Research Article

In Vivo Longitudinal Imaging of Experimental Human Papillomavirus Infection in Mice with a Multicolor Fluorescence Mini-Endoscopy System

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Abstract

Human papillomavirus (HPV) infection is the most common sexually transmitted infection. Vaccines for HPV infection can reduce the risk of developing cervical cancer. To further improve such vaccines and to explore other methods of preventing or treating viral infection, longitudinal studies in experimental animals are desirable. Here, we describe a newly developed multicolor endoscopic fluorescence imaging system to visualize early HPV infection with fluorescent protein–encoded pseudoviruses (PsV) in the female genital tract of living mice. With this imaging method, the course of HPV PsV infection and the effects of intervention to prevent infection can be monitored in a single mouse over time. Female immunocompetent or athymic mice were pretreated with a vaginal spermicide and then HPV PsV composed of an authentic viral capsid and encapsidating green or red fluorescent protein (GFP or RFP) reporter gene was intravaginally instilled. Expression of GFP or RFP was detected 1 day after PsV challenge, which peaked after 2 or 3 days and decreasing thereafter. No fluorescence was detected in vaccine-treated immunocompetent mice. By using serial infection of the same PsV type (HPV16) encoding either GFP or RFP, different infection patterns of repeated exposure can be monitored. This method offers the ability to monitor experimental virus infections before and after intervention, thereby accelerating the development of appropriate prevention and therapy. Cancer Prev Res; 4(5); 767–73. ©2011 AACR.

Introduction

Human papillomavirus (HPV) infection is usually acquired by sexual contact, and up to 80% of sexually active women show evidence of HPV infection at some stage of their lives. Although HPV infection is usually asymptomatic, more than 90% of cervical cancers are associated with HPV infection (1, 2). HPV16 is detected in approximately 50% of cervical cancers worldwide. The recently developed and U.S. Food and Drug Administration-approved HPV vaccines, Gardasil (Merck) and Cervarix (GlaxoSmithKline), markedly reduce the risk of infection by the vaccine-targeted HPV types, thereby preventing cervical precancerous lesions, provided the vaccination occurs prior to HPV exposure (3, 4). To develop the next generation of vaccines with maximal potency and to explore interventions that prevent infection after exposure to HPV, studies using large numbers of mice, euthanized at various time points, must be conducted. Alternatively, a smaller number of animals can be serially studied over time, using each mouse as its own control. Although both methods are valid, the latter approach may be more cost-effective and realistic. We have developed a multicolor fluorescence mini-endoscopic imaging system with a highly sensitive charge-coupled device (CCD) camera and multicolor fluorescence imaging capabilities for real-time monitoring of the viral infection with HPV pseudovirus (PsV) encapsidating green or red fluorescent protein (GFP or RFP). Animals were observed longitudinally after experimental interventions to determine the effectiveness in reducing or preventing infection.

Materials and Methods

Reagents

Depo-Provera was purchased from Pfizer and diluted to a final concentration of 30 mg/mL in PBS. Carboxymethylcellulose (CMc) was purchased from Sigma-Aldrich and dissolved in deionized water. Conceptrol, which contains 4% of nonoxynol-9 (N-9), was purchased from McNeil-PPC Inc. Gardasil was purchased from Merck. Intron A was purchased from Schering-Plough and dissolved in distilled...
water. It was mixed with CMC to make a final concentration of 2% CMC.

Pseudovirus

HPV16 pseudovirus (PsV) with GFP or tdTomato (RFP) fluorescent reporter constructs were prepared as previously described (5–7).

HPV PsV challenge in vivo

All procedures were conducted in compliance with the Guide for the Care and Use of Laboratory Animal Resources (1996), National Research Council, and approved by the local Animal Care and Use Committee. Six- to 8-week-old female Balb/c or homozygote athymic mice were purchased from Charles River (NCI-Frederick). During the procedure, mice were anesthetized with isoflurane. HPV PsV challenge was conducted according to a previous report (5) with a slight modification. In brief, mice were injected subcutaneously with 3 mg of Depo-Provera 4 days before HPV PsV challenge. For chemical disruption, mice were pretreated with intravaginal administration of 50 μL of N-9. For mechanical disruption, a commercially available model BF XP-60 bronchoscope (Olympus Co.), 2.8 mm in diameter with a single biopsy channel, was inserted into the mice vagina with gentle insufflation of carbon dioxide. With white light endoscopic imaging (CLV-180; Olympus Co.), a cervical specimen was disrupted using biopsy forceps (FB-56D-1; Olympus Co.). Four hours after chemical or mechanical disruption, a HPV PsV titer of ~2.5 x 10^7 infectious units in 20 μL of 2% CMC was administered intravaginally. For vaccination, Balb/c mice were injected intramuscularly with 6 μg of Gardasil once or twice, 3 weeks apart. Seven days after the last immunization, mice received HPV PsV challenge. For interferon (IFN) treatment, 1 MIU (million international units) of Intron A dissolved in 20 μL of 2% CMC was administered intravaginally.

