Dominance of highly divergent feline leukemia virus A progeny variants in a cat with recurrent viremia and fatal lymphoma

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Abstract

Background: In a cat that had ostensibly recovered from feline leukemia virus (FeLV) infection, we observed the reappearance of the virus and the development of fatal lymphoma 8.5 years after the initial experimental exposure to FeLV-A/Glasgow-1. The goals of the present study were to investigate this FeLV reoccurrence and molecularly characterize the progeny viruses.

Results: The FeLV reoccurrence was detected by the presence of FeLV antigen and RNA in the blood and saliva. The cat was feline immunodeficiency virus positive and showed CD4⁺ T-cell depletion, severe leukopenia, anemia and a multicentric monoclonal B-cell lymphoma. FeLV-A, but not -B or -C, was detectable. Sequencing of the envelope gene revealed three FeLV variants that were highly divergent from the virus that was originally inoculated (89-91% identity to FeLV-A/Glasgow-1). In the long terminal repeat 31 point mutations, some previously described in cats with lymphomas, were detected. The FeLV variant tissue provirus and viral RNA loads were significantly higher than the FeLV-A/Glasgow-1 loads. Moreover, the variant loads were significantly higher in lymphoma positive compared to lymphoma negative tissues. An increase in the variant provirus blood load was observed at the time of FeLV reoccurrence.

Conclusions: Our results demonstrate that ostensibly recovered FeLV provirus-positive cats may act as a source of infection following FeLV reactivation. The virus variants that had largely replaced the inoculation strain had unusually heavily mutated envelopes. The mutations may have led to increased viral fitness and/or changed the mutagenic characteristics of the virus.
The development of sensitive and specific real-time TaqMan polymerase chain reaction (PCR) assays [13-15] led to the reassessment of FeLV infection outcomes [14,16-18]. In these studies, cats with progressive infection became persistently positive for the provirus and viral RNA and had high viral loads. Cats with regressive infection had lower provirus and viral RNA loads than cats with progressive infection. The provirus became undetectable over time only in a few cats with regressive infection [14].

We now report on a specific pathogen-free (SPF) cat that had been part of an early FeLV vaccination study, which was performed to test the first recombinant FeLV vaccine and to examine the influence of a preexisting FIV infection on the immune response and vaccine efficacy [19]. The cat had been infected with FIV prior to FeLV vaccination and exposure to FeLV-A/Glasgow-1. The cat developed transient FeLV viremia but was FeLV negative thereafter. After being healthy for 8.5 years, the FeLV antigen and viral RNA reappeared in the blood; FeLV was shed via the saliva; and the cat developed a multicentric FeLV-positive lymphoma. The goals of the present study were to investigate the recurrence of FeLV in this cat and to determine the molecular characteristics of the progeny viruses and their distribution in order to provide further knowledge on the molecular determinants of FeLV pathogenicity and to deepen our understanding of the host-retrovirus interaction.

Methods

Animal, virus exposures and long-term follow-up
A female SPF cat (cat #261; Ciba Geigy, Basel, Switzerland), which was kept under barrier conditions and housed in a group, was intraperitoneally infected with FIV at the age of 17 weeks (Fig. 1A), as described [19]. The cat was vaccinated with a recombinant FeLV p45 protein vaccine (Leucogen, Virbac, Nice, France) at the age of 41 weeks (Fig. 1B), and it was exposed intraperitoneally to FeLV-A/Glasgow-1 18 weeks later (Fig. 1B) [19]. At the age of four years (2.9 years post FeLV infection [p.i.]), the cat was revaccinated twice with the FeLV vaccine (Fig. 1B). The cat was observed for 8.5 years p.i., for a total observation period of 9.6 years. It was co-housed with FeLV p27-positive cats during the first seven years p.i., after which it was kept with p27-negative cats. The study was officially approved by the veterinary office of the Swiss Canton of Zurich (197/89, 43/90, 66/91, 131/91, 329/91, 56/95). Complete hemograms and, at selected time points, serum biochemistry analyses were performed. CD4+ and CD8+ cell subsets were determined by flow cytometry as described [20], starting twenty months after FIV infection at the age of two years.

Serological assays and virus isolation
ELISA was used to detect the levels of the FeLV p27 antigen [21] and antibodies to the FIV transmembrane protein [22], total FeLV, and FeLV p45 [19,23]. ELISA results were calculated as a percentage after normalization to the positive control, which was assayed on every plate. FeLV-neutralizing antibodies were measured by a focus-inhibition assay [19]. Virus isolation was performed for FIV using blood lymphocytes [19,23] and for FeLV using heparinized plasma [19]. To detect FeLV latency, bone marrow that was collected 24 weeks p.i. at the age of 1.6 years was cultured in the presence of hydrocortisone [19].

Necropsy
The cat underwent histopathological examination, and samples from 27 tissues (Table 1) were collected. Tissues for histology were fixed in 10% buffered formalin and processed by standard procedures. Samples for PCR analyses were snap-frozen in liquid nitrogen and stored at -70°C.

Immunohistology
FeLV proteins were detected in formalin-fixed paraffin-embedded (FFPE) tissue sections by an indirect immunoperoxidase assay (IPA) using antibodies directed against p27, gp70, and p15E as previously described [24,25]. Controls were established with a monoclonal antibody directed against an unrelated antigen. FFPE lymphoma-positive tissue sections (sternal and mesenteric lymph nodes, large intestine, spleen and liver) were tested to identify B and T cells using a CD3 T-cell marker (M7051, DAKO), CD20 (RB-90-13-P, Labvision, Thermo Fisher Scientific, Fremont, USA) and CD45R [26] (clone B220 [Ly5]; Linaris, Wertheim-Bettingen, Germany) together with the ChemMate detection kit (K5003, DAKO).

