British Columbia. Six- to 12-week-old female mice were used in the experiments. Fetal liver was split, and 1-mm<sup>3</sup> pieces of liver and thymus were implanted under the kidney capsule. Autologous fetal liver tissue was used for CD34<sup>+</sup> cell isolation using a CD34<sup>+</sup> cell positive selection kit from Stemcell Technology Inc. according to the manufacturer’s instructions, and cells were frozen in 90% human serum–10% dimethyl sulfoxide. At 3 weeks following the surgical implantation, mice received 225 Gy of irradiation from an X-ray source, followed by intravenous injection of 100,000 to 200,000 CD34<sup>+</sup> cells within 30 h. The mice were infected with Cc-bl virus before blood was characterized for human immune cell reconstitution.

**Flow cytometry.** One hundred microliters of blood was collected from BLT mice and treated with EDTA. Red blood cell lysis buffer was added, and samples were incubated for 15 min before washing with fluorescence-activated cells sorting buffer (phosphate-buffered saline [PBS] plus 2% FBS). Cells were stained with anti-mouse CD45 Pacific Blue (eBioscience), mouse anti-human CD45 phycoerythrin (PE)-Cy7 (BD Pharmingen), mouse anti-human CD3 Alexa Fluor 700 (eBioscience), mouse anti-human CD4 PE (BD Pharmingen), mouse anti-human CD8 PE-Cy5 (BD Pharmingen), mouse anti-human CD14 allophycocyanin (APC)-Cy7 (BD Pharmingen), mouse anti-human CD19 APC (BD Pharmingen), and mouse anti-human CD56 fluorescein isothiocyanate (FITC) (BD Pharmingen). Data were acquired on an LSR II flow cytometer and analyzed using FlowJo software (TreeStar). Mouse versus human CD45<sup>+</sup> cells were examined for expression of CD34<sup>+</sup> cells and include human CD45<sup>+</sup> cells. The human CD45<sup>+</sup> cells were gated for expression of CD3. CD3<sup>+</sup> cells were examined for expression of CD4 and CD8. The CD3<sup>+</sup> cells were examined for expression of CD19 (B cells) versus CD56 (NK cells). Unstained and single-stained human PBMCs were used as controls.

**HIV-1<sub>r,cres</sub> infection.** Mice were anesthetized before atraumatic vaginal infection with 10,000 TCID<sub>50</sub> of HIV-1<sub>r,cres</sub> in the presence or absence of 10<sup>6</sup> *C. crescentus* bacteria. The *C. crescentus* bacteria were mixed with HIV-1<sub>r,cres</sub> less than 5 min before inoculation into the vaginal tract in a volume of 20 μl using a sterile p200 pipette tip, and then the entire volume was inoculated intravaginally. Sixteen independent cohorts of BLT mice were created for these studies. Each cohort contained at least one mouse infected with HIV-1<sub>r,cres</sub> only and one mouse infected with HIV-1<sub>r,cres</sub> and treated with Cc-control. Each recombinant *C. crescentus* bacterium was tested in mice from at least 2 different cohorts.

**HIV-1<sub>r,cres</sub> infection analysis.** Blood was collected from the retro-orbital sinus on days 0, 14, 28, and 42 postinfection. Blood was concentrated at 8,000 rpm for 15 min, and serum was aliquoted and frozen at −80°C before being analyzed by p24 ELISA (ZeoMetrix) and RT-qPCR. The p24 ELISA was performed according to the manufacturer’s instructions and had a limit of detection of 3.9 pg/ml. Viral RNA was extracted from equal volumes of serum using a QiAamp viral RNA minikit (Qiagen). Equal volumes of RNA were converted to cDNA using an Applied Biosystems cDNA kit (Thermo Fisher Scientific). Equal volumes of cDNA were run in duplicate on an Mx3005p PCR multiplex quantitative PCR instrument (Stratagene) with Sybr green (Bio-Rad), forward primer ATCAAGCAGCTATGCAAATGCT, and reverse primer GTGAAG GGTACTAGTAGTCCCTGCTATGTC; the settings were 50°C for 2 min, 95°C for 10 min, and 40 cycles at 95°C for 15 s and 60°C for 1 min. The limit of detection was 500 copies/ml. Mice were considered HIV-1 negative if HIV-1 was not detected by both the p24 assay and the RT-qPCR at all time points.

**Statistics.** Statistical analysis was performed with GraphPad Prism software. Viral blocking assay results are reported as the mean ± standard error of the mean (SEM). Experiments were conducted in quadruplicate wells and repeated in a minimum of three independent experiments, unless indicated otherwise. An unpaired two-tailed t test (for comparison of the results for mice receiving the Cc-control and those receiving 1 recombinant *C. crescentus* bacterium) or analysis of variance (ANOVA) with Bonferroni’s correction for multiple comparisons (for comparison of the results for mice receiving the Cc-control and >2 groups receiving recombinant *C. crescentus* bacteria) was used as appropriate to evaluate the significance of differences between groups. Statistical analysis for BLT mice was performed using the log-rank (Mantel-Cox) test for survival curves and the Kruskal-Wallis test with Dunn’s multiple-comparison test for p24 ELISA and RT-qPCR analysis. The reported statistics represent the comparison between mice infected with HIV and receiving Cc-control and mice infected with HIV and receiving a recombinant *C. crescentus* bacterium. A P value of <0.05 was considered statistically significant.

**Study approval.** All animal work was approved by the University of British Columbia Animal Care Committee (protocols A13-0055, A13-0234, and A12-0245). Human ethics approval was obtained from the University of British Columbia Clinical Ethics Board (H12-02480). Human PBMCs were obtained from healthy donors after they provided informed written consent. Second-trimester human fetal liver and...
thymus tissue was obtained after informed written consent from women undergoing elective abortion and procured by Advanced Bioscience Resources Inc. (Alameda, CA).

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