Severe Restriction of Xenotropic Murine Leukemia Virus-Related Virus Replication and Spread in Cultured Human Peripheral Blood Mononuclear Cells*

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Received 7 January 2011/Accepted 8 February 2011

Xenotropic murine leukemia virus-related virus (XMRV) is a gammaretrovirus recently isolated from human prostate cancer and peripheral blood mononuclear cells (PBMCs) of patients with chronic fatigue syndrome (CFS). We and others have shown that host restriction factors APOBEC3G (A3G) and APOBEC3F (A3F), which are expressed in human PBMCs, inhibit XMRV in transient-transfection assays involving a single cycle of viral replication. However, the recovery of infectious XMRV from human PBMCs suggested that XMRV can replicate in these cells despite the expression of APOBEC3 proteins. To determine whether XMRV can replicate and spread in cultured PBMCs even though it can be inhibited by A3G/A3F, we infected phytohemagglutinin-activated human PBMCs and A3G/A3F-positive and -negative cell lines (CEM and CEM-SS, respectively) with different amounts of XMRV and monitored virus production by using quantitative real-time PCR. We found that XMRV efficiently replicated in CEM-SS cells and viral production increased by >4,000-fold, but there was only a modest increase in viral production from CEM cells (<14-fold) and a decrease in activated PBMCs, indicating little or no replication and spread of XMRV. However, infectious XMRV could be recovered from the infected PBMCs by cocultivation with a canine indicator cell line, and we observed hypermutation of XMRV genomes in PBMCs. Thus, PBMCs can potentially act as a source of infectious XMRV for spread to cells that express low levels of host restriction factors. Overall, these results suggest that hypermutation of XMRV in human PBMCs constitutes one of the blocks to replication and spread of XMRV. Furthermore, hypermutation of XMRV proviruses at GG dinucleotides may be a useful and reliable indicator of human PBMC infection.

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† Published ahead of print on 16 February 2011.
and named DIG cells (D by the SV40 terminator. Over 8,000 colonies of transfected cells were selected in vening MLV sequences. The R and U5 in the 3'/H11032 a simian virus 40 (SV40) promoter, is located after MLV RNA. The hygromycin phosphotransferase B gene (previously (20).

85-bp stretch of homology that is able to direct minus-strand transfer, resulting - and 3' after U3 in the 5'-LTR, respectively. The two GFP fragments share a long terminal repeats (LTRs) in which the sequences coding for C- and N-ter- canine osteosarcoma cells were transfected with the previously described MLV- reverse transcription-PCR (RT-PCR) for detection of XMRV 

XMRV RNA copy numbers were determined by using quantitative real-time 
m filter, and viral RNAs were isolated using a QIAamp viral RNA minikit (Qiagen). To avoid DNA contamination, the isolated RNA was treated for 30 min with DNase I, using the Turbo DNA-free kit (Ambion). To quantify the RNA copy numbers, quantitative real-time RT-PCR was performed using the LightCycler 480 RNA master hydrolysis probes reaction mixture (Roche) and the primers and probes described previously for detection of XMRV env (20). A serial 10-fold dilution of VP62 plasmid (9), kindly provided by Robert Silverman, was used to generate a standard curve.

to determine XMRV proviral copy numbers, infected CEM and CEM-SS cells were harvested at day 3 and day 13 after infection, and genomic DNA was isolated using a DNeasy blood and tissue kit (Qiagen). Genomic DNA (500 ng) was used to quantify the proviral copy numbers by using primers that detect XMRV gag as previously described (20). Quantitative real-time PCR was performed in duplicate and normalized to the CCR5 gene by using primers and probes as previously described (35).

Recovery of infectious XMRV from PHA-activated PBMCs. To recover infec- tious virus from XMRV-infected PBMCs, 8 × 10⁷ DIG cells were cocultivated with PBMCs (1 × 10⁶ cells) that were infected with culture supernatant containing 10⁴, 10⁵, 10⁶, 10⁷, or 10⁸ XMRV RNA copies and harvested 7 days postinfection. Cells were transferred to a 10-cm dish 3 days postinfection and maintained for 25 days. After 14 and 25 days, cells were harvested and the percentage of GFP-expressing cells was determined by fluorescence-activated cell sorting (FACS) analysis.

G-to-A hypermutation of XMRV proviruses in infected PBMCs. Genomic DNA from XMRV-infected PBMCs was isolated with the DNeasy blood and tissue kit (Qiagen), and a 1.2-kb XMRV fragment was amplified, cloned, and sequenced as described previously (37).

Western blot analysis of A3G and A3F expression in PHA-activated PBMCs. A3G and A3F expression was analyzed by Western blotting as previously de- scribed (37). Endogenous A3G and A3F proteins were detected by using an anti-A3G apoC17 antibody (1:5,000 dilution) and rabbit anti-human A3F (C18; 1:500 dilution; AIDS Research and Reference Reagent Program, Division of AIDS, NIAID, NIH), anti-apoC17 was from Klaus Strebel, and anti-human APOBEC3F was obtained from Michael Malim. Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was detected using anti-GAPDH antibody (Abcam) and served as a loading control.

RESULTS

XMRV replication is restricted in CEM cells but not in CEM-SS cells. The protocol used to compare the kinetics of replication and spread of XMRV in A3G/A3F-expressing CEM and A3G/A3F-deficient CEM-SS cells is outlined in Fig. 1A. Infectious XMRV was harvested from prostate carcinoma cell line 22Rv1. The 22Rv1 cell line has been shown to produce large amounts of a virus that is virtually identical to XMRV, originally found in prostate cancer samples (22, 56). We quanti- fied the amount of XMRV harvested from the 22Rv1 cell line by using real-time RT-PCR to detect XMRV env RNA as previously described (20).

Generation of canine indicator cells for detection of infectious XMRV. D17 canine osteosarcoma cells were transfected with the previously described MLV-based plasmid pMS2-FP-GF-no3R (6). Briefly, the plasmid contains modified long terminal repeats (LTRs) in which the sequences coding for C- and N-terminal fragments of green fluorescent protein (GFP, FP and GP) were inserted after U3 in the 5'- and 3'-LTR, respectively. The two GFP fragments share a 85-bp stretch of homology that is able to direct minus-strand transfer, resulting in the reconstitution of full-length gfp upon reverse transcription of this packaged RNA. The hygromycin phosphotransferase gene (hygro), under the control of a simian virus 40 (SV40) promoter, is located after MLV ², replacing all inter- vening MLV sequences. The R and U5 in the 3' untranslated region is replaced by the SV40 terminator. Over 8,000 colonies of transfected cells were selected in medium containing hygromycin at a final concentration of 120 μg/ml, pooled, and named DIG cells (D17 indicators of gammaretroviruses).

XMRV infection and replication. CEM and CEM-SS cells (1 × 10⁶ cells) as well as PHA-activated PBMCs (6 × 10⁶ or 1 × 10⁶ cells) were infected with XMRV obtained from 22Rv1 cells at different dilutions. The XMRV RNA copies were quantified in the virus stock by using quantitative real-time RT-PCR, and different viral RNA copy numbers (10⁴, 10⁵, 10⁶, 10⁷, 10⁸, or 10⁹) were used to infect cells. The cells were washed 6 h after infection and maintained in 6 ml of medium. Two milliliters of culture medium was harvested every day, frozen at −80°C, and replaced with fresh medium.

Quantitative real-time PCR. To quantify XMRV RNA copy numbers, culture supernatants containing virus were collected and filtered through a 0.45-μm filter, and viral RNAs were isolated using a QIAamp viral RNA minikit (Qiagen). To avoid DNA contamination, the isolated RNA was treated for 30 min with DNase I, using the Turbo DNA-free kit (Ambion). To quantify the RNA copy numbers, quantitative real-time RT-PCR was performed using the LightCycler 480 RNA master hydrolysis probes reaction mixture (Roche) and the primers and probes described previously for detection of XMRV env (20). A serial 10-fold dilution of VP62 plasmid (9), kindly provided by Robert Silverman, was used to generate a standard curve.

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The amounts of XMRV produced from the infected CEM and CEM-SS cells are shown in Fig. 1B to G. Neither CEM nor CEM-SS cells infected with 22Rv1 culture supernatant containing 10⁴ RNA molecules produced detectable amounts of XMRV (detection limit, ~1.1 × 10⁴ copies/8.4 μl) (Fig. 1B). XMRV particles were detected in both CEM and CEM-SS cells infected with 10⁵, 10⁶, 10⁷, or 10⁸ viral RNA copies (Fig. 1C to G, respectively); virus production at the beginning
FIG. 1. Replication of XMRV in CEM and CEM-SS cells. (A) Schematic overview of the experimental protocol. Supernatant containing XMRV was harvested from 22Rv1 cells, and XMRV RNA copies in the supernatant were determined by quantitative real-time RT-PCR using primers and a probe for a region of XMRV env. Culture supernatants containing different amounts of XMRV RNA copies were then used to infect 1 × 10^6 CEM or CEM-SS cells. At 6 h postinfection, cells were extensively washed, and culture supernatants were harvested every day from days 0 to 14. XMRV RNA copy numbers were determined by quantitative real-time RT-PCR. (B to G) Replication kinetics of XMRV 22Rvl in CEM and CEM-SS cells infected with 22Rv1 culture supernatant containing 10^4 XMRV RNA copies (B), 10^5 XMRV RNA copies (C), 10^6 XMRV RNA copies (D), 10^7 XMRV RNA copies (E), 10^8 XMRV RNA copies (F), or 10^10 XMRV RNA copies (G). The viral RNA in the supernatant of infected cells was isolated, and XMRV RNA was detected by quantitative real-time RT-PCR. The results are plotted as copies of viral XMRV RNA per 8.4 µl of culture supernatant of infected cells. The detection limit for quantitative RT-PCR was 1.1 × 10^2 copies/8.4 µl (indicated as a solid line). A representative experiment from two independent experiments is shown. Error bars indicate standard deviations. (H) Determination of increases in viral RNA copies in CEM and CEM-SS cells 14 days postinfection. The increase in viral RNA copies produced from CEM and CEM-SS cells infected with different amounts of 22Rvl XMRV are shown in panels B to G. The average number of RNA copies per 8.4 µl for days 12 to 14 was divided by the average number of RNA copies per 8.4 µl from days 1 to 3.
of infection (days 1 to 3) was dependent on the amount of XMRV used to infect the cells, and there was no difference in the virus production between CEM and CEM-SS cells (Fig. 1B to G).

Virus production increased to detectable levels in infected CEM-SS cells starting around days 5 to 7, whereas viral production either did not increase or increased only slightly in CEM cells infected with \(10^5\) to \(10^{10}\) XMRV RNA copies (Fig. 1C to G). Virus production did not increase in either CEM-SS or CEM cells infected with \(10^4\) RNA copies, indicating that exposure to culture supernatant containing \(10^4\) XMRV RNA copies did not result in an infection event (Fig. 1B). In contrast, while no virus production could be detected in CEM cells infected with \(10^5\) RNA copies, infection was clearly established in the CEM-SS cells, as evidenced by the dramatic increase in virus production from less-than-detectable levels at day 6 to \(2.5 \times 10^5\) copies per 8.4 \(\mu\)L at day 14 (Fig. 1C). A similar increase in virus production was observed in CEM-SS cells infected with \(10^6\), \(10^7\), and \(10^8\) RNA copies, whereas less than a 10-fold increase was observed in CEM cells (Fig. 1D, E, and F). In CEM-SS cells infected with \(10^{10}\) RNA copies, a modest 18-fold increase in virus production was observed, and no increase was observed in CEM cells (Fig. 1G).

To compare the increases in virus production, we compared the average number of viral RNA copies in culture supernatants from days 1 to 3 and days 12 to 14 to determine the increase in RNA copies (Fig. 1H). The results showed a remarkable increase in virus production in CEM-SS cells infected with \(10^5\), \(10^6\), \(10^7\), and \(10^8\) RNA copies; the increase in virus production ranged from \(~863\)-fold to \(~4,276\)-fold. In contrast, there was only a modest increase in virus production in CEM cells, which reached a maximum of \(~14\)-fold in cells infected with \(10^8\) RNA copies. Only an \(~18\)-fold increase in virus production was detected in CEM-SS cells infected with \(10^{10}\) RNA copies, suggesting that a large proportion of the cells were infected with the input virus, and a plateau was reached after all the cells were infected, resulting in superinfection interference. This result is consistent with the estimation that exposure of CEM or CEM-SS cells to \(10^5\) RNA copies results in 1 to 10 infection events (compare Fig. 1B and C). Thus, infection with \(10^{10}\) RNA copies of input virus would result in up to \(~1\times10^6\) infection events, and a large proportion of the \(~1\times10^6\) CEM-SS cells would be infected with input virus; consequently, most of the cells would be infected after a small increase in virus production, and superinfection interference would be established, suppressing any additional increase in virus production.

An increase in virus producer cells does not account for the increase in virus production. To determine whether an increase in virus producer cells could have contributed to the modest (\(~14\)-fold) increase in virus production observed in CEM cells (Fig. 1H), we repeated the experiment described for Fig. 1 and quantified the virus producer cells every 2 days. The fold increase in virus production was determined by dividing the average RNA copies present on days 13 to 15 with the average RNA copies present on days 1 to 3 (Fig. 2A). The results were similar to those observed in Fig. 1H; the increase
in viral production ranged from ~6- to ~3,936-fold in CEM-SS cells and up to 34-fold in CEM cells. In this experiment, the increase in viral load in CEM-SS cells after infection with 10^5 RNA copies was 77-fold, compared to 1,050-fold in Fig. 1H. Since <10 cells were likely to be infected after infection with 10^5 RNA copies, the variability in viral production may be due to stochastic events that led to greater variation in the number of initially infected cells or loss of virus production from some of the initially infected cells.

To determine the extent to which the number of virus-producing cells affected virus production, the increases in virus production were adjusted for the numbers of cells in culture (Fig. 2B). The increase in virus production in CEM and CEM-SS cells was generally reduced by 2- to 4-fold after adjusting for the number of virus-producing cells, but the CEM cells still exhibited modest increases in virus production (up to 10-fold). Therefore, despite the presence of A3G and A3F, XMRV replication was modestly increased in CEM cells infected with 10^6, 10^7, and 10^8 RNA copies. Proviral copy numbers per cell did not increase in PBMCs from day 1 to day 15, indicating that there was little or no virus replication and spread. To quantify viral replication and spread, we compared the average RNA copy numbers from days 1 to 3 and days 13 to 15 (Fig. 3C). Viral RNA copy numbers did not increase, since the ratios of the viral RNA copy numbers on days 13 to 15 to days 1 to 3 were less than 1 (0.14 to 0.5), indicating that XMRV replication and spread in activated human PBMCs is severely restricted.

We also compared the proviral copy numbers per cell in PBMCs infected with 10^10 XMRV RNA copies from day 1 and day 15 (Fig. 3D). The proviral copy numbers per cell in the three donors on day 1 averaged ~0.07, compared to ~2 proviruses per CEM cell after infection with 10^10 XMRV RNA copies; since 10-fold more PBMCs (1 × 10^7) were infected compared to CEM cells (1 × 10^6), infection of PBMCs was ~3-fold less efficient (~2 × 10^4 proviruses in CEM cells versus ~7 × 10^4 proviruses in PBMCs). The proviral copy numbers per cell did not increase in the PBMCs from day 1 to day 15 and actually decreased from 0.07 to 0.03 proviral copies/cell, but the decrease was not statistically significant (P = 0.15; t-test); these results indicated that XMRV replication and spread is severely restricted in PBMCs. Viral production from the PBMCs was approximately 10-fold lower than in CEM cells, which is in general agreement with the ~3-fold reduction in infection efficiency.

Murine leukemia viruses require cell division to integrate into the target cell chromosomes and complete viral replication (25, 41). To determine whether XMRV replication was restricted in PBMCs because of inefficient cell division, we infected PBMCs on days 1, 5, and 9 after activation and monitored virus production (Fig. 3E). Similar virus production was observed after infection on all 3 days, indicating that equivalent numbers of cells on days 1, 5, and 9 were susceptible to XMRV infection. Thus, the severe restriction to virus replication and spread in the PBMCs was not because of the absence of cells susceptible to XMRV infection.

**XMRV replication is severely restricted in PHA-activated human PBMCs.** Since XMRV replication and spread was restricted in A3G/A3F-expressing CEM cells, we sought to determine the extent to which XMRV can replicate and spread in A3G/A3F-expressing human PBMCs. We first compared the steady-state levels of A3G and A3F proteins in PHA-activated PBMCs obtained from three different donors to the levels present in human T cell line and prostate cancer cell lines (Fig. 3A). As expected, A3G and A3F expression was clearly detected in CEM and H9 cells but not CEM-SS cells, and A3G was not detectable in prostate cancer cell lines 22Rv1, LNCaP, and DU145. Compared to CEM cells, A3F expression was similar in H9 cells and substantially lower in LNCaP, 22Rv1, and DU145 cells. A3G and A3F were expressed in PBMCs from all three donors at levels similar to or higher than those in CEM cells.

To determine the extent of XMRV replication and spread in A3G/A3F-expressing PBMCs, 22Rv1 culture supernatants containing 10^5, 10^6, 10^7, and 10^10 XMRV RNA copies were used to infect 1 × 10^7 PBMCs from donor 2 (Fig. 3B). Consistent with the results obtained with CEM cells, no virus was detected in PBMCs infected with culture supernatants containing 10^7 XMRV RNA copies, suggesting that this amount of virus is not sufficient to establish a productive infection in PBMCs. In culture supernatants of PBMCs infected with 10^6 XMRV RNA copies, a low level of virus was detected, which became undetectable after 7 days. In culture supernatants of PBMCs infected with 10^7, 10^8, and 10^10 XMRV RNA copies, detectable levels of virus were present early in infection, which ranged from 2 × 10^5 to 2 × 10^7 RNA copies per 8.4 μl of culture supernatant. Importantly, we did not observe any increase in viral RNA copies in the infected PBMCs during the next 15 days, indicating that there was little or no virus replication and spread. We also compared the proviral copy numbers per cell in PBMCs infected with 10^10 XMRV RNA copies from day 1 and day 15 (Fig. 3D). The proviral copy numbers per cell in the three donors on day 1 averaged ~0.07, compared to ~2 proviruses per CEM cell after infection with 10^10 XMRV RNA copies; since 10-fold more PBMCs (1 × 10^7) were infected compared to CEM cells (1 × 10^6), infection of PBMCs was ~3-fold less efficient (~2 × 10^4 proviruses in CEM cells versus ~7 × 10^4 proviruses in PBMCs). The proviral copy numbers per cell did not increase in the PBMCs from day 1 to day 15 and actually decreased from 0.07 to 0.03 proviral copies/cell, but the decrease was not statistically significant (P = 0.15; t-test); these results indicated that XMRV replication and spread is severely restricted in PBMCs. Viral production from the PBMCs was approximately 10-fold lower than in CEM cells, which is in general agreement with the ~3-fold reduction in infection efficiency.

**Hypermutation of XMRV proviruses in PHA-activated PBMCs.** To analyze the effects of A3G/A3F expression on XMRV replication in PBMCs, we harvested XMRV-infected cells at day 15, isolated total cellular DNA, PCR amplified and cloned a 1.2-kb fragment from the pol region of XMRV proviral DNA, and sequenced the cloned PCR products (Fig. 4).
Proviral sequences were recovered from PBMC cultures infected with 10^{10} RNA copies from the three different donors described for Fig. 3; of the 38, 40, and 46 sequences obtained at day 15 from donors 1, 2, and 3, respectively, 2, 6, and 18 (2/39 to 39%) sequences were hypermutated. Furthermore, most of the G-to-A substitutions occurred in the GG dinucleotide context, which is characteristic of A3G-mediated hypermutation. The detection of these hypermutated proviruses in PBMCs indicated that some of the PBMCs were infected with virus that was produced from A3G-expressing cells. A few of the sequences also had multiple G-to-A mutations in the GA dinucleotide context, which is characteristic of A3F-mediated hypermutation. These results indicated that XMRV produced from the PBMCs packaged A3G or A3F into the virions and reinfected the PBMCs, resulting in G-to-A hypermutation in the context of GG or GA dinucleotides, respectively.

