Rapid evolution of the \textit{env} gene leader sequence in cats naturally infected with feline immunodeficiency virus

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Analysing the evolution of feline immunodeficiency virus (FIV) at the intra-host level is important in order to address whether the diversity and composition of viral quasispecies affect disease progression. We examined the intra-host diversity and the evolutionary rates of the entire \textit{env} and structural fragments of the \textit{env} sequences obtained from sequential blood samples in 43 naturally infected domestic cats that displayed different clinical outcomes. We observed in the majority of cats that FIV \textit{env} showed very low levels of intra-host diversity. We estimated that \textit{env} evolved at a rate of $1.16 \times 10^{-3}$ substitutions per site per year and demonstrated that recombinant sequences evolved faster than non-recombinant sequences. It was evident that the V3–V5 fragment of FIV \textit{env} displayed higher evolutionary rates in healthy cats than in those with terminal illness. Our study provided the first evidence that the leader sequence of \textit{env}, rather than the V3–V5 sequence, displayed the highest intra-host diversity and the highest evolutionary rate of all \textit{env} fragments, consistent with this region being under a strong selective pressure for genetic variation. Overall, FIV \textit{env} displayed relatively low intra-host diversity and evolved slowly in naturally infected cats. The maximum evolutionary rate was observed in the leader sequence of \textit{env}. Although genetic stability is not necessarily a prerequisite for clinical stability, the higher genetic stability of FIV compared with human immunodeficiency virus might explain why many naturally infected cats do not progress rapidly to AIDS.

INTRODUCTION

Deciphering the evolution of feline immunodeficiency virus (FIV) is essential for understanding why some infected cats have normal lifespans, whilst others progress rapidly to AIDS. The generation of polymorphic populations (Eigen, 1993) and the potentially high intra-host diversity of RNA viruses are features attributable to their biology as well as the unique characteristics of retroviral reverse transcription (Domingo, 2000). Improved understanding of the dynamics of viral populations at the intra-host level will lead to the development of more effective diagnostic tests and control strategies, and ultimately, will inform the design of the next generation of FIV vaccines, as the limited efficacy of the current commercial vaccine (Dunham \textit{et al.}, 2006) is likely attributable to the high genetic diversity of naturally occurring viruses.

The majority of FIV phylogenetic analyses have focused on sequence variation at the population level (Carpenter \textit{et al.}, 1998; Olmsted \textit{et al.}, 1992; Pistello \textit{et al.}, 1997; Samman \textit{et al.}, 2011; Teixeira \textit{et al.}, 2010) and highlighted the high variability of the V3–V5 region of the \textit{env} gene, which has been used to classify FIV into six distinct subtypes (A–E and putative subtype F) (Marcola \textit{et al.}, 2013; Sodora \textit{et al.}, 1994). Employing GARD (Kosakovsky Pond \textit{et al.}, 2006a) and jPHMM (Schultz \textit{et al.}, 2006) recombination detection methods, we recently observed an abundance of recombinant \textit{env} sequences in natural FIV infection, emphasizing the important role of recombination in generating viral
diversity at the population level and highlighting the limitations of the current phylogenetic classification of FIV (Bęczkowski et al., 2014a).

Mutations also contribute to viral diversity, although little is known about the evolutionary rate and the within-host viral diversity of the entire FIV env in natural infection (Table 1). In human immunodeficiency virus type 1 (HIV-1)-infected individuals, quasispecies composition varies greatly, with some studies reporting 10% diversity in long-term infected individuals (Delwart et al., 1997). It has also been proposed that intra-host diversity could influence the outcome of disease and that the rate of viral evolution decreases during disease progression (Delwart et al., 1997; Shankarappa et al., 1999).

Selection measures have been applied previously to assess forces acting on the evolution of different retroviral genes and at different regions within the same gene (Choisy et al., 2004). For example, in HIV-1 infection, gag is under negative selection pressure (Kils-Hütten et al., 2001), whilst env evolution is generally shaped by positive selection pressure (Wolfs et al., 1990). As gag encodes structural proteins, its conserved nature is essential to maintain viral integrity. In contrast, HIV-1 env encodes a heavily glycosylated envelope glycoprotein in which mutations in N-linked glycosylation sites are important for immune evasion (Yamaguchi & Gojobori, 1997). By analogy with HIV, we predict that FIV env is under a similar positive selection pressure as its human counterpart, but relatively little is known about the glycosylation pattern or selection forces acting on FIV Env in naturally infected cats.

