A study of multiple Felis catus Papillomavirus types (FcaPV-1, 2, 3, 4) in cat skin lesion in Italy by q-PCR

Journal:	Journal of Feline Medicine and Surgery
Manuscript ID	JFMS-17-0031.R2
Manuscript Type:	Original Article
Date Submitted by the Author:	n/a
Complete List of Authors:	Mazzei, Maurizio; University of Pisa, Department of Veterinary Science Forzan, Mario; University of Pisa, Department of Veterinary Science Carlucci, Vito; University of Pisa, Department of Veterinary Science Anfossi, Antonio; Universita degli Studi di Sassari, Dipartimento di Medicina Veterinaria Alberto; Universita degli Studi di Sassari Dipartimento di Medicina Veterinaria Albanese, Francesco; Private veterinary laboratory "LaVallonea" Binanti, Diana; Private veterinary laboratory AbLab Millanta, Francesca; University of Pisa, Department of Veterinary Science Baroncini, Lisa; University of Pisa, Department of Veterinary Science Pirone, Andrea; University of Pisa, Department of Veterinary Science Abramo, Francesca; University of Pisa, Department of Veterinary Science
Keywords:	feline, papillomavirus, skin, viral plaque, in situ Bowenoid carcinoma, squamous cell carcinoma, actinic keratosis, q-PCR
Abstract:	Abstract Objective: Aim of the study is to investigate by q-PCR the presence of Papillomavirus in feline viral plaques (VP), Bowenoid in situ carcinoma (BISC), squamous cell carcinoma (SCC), and actinic keratosis (AK). Methods: Twenty-nine cases with previously established diagnosis of feline VP, BISC, invasive SCC, and AK were selected from a dermatopathological database. A critical re-evaluation of diagnosis was performed by defining clear criteria toward carcinomatous vs non carcinomatous, in situ vs invasive (if carcinomatous) and viral vs actinic. Cases were evaluated for p16 immunolocalisation. The presence of the target viral genes for FcaPV- 1, 2, 3 and 4 was determined by q-PCR. Data generated Cq values, which represent a normalized measure of DNA viral quantity. Samples with a positive Cq values were submitted to sequence analysis. Results: 4 VPs, 19 BISCs 4 SCCs and 1 AK were included. By Cq analysis we found that all VPs were positive for FcaPV-1 or FcaPV-2; and 8 BISCs for FcaPV-1, 2, 4. FcaPV-2 was the most prevalent among the group of VPs and BISCs. Conclusion and relevance: Using Cq method we report the first evidence of FcaPV-1, 2 and 4 in Italy, FcaPV-2 was the most frequently detected. To a lesser extent also FcaPV-1 and FcaPV-4 has been detected in our examined samples while FcaPV-3 was never associated to viral induced lesions by Cq investigation. Compared to conventional PCR the Cq method has the advantage to establish a possible role of the virus in the outcome of infection.



1	A study of multiple Felis catus Papillomavirus types (FcaPV-1, 2, 3, 4) in
2	cat skin lesion in Italy by q-PCR.
3	
4	Mazzei M.*, Forzan M.*, Carlucci V.*, Anfossi A.G.§, Alberti A.§, Albanese F.°,
5	Binanti D.^, Millanta F.*, Baroncini L.*, Pirone A.*, Abramo F.*
6	* Department of Veterinary Sciences, University of Pisa, Viale delle Piagge 2, 56124 Pisa,
7	Italy;
8	[§] Department of Veterinary Medicine, University of Sassari, via Vienna 2, 07100 Sassari,
9	Italy;
10	° Private Veterinary Laboratory "LaVallonea", via G. Verdi 39, 73031 Alessano (LE), Italy;
11	^ Private Veterinary Laboratory "AbLab", via Privata Massa Neri 13, 19038 Sarzana (SP),
12	Italy.
13	
14	Corresponding author: Francesca Abramo, associate professor.
15	Department of Veterinary Sciences, University of Pisa, Viale delle Piagge 2, 56124 Pisa, Italy
16	Phone number +39 0502216988 – Mobile +39 3287119710; e-mail: francesca.abramo@unipi.it
17	

Key words: feline papillomavirus, skin, viral plaque, *in situ* Bowenoid
 carcinoma, squamous cell carcinoma, actinic keratosis, q-PCR

20 Abstract

Objective: Aim of the study is to investigate by q-PCR the presence of
Papillomavirus in feline viral plaques (VP), Bowenoid *in situ* carcinoma (BISC),
squamous cell carcinoma (SCC), and actinic keratosis (AK).