Nucleic acid extraction
DNA from 200 μL of saliva oruffy coat that was collected from EDTA-anticoagulated blood was extracted using the QiAamp Blood Mini Kit (Qiagen, Hombrechtikon, Switzerland). RNA from serum and saliva samples that were collected at the time of euthanasia was extracted using the viral RNA Mini Kit (Qiagen). Tissue samples were homogenized as described [27], and DNA was extracted using the QiAamp DNA Tissue Kit (Qiagen). RNA from tissues was purified using the ABI Prism 6700 Automated Nucleic Acid Workstation (Applied Biosystems, Rotkreuz, Switzerland) or the RNeasy Mini Kit (Qiagen). Nucleic acids were extracted from urine (200 μL) and feces (~5 mg) collected at the time of euthanasia as described [28,29]. Negative
Figure 1 Time course of FeLV infection in cat #261. A) FIV transmembrane (TM) specific antibody levels as determined by ELISA. B) Total anti-FeLV antibodies (black squares) and anti-FeLV p45 antibodies (black triangles) as determined by ELISA. C) Plasma FeLV p27 antigen levels as determined by ELISA. D) FeLV provirus loads as determined by real-time PCR. Two samples, indicated by asterisks, were positive for FeLV by non-quantitative PCR. Time points of FIV infection (at the age of 17 weeks), FeLV vaccinations (41 weeks, 4 years) and FeLV exposure with FeLV-A/Glasgow-1 (59 weeks) are indicated.
extraction controls consisting of phosphate buffered saline were included with each batch. In those experiments for which complementary DNA (cDNA) levels are given, the isolated RNA was reverse transcribed into cDNA using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems) prior to real-time PCR.

Total FeLV provirus and viral loads

Total FeLV provirus loads were quantified by TaqMan real-time PCR (U3 region) [15]. The number of provirus copies per cell was calculated using feline glyceraldehyde-3-phosphate dehydrogenase (GAPDH) copy numbers [30,31]. Ten blood samples that were collected from 7.3 to 8.5 years p.i. (at the age of 8.4 to 9.6 years) were available for quantitative analyses. Two samples that were collected at 4.5 and 6.4 years p.i. (at the age of 5.6 and 7.5 years) were analyzed by nested FeLV PCR [13]. Tissue viral loads were determined from cDNA using the U3 real-time PCR assay and were normalized to GAPDH and ribosomal protein S7 (RPS7) cDNA copy numbers [27]. Viral RNA loads in the serum and saliva were calculated as copies per mL.

FIV provirus and viral RNA loads

FIV provirus loads were determined by quantitative TaqMan real-time PCR [32]. FIV RNA loads were quantified using a protocol [15] and oligonucleotides previously described [32]. They were normalized according to GAPDH mRNA loads as determined by real-time PCR using a protocol [15] and previously described oligonucleotides [31]. For absolute quantification, standard RNA templates were prepared [33] from plasmids containing either FIV [34] or GAPDH [31] sequences. The standard RNA was quantified and aliquoted as described [35].

Table 1 Detection of lymphoma and FeLV in tissues collected upon necropsy from cat #261.

| Tissue                | Lymphoma (Histology) | In situ Hybridization | Immunohistology |
|-----------------------|----------------------|-----------------------|-----------------|
|                       | gp70 | p27 | gp70 | p27 | p15E |
| Salivary glands:      |      |     |      |     |      |
| - Mandibular          |      |     |      |     |      |
| - Parotid             |      |     |      |     |      |
| Duodenum              |      |     |      |     |      |
| - in GALT             | +/++ | +/++ |       |      |      |
| Jejunum               |      |     |      |     |      |
| - in GALT             | +/++ | +/++ |       |      |      |
| Ileum                 |      |     |      |     |      |
| - in GALT             | ++   | GALT |       |      |      |
| Colon                 |      |     |      |     |      |
| - in GALT             | +/++ | +/++ |       |      |      |
| Rectum                |      |     |      |     |      |
| Liver                 |      |     |      |     |      |
| - (mainly centro-acinous) | ++ | ++ |       |      |      |
| Spleen                |      |     |      |     |      |
| Thymus                |      |     |      |     |      |
| Tonsil                |      |     |      |     |      |
| Lymph nodes:          |      |     |      |     |      |
| - Sternal             |      |     |      |     |      |
| - Popliteal           |      |     |      |     |      |
| - Submandibular       |      |     |      |     |      |
| - Mesenteric          |      |     |      |     |      |
| Bone marrow           |      |     |      |     |      |
| Kidney                |      |     |      |     |      |
| Urinary bladder       |      |     |      |     |      |
| Brain                 |      |     |      |     |      |
| Spinal cord           |      |     |      |     |      |
| Ischiatic nerve       |      |     |      |     |      |
| Muscle upper hind leg |      |     |      |     |      |
| Lung                  |      |     |      |     |      |
| Myocardium            |      |     |      |     |      |
| Aorta                 |      |     |      |     |      |
| Diaphragm             |      |     |      |     |      |
| Thyroid and parathyroid glands |      |     |      |     |      |

Lymphoma diagnosed by macroscopic and histological examination and FeLV detected by in situ hybridization and immunohistology. GALT = gut associated lymphoid tissue; nt = not tested; - = negative reaction; + = positive reaction; ++ = strong positive reaction; +/- = positive or strong positive reaction depending on tissue section/cell type.
FeLV envelope gene specific real-time PCR assays

FeLV-A/Glasgow-1 envelope gene (env) was quantified by TaqMan real-time PCR assay as described [5]. In addition, primers and probes for an env variant-specific assay were designed using Primer Express software (version 3, Applied Biosystems; Table 2). The PCR reactions were performed as described [27] using 400 nM primers, and 200 nM of fluorogenic probe (Microsynth, Balgach, Switzerland).

Linearized plasmid DNA containing the appropriate envelope gene sequence (Table 2) was used as a standard template to test the specificity and sensitivity of the two env-specific real-time TaqMan assays and for absolute quantification. Copy numbers were determined spectrophotometrically (Nanodrop® ND-1000, Witec, Littau, Switzerland), and ten-fold serial dilutions were prepared as described [33]. The sensitivity of the system was determined by an endpoint dilution experiment [33]. The specificity was tested further with an endogenous FeLV sequence standard containing 10^8 copies/reaction [36] and with DNA from three SPF cats.

Detection of FeLV subgroups

FeLV-subgroups were investigated in the kidney, spleen, rectum, diaphragm, thymus, mandibular gland and myocardium by conventional PCR using the FeLV-A specific primers RB59 and RB17, the FeLV-B specific primers RB53 and RB17 and the FeLV-C specific primers RB58 and RB47 as described [37,38].

In situ hybridization

Digoxigenin-labeled RNA probes recognizing gp70 and p27 were used for in situ hybridization [39]. The probes were constructed from FeLV-A [GenBank: M18247] [40] using the primers listed in Table 2. The PCR products were cloned using the TOPO TA Cloning kit (Invitrogen BV, Groningen, The Netherlands). In vitro reverse transcription of the linearized plasmids and digoxigenin labeling was performed using the DIG RNA Labeling Kit (Roche Diagnostics GmbH, Mannheim, Germany). Positive strand RNA was used as a negative control. Hybridized digoxigenin was visualized with 150 U of anti-digoxigenin AP Fab fragments (Roche Diagnostics GmbH) and nitroblue tetrazolium chloride/5-bromo-4-chloro-3-indolyl phosphate.

Sequencing of the env and long terminal repeat (LTR) regions of FeLV progeny viruses

For the analysis of the full-length FeLV env sequences, DNA from the kidney and spleen was amplified as described [29] using env variant primers yielding a 2664 bp product (Table 2). PCR products were either sequenced directly (Synergene Biotech GmbH, Schlieren, Switzerland) or after TOPO TA cloning. Three env variants were identified: KI261-I from the kidney (direct sequencing), KI261-II from the kidney (direct sequencing) and spleen (2 clones) and SP261-III from the spleen (19 clones).

Full-length U3 regions of the 3’ LTR were amplified using the forward primer CAA TAC GAT CCG GAC CGA CCA TG and the reverse primer CGG GGC GGT CAA GTC TCG GCA AAG (adapted from [41]). PCR products (446 bp) were cloned as above. A total of 18 FeLV LTR clones, including 7 from the kidney, 3 from the bone marrow, 3 from the liver and 5 from the spleen, were sequenced (Microsynth).

| Table 2 Oligonucleotides used in this study |
|------------------------------------------|
| **Assay/Application** | **Oligonucleotide** | **Sequence** | **Amplicon size (bp)** | **Nucleotide position (bp)** |
|----------------------|---------------------|--------------|------------------------|----------------------------|
| TaqMan® PCR assay    |                     |              |                        |                            |
| env variant^1        | Forward             | GAT CCG GAC CGA CCA TAA TTA A | 105 | 1,912 - 1,933 |
|                      | Probe               | TGT ATG ATT CCA TTT AGT CCC^6 | 1,935 - 1,955 |
|                      | Reverse             | ACA CCA CTG CAG TAG CTG GCT AA | 2,017 - 1,995 |
| Production of standard | Forward             | TGG GCC CAA AGG GAA CAC AT | 598 | 456 - 475 |
|                      | Reverse             | GTT ACC TAA GAT TGC AAT CCC TTC G | 1,054 - 1,030 |
| env variant^3        | Forward^5           | CCT ATG GCT CAC TTC TTG GAT ACT GAT ATC TCT A | 2,664^7 | 5,617 - 5,650 |
|                      | Reverse             | TTA TAG CAG AAA GGG CCC G | 8,281 - 8,263 |
| In situ hybridization^4 | p27 Forward         | TAC GCC TTT ATC GCC AGT TG | 342 | 1,840 - 1,859 |
|                      | Reverse             | ATC TTT CCT CCC TTG CCT CTG G | 2,181 - 2,160 |
|                      | gp70 Forward        | AGG GAT TGC AAT CTT AGG TA | 219 | 6,952 - 6,971 |
|                      | Reverse             | TTA CAG GCC CAA TAG GTG | 7,170 - 7,153 |

^1 based on FeLV SP261-III [GenBank: EU359305]; ^2 based on FeLV-A/Glasgow-1 [GenBank: M12500]; ^3 based on FeLV-A Rickard strain [GenBank: AF052723]; ^4 based on FeLV-FADS [GenBank: M18247]; ^5 5’ FAM/3’ TAMRA; ^6 the PCR product included the first 300 nucleotides of the LTR in addition to env

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Phylogenetic analyses

Phylogenetic and molecular evolutionary analyses were conducted using MEGA version 4 [42]. The FeLV surface unit (SU) and the LTR sequences were aligned using CLUSTAL W [43]. For SU sequences, bootstrap support (1,000 replicates) was calculated by the neighbor-joining (NJ), minimum evolution (ME) and maximum parsimony (MP) methods, and results > 70% were considered to be significant [44]. MP trees were obtained using the Close-Neighbor-Interchange algorithm [45] with search level three [44], in which initial trees were obtained by the random addition of sequences (10 replicates). All positions containing gaps and missing data were eliminated from the dataset (complete deletion option).

Assessment of clonality in the lymphoma

To assess the clonality in the lymphoma from cat #261, the variable region genes of the feline immunoglobulin heavy chain (IGHV) and the T-cell receptor gamma (TCRG) were analyzed using PCR for antigen receptor gene rearrangements (PARR). To this end, DNA extracted from snap-frozen tissues was evaluated as described [46,47]. In addition, FFPE samples were analyzed without preceding DNA extraction: single 10 μm sections were treated with 20 mg/mL of proteinase K in 150 μL 1x Phusion HF Reaction Buffer (BioCat, Heidelberg, Germany) at 60°C overnight. After inactivation of the proteinase and centrifugation at 16,000 × g for 2 minutes, the fluid phase was used to confirm the DNA quality as described [48] with Phusion DNA Polymerase (BioCat). The samples were assayed by IGHV PARR analysis as described [48] with the following modifications: sense primers with annealing sites within the same framework region (1 and 3) were combined in one reaction (V1FR1 and V3FR1; V1FR3 and V3FR3).

Statistics

Statistical analyses were performed with GraphPad Prism for Windows (version 4.03, GraphPad software, San Diego, CA). Differences among three or more groups were analyzed by Kruskal-Wallis one-way ANOVA by Ranks (pKW) and Dunn’s post test. Differences between two groups were tested for significance using the Mann-Whitney U-test (pMWU). Differences were considered significant if p < 0.05. Observed agreement, expected agreement and Cohen’s kappa coefficient were calculated as described [49], with kappa values interpreted as suggested [50].

Nucleotide sequence accession numbers

The sequences described were submitted to GenBank: env [EU359303 to EU359305] and LTR [FJ613291 to FJ613296].

Results

Long-term follow-up

We analyzed blood samples collected from cat #261 over an observation period of 9.6 years as well as tissue samples collected at necropsy to investigate the FeLV recurrence and the development of virus variants in a long-term FIV-infected, FeLV provirus-positive, antigen-negative cat. The cat remained FIV-infected throughout the entire study (Fig. 1A), and all blood samples tested were FIV provirus-positive. Subsequent to the FeLV exposure at the age of 1.1 year, the cat developed transient FeLV viremia. FeLV virus was isolated from the blood in week 2 p.i., and the cat was p27-positive in week 3 p.i. (Fig. 1C). Transiently decreased white blood cell (WBC) counts were observed in weeks 1 and 4 p.i. (Fig. 2A), and the cat developed neutropenia (Fig. 2B). High neutralizing antibody titers were demonstrated by 12 weeks p.i., and total anti-FeLV antibody levels were persistently high (Fig. 1B). The virus could not be isolated from the bone marrow. Revaccination against FeLV at the age of four years yielded a marked boost in anti-p45 antibodies (Fig. 1B) and temporary increases in the WBC count and the number of neutrophils (Fig. 2A and 2B). There was no increase in the level of FeLV p27 (Fig. 1C). All tested blood samples were FeLV provirus-positive (Fig. 1D).

Development of disease

At 8.5 years p.i., at the age of 9.6 years, cat #261 became anorexic and lost weight, which was followed by dyspnea, dehydration, pale mucous membranes and a painful abdomen upon palpation. Severe non-regenerative anemia (packed cell volume: 12%; Fig. 2E; hemoglobin 3.8 g/dL) and severe leukopenia (100 cells/μL; Fig. 2A) were observed. Due to the low WBC count no differential was possible at this time. The last WBC differential performed three weeks prior to sacrifice revealed neutropenia (1,066 cells/μL Fig. 2B) and severe lymphopenia (182 cells/μL; Fig. 2C). In addition, CD4⁺ T cells were depleted (9 cells/μL; Fig. 2D). At the time of euthanasia, the animal had elevated levels of blood urea (37.6 mmol/L), creatinine (227 μmol/L), calcium (3.8 mmol/L), potassium (6.3 mmol/L) and phosphorus (2.9 mmol/L). The urine had a specific gravity of 1.016, and the protein-creatinine quotient (3.50) was elevated. Upon necropsy, lymphoma was detected in 18 out of 27 tissues by histological examination (Table 1, Additional file 1).

Reoccurrence of FeLV

Simultaneously with disease development, the cat became FeLV p27 antigen-positive (Fig. 1C; at the age of 9.6 years). A marked decrease in FeLV-specific antibodies was noted (Fig. 1B). FeLV RT-PCR analysis of
Figure 2 Time course of hematological parameters for cat #261. A) White blood cell counts. B) Neutrophil granulocyte counts. C) Lymphocyte counts. D) CD4⁺ T-cell counts. E) Packed cell volume. The reference ranges (5th to 95th percentiles) are indicated by the shaded areas (A to C, and E). No reference range was available for the absolute numbers of CD4⁺ T cells. In panel D, the dotted line indicates a CD4⁺ T-cell count of 200 CD4⁺ T cells/μL. No differential was possible at the time of sacrifice due to the low WBC number (100 cells/μL).
the serum was positive (mean load: $2.8 \times 10^4$ copies/mL serum), and a saliva sample was positive for FeLV p27 (11%; values above 4% are considered positive [28]) and FeLV viral RNA ($1.2 \times 10^7$ copies/mL saliva).

**Characterization of lymphoma**

A diffuse proliferation of small to predominantly medium sized lymphatic cells was observed. The nuclei were mostly round, and had a finely to coarsely stippled chromatin. One single central nucleolus or multiple randomly distributed nucleoli were seen (Additional file 1C). Between 10% (liver) and 100% (sternal lymph node) of the assessed tissues were affected. Based on the histological appearance and according to the WHO Histological Classification of Hematopoietic Tumors of Domestic Animals [51] a diffuse large B-cell lymphoma was diagnosed. The B-cell lineage was confirmed by positive staining in immunohistochemistry using the anti-CD20 and anti-CD45R B-cell antibodies (Additional file 1D). Anti-CD3 T-cell and anti-CD79 B-cell staining were negative. PARR analysis of the IGHV using FFPE tissues (liver, intestine, lymph node node) with the framework region 3-specific primers showed a reproducible intensive band in the expected size range (~180 bp) upon heteroduplex analysis; a second weak band was non-reproducible. These results were corroborated by the investigation of snap-frozen tissues from spleen for rearrangements in the IGHV; a reproducible clonal PCR product of the expected size (~140 bp) within a background polyclonal smear was detected. No rearrangement of the TCRG gene was observed. Accordingly, the lymphoma was categorized as a monoclonal proliferation of B cells.

**Presence of FeLV and FIV in the lymphoma**

Using IPA, FeLV p27-specific reactions were detected in 2 out of 25 tested samples, while FeLV p27 and gp70-specific gene sequences were found in 16 out of 25 samples by the more sensitive in situ hybridization (Table 1). When the results of the histological detection of lymphoma and FeLV in situ hybridization were compared, we observed an agreement of 96% (expected agreement 55%) and a Cohen’s kappa value of 0.91, indicating almost perfect agreement between these two analyses.

All 27 tissues examined were FeLV and FIV provirus-positive. FeLV provirus loads were significantly higher in tissues than in the blood ($p_{MWU} < 0.0001$), and they were significantly higher in tissues with lymphoma than in those without lymphoma (Fig. 3A). FIV provirus loads in tissues with lymphoma were higher when compared to the loads in healthy tissues, but they were not higher than the loads in the blood (Fig. 3B).

