Infectious DNA of the Human spumaretrovirus

Axel Rethwilm, Gerald Baunach, Kai-Olaf Netzer, Bernd Maurer, Bettina Borisch and Volker ter Meulen
Institut für Virologie and 1Institut für Pathologie, Universität Würzburg, Versbacher Strasse 7, 8700 Würzburg, FRG

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ABSTRACT
An infectious molecular clone (pHSRV) of the human Spumaretrovirus (HSRV) was constructed using viral DNA and cDNA clones. The infectivity of pHSRV was proven by transfection of cell cultures and subsequent infection of susceptible cultures with cell free transfection derived virus. pHSRV derived virus produced foamy virus typical cytopathic effects in susceptible cultures. Infected cells could be stained specifically with foamy virus antisera by means of indirect immunofluorescence. Radioimmuno-precipitation revealed the presence of characteristic HSRV structural proteins in pHSRV infected cultures. By cotransfection of pHSRV and an indicator plasmid it was found that pHSRV is able to transactivate the viral LTR. Viral transcripts were found to be approximately 200 bases longer in pHSRV infected cultures compared to wildtype infected cultures. This difference is most likely due to an insertion of DNA of non-viral origin in the U3 region of the 3'LTR of the infectious clone.

INTRODUCTION
Retroviruses have been subdivided into three major groups: oncoviruses, lentiviruses and spumaviruses (1). While oncoviruses and more recently lentiviruses have been characterized intensively there is only little knowledge on the biology of spumaviruses. Spumaviruses are widespread in wild non-human primates, in bovines, and felines, and more recently lentiviruses have been associated with retroviruses (11, 12). Moreover, it was shown recently that transcription from the HSRV LTR is markedly enhanced in virus infected cells (15). To conduct more detailed studies on the molecular biology of HSRV we have constructed an infectious molecular clone. This plasmid will be invaluable in investigating the proposed regulatory genes of HSRV.

MATERIALS AND METHODS
Cells and virus
Baby hamster kidney cells (BHK-21) were obtained from Dr. D. Neumann-Haefelin (Freiburg, FRG), human glioblastoma cells U-251 MG from Dr. D.D. Bigner (Durham, USA), human hepatoma cells HEP G2 from Dr. H. Will (Munich, FRG), VeroA and primary human embryonic lung cells (HEL) from Dr. A. Pohl-Koppe (Würzburg, FRG). Cells were grown in modified Eagles medium supplemented with 5% fetal calf serum, glutamine and antibiotics. Wildtype HSRV infected HEL cells were a gift of Dr. R.M. Flügel (Heidelberg, FRG). Virus was first passaged on HEL cells and then adapted to different cell cultures by cocultivating recipient cells with a freeze/thaw lysate of infected HEL cells. For transmission of cell free virus supernatant from infected cultures was cleared by low speed centrifugation and passed through a 0.2 µm filter (Millipore). Infection was carried out by incubating the cells with virus containing supernatant for one hour at 37°C, washing once with medium and feeding with fresh medium. Titration of virus was done as described previously (16).

Plasmids and DNA constructions
The infectious clone pHSRV was constructed from viral DNA clones pHSRV-B-C11 (C11) and pHSRV-B52 (B52) and cDNA clone pHSRV-H-C55 (C55), which together represent the whole viral genome. Molecular cloning and nucleotide sequence analysis of the clones have already been described (11,12,17). Cloning procedures were according to established methods (18), using restriction enzymes and DNA modifying enzymes from Boehringer-Mannheim and New England Biolabs.
Oligonucleotide linkers were from New England Biolabs. All plasmid constructions were verified by restriction enzyme analysis. Molecular cloning of pHSRVcat which bears the bacterial chloramphenicol acetyltransferase gene has been described recently (15).

Transfection and CAT assay
DNA was transfected into adherent cells as a 1:2 mix with 15% glycerol in transfection buffer for 1-2 min. Assay (Biorad), various amounts of protein as determined by a commercial protein assay (Bio-Rad) was transfected into adherent cells as a 1:2 mix with 15% glycerol in transfection buffer for 1-2 min. After overnight incubation at 37°C, cultures were harvested 48 hrs after transfection and processed as described (20). CAT assays were performed with various amounts of protein as determined by a commercial protein assay (Biorad), 0.1 μCi 3H-chloramphenicol (Amersham), and 1mM acetyl-CoA (Sigma) in 0.25M TRIS, pH 8.0. Assays were incubated for 60 min. at 37°C.

RNA extraction and Northern-blot hybridization
RNA was extracted by the guanidinium-isothiocyanate method followed by centrifugation through a CsCl cushion (21). Poly A+ RNA was selected on oligo (dT) cellulose (Boehringer-Mannheim). RNAs were run on formaldehyde containing 1% SDS and t-RNA at 48°C. Hybridization was carried out in the same buffer containing DNA probe and reduced concentrations of sodium phosphate (0.02M) and t-RNA (0.05g/l) at 48°C. Filters were washed 3 x 30min. in 0.5 x SSC, 0.1% SDS at 75°C, dried and exposed to DuPont Cronex films overnight at 70°C using an intensifying screen. Probes were labeled with 32P-dCTP (Amersham) as described (22) to a specific activity of 5 x 108 cpm/μg.

Table 1: Appearance of foamy virus c.p.e. in cultures infected with wildtype virus or transfected with plasmid DNA

<table>
<thead>
<tr>
<th>Viral stock</th>
<th>BHK-21</th>
<th>U251-MG</th>
<th>VeroA</th>
</tr>
</thead>
<tbody>
<tr>
<td>HSRVwt</td>
<td>d 2 +</td>
<td>d 1 +</td>
<td>d 2 +</td>
</tr>
<tr>
<td>pHSRV</td>
<td>d 4 +</td>
<td>d 8 +</td>
<td>d 10 +</td>
</tr>
<tr>
<td>pSLG/pEB3L</td>
<td>d 6 +</td>
<td>d14 +</td>
<td>d20 +</td>
</tr>
<tr>
<td>pSLGE</td>
<td>n.d.</td>
<td>d21</td>
<td>d14</td>
</tr>
<tr>
<td>pBR322</td>
<td>d15</td>
<td>d21</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

Cells were infected with cell free wildtype virus or were transfected as indicated and observed for multinucleated giant cell c.p.e. Given are the days after infection/transfection with minimal c.p.e. (+ < 10% multinucleated cells), medium c.p.e. ( + + = 10-50% multinucleated cells), and maximal c.p.e. ( + + + 50-100% multinucleated cells), = no c.p.e. visible, n.d. not done.

Indirect immunofluorescence (IFA) and radioimmuno-precipitation (RIPA)
Cells were grown on coverslips, were fixed in cold acetone and IFA was performed as described (23), using HSRV positive and negative human sera at a 1:10 dilution and FITC conjugated anti human IgG (Dakopatts). RIPA was essentially done as described recently (24). Briefly, cultures were metabolically labeled with 35S-methionine for 16hrs at 37°C. Cells were lysed in RIPA detergent buffer (0.02M TRIS (pH7.4), 0.3M NaCl, 1% Triton X-100, 1% deoxycholate, 0.1% SDS, 0.001M phenylmethyl-sulfonylfluoride), and lysates were cleared by brief ultracentrifugation. 50μl of lysate were reacted with 2.5μl of serum at 4°C for 60min. Immunocomplexes were precipitated with protein-A-sepharose beads (Pharmacia), washed with RIPA detergent buffer and resolved on 7.5% polyacrylamid-SDS gels (25). Gels were fixed, washed in fluorographic reagent (Amersham), dried and exposed to X-ray films at -70°C.

Figure 1: Genome organization of HSRV and cloning strategy of pHSRV (modified from (11) and (12). A (Asp718), Aa (Aattl), B (BamHI), E (EcoRI), H (HindIII), N (Ncol), Ns (NsiI), Pv (PvuII), X (XbaI).
Figure 2: (a) Multinucleated giant cell c.p.e. in U251-MG glioblastoma cells infected with pHSRV derived cell free virus. (b) Indirect immunofluorescence of pHSRV infected hepatoma cell line Hep G2 using a human α HSRV serum.