To our knowledge, there are no data available concerning rates of evolution of the entire ORF of FIV env which, in contrast to primate retroviruses, contains an unusually long leader/signal region (Verschoor et al., 1993). Evolutionary studies of HIV-1 infection (Korber et al., 2000; Leitner & Albert, 1999; Lukashov et al., 1995; Shankarappa et al., 1999; Zhang et al., 1997) suggested that within-host evolution is influenced by both viral- and host-dependent factors. Whereas some studies reported increased evolutionary rates during the early stages of infection, declining towards terminal stage disease (Delwart et al., 1997; Shankarappa et al., 1999), others did not observe a similar pattern (Mikhail et al., 2005). Nevertheless, the assumption that the virus is under constant pressure in immunocompetent individuals and evolves at a high rate in response to a changing environment provides a plausible explanation for differences in evolutionary rates at different stages of disease (Lukashov et al., 1995). It is apparent that rates of evolution are dependent on the combined effects of host and viral factors, demonstrated by comparisons of the evolutionary tempo of different HIV-1 subtypes (Abecasis et al., 2009), and differences in replication rates of HIV-1 and -2 (MacNeil et al., 2007b).

Our ability to understand immune evasion, and to predict the outcome of FIV infection, depends on understanding how the intra-host evolution of FIV differs within and among cats. For this reason, we examined env sequences collected serially over 12 months from two distinct cohorts of naturally infected cats that developed different clinical outcomes. Our specific aims were to (i) quantify viral diversity, selection and rates of evolution in naturally infected cats, and (ii) determine whether the evolutionary rate correlated with the clinical outcomes of FIV infection.

**RESULTS**

**Phylogenetic inference**

The maximum-likelihood (ML) tree constructed using the entire dataset (Additional file 1, available in the online Supplementary Material) revealed high genetic diversity in FIV env at the population level (overall mean pairwise distance of 14.1%). Although >41% of cats were infected with circulating recombinant viruses, we found no evidence of within-host recombination (Bęczkowski et al., 2014a). At the intra-host level, sequential sequences amplified from 93% (40/43) of cats clustered together in monophyletic groups, whilst sequences from three cats (M5, P8 and P21), amplified at different time points, showed incongruent phylogenetic assignment (Fig. 1). Sequences from a parent/
offspring pair (cat M1 was an offspring of female M11) were classified as non-recombinant clade B and clustered closely together, with an overall mean pairwise distance of 0.3 % (Fig. S1).

**Intra-host diversity**

Having excluded sequences from the three cats that formed non-monophyletic groups, the mean pairwise distances for env within cats ranged from 0 to 2.9 % (median 0.13 %), 0 to 1.9 % (median 0.11 %) and 0 to 1.9 % (median 0.17 %), for early (A), intermediate (B) and late (C) time points, respectively (Table S1). When we compared the diversity of structural fragments of env, although not statistically significant ($P=0.29$), the leader region displayed the highest variation at the intra-host level; the median overall mean pairwise distances for each time point were 0.2, 0.2 and 0.27 %, respectively (Fig. 2, Table S1).

**Selection**

The mean $\omega$ value (dN/dS ratio) determined by SLAC (single likelihood ancestor counting) for the env sequences from both cohorts was 0.34. Examining the entire dataset, the SLAC analysis indicated 14 (1.6 %) positively and 201 (23 %) negatively selected sites. FEL (fixed effects likelihood) reported 22 (2.5 %) positively and 391 (44.8 %) negatively selected sites, whilst IFEL (internal fixed effects likelihood) indicated 24 (2.7 %) and 377 (43.2 %) sites under positive and negative selection, respectively. Positively selected sites, consistently identified by the three methods (Table S2), were restricted to specific regions of
**env**, with a high proportion (5/10, 50 %; sites 8, 62, 96, 156 and 157) lying within the leader region (Fig. 3). The selection hot spots at positions 400 and 750 were located within B-cell epitopes identified in previous studies (Lombardi et al., 1993; Pancino et al., 1993) (Fig. 3). An investigation of the global distribution of potential N-linked glycosylation sites (PNGSs) for the presence of fixed and shifting sequons revealed that, regardless of the high genetic variability of the analysed sequences (n=329), conserved PNGSs did exist (Fig. 3). However, none of the predicted PNGSs fell within sites identified as being under positive selection.

![Fig. 2.](image1.png) **Fig. 2.** Comparison of mean pairwise distances between **env** fragments and entire **env** sequences amplified from three samplings (A, B and C on the x-axis). Median values for the leader, V3–V5, NV and **env**, respectively, for time points A (0.20, 0.11, 0.16 and 0.13 %), B (0.20, 0.12, 0.12 and 0.11 %) and C (0.27, 0.19, 0.14 and 0.17 %).

![Fig. 3.](image2.png) **Fig. 3.** Frequency and location of predicted PNGSs (a) and sites under positive selection (b) with corresponding dN/dS (ω) values. PNGSs at positions 303, 335, 347, 536, 553 and 738 represent fixed sequons among all clades of FIV from different geographical origins (n=329). Sequons at positions 263, 279, 423, 427, 504, 734 and 341, 523 and 746 were also fixed, and were present in >99 % of sequences. Positively selected sites (n=10) were consistently identified by three different detection methods (for details, see Table S2). Leader sequence and B-cell epitope regions identified in previous studies are highlighted in grey. The putative leader sequence cleavage site is located at the position 528 bp from the start codon of the **env** ORF of reference FIV Petaluma (GenBank accession number M25381.1). Location of V1–V8 regions is highlighted below the graphs.
Rates of molecular evolution

For all cats in the study, env was estimated to evolve at a mean rate of $1.16 \times 10^{-3}$ [0.7–1.67 95% highest posterior density (HPD)] substitutions per site per year under a relaxed log-normal clock model (Table S3) with the first and second codon positions evolving at a lower rate than the third codon position [0.95 (0.90–0.98 95% HPD) versus 1.1 (1.02–1.19 95% HPD)]. To determine whether specific regions of the env gene evolved at different rates, we estimated the clock rate for the leader, the V3–V5 and the non-variable (NV) regions. The leader region evolved at the highest rate [$3.43 \times 10^{-3}$ (1.55–5.6 95% HPD) substitutions per site per year under a relaxed log-normal clock model] in comparison with other fragments, whilst the evolutionary rate of the V3–V5 region, commonly regarded as highly variable, was over three times lower [$1.08 \times 10^{-3}$ (0.5–1.74 95% HPD) substitutions per site per year under a relaxed log-normal clock model] (Fig. 4, Table S3).

As the individually housed cats from the Chicago cohort were generally healthy, whilst the majority of group housed cats from the Memphis cohort had high morbidity and 63% mortality over the study period, we compared the evolutionary rates between and within the cohorts. Median rates of evolution of the entire env and its fragments were higher for the sequences from cats in the Chicago cohort (Fig. 4). We hypothesized that env evolution in healthy cats that remained alive throughout the study period would be faster than in terminally ill cats that died during the study period. Fig. 5 illustrates a comparison of the evolutionary rates for sequences from (i) 11 deceased cats, and (ii) six cats that remained alive and generally free of clinical signs during the study. The entire env, leader and NV fragment evolved at higher rates in terminally ill animals. This relationship, however, was reversed for the V3–V5 region, which evolved at a higher rate in cats that remained alive during the study period. Recombinant env sequences ($n=123$) evolved three times faster than sequences without any detectable prior history of recombination ($n=184$) (Table S3).

**DISCUSSION**

Analyses of serial env sequences collected over 12 months from naturally infected cats offered a rare opportunity to examine the intra-host dynamics and evolution of FIV. Despite considerable variation among individuals, the overall intra-host diversity of env (up to 3%) was low compared with primate immunodeficiency viruses (commonly $\geq$5%) (Rambaut et al., 2004; Salemi, 2013). Furthermore, the overall mean distance between sequences obtained from two closely related animals (M1 was the offspring of M11), following at least one postulated transmission event, was remarkably low. These results suggest that the env sequences were stably maintained and evolved slowly following vertical transmission. This is consistent with an earlier study examining V3–V5 env sequence diversity of the Aomori 2 strain of FIV following vertical transmission; 100% homology was observed between viruses isolated from a queen and her kitten 48 weeks post-transmission (Motokawa et al., 2005).

The apparent discrepancy between low, within-host viral diversity and relatively high diversity of FIV at the population level is intriguing, especially in light of differences in evolutionary rates demonstrated at the within-host and epidemiological levels in HIV infection (Alizon & Fraser, 2005; Lythgoe & Fraser, 2012). Analysis of FIV evolutionary rates at the epidemiological level are constrained by the low
number of longitudinal full-length FIV env sequences that are available and comprehensive datasets would be needed to quantify these reliably.

Low diversity and rates of evolution in lentiviruses are commonly associated with lower pathogenicity. Although HIV-1 and -2 achieve similar proviral loads in infected individuals, HIV-2 replicates to lower titres in the plasma (Popper et al., 2000), and displays a lower rate of sequence evolution (MacNeill et al., 2007a) and lower pathogenicity, reflected by a slower decline in CD4⁺ T-cell numbers and slower disease progression (Marlink et al., 1988, 1994; Popper et al., 1999; Whittle et al., 1994). Similarly, bovine immunodeficiency virus, which has low pathogenicity, is more closely related to FIV than to primate lentiviruses (Olmsted et al., 1989) and also exhibits very little sequence variation (Carpenter et al., 2000).

The genetic stability of some lentiviruses could be associated with the fidelity of their reverse transcriptase (Lewis et al., 1999). Mutations in the catalytic YMDD motif of HIV-1 and the V148S mutation in simian immunodeficiency virus (SIV) (Huisman et al., 2008) have been implicated in increasing reverse transcriptase fidelity (Olmsted et al., 1989; Operario et al., 2005). It is likely that the reverse transcriptase of FIV has enhanced fidelity compared with the reverse transcriptase of HIV-1 (Huisman et al., 2008). The presence of almost identical sequences in our study might further indicate that FIV exhibits a low replication rate during persistent infection or that most infections occur via cell-to-cell transfer owing to feline tetherin/BST-2 (Dietrich et al., 2011). It is also possible that viral evolution is constrained due to the high fitness cost associated with divergence from the parental virus (Domingo & Holland, 1997). Whatever the specific mechanism, the low sequence variation of FIV in comparison with HIV-1 could be indicative of co-adaptation between FIV and its host, which are thought to have co-existed together for longer than HIV-1 and humans (Pecon-Slattery et al., 2008). Furthermore, the analysis of viral sequence divergence and disease pathogenesis amongst the Felidae suggests different periods of virus–host co-evolution (Troyer et al., 2008). In general, the least pathogenic viruses appear to have co-evolved with their hosts for longer periods of time, whereas more virulent strains have more recent origins. This pattern is reflected in the disease-inducing potential of different species-specific FIVs, with FIV-Pco (FIV of the puma) being the least pathogenic, whilst FIV-Fca (FIV of the domestic cat) is the most pathogenic. Although FIV infection in pumas and lions was previously reported to be asymptomatic, recent evidence suggests that infected lions do indeed exhibit mild clinical sings of immunodeficiency, subsequently affecting the lifespans of infected animals (Roelke et al., 2009). Similar observations have been made recently in SIV-infected chimpanzees, previously regarded as asymptomatic virus carriers (Terio et al., 2011).

It has been suggested that the slow rate of evolution documented in terminal HIV-1 infection might be linked to the weakened host immune system (Delwart et al., 1997; Shankarappa et al., 1999). Consistent with this theory, we found that the V3–V5 fragment, which contains neutralization epitopes and is under immune system surveillance (Lombardi et al., 1993; Pancino et al., 1993), evolved faster in healthy cats than in terminal ill cats. In addition, the rates of evolution across all datasets from the individually housed cats from Chicago (which were generally in good health) were consistently higher than those estimated for sequences from the cats from Memphis (which displayed more clinical signs of illness).

Recombination is a potential driver of rapid evolutionary change and plays a significant role in generating viral diversity among retroviruses (Onafuwa-Nuga & Telesnitsky, 2009). In the present study, viruses with a prior history of recombination (Bęczkowski et al., 2014a) exhibited higher evolutionary rates, which could be the result of strong immune selection pressure acting on the recombinant Envs compared with (presumably) more host-adapted, evolutionary older non-recombinants. Nonetheless, the mechanisms responsible for the faster evolution of recombinants might not be related to the immune system, and could be attributed to their higher replication tempo and increased infectivity (Tebit et al., 2007).

This is the first study to demonstrate that against a background of low diversity and slow evolution for env as a whole, the leader region exhibited the highest proportion of positively selected sites and the highest evolutionary rate among all fragments examined, but more data are required to confirm this. The N-terminal signal peptide of FIV plays an important role in post-translational targeting of the Env precursor and its translocation through the endoplasmic reticulum (Verschoor et al., 1993). Studies of viral leader sequences highlighted their enormous complexity, suggesting a role not only in post-translational events, but also in post-cleavage events (Hegde & Bernstein, 2006; Lemberg & Martoglio, 2002). Leader-encoded peptides are involved in self-antigen presentation (Borrego et al., 1998) – a property exploited by human cytomegalovirus. This virus has a signal peptide with nine residues identical to the MHC class I signal peptides (Tomasec et al., 2000), which helps it to evade detection by NK cells presenting a virus-encoded MHC class I molecule (Ulbrecht et al., 2000). Furthermore, sequence variation among the signalling peptides of acute and chronic isolates of HIV has also been demonstrated (Gnanakaran et al., 2011). These findings suggest that the leader may play an important role in regulating env expression on the viral surface (Hegde & Bernstein, 2006) and therefore viral infectivity. This hypothesis warrants further research and validation in functional studies.

By comparing amino acid sequences and global patterns of PNGSs, 19 sites were identified which were conserved in >97% of the examined sequences, with six PNGSs displaying 100% fixed cross-clade pattern. This striking consistency at the population level suggests that glycosylation at these sites is likely to play an essential role in the
folding and integrity of the viral Env. Patterns of fixed N-linked glycosylation sites have also been noted in infections with SIV (Chackerian et al., 1997) and HIV-1 group M subtypes A–G (Gao et al., 1996).

Statistical characterization of a relationship between within-host evolutionary dynamics and the clinical outcome of retroviral infection remains problematic. Although HPDs permit a comparison of evolutionary rates between groups of patients, the estimate uncertainty tends to be high due to limited signal within intra-host datasets (Carvajal-Rodríguez et al., 2008). We have considered hierarchical phylogenetic models to test for rate differences between cats, but preliminary analyses suggested limited power to detect effects in our data using these parameter-rich models, as also suggested by others (Edo-Matas et al., 2011).

The conclusions drawn from the intra-host diversity and rates of evolution are restricted by (i) the 63 % mortality rate within the Memphis cohort, (ii) the relatively low number of sequences available from some cats at selected time points and (iii) the short 12-month study period (Seo et al., 2002), and indeed we identified almost identical sequences over the three different sampling times. However, despite this short time frame and the potential limitations of the methodology used, we reliably quantified within-host evolution. The presence of identical amplicons in cats at three different sampling times suggest that factors which tend to generate false diversity (polymerase template switching, PCR-induced errors) did not affect our results. Furthermore, the results presented here are in agreement with previous FIV studies employing ‘bulk’ PCR (Ikeda et al., 2004; Motokawa et al., 2005), end-point dilution proviral DNA PCR (Kraase et al., 2010) as well as a study examining sequences from plasma viral RNA (Huisman et al., 2008), and we observed low sequence variation and high genetic stability of FIV in a relatively large sample of naturally infected cats.

CONCLUSION

We observed relatively low intra-host diversity and a low rate of evolution of the entire env in cats naturally infected with FIV. The greater overall genetic stability of FIV compared with HIV-1 might explain why many naturally infected cats do not progress rapidly to AIDS. To the best of our knowledge, this is the first study to demonstrate that the leader sequence is the fastest evolving region of env. As the majority of previous studies focused on the V3–V5 region, these results indicate that the role of the unusually long leader sequence of FIV env in viral infectivity and immune evasion might have been underestimated.