Out of the 27 tissues tested, 93% were positive for FeLV transcription, and 89% were positive for FIV transcription. Tissues with lymphoma had significantly higher FIV and FeLV viral loads than healthy tissues (Fig. 3C and 3D).

**Characterization of FeLV progeny viruses**

No FeLV-B (recombination with endogenous FeLV sequences) and FeLV-C subtypes were detected by conventional PCR. Three heavily mutated FeLV-A env variants were identified that showed 89-92% amino acid identity with each other and the highest, albeit modest, resemblance to FeLV-A/Glasgow-1 (for details see Table 3). Sequence variations were scattered throughout env and included point mutations in functional domains, such as the variable regions (VRA, VRB and VRC) and the proline rich region (PRR; [52]; for details see Additional file 2). All potential disulfide bonds were conserved in the three env variants compared to FeLV-A/Glasgow-1, as was the PHQ motif that is located in the N terminus of the receptor-binding domain and is critical for triggering virus fusion [52,53]. In addition, 11 out of 13 potential N-linked glycosylation sites were conserved when compared to FeLV-A/Glasgow-1, with two sites being lost. In addition, three new potential N-linked glycosylation sites were identified.

Phylogenetic analysis conducted by the MP, NJ and ME methods revealed the presence of a cluster of the three env progeny variants. In the trees based on nucleotide as well as protein sequences, the env variant sequences were most closely related to the original challenge strain FeLV-A/Glasgow-1 (Fig. 4). Sequencing of U3 led to nine groups of LTR sequences (Additional file 3). An overall U3 sequence conservation of 95-97% was found when the full-length progeny LTR sequences were compared to FeLV-A/Glasgow-1. Point mutations were found at 31 locations in the LTR sequence between the start of U3 and the TATA box. Several changes were found within the enhancer framework that comprises the binding sites for the transcription factors, the leukemia virus factor b (LVb), simian virus 40 core enhancer (CORE), nuclear factor 1 (NF1), glucocorticoid response element (GRE) and the FeLV-specific binding motif (FLV-1). There was one insertion and one transition found in the LVb binding site (Additional file 3). Four clones had a point mutation in the CORE. Mutations were found at two locations within the NF1 binding site, and one of these was detected in all of the clones. One mutation was located in the GRE binding motif, and two clones had a mutation in the FLV-1. Additional mutations were detected at 24 locations outside of these domains, with the majority of these mutations located upstream of the LVb site. No duplications of the enhancer or the upstream region...
**Figure 3** FeLV and FIV provirus and viral loads in blood and tissue samples from cat #261. FeLV and FIV provirus and viral loads in cat #261, quantified in the various tissues collected upon necropsy and in blood samples collected over the course of the last 14 months prior to sacrifice. Provirus and cDNA loads were determined using TaqMan real-time PCR and viral RNA loads were measured by TaqMan real-time RT-PCR. The tissues were classified according to the absence (n = 9) or presence (n = 18) of apparent lymphoma, as determined by histological examination. A) FeLV provirus loads. B) FIV provirus loads. C) FeLV viral (cDNA) loads in the tissues. D) FIV viral loads in the tissues. Viral tissue loads were normalized using GAPDH. Provirus loads (A and B) were tested for statistical differences by Kruskal-Wallis one-way ANOVA by Ranks (p_{KW} as indicated) and subsequently by Dunn's post test: * = p < 0.05; ** = p < 0.01; *** = p < 0.001. Viral loads (C and D) were tested for statistically significant differences using the Mann-Whitney U-test (p_{MWU} as indicated).

**Table 3** Sequence comparison of the env variants with prototype FeLV-A, -B, and -C

| Amino acid identity (%) | FeLV-A/Glasgow-1 [GenBank: M12500] | FeLV-B/Gardner-Arnstein [GenBank: K01209] | FeLV-C/Sarma [GenBank: M14331] |
|-------------------------|-------------------------------------|----------------------------------------|---------------------------------|
| env                     | 89-91                               | 76-77                                  | 85-86                           |
| SU                      | 86-89                               | 68-69                                  | 81-83                           |
| VRA                     | 68-74                               | 29-34                                  | 47-53                           |
| VRB                     | 48-69                               | N.A.                                   | 40-63                           |

N.A. = Not applicable (< 20%)
of the enhancer (URE) were detected in any of the clones that were investigated.

**Dominance of env variants**

Using standard DNA templates, real-time PCR assays for the variant and FeLV-A/Glasgow-1 env were shown to be specific for the respective sequences, without amplifying endogenous FeLV sequences. The detection limit of both assays was one copy/PCR, and the amplification efficiency was 99% for FeLV-A/Glasgow-1 and 98% for the variant.

Provirus of the env variants was identified in all 27 tissues, and the FeLV-A/Glasgow-1 env provirus was found in 26 out of 27 tissues (Fig. 5A; for details see Additional file 4A). With the exception of the duodenum, the provirus loads of the env variants were higher in every tissue than the FeLV-A/Glasgow-1 env provirus loads, and, when the results from all tissues were combined, a significant difference was found (p<0.0001, Fig. 5A). Remarkably, the provirus loads of the env variants were also significantly higher in tissues with

**Figure 4** Evolutionary relationship of the three SU variants found in cat #261. Phylogenetic trees were constructed by the MP method. Trees were drawn to scale, with the length being relative to the number of changes over the entire sequence. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) is shown next to the branches. A) Relationships at the DNA level. MP tree length: 765. The codon positions included were 1st + 2nd + 3rd + noncoding. There were a total of 1,363 base positions in the final dataset of which 349 were parsimony-informative. B) Relationships at the protein level. MP tree length: 329. There were a total of 453 amino acid positions in the final dataset of which 121 were parsimony-informative. GenBank accession numbers of the sequences included in the phylogenetic analyses are noted in square brackets following the virus identity. SP261-III, KI261-I and KI261-II (depicted in bold) were derived from cat #261.
lymphoma than in tissues without lymphoma ($p_{MWU} = 0.0168$, Fig. 5B), while no significant difference was found in provirus loads between these two groups for FeLV-A/Glasgow-1 (data not shown).

Concordant with these observations, the viral RNA loads of the $env$ variants were higher than the FeLV-A/Glasgow-1 viral RNA loads in all but three tissues, and overall the variant $env$ RNA levels were significantly higher than those of FeLV-A/Glasgow-1 ($p_{MWU} < 0.0001$, Fig. 5C and for details see Additional file 4B). Moreover, viral RNA levels of the $env$ variants were significantly higher in tissues with lymphoma than in healthy tissues ($p_{MWU} = 0.0019$, Fig. 5D), whereas no difference was detectable for the levels of FeLV-A/Glasgow-1 $env$ RNA (data not shown and Additional file 5D). These results were confirmed using the ribosomal protein S7 (RPS7) as an additional reference gene for the normalization of expression (Additional file 5). RPS7 was among the most stable reference genes tested previously [27].

When blood samples collected over the course of the last 14 months prior to sacrifice were analyzed, all samples tested provirus positive for the $env$ variant and FeLV-A/Glasgow-1. An increase in the $env$ variant provirus load (>100×) was observed at the time of FeLV reoccurrence, while the FeLV-A/Glasgow-1 provirus load remained low (Fig. 5E). No RNA samples from blood were available for detailed RT-PCR analyses. The FeLV-positive saliva sample collected at the time of sacrifice, tested viral RNA positive for FeLV-A/Glasgow-1 but negative for the $env$

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**Figure 5** FeLV-A/Glasgow-1 and $env$ variant loads in blood and tissue samples from cat #261. Loads of FeLV-A/Glasgow-1 and the $env$ variants of cat #261 quantified by real-time PCR in the tissues collected upon necropsy (A to D) and in blood samples collected over the course of the last 14 months prior to sacrifice (E). A) Provirus loads of FeLV-A/Glasgow-1 and $env$ variants in all tissues. B) $env$ variant provirus loads in tissues without (n = 9) and with (n = 18) apparent lymphoma. C) Viral (cDNA) FeLV-A/Glasgow-1 and $env$ variant loads in all tissues. D) Viral (cDNA) $env$ variant loads in tissues with and without apparent lymphoma. E) Time course of provirus loads of FeLV-A/Glasgow-1 and $env$ variants in the blood of cat #261. Viral loads (C and D) were normalized using GAPDH. Provirus and viral loads (A to D) were tested for statistical significance using the Mann-Whitney U-test ($p_{MWU}$ as indicated).
variants. The feces and urine collected at the same time point were positive for env variants.

Discussion

We demonstrate the reappearance of FeLV in an ostensibly recovered cat 8.5 years after the initial virus exposure. The SPF cat under investigation was kept in a controlled environment and was monitored closely throughout the entire observation period. It was exposed to a specified prototype FeLV-A strain, FeLV-A/Glasgow-1, after which a regressive infection with clearance of viremia was demonstrated. Latent FeLV infection, which is defined as replicating virus in bone marrow cell cultures supplemented with corticosteroids [8-10], was undetectable 24 weeks p.i. (at the age of 1.6 years); additional samples for testing were unavailable. All blood samples collected during the last 14 months before sacrifice were FeLV provirus-positive and the provirus loads in the peripheral blood were low, a characteristic of FeLV provirus-positive antigen-negative cats [13]. Only at the age of 9.6 years and concurrent with the reoccurrence of FeLV antigenemia, the blood FeLV provirus loads increased.

The cat was co-infected with FIV. This virus infection is frequently used as an animal model for retroviral diseases and in particular AIDS due to the common genus and the similarities with the human and simian immunodeficiency virus (HIV, SIV). The Swiss FIV isolate used in this study, FIV Z Ga, led to an AIDS-like syndrome characterized by complete CD4⁺ depletion, similar to what has been described for FIV Petaluma [2] and HIV. The breakdown of the immune system in cat #261, as a result of end-stage FeLV infection, and the resulting loss of the surveillance of FeLV replication was probably crucial for the reoccurrence of FeLV. Reactivation of FeLV infection upon experimental immune suppression in aviremic cats has been reported [9-11,17]. Some of these cats received high doses of corticosteroids for several weeks to provoke FeLV reactivation. The potential for reactivation seemed to be associated with the FeLV isolate [11], and the probability of reactivation decreased with increasing duration after FeLV exposure. Most cats were free of reactivable virus several months after clearance of FeLV viremia or after the removal from constant challenge [11,12]. In cat #261, the last potential exposure to FeLV was 18 months prior to FeLV reactivation. So far, in the absence of iatrogenic/experimental immune suppression, only a limited number of cases of reactivation of FeLV infection have been reported [7,11,14,17]. In some of these cats, natural stress was thought to be the initiator of reactivation [11]. To the best of our knowledge, this is the first documentation of FeLV reoccurrence in a cat with immunosuppression due to FIV infection.

Concurrent with the reappearance of FeLV, the FIV-infected cat had developed a fatal multicentric lymphoma. In general, lymphoma is a well-known manifestation of FeLV infection and is commonly associated with variant viruses [1,54-57]. In cat #261, the emergence of several distinct FeLV variants has been demonstrated. Moreover, lymphoma has also been described in FIV and FeLV co-infected cats [54,58]. As a result of the co-infection in cat #261, the role of FeLV and FIV in the tumorigenesis could not be easily assessed. Using specific real-time PCR assays, the FeLV progeny env variant loads were quantified relative to the load of the originally inoculated FeLV-A/Glasgow-1. At the time of euthanasia, the progeny viruses were more abundant than FeLV-A/Glasgow-1, thereby largely replacing the originally inoculated virus. Moreover, the env variant viruses were present in higher numbers in lymphoma-positive tissues than in healthy tissues. This was not the case for FeLV-A/Glasgow-1. These results may indicate that the FeLV progeny viruses found in cat #261 were relevant to the course of disease progression. Nonetheless, a strong association was also demonstrated between the presence of lymphoma and the provirus and viral RNA loads of FIV in the affected tissues.

The lymphoma was subsequently further characterized using immunohistology and PARR analysis and was, based on these results, categorized as a monoclonal proliferation of B cells. B-cell lymphomas have been reported to be the predominant form occurring during infections with FIV [59-62]. This is in parallel to HIV and SIV, where the majority of the lymphomas occurring in infected individuals were of B-cell origin [63,64]. In FeLV infection both T and B-cell lymphomas, with a predominance of T-cell lymphomas, have been described [65,66]. In the present case, FIV may have played an indirect role in tumorigenesis via immune suppression, which may have led not only to reactivation of FeLV infection but, independently, favored the emergence of a tumor cell clone. The role of the immune dysfunction induced by FIV for lymphomagenesis has been recognized earlier [67]. FIV-positive cats with lymphoma were reported to have substantially decreased CD4⁺ T cells, a reduced cell-mediated immunity and B-cell hyperactivity [60,67]. This parallels observations for other lentiviruses. HIV-1 is generally considered to play an indirect role in tumor development via immune suppression and co-factors, such as polyclonal B-cell activation; and the involvement of other infectious agents, notably the Epstein-Barr virus, have been identified [68]. Only rarely FIV seemed to have a direct mutagenic potential [62]. Accordingly, FIV as well as HIV and SIV proviruses have been reported to be only seldom
integrated in the host genome [62,67]. This is in contrast to directly oncogenic retroviruses including FeLV. In the present study, tumor tissues were positive for FIV, but since the cat was viremic, the PCR signal could have resulted at least partially from FIV positive circulating blood cells.

The outcome of an FeLV infection may depend on the age and genetic disposition of the cat, as well as on the virus genotype [4]. FeLV-A is thought to be minimally pathogenic but highly infectious and present in all natural FeLV infections [69,70]. FeLV-B, -C and -T develop within the FeLV-A infected cat via mutation, insertion or recombination events and are associated with the development of disease [71]. Using subgroup-specific PCR, only evidence of FeLV-A, but not of any other FeLV subgroups was found in cat #261. The inability to detect FeLV-B indicates that no recombination with endogenous FeLV sequences had taken place. So far reported FeLV-A isolates obtained from domestic and wild cats over several decades and across the world were found highly conserved, sharing over 97% amino acid sequence identity in the SU gene [29,40]. Remarkably, the SU sequences from the FeLV in cat #261 showed only 86-89% amino acid identity to FeLV-A/Glasgow-1. This divergence clearly exceeds that so far found among the members of the FeLV-A subgroup. Phylogenetic analysis confirmed that the closest relationship was to the original challenge strain, FeLV-A/Glasgow-1. At the time of sacrifice, virus isolation for the detection of replicating virus was not performed; therefore, receptor interference was not tested. The VRA sequence predominantly determines the receptor specificity [72]. The VRA of the progeny viruses revealed the highest resemblance to the VRA of FeLV-A/Glasgow-1; nonetheless a remarkable number of changes were observed. Recently, small changes in the VRA were shown to yield an envelope protein that was capable of using a different receptor [73,74]. However, these studies were conducted using constrained peptides in the VRA that were different from the naturally occurring FeLV isolates, and the engineered viruses were capable of using receptors outside of those traditionally used by the FeLV-A, -B and -C viral interference groups [73-75].

In gamma retroviruses, the U3 region of the LTR is a particularly potent viral determinant for pathogenicity. Changes in the FeLV U3 sequence, including the URE and the enhancer itself, drastically influence viral pathogenicity [41,76]. In the case of FeLV-related non-T-cell malignancies or non-neoplastic diseases, only one copy of the enhancer is typically found, but multiple copies of other regions of the LTR, such as the URE, have been reported [41,77]. FeLV that has been isolated from thymic lymphomas, however, contained duplications of either a partial or the entire enhancer sequence [78-80]. In FeLV-945, a natural isolate from a cat with a multicentric lymphoma, a 21-bp tandem triplication downstream of a single copy of the enhancer was shown to confer a replication advantage and to accelerate its disease onset [81]. The U3 sequences detected in cat #261 did not contain any duplication of the enhancer sequence or any repeats in the URE. However, point mutations at 31 locations were identified, including mutations in the enhancer region. Interestingly, 12 mutation sites had been described previously in FeLV-infected cats with lymphoma [41,77-79], and three additional mutations had been found in other kinds of tumors [41,77]. These mutations included one point mutation each in the CORE, the NF1 and the FLV-1 sequences. Thus, the mutations found in the U3 region of the progeny viruses in cat #261 may be causatively linked to the induction of the neoplastic disease.

The particularly high divergence of the progeny viruses from the originally inoculated FeLV-A/Glasgow-1 found in cat #261 may be explained by the long period (8.5 years) during which the virus had time to evolve in this cat. This, in turn, may indicate that minimal viral replication, at a level below the detection limit, had occurred in cat #261. In cats that have ostensibly recovered from FeLV viremia, we have found an association between plasma viral RNA, as a probable indicator of minimal viral replication in a sequestered tissue, and FeLV reactivation and tumor development [17]. No samples were available from cat #261 to determine plasma viral RNA throughout the infection. However, antigens expressed at very low levels would have constantly boosted the specific immunity, which may be in agreement with the persistently high anti-FeLV antibodies that were detected in cat #261. The long observation period in cat #261 exceeds the lifespan of most FeLV-infected pet cats, which often succumb within a few years, e.g. due to secondary infections. Since cat #261 was kept under barrier conditions, the risk of opportunistic infections or accidents was reduced.

At the time of FeLV reoccurrence, cat #261 was shedding FeLV via the saliva and had, therefore, become a potential source of infection for other cats. Shedding of FeLV RNA in the saliva has been demonstrated to be a consistent feature in antigenemic cats [28], and is thought to be the main transmission route for FeLV [82]. Interestingly, the saliva sample tested negative for the presence of env variants. In contrast, urine and feces were positive, albeit at a low level, for the env variants, indicating that the progeny viruses were shed via these secondary routes. It needs to be noted that the gastrointestinal and urinary tract, but not the salivary glands, were lymphoma positive and particularly the urinary
tract showed high env variant provirus and viral loads whereas the mandibular gland had high FeLV-A/Glasgow-1 viral loads (Additional file 4). We subsequently tested the cats from the same cohort and co-housed with cat #261, but none of them tested provirus positive for the env variants. The fact that no transmission of the env variants was detectable could be due to the low loads of the env variants in urine and feces and the presumably only very short duration of shedding. Alternatively, the variants may only inefficiently or not be transmittable. Viruses selected for their replication fitness within a host during long-term infections may differ from those that are efficaciously transmitted between hosts.

Conclusions
This is the first study to document the reoccurrence of FeLV in a cat with immunosuppression due to FIV infection many years after the initial virus inoculation. Since FIV is highly prevalent in some countries, we postulate that this phenomenon could also be observed in the field. Our results indicate that both FeLV and FIV were important for the disease development in the cat under investigation. While FIV may have mainly contributed via an immunosuppressive effect, FeLV and/or FIV may have been causally linked with the tumorigenesis. Cat #261 developed an active FeLV infection and was shedding FeLV at the time of euthanasia; thus at least part of the FeLV provirus had remained full-length and replication competent throughout the long aviremic phase. Moreover, we quantified for the first time viral loads of the FeLV challenge strain and the evolved progeny variants using sensitive, discriminating real-time PCR assays. The virus variants had largely replaced the inoculated prototype FeLV-A over time. Molecular characterization of the progeny viruses revealed a high variance in env not commonly found in the otherwise highly conserved FeLV-A subgroup. The large number of mutations may have led to increased viral fitness and/or changed the mutagenic characteristics of the virus.

Additional file 1: Multicentric lymphoma. Lymphoma detected upon necropsy. A) Kidney. B) Spleen. C) Histology of sternal lymph node: diffuse proliferation of mainly medium sized lymphatic cells with round nuclei, coarsely stippled chromatin and one to multiple medium sized nucleoli. Hematoxylin and Eosin, bar = 10 μm. D) Liver: positive immunohistochemical labeling of perportal infiltrating neoplastic cells for CD45R. Avidin-biotin complex method; Papanicolaou’s hematoxylin counterstain, bar = 50 μm.
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Additional file 2: Amino acid alignment of FeLV env sequences. Amino acid alignment of the env coding region from three FeLV subtypes (FeLV-A/Glasgow-I [GenBank: M12500], FeLV-B/Gardner-Arnstein [GenBank: K01209] and FeLV-C/Sarma [GenBank: M14331]) and the three env variants (KI261-I, KI261-II and SP261-III). The start of the SU region, the transmembrane domain (TM), the variable regions VRA, VRB and VRC, the PRR and the C2 disulfide-bonded loop (S-S) are labeled (according to [52]). Circles and stars represent amino acid sequences containing Asn-X-Ser/Thr, which indicate possible sites of N-glycosylation, as previously described [5383]. Potential N-linked glycosylation sites that are conserved in FeLV-A/Glasgow-1 and all env variants are represented by filled circles. New potential N-linked glycosylation sites in the env variants that were not present in the challenge strain FeLV-A/Glasgow-1 are labeled with filled stars; those that were present in FeLV-A/Glasgow-1 but lost in the env variants are marked with empty stars. Dots represent identical residues, and dashes represent spaces, which were introduced for proper alignment.
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Additional file 3: Comparison of nucleotide sequences of the FeLV US region. Nucleotide sequence comparison of the US region from FeLV-A/Glasgow-1 [GenBank: M12500] and the progeny virus variants retrieved from cat #261. Sequences found in multiple clones were depicted once. KI261-I was found in clones from the kidney. SP261-III, pKH11.8 and pKH11.9 were found in clones from the spleen. All other listed sequences were found in at least two different tissues, including the kidney, liver, spleen, and bone marrow. Enhancer elements and their corresponding nucleotide sequences are marked in the reference strain FeLV-A/Glasgow-1 [GenBank: M12500]. Nucleotides differing from the originally inoculated strain are indicated. Primer sequences for PCR [41] were located at positions -415 to -392 and 9 to 32 (not included in the figure). Previously described mutations are indicated by numbers in brackets: (1) Jackson et al., 1996; (2) Nishigaki et al., 1997; (3) Fulton et al., 1990 and (4) Matsumoto et al., 1990. Numbers at the top of the sequence indicate nucleotide positions relative to the presumptive RNA cap site in the FeLV/Glasgow-1 LTR [53]. Grey dashed lines indicate unknown sequences.
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Additional file 4: FeLV-A/Glasgow-1 and env variant provirus and viral loads in the tissues from cat #261. A) Provirus loads of FeLV-A/Glasgow-1 and the env variants. B) Viral (cDNA) loads of FeLV-A/Glasgow-1 and env variants. Viral tissue loads were normalized to GAPDH (top) and to RPS7 cDNA copy numbers (bottom). Tissues with apparent lymphoma are indicated by shaded areas.
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Additional file 5: FeLV viral loads in tissues from cat #261 normalized to RPS7. A) Total FeLV viral (cDNA) loads (US region PCR) in tissues with and without apparent lymphoma (analogous to Fig. 3C). B) Viral (cDNA) loads of FeLV-A/Glasgow-1 and env variants (analogous to Fig. 3C). C) Viral (cDNA) loads of env variants in tissues with and without apparent lymphoma (analogous to Fig. 3D). D) Viral (cDNA) loads of FeLV-A/Glasgow-1 in tissues with and without apparent lymphoma. Viral loads were normalized to RPS7 cDNA copy numbers determined by TaqMan real-time PCR, as described [27]. Viral loads were tested for statistically significant differences using the Mann-Whitney U-test (pMWU) as indicated.
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Authors’ contributions

AKHH performed and analyzed the research and drafted the manuscript. VC participated in the assay design and data analysis and revised the manuscript. FSB was the veterinarian in charge for this cat and conducted ex vivo sample collections. PO performed the necropsy, the macroscopic and histological analysis, the sample collection and revised the manuscript. FG performed histological and part of the immunohistochemical analyses. JR performed the IPA and in situ hybridization. MH performed part of the immunohistochemical and PARR analyses. EB performed molecular assays. KBP was responsible for the housing and the animal care. EN and EH performed the flow cytometry experiments. HL contributed to the study design and supervised the initial study. RHL conceived and supervised the follow-up study and edited the manuscript.

Competing interests

The authors declare that they have no competing interests.

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