RESULTS

Construction of pHSRV

The infectious plasmid pHSRV was constructed using the three viral clones C11, C55 and B52. Plasmid C11 harbours a 15.4 kb BamH1 insert where viral genes are organized in the order: 5'-Δbell-env-S1-Δpol-5'LTR-gag-pol-S1-env-Δbell-3'. Viral genes in the 5.5kb HindIII insert of C55 appear in the order: 5'-Δpol-S1-env-bell1-bell2-Δbell3-3', and in the 2.2 kb insert of B52 in the order: 5'-Δbell1-bell2-bell3-3'LTR-3'. Thus, the whole viral genome of HSRV is represented by these molecular clones. The strategy for generation of pHSRV is outlined in Fig.1. In general, two overlapping viral clones were constructed by stepwise addition of viral DNA fragments using enzymes that cut only once or twice in the portion to be cloned. Following this principle, Asp718 linkers were added to a 3.0
Figure 3: Northern-blot of poly A + RNA from wildtype infected (a), uninfected (b), and BHK21 cells infected with pHSRV derived cell free virus (c). RNAs were hybridized to 32P-labeled pHSRV DNA.

Figure 4: SDS-gel electrophoresis of protein precipitated from wildtype (a,b) and pHSRV (c,d) infected BHK21 cells using HSRV negative (a,c) and HSRV positive (b,d) human serum.

kb fragment of C11, running from a PvuII site in the flanking sequence 43bp 5' to the 5' LTR to a PvuII site in the gag ORF, and the fragment was cloned into pUC19 vector, leading to p5LG. A 3.6 kb NsiI/EcoRI fragment from C11 was inserted into NsiI/EcoRI cut p5LG, leading to p5LGP. Plasmid p5LGP was derived by inserting a 3.5 kb EcoRI fragment of C55 into EcoRI cut p5LG. The 3' third of the viral genome was reconstituted by (i) modifying a single BamHI site of B52 and a single XbaI site of C55, respectively, into Asp718 sites and (ii) inserting a 1.7 kb Asp718/NcoI fragment of the latter into the respective sites of the first, leading to pEB3L. Plasmids p5LGP and pEB3L overlap by 1.0 kb and have a single AatII site in the overlap. Plasmid pHSRV was constructed by cloning the 9.3 kb Asp718/AatII fragment of p5LGP into Asp718/AatII cut pEB3L. Restriction enzyme mapping revealed all restriction enzyme sites used during the cloning procedure were unchanged in pHSRV (data not shown).

Transfection experiments
BHK-21, U-251 MG and Vero A cells were transfected with 20 μg of DNA. DNAs used for transfection were pHSRV (20 μg), p5LGP (20 μg), p5LGP/pEB3L (10 μg/10 μg), and pBR322 (20 μg). Parallel cultures were infected with cell free wildtype virus at a multiplicity of infection (m.o.i.) of 0.1. Cultures were passaged and examined for foamy virus typical cytopathic effect (c.p.e.) daily by light microscopy. The result of a typical experiment is shown in Tab.1. In wildtype infected cells multinucleated giant cell c.p.e. first appeared after one (U-251 MG) or two (BHK-21 and Vero A) days and was maximal within 7–9 days. In cultures transfected with pHSRV c.p.e. occurred 4–10 days post transfection and in cultures transfected with the two overlapping plasmids p5LGP and pEB3L 6–14 days after transfection. Once c.p.e. had become visible in the transfected cultures there was no delay in the progression of infection until c.p.e. reached maximal levels relative to wildtype infected cultures. No c.p.e. was observed in cells transfected with p5LGP only or with pBR322 even after 3 weeks of cultivation. All following experiments were carried out with cell free transfection derived virus. As shown in Fig.2a cells infected with
Acetylation of chloramphenicol could clearly be observed in assays cotransfected with pHSRV (lane b: different molecular weights we performed radioimmuno-
acetylation at p. g.

analysis of viral proteins

Poly A+ RNA from pHSRV and wild type infected cultures was run on denaturing gels, blotted and hybridized to 32P-labeled pHSRV DNA. As shown in Fig.3 three bands were detected in both RNA preparations (compare lanes a and c). In lane a the smallest band corresponds to RNA of ca. 2.5 kb, the middle band to RNA of ca. 6.3 kb and the largest band to ca. 11.2 kb RNA. While the pattern of HSRV transcripts was clearly the same in pHSRV and wildtype infected cultures, the pHSRV infected cultures yield transcripts slightly longer (ca. 0.2 kb) than the wildtype infected cultures.

analysis of viral proteins

To test whether differences found in viral transcript lengths between pHSRV and wildtype virus resulted in viral proteins of different molecular weights we performed radioimmuno-
precipitation assays. Proteins from lysates of infected BHK-21 cultures were precipitated with HSRV positive human sera detecting the main structural viral proteins. The precipitates were analysed on SDS polyacrylamide gels. As shown in Fig.4 the relative mobility of precipitated proteins were the same in pHSRV and wildtype infected cultures (compare lanes b and d). Proteins identified in both lysates were the env related glycoproteins gp170, gp130 and gp47 (24). Further virus specific but so far uncharacterized proteins detected in both preparations were of the apparent molecular weight of 80, 70, and 55 kD.

transactivation assay

Since transcription of a reporter gene under control of the HSRV LTR is markedly enhanced in HSRV infected cells compared to uninfected cells (15), we tested whether cotransfection of pHSRV and an indicator plasmid leads to transactivation. As an indicator plasmid we used pHSRVcat, a plasmid harbouring the 5'LTR to the cat gene. Fig.5 shows that pHSRVcat is silent upon cotransfection with pBR322 DNA assaying 250 µg of protein from prepared lysates (lane a). In contrast, acetylation of chloramphenicol could clearly be observed in assays with increasing protein amounts of lysates prepared from cultures cotransfected with pHSRV (lane b: 25 µg, lane c: 50 µg, lane d: 75 µg, lane e: 100 µg). As positive control 100 µg protein of a lysate from a culture transfected with pSV2CAT were run (lane f).

discussion

Here we have described the construction and characterization of an infectious molecular clone of the human spumaretovirus. Transfection of susceptible cultures with the construct pHSRV leads to cell free infectious viral particles as does transfection with the two overlapping plasmids pSLGPE and pEB3L. pHSRV derived virus was found to be identical to wildtype virus with respect to c.p.e. formation, apparent molecular weight of viral proteins and transactivation of the viral LTR. The only differences found were in the lengths of viral transcripts. In both RNA preparations three predominant viral transcripts were found, corresponding most likely to genomic RNA, spliced env message and probably double spliced messages for the regulatory proteins as described for related retroviruses (26,27). pHSRV messages were ca. 0.2 kb longer than wildtype messages. Interestingly, 5'LTR and 3'LTR of pHSRV differ by a stretch of 15B nucleotides flanked by a 4 bp repeat that is present in the U3 region (at nucleotide position 11336 of pHSRV) of the 3'LTR (derived from clone B52) but missing in the 5'LTR (derived from clone C11) (11,12). Since all retroviral transcripts terminate at the R-U5 border of the 3'LTR one would expect, that the 15B nucleotide stretch is present in all pHSRV derived viral transcripts. It has been speculated that the 15B bp stretch is an insertion of non-viral origin in the 3' LTR of clone B52 (12). The ca. 0.2 kb smaller wildtype transcripts found in the present study strongly support this view. Whether this insertion will influence the biological properties of the virus remains to be elucidated. Obviously, it does not interef with viral replication.

HSRV shares some striking features with the pathogenic human retroviruses especially of the lentiviral subgroup (13). Like HIV HSRV uses RNA-lys as primer for first strand cDNA synthesis (12), has an extraordinary long ORF for heavily glycosylated env proteins (12,24), and encodes for several additional gene products to gag, pol and env (11,12,13). One of which probably represents a viral transactivator (15). The availability of an infectious molecular clone for HSRV will allow to characterize the function of these genes in relation to the function of the regulatory genes of HIV and HTLV.

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references

8. Muller, H.K., Ball, G., Epstein, M.A., Achong, B.G., Lensor, G. and Levin,