METHODS

Cats, FIV env sequences and alignments. In total, 44 privately owned, neutered domestic cats, of various ages, breeds and conditions of health, were enrolled into the study (Bęczkowski, 2013) based on a history of a positive FIV antibody test result (SNAP FIV/FelV Combo Test; IDEXX Laboratories). All cats were feline leukemia virus antigen-negative and their FIV-positive status was confirmed by virus isolation (Hosie et al., 2009). In the study group, 27 cats lived together in a large multi-cat household in Memphis, TN, USA, where FIV-positive and FIV-negative cats were housed indoors with unrestricted access to one another. The remaining 17 cats lived in individual households in Chicago, IL, USA, with exception of five cats: two cats (P7 and P4) had been rehomed together and were living in the same household, one cat (P9) had been rehomed with another FIV-positive cat not enrolled in the study, one cat (P13) had been rehomed with another FIV-negative cat, and one cat (P21) was housed with another two FIV-positive cats in the rehoming centre. Cats were classified, based on a 6-monthly clinical examination by a registered specialist in feline medicine (A.L.), into two groups: (i) healthy (cats with no abnormalities found on clinical examination) and (ii) unhealthy (cats with any abnormalities detected on clinical examination). At the time of enrolment, there were 10 healthy (59 %) and seven unhealthy (41 %) cats in the Chicago cohort. In the Memphis cohort, 12 cats were classified as healthy (44 %) and 15 cats were classified as unhealthy (56 %) at the time of enrolment. During the study period, there was a 63 % mortality rate in the Memphis cohort (17/27), whilst only one cat (5.9 %) from the Chicago cohort died during the same time frame (Bęczkowski, 2013).

Multiple full-length FIV env genes (~2500 bp) were amplified directly from whole blood collected at 6-monthly intervals starting in January and May 2010 for Memphis and Chicago, respectively (time points A, B and C), using a nested PCR protocol (Table S4). First-round PCR products were amplified directly from blood, without genomic DNA extraction by Phusion Blood Direct II Polymerase (Thermo Fisher Scientific) followed by direct nucleic acid sequence determination. Phusion Blood Direct II Polymerase is a proofreading polymerase with 25-fold greater accuracy than Taq DNA polymerase, determined with a modified lacd-based method (Frey & Suppmann, 1995). The nucleic acid sequence of the first-round PCR product informed the primer design for the second-round PCR, which was performed using High Fidelity PCR Master (Roche), which amplifies with threefold greater accuracy than Taq DNA polymerase (Roche). Strain-specific primers for the second-round PCR incorporated restriction sites. Amplified env sequences were cloned into the eukaryotic expression vector VR1012 (Hartikka et al., 1996) and transformed into Escherichia coli MAX Efficiency DH5x Competent Cells (Invitrogen). Thus constructed VR1012 plasmids expressing FIV env were sequenced (Table S5) using a Big Dye Terminator v1.1 kit (Applied Biosystems) for the purpose of the present study, before being assessed in functional studies (Bęczkowski et al., 2014b, c). Special measures were taken to avoid the possibility of contamination, in both the clinical and laboratory settings: cats were double identified prior to blood sampling, PCRs were prepared in a designated UV-treated room and fresh, unopened reagents were used at each separate time point throughout the study.

There were 355 serial env sequences from 43 cats available for analysis from the two cohorts (Table S5). The number of sequences varied according to the availability of follow-up samples, being largely influenced by the 63 % mortality rate within the Memphis cohort. Multiple sequence alignments were conducted in MEGA5 (Tamura et al., 2011) and curated manually to ensure homology of gaps in sequences of variable length. Analyses were performed using the entire env DNA sequences and the three structural fragments: (i) leader/signal region (~509 bp in length), (ii) variable V3–V5 region (~630 bp) and (iii) remaining concatenated fragments of the entire env after exclusion of the V3–V5 region, denoted NV (~1900 bp). The temporal signal and ‘clocklikeness’ of the env phylogenies were tested in Path-o-Gen (http://tree.bio.ed.ac.uk) (Table S6).
Phylogenetic trees. ML trees were constructed in MEGAS (Tamura et al., 2011) under the HKY nucleotide substitution model, selected through jMODELTEST analysis (Posada, 2008). Statistical support for ML trees was estimated using bootstrapping analysis with 1000 replicates (Efron et al., 1996). The ML tree constructed using the entire dataset was carefully examined for evidence of non-monophyletic clustering of multiple sequences amplified from each individual. Sequences from three animals (M5, P8 and P21) amplified at different time points did not form monophyletic groups, and were excluded from further intra-host and evolutionary rate analyses.

Intra-host diversity. Intra-host sequence variation among the env sequences and three fragments of the gene (leader, V3–V5 and NV) at each time point were calculated as mean and highest pairwise distances under the HKY nucleotide substitution model in PAUP 4.0 v10 (Wilgenbusch & Swofford, 2002). To explore the variation in mean pairwise distances of three structural fragments of env, a general linear model (Khan, 2013) was used with ‘structural fragment of the env’ and ‘time point’ as fixed effects and ‘cat’ as a random effect. No evidence of recombination within hosts was found using rigorous fivefold recombination testing (Bęczkowski et al., 2014a).

Selection and PNGSs. Nucleotide sites under diversifying or purifying selection were identified and dN/dS ratios (o) for every codon in the alignment were estimated using three different methods: SLAC (Kosakovsky Pond & Frost, 2005) at P<0.01, FEL (Kosakovsky Pond & Frost, 2005) at P<0.1 and IFEL (Kosakovsky Pond et al., 2006b) at P<0.1. To focus on the most strongly supported positively selected sites, we reported those sites which were consistently identified by all three methods.

PNGSs and their position in the protein alignment were identified by the N-GlycoSite tool available at the Los Alamos National Laboratory web server (http://www.hiv.lanl.gov/content/index) and the number of sequences with glycosylation sites was counted. The analysis included reference sequences from GenBank representing clades A–D: Aomori 1 (GenBank accession number D37816), Aomori 2 (GenBank accession number D37817.1), FIV C (GenBank accession number AF474246.1), Dixon (GenBank accession number L00608.1), Dutch (GenBank accession number X60725), Fukuoka (GenBank accession number D37815.1), Sendai 1 (GenBank accession number D37813.1), Shizouka (GenBank accession number D37811.1), UK2 (GenBank accession number X69494.1), UK8 (GenBank accession number X69496.1), USL2489 (GenBank accession number U11820.1), Yokohama (GenBank accession number D37812.1), Petaluma (GenBank accession number M25381.1), PPR (GenBank accession number M36968.1) and sequences from all cats excluding identical, duplicate sequences (n=329). Alignments were numbered according to positions in the M49C C76 sequence.

Rate of molecular evolution. Rates of evolution of the entire env gene and individual fragments from each Memphis and Chicago cohorts and all cats were estimated using BEAST v1.7.1 (Drummond & Rambaut, 2007), based on sampling date information. Additionally, rate comparisons were made between non-recombinant or recombinant sequences from healthy and sick cats, and cats from Memphis that remained alive and those that died during the study period.

Sequence alignments from animals from which data were available from more than one time point were included in the analysis: 17 cats from Memphis (197 sequences) and 12 cats from Chicago (108 sequences). The analysis of differences in evolutionary rate between sequences from alive (n=6) and deceased cats (n=11) was based on sequences from Memphis cohort. The HKY evolutionary model of substitution with four-category Γ distribution was selected with codon positions (1 +2) and 3 as partitions (Shapiro et al., 2006). The analyses in BEAST were performed estimating independent trees for each cat, with a linked clock rate parameter across individual cats, under strict and relaxed log-normal clock models with uniform distribution clock rate priors, informed by previous estimates of evolutionary rates for FIV and HIV (Table 1). The length of the Markov chain Monte Carlo was set for 2 × 10^8 iterations with 2 × 10^7 burn-in, and was run until convergence and effective sample sizes >100 were obtained.

The BEAST-generated output log file was analysed in TRACER v1.5. The coefficient of variation in rates among branches was used to determine whether a relaxed molecular clock was more appropriate for these data. This was assumed to be the case if the estimate for the coefficient of variation excluded zero.

Statistical support was provided in the form of parameter estimates and their HPD values.

Ethics and consent. The study and its aims were reviewed and approved by the University of Glasgow Ethics Committee and the Purdue Animal Care and Use Committee. Cat owners provided written informed consent for their participation in the study.

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REFERENCES