To determine whether infectious XMRV can be recovered from infected PBMCs or from other human cells, we developed a sensitive assay for recovery and detection of replication-competent XMRV. The strategy for XMRV detection and the construction of the DIG cells is illustrated in Fig. 5A. D17 canine osteosarcoma cells were transfected with the previously described MLV-based plasmid pMS2-FP-GF-no3R. This vector contains modified LTRs, in which partially redundant portions of the *gfp* gene are located...
in each LTR such that the gfp gene is reconstituted upon reverse transcription. The plasmid includes the MLV packaging signal (Ψ), the hygromycin phosphotransferase B gene (hygro), which is expressed from the SV40 promoter, and the FP portion derived from the 3' end of gfp in the left LTR and the GF portion derived from the 5' end of gfp in the right LTR. In addition, the vector contains all other cis-acting sequences required for retroviral replication but lacks gag, pol, and env, and thus can only be packaged by a helper virus that has infected the same cell. XMRV and MLV packaging sequences (Ψ) share ~70% homology, and XMRV has previously been shown to package MLV-based vectors. During reverse transcription of the MS2 vector, the middle F portion of gfp, which is present in both LTRs and has replaced the R region of the MLV LTR, is used to carry out the minus-strand DNA transfer step of reverse transcription. This results in functional reconstitution of the gfp gene and expression of functional GFP. A D17 cell line that was constructed to stably express the MS2 vector was named the DIG cell line. When XMRV infects DIG cells, many of the newly assembled XMRV particles will package the DIG RNA instead of XMRV RNA, resulting in GFP-positive cells in subsequent rounds of infection.

The protocol used to demonstrate the sensitivity of XMRV detection using the DIG cells is outlined in Fig. 5B. Briefly, we infected the DIG cells with culture supernatants from 22Rv1 cells containing $6 \times 10^6$ to $6 \times 10^7$ RNA copies. After 5 h, the virus-containing medium was removed, and the cells were washed with fresh medium. The infected cells were further incubated for 12 to 16 h, at which point $3 \times 10^5$ LNCaP cells were added (day 1), to ensure that highly infectible cells were present in the culture. Other experiments have indicated that the D17 cells can be efficiently infected with XMRV (14) (data not shown). The infected cells were monitored every day for 5 days (days 2 to 6) by using high-content imaging (ImageXpress; Molecular Devices), and the percentage of GFP-positive cells was determined. A representative result of three independent experiments (Fig. 5C) shows that XMRV replication and spread were detectable in all cultures that were infected with XMRV.

Recovery of replication-competent XMRV from infected PBMCs. To detect replication-competent XMRV from infected PBMCs, we cocultured DIG cells with PBMCs that were infected with different amounts of XMRV and monitored GFP expression in the DIG cells after 14 and 25 days (Fig. 5D). Activated PBMCs ($6 \times 10^6$) from donor 2 were infected with different amounts of XMRV RNA copies, as described earlier (Fig. 3), and virus replication was monitored (Fig. 5E). Similar to donor 2 (Fig. 3B), no increase in virus production was observed after 8 days (Fig. 5E). In fact, the virus production slightly decreased over the course of the experiment, which corresponded with a decrease in the total number of live cells in the cultures. Virus production in the PBMCs infected with $10^5$ RNA copies decreased to undetectable levels 5 days after infection. At 7 days postinfection, $1 \times 10^5$ infected PBMCs were cocultured with $8 \times 10^5$ DIG cells for 25 days. After 14 days (data not shown) and 25 days, DIG cells were harvested and the percentage of GFP-positive cells was determined by FACS analysis (Fig. 5F). Additional studies are required to determine the minimum amount of time required for recovery of infectious virus using the DIG cells. PBMCs infected with $10^5$, $10^6$, and $10^7$ XMRV RNA copies generated XMRV that infected the DIG cells. Subsequent spread of the MLV reporter vector in the DIG cells increased the proportion of GFP-positive cells to 31 to 41%. However, infectious XMRV could not be recovered from PBMCs infected with $10^4$, $10^5$, or $10^6$ XMRV RNA copies.

**DISCUSSION**

Recently, we and others showed that XMRV replication is restricted by A3G, A3F, and A3B as well as human tetherin/BST2/CD317 in a single-cycle assay (5, 14, 37, 52). On the
other hand, Lombardi et al. reported detection and recovery of infectious XMRV from PBMCs of chronic fatigue syndrome patients by cocultivation with LNCaP cells (28). Here, we compared viral replication in human PBMCs and T cell lines and observed that XMRV replication and spread was severely restricted in A3G/A3F-positive PBMCs and CEM cells, but not in A3G/A3F-negative CEM-SS cells. In fact, XMRV replication was more potently restricted in the PBMCs than in the

FIG. 5. Characterization of DIG cells and isolation of replication-competent XMRV from infected PBMCs. (A) Structure of MLV-based vector pMS2. FP, the 3' 462-bp fragment of GFP; GF, the 5' 350-bp fragment of GFP; Y+, the extended MLV packaging signal; SV-hygro, hygromycin phosphotransferase B gene under the control of the SV40 promoter; ter, SV40 termination signal. GFP was reconstituted upon F region-mediated minus-strand strong-stop transfer during reverse transcription. (B) Overview of the experimental procedure used to test the DIG XMRV reporter cell line. D17 cells stably transfected with the pMS2 vector were infected with serial 10-fold dilutions of XMRV or no XMRV. LNCaP cells were added 1 day postinfection. The cells were imaged and scored for GFP daily beginning at 2 days postinfection. (C) DIG cells were infected with serial 10-fold dilutions of XMRV or no XMRV. LNCaP cells were added 1 day postinfection. The cells were analyzed by high-content imaging and scored for GFP daily beginning at 2 days postinfection. Spread of XMRV through coculture of DIG and LNCaP cells is represented by the percentage of cells that expressed GFP. RNA copy numbers of the XMRV stock used to infect the DIG cells were determined by quantitative real-time RT-PCR. A representative experiment of three independent experiments is shown. (D) Protocol for recovering replication-competent XMRV from infected PBMCs in DIG cells. PBMCs (6 x 10^6) were infected with different XMRV as described for Fig. 3A, and virus production was quantified by real-time RT-PCR. At 7 days postinfection, 1 x 10^5 PBMCs were cocultured with 8 x 10^5 DIG cells. Virus recovery was detected by the percentage of GFP-positive cells. (E) Virus production in PBMCs infected with different amounts of XMRV. At 7 days postinfection, 1 x 10^5 infected PBMCs were collected from each infection and added to 8 x 10^5 DIG cells in six-well dishes. (F) At 3 days postcocultivation, the cells were transferred to a 10-cm dish, and the percentages of GFP-positive cells were determined after 25 days using FACS analysis. The averages of two independent experiments using two different donors are shown. Error bars represent the standard deviations.
CEM cells, since virus production increased modestly in CEM cells (up to 14-fold) but did not increase in the PBMCs, suggesting that other mechanisms of restriction, such as tetherin-mediated inhibition of virus release, may contribute to more potent restriction of viral replication (14, 34, 57). Although A3B was shown to inhibit XMRV replication in single-cycle assays (14), two recent studies have shown that PHA-activated PBMCs express little or no A3B (23, 38). Therefore, it is unlikely that A3B is contributing to the restriction of XMRV replication in PBMCs. However, it is also possible that XMRV infection is less efficient in PBMCs than in CEM cells, and as a consequence, the inhibitory effects of A3G are more potent in PBMCs than in CEM cells.

Our observation that XMRV replication and spread is severely restricted in PHA-activated PBMCs is a novel finding that has not so far been reported. These results are also inconsistent with the report by Lombardi et al. (28), which showed that after activation with PHA and IL-2, nearly all of the PBMCs from XMRV-positive patients become reactive to a monoclonal antibody against the MLV p30 Gag protein; these studies suggested that XMRV replicated and spread efficiently in CFS patients and was present in most of the activated PBMCs (28). In future studies, it will be important to compare the replication and spread of XMRV in PBMCs isolated from normal control donors and CFS patients.

Despite the potent restriction, the PBMCs could be infected with XMRV, as recently shown (17), and the infected PBMCs could produce virus particles; this is consistent with the mechanism of A3G/A3F-mediated inhibition of viral replication, which does not block virus entry or production but greatly reduces the infectivity of the virions (47, 61). We also observed G-to-A hypermutation of XMRV proviruses in PBMCs, indicating that virus produced from the PBMCs could reinfect the cells and generate proviruses; since there was no detectable increase in virus production, most of these reinfection events probably resulted in abortive infection or the generation of highly mutated proviral genomes that were unable to produce viral particles. The extent of hypermutation was variable between different donors and very low for donor 1, suggesting that replication and spread of XMRV in PBMCs from some donors may be very inefficient. These results indicate that hypermutation of XMRV proviral genomes at GG dinucleotides, which indicate A3G substrate specificity, may serve as a useful marker for infection and replication of XMRV in human PBMCs. We recovered replication-competent XMRV from 1 × 10^5 infected human PBMCs that were initially infected with 10^{10}, 10^{9}, or 10^7 XMRV RNA copies by cocultivation with the DIG cells. We observed that there were approximately 0.07 XMRV proviral copies per cell in PBMCs infected with 10^{10} RNA copies; we estimate that of the 1 × 10^5 PBMCs that were cocultivated with the DIG cells, 7,000, 70, and 7 cells were initially infected upon infection with 10^{10}, 10^9, and 10^7 XMRV RNA copies, respectively. This estimate suggests that cocultivation with as few as ~7 PBMCs that were initially infected resulted in the rescue of replication-competent XMRV. Thus, the DIG cell line can be used for sensitive detection and isolation of XMRV or other gammaretroviruses that can serve as a helper virus for the MLV-based vector in the DIG cells.

Although XMRV replication and spread in PHA-activated PBMCs was severely restricted, PBMCs could serve as a reservoir of replication-competent XMRV and facilitate infection of A3G/A3F-deficient cells. A3G/A3F mRNA levels are similar in different T cell subsets (23, 38), but their expression levels were shown to be lower in monocytes; high levels of A3G mRNA expression have also been reported in lung, ovaries, spleen, and thymus (23, 38). However, as shown here and in previous studies (37, 52), A3G expression levels are very low in prostate cancer cells, suggesting that XMRV might replicate more efficiently in these cells. Although 22Rv1, LNCaP, and DU145 cells all expressed detectable levels of A3F, these levels were generally lower than those in CEM cells and did not appear to be sufficient to restrict XMRV replication. Although the XMRV promoter activity in lymphocytes has not been determined, we were able to readily detect virus production from the infected cells, indicating that XMRV RNA and proteins are expressed in the PBMCs. It has been reported that the XMRV promoter is responsive to androgen stimulation, and XMRV transcription is robust in LNCaP prostate tumor cells, suggesting that XMRV transcription may be higher in prostate tissues than in PBMCs (40).

We and others have shown that the anti-HIV-1 drugs zidovudine, tenofovir, and raltegravir are potent inhibitors of XMRV replication (20, 37, 43, 49, 51). Although there is no evidence to indicate that XMRV infection and replication contribute to the etiology of chronic fatigue syndrome or prostate cancer, clinical trials are being contemplated to determine the effectiveness of antiviral drugs for the treatment of these diseases. We believe it is necessary to have reliable and sensitive assays for XMRV detection and replication before the efficacy of any antiviral therapy can be assessed. Our results show that monitoring virus production from PBMCs is not a reliable indicator of virus replication and spread in PBMCs; moreover, the fact that we observed little or no replication in PBMCs implies that it may be very difficult to monitor any potential inhibitory effects of antiviral drugs by evaluation of XMRV levels in peripheral blood.

In summary, our results show that XMRV replication and spread is severely restricted in PBMCs, but these cells can serve as a reservoir for generation of infectious virus that can potentially spread to cells that express low levels of these restriction factors. Additionally, hypermutation of XMRV at GG dinucleotides may be a useful marker for detection of human PBMC infection.

ACKNOWLEDGMENTS

This research was supported in part by the Intramural Research Program of the NIH, National Cancer Institute, Center for Cancer Research.

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