Methods: Twenty-nine cases with previously established diagnosis of feline VP, 24 BISC, invasive SCC, and AK were selected from a dermatopathological 25 database. A critical re-evaluation of diagnosis was performed by defining clear 26 criteria toward carcinomatous vs non carcinomatous, in situ vs invasive (if 27 carcinomatous) and viral vs actinic. Cases were evaluated for p16 28 immunolocalisation. The presence of the target viral genes for FcaPV-1, 2, 3 29 and 4 was determined by q-PCR. Data generated $\Delta\Delta$ Cq values, which 30 represent a normalized measure of DNA viral quantity. Samples with a positive 31 $\Delta\Delta$ Cg values were submitted to sequence analysis. 32

Results: 4 VPs, 19 BISCs 4 SCCs and 1 AK were included. By $\Delta\Delta$ Cq analysis we found that all VPs were positive for FcaPV-1 or FcaPV-2; and 8 BISCs for FcaPV-1, 2, 4. FcaPV-2 was the most prevalent among the group of VPs and BISCs.

Conclusion and relevance: Using $\Delta\Delta Cq$ method we report the first evidence of 37 FcaPV-1, 2 and 4 in Italy, FcaPV-2 was the most frequently detected. To a 38 lesser extent also FcaPV-1 and FcaPV-4 has been detected in our examined 39 samples while FcaPV-3 was never associated to viral induced lesions by $\Delta\Delta$ Cq 40 investigation. Compared to conventional PCR the $\Delta\Delta$ Cq method has the 41 Je ro. advantage to establish a possible role of the virus in the outcome of infection. 42

43

http://mc.manuscriptcentral.com/jfms

44 Introduction

Detection of Feline papillomaviruses (FcaPVs) from feline cutaneous 45 lesions has been reported worldwide^{1,2} and these viruses are considered liable 46 of causing several feline skin conditions.¹⁻⁵ These include a wide spectrum of 47 proliferative skin lesions ranging from non-neoplastic viral plaques (VP) to pre-48 neoplastic Bowenoid in situ carcinoma (BISC), and invasive squamous cell 49 carcinoma (SSC).⁶ A progression from non neoplastic viral induced lesions to 50 overt neoplasia has also been documented to occur.⁷ Four FcaPV types 51 (FcaPV-1 to FcaPV-4) have been so far described in cats, having different 52 anatomical distribution and presentation.^{1,8-11} The most frequent FcaPV 53 (FcaPV-2, belonging to Dyothetapapillomavirus genus) was isolated from a 54 cutaneous pigmented plaque,¹² and later related to cutaneous SCC.^{13–15} 55 FcaPV-2 seems to play a role in a high proportion of nasal planum squamous 56 cell carcinomas and, animals with tumours associated to papillomaviral 57 aetiology with p16 upregulation, show increased survival compared with those 58 attributable to UV radiation.¹⁶ The relevance of FcaPV-2 in the pathogenesis of 59 both premalignant and malignant lesions has been recently investigated by 60 concurrently assessing number of DNA viral copies and viral gene expression.^{5–} 61 ⁷ However, the effective role of FcaPV-2 in the development of malignant 62 lesions is still a debated question, especially in view of recent results indicating 63