Fluorescence endoscopy and image analysis

Mice were examined with fluorescence endoscopy every day for up to 14 days after HPV PsV challenge. A model BF XP-60 bronchoscope system was inserted intravaginally while the animal was under anesthesia, and cervicovaginal epithelium was observed with white light imaging and dual fluorescence imaging by switching back and forth between the blue (465–500 nm) or green (530–555 nm) excitation filters. Endoscopic images were obtained via a dual fluorescence imaging by switching back and forth between the blue (465–500 nm) or green (530–555 nm) excitation filters. This enabled the simultaneous, side-by-side imaging of white, or excitation, light imaging; and a highly sensitive EM-CCD camera for fluorescence imaging of the cervicovaginal epithelium (Fig. 1A). Camera gain, exposure time, and binning for the fluorescence images were held constant in each fluorescent protein throughout the study.

To compare fluorescence intensities during different HPV PsV challenges, the distance between the cervical opening and the endoscope head was maintained using the biopsy forceps as a guide. Snapshot images obtained from the hard disk recorder were used for calculating fluorescence intensities. All fluorescence images were analyzed with Image J software [http://rsweb.nih.gov/ij/]. Circular regions of interest (ROI) were placed in the same regions of cervical epithelium in each mouse to determine the change of fluorescence intensities over time. The pixel intensities, varying between 0 and 255, were obtained from each ROI. Normal background signal was determined by measuring signal intensity prior to HPV PsV challenge to remove autofluorescence and camera noise.

Validation of HPV PsV infection

Mice were examined with fluorescence endoscopy 2 days after HPV PsV challenge. The green fluorescence filter set was used to do real-time biopsies with biopsy forceps. Smears were prepared from the fresh biopsy specimens and examined with fluorescence microscopy (BX51; Olympus America) equipped with the following filters: excitation wavelength, 450 to 490 nm; emission wavelength, 500 to 550 nm. Transmitted light differential interference contrast (DIC) images were also acquired.

Statistical analysis

Data are expressed as mean ± SEM. Analyses were carried out using a statistics program (GraphPad Instat, version 3.06; GraphPad Software). For comparison of groups, the Mann–Whitney rank-sum test was used. A level of P < 0.05 was considered statistically significant.

Results

Monitoring the HPV PsV infection by fluorescence mini-endoscopy in vivo

Quantitative analysis of the initial phase of HPV has previously been carried out by intravaginal infection of mice with HPV PsV encapsidating plasmids encoding genes for RFP. To assess the kinetics of gene expression, large numbers of animals were required to obtain ex vivo imaging of the vaginal tracts over time (5). Here, we sought to repeatedly monitor localized HPV PsV infection kinetics in the same animal by employing a multicolor mini-endoscope based on a commercial clinical endoscope, which allows for the independent detection of GFP and RFP fluorescence. This mini-endoscope uses a clinically available light source and fiber-optic endoscope in addition to in-house designed excitation filters, a dichroic splitter, emission filters, and 2 cameras; a color camera for white light imaging; and a highly sensitive EM-CCD camera for detecting fluorescence. This enabled the simultaneous, side-by-side imaging of white, or excitation, light imaging and fluorescence imaging of the cervicovaginal epithelium.
This system, therefore, has the ability to image green, red, and near infrared (NIR) fluorescence with the appropriate filter set (Fig. 1B). For quantitating fluorescence intensity over time, the distance between the cervical opening and the endoscope head was held constant at 3.0 mm (Fig. 1C).

After pretreatment with a vaginal spermicide, N-9, an HPV16 PsV encapsidating either a GFP or an RFP expression plasmid was administered intravaginally to the animals (5). Expression of GFP or RFP was detected as early as 1 day after PsV challenge ($n = 6$ mice each), peaked at 2 or 3 days, and then decreased thereafter to the preinfected baseline (Fig. 2A–D). Both GFP and RFP encoding HPV PsVs behaved in the same manner. Settings of camera gain, exposure time, and binning for the fluorescence images were held constant throughout the study. There was almost no background signal arising from the cervicovaginal epithelium, as only minimal autofluorescence could be detected prior to infection. In addition, no cross-talk was observed between GFP and RFP fluorescence and the sensitivity of the fluorescence did not vary significantly between the two (Fig. 2A and C). Moreover, HPV PsV, which did not encapsidate a fluorescence reporter plasmid, or no HPV PsV challenge showed no detectable fluorescence (Supplementary Fig. S1). These results indicate that it is possible to detect and monitor HPV infection over time in the cervicovaginal epithelium in living mice, using this fluorescence mini-endoscopy system (Supplementary Video S1A and B).

To assess the infection efficiency of HPV PsV, simultaneous coinfection with approximately equal titers of GFP and RFP encoding HPV PsVs was carried out. Both GFP and RFP expression were detected by day 1 and peaked on day 2, indicating that there were no significant differences in infectivity between the GFP- and RFP-expressing PsVs (Supplementary Fig. S2A). Merged green and red fluorescent images obtained 2 days after infection revealed colocalization of GFP and RFP fluorescence ($n = 3$ mice; Supplementary Fig. S2B). Next, biopsy forceps were used to mechanically disrupt the cervical epithelium, thereby promoting infection with HPV PsV. During endoscopy, the dorsal cervix was selectively disrupted and then mice received the HPV PsV challenge. Fluorescence was observed only in the rim and surrounding mucosa of the disrupted lesion, and peak expression was observed on day 2 after the HPV PsV challenge, which was similar to the results with chemical disruption ($n = 3$ mice; Supplementary Fig. S3). These results confirm the findings of Roberts and colleagues that epithelial abrasion, whether induced chemically...

**Figure 1.** Multicolor fluorescence mini-endoscopic imaging of the mouse cervix. A, multicolor fluorescence imaging system is based on a clinically available fiber-optic endoscope and light source. Excitation light is provided by in-house designed excitation filters, which are switchable to either the visible light filter (white light) or band-pass filter. Endoscopic images were obtained through a dichroic splitter, wherein the excitation light images were displayed with an image processor, and the fluorescence images were filtered by multicolor emission filters before reaching the EM-CCD camera. Both images are displayed side-by-side on the monitor. B, ex vivo images of the phantoms (lymph nodes) incorporating fluorescent tracers [Rhodamine green (close to GFP), TAMRA, RhodamineX (close to RFP), Cy5.5, ICG] showed the capability of multicolor fluorescence detection. Excitation light image (EL) and the fluorescence image (FL) were simultaneously displayed on the screen side by side. The emission filter spectrums are indicated. C, white light endoscopic imaging of the mouse cervix. Biopsy forceps (tip = 1-mm diameter) were used to gauge the distance to the cervix that was held constant.
or mechanically, is required for HPV PsV infection (ref. 5; Supplementary Figs. S2 and S3).

Next, biopsy specimens (n = 3) were obtained from fluorescing lesions of the cervicovaginal epithelium during endoscopy. Fluorescent lesions were selectively sampled using biopsy forceps. Cells exhibiting GFP expression at the site of biopsy was confirmed with fluorescence microscopy (Supplementary Fig. S4 and Video S2).

Detection of HPV PsV infection after intervention

Previous studies report that intramuscular vaccination with native L1 virus-like particles (VLP) can induce antibody-mediated, type-restricted protection against experimental papillomavirus infection (8, 9). To test whether intravaginal instillation of HPV capsids can generate protective immunity, 14 days after the initial vaginal infection, immunocompetent Balb/c and immunocompromized homozygote athymic mice were vaginally infected using an HPV PsV encapsidating the opposite fluorescent reporter plasmid from what was employed during the first HPV PsV challenge (e.g., GFP encoding HPV PsV followed by RFP encoding HPV PsV, and vice versa). The use of 2 different probes allowed the second infection to be distinguished from the first, but it also abrogated the possibility of immunity generated against the first fluorescent protein. There was no detectable fluorescence after the second HPV PsV challenge in immunocompetent Balb/c mice (Fig. 3A), whereas fluorescence could be detected in immunocompromised athymic mice (Fig. 3B; n = 3 mice each), suggesting that acquired immunity against the HPV capsid played a role in the inhibition of HPV PsV infection.

Next, Balb/c mice were immunized once or twice at an interval of 3 weeks with 6 µg of Gardasil, a quadrivalent vaccine composed of L1 VLPs from HPV types 6, 11, 16, and 18, to confirm the effect of vaccination. Seven days after the last immunization, mice received an HPV16 PsV challenge. Consistent with recent studies showing immunization with HPV VLPs protects mice from vaginal PsV challenge (10, 11), there was no apparent fluorescence after HPV PsV challenge at any time point in both vaccination groups (n = 4 mice each), indicating that the vaccine induced protection from infection (Fig. 3C shows results).

Topical IFN was then tested as an antiviral therapy. Intron A (10^6 MIU), a recombinant human IFNα-2b (rhIFNα-2b) that induces antiviral activity by suppressing...
the transcription and translation of viral genes, was administered intravaginally to Balb/c or athymic mice every day starting 3 days before HPV PsV challenge. Fluorescence was greatly reduced in both immunocompetent and immunocompromised mice pretreated with IFN (n = 4 mice each), indicating that topical IFN inhibits HPV PsV infection and/or suppresses the transcription of the reporter gene (Fig. 3C).

Collectively these results suggest that various strategies to inhibit HPV PsV cervicovaginal infection can be assessed in vivo using HPV PsVs encapsidating different fluorescent protein genes. The imaging technique described here allows for the monitoring of the early stages of HPV infection over time by minimally invasive fluorescence endoscopy in the cervicovaginal tract of living immunocompetent or immunocompromised mice.

Discussion

The pathologic consequences of HPV infection are strictly species- and tissue-restricted to humans. However, the early events of binding, entry, and genome release seem not to be species-restricted and therefore cervicovaginal infection with HPV PsV in mice provides a platform for the examination of the initial phase of papillomavirus infection. Infection is promoted by trauma of the cervicovaginal epithelium, which exposes the basement membrane and the basal epithelial cells to virus. Fluorescent proteins are used as a robust tool for investigating the viral infection, as well as the cancer research in vivo (12–16). A previous study showed that the spermicide N-9 increased the likelihood of HPV infection, which was shown with ex vivo optical fluorescence utilizing GFP and RFP encapsidated by the
PsV (5). This experiment required multiple mice at each time point to reach statistical end points. The key technology advance reported herein is the development of a mini-endoscope that permits fluorescence imaging using a highly sensitive EM-CCD camera. By changing the filter sets, multicolor imaging becomes possible. The advantages of this are (i) the infection in a single living mouse can be followed over time and (ii) multicolor fluorescence signals can be analyzed to separately test the impact of a second HPV PsV infection by using a PsV encapsidating a different reporter gene. Furthermore, the use of this method provides a means to accurately induce and image pinpoint infections by creating specific mini-abrasions or sample the infected tissue with fluorescence-guided biopsy under endoscopic guidance (Supplementary Fig. S3 and Video S2).

As there was no cross-talk between GFP and RFP fluorescence (17), preventive and early therapeutic interventions could be used to determine resistance to reinfec- tion by using serial infections with different HPV PsVs. Therefore, infecting with the same type of HPV PsV but with different color reporters can be useful in the investigation of the duration and intensity of the immune response against the primary infection or vaccination (Fig. 3A and B). For instance, in this study, infection was prevented or ameliorated by prior vaccination with Gardasil and topical administration of rhIFN (Fig. 3C), as shown by longitudinal observations of reduced fluorescent gene expression after intervention.

In vivo observation of HPV PsV infection has also been successfully done using a bioluminescence reporter imaging system (Supplementary Fig. S5; refs. 18, 19). There are 2 technical advantages of our method over the bioluminescence imaging: (i) real-time multiple color capability and (ii) direct access to the tissue for localizing infection or sampling infected cells, resulting in much higher resolution imaging. From the photophysical point of view, the endoscope can minimize light scattering and absorbance caused by overlapping tissue, resulting in more accurate depiction of the lesion and, theoretically, allowing better quantitative measurements of the infection. Moreover, this method does not require administration of an exogenous substrate, luciferin, which is necessary for bioluminescence (20). Furthermore, GFP and RFP are readily translated to ex vivo histopathology whereas this is more difficult with bioluminescence.

Until recently, it has not been possible to directly visualize HPV PsV infection in vivo. Improvements in optical technology allow placement of mini-endoscopes in small cavities with full fluorescence capabilities in real time. These tools provide an opportunity to investigate the effects of preventive and therapeutic interventions for HPV PsV infections in small rodents, much as they would be investigated in humans.

Disclosure of Potential Conflicts of Interest

No conflict of interest was disclosed.

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