a high FcaPV-2 prevalence in healthy domestic cats.^{7,17} The role of other feline 64 PV types, namely FcaPV-1, 3, and 4, remains obscure. FcaPV-1, belongs to the 65 Lambdapapillomavirus genus and has been rescued from a hyperkeratotic 66 cutaneous lesion in a Persian cat by Tachezy et al.⁹ In the same year another 67 FcaPV was also identified in a shorthair domestic cat with papillomatosis.¹¹ 68 More recently the virus was detected in multiple small sessile raised lesions on 69 the ventral surface of the tongue in two 13-year-old domestic cats.¹⁸ FcaPV-3 is 70 a Taupapillomavirus, isolated for the first time from a cutaneous in situ 71 carcinoma.³ Recently FcaPV-3 has been detected from a BISC with a novel 72 behaviour.¹⁹ same histological feature and a benign clinical The 73 histopathological changes were observed in a feline basal cell carcinoma, from 74 which DNA sequences of a novel PV closely related to FcaPV-3 were detected 75 by the authors.⁴ FcaPV-4 belongs to genus *Taupapillomavirus*, and has been 76 detected in oral cavity following a severe gengivitis.¹⁰ Pathogenicity of FcaPV-1 77 and FcaPV-4 is still unclear and these viruses are rarely detected in oral 78 inflammatory as well as neoplastic lesions. Therefore, an active role of FcaPV-1 79 and FcaPV-4 in carcinogenesis is still debated. 80

Based on literature, it emerges that most cases of FcaPVs infection have been documented in domestic felids from New Zealand, North and South America and, to a lesser extent from Europe. This might just reflect the active research

http://mc.manuscriptcentral.com/jfms

done in these countries without any epidemiological meaning. In Europe PVs 84 infections have been reported in Switzerland in nine cases,^{12,20} in one case in 85 Germany²¹ and in one case in Italy.²² Here we assess by a combination of 86 histology, immunohistochemistry and q-PCR the concurrent presence of 87 FcaPV-1, 2, 3 and 4 viral DNA and p16 immunostaining in 29 feline lesions 88 89 collected in Italy including non neoplastic and preneoplastic, viral and non viral induced skin lesions namely: VP, BISC, invasive SSC and actinic keratosis 90 (AK). Two methods were applied to analyse the viral load data: absolute 91 quantification (AQ) and relative quantification (RQ). The AQ was applied to 92 determine the viral copy number per μg of DNA; however, by this method wide 93 range of values can be obtained without a cut-off value that would allow to 94 discriminate FcaPVs presence as innocent bystander or as responsible of viral 95 lesion. To address this problem we approached the RQ by $\Delta\Delta$ Cq calculation. ²³ 96 This method measures viral genome presence by relating the obtained values 97 of each of the 4 FcaPVs in each lesion classified as VP or BISC to those of 98 SCC and AK. 99

100 Materials and methods

101 Biopsies

¹⁰² Feline skin biopsies have been identified from dermatopathological databases

and 29 cases with a diagnosis of VP, BISC, invasive SCC and AK were 103 selected. The retrieved formalin fixed cases were routinely processed for 104 histopathology and 4 µm thick paraffin embedded tissues sections were stained 105 with haematoxylin and eosin (HE). Before inclusion in the study, the selected 106 samples were subjected to critical re-evaluation of diagnosis by defining clear 107 108 distinctive criteria, such as carcinomatous vs non carcinomatous, in situ vs 109 invasive (if carcinomatous), and viral vs actinic. The established criteria were 110 defined as described in the scientific literature and are summarized in table 1.^{14,20,24} Data from signalment and lesion distribution, when available, were 111 included. 112

113 *P16 immunohistochemistry*

Cases were evaluated for the detection of p16 by immunohistochemistry 114 employing an ABC system (Vector Laboratories, Burlingame, USA). Antigen 115 unmasking was carried out at 120°C for 3 min in a pressure cooker with a TRIS-116 EDTA buffer (1,2 g/L Tris, 0,36 g/L EDTA, pH 9.0). Sections were pre-treated 117 for 10 min with 1% H₂O₂ in 0.1 M phosphate buffered saline (PBS), pH 7.4 to 118 quench endogenous peroxidase activity, and blocked for 30 min at room 119 120 temperature (rt) in PBS with a 2% normal horse serum (PK-7200, Vector 121 Laboratories, Burlingame, USA) and 0.05% TritonX-100. Sections were then 122 incubated overnight at 4°C with a 1:100 diluted monoclonal primary antibody

p16-INK4a (mouse monoclonal IgG, BiorByt, San Francisco, USA). After
washing in PBS, an incubation with a secondary universal biotinylated antimouse/rabbit antibody (PK-7200, Vector Laboratories, Burlingame, USA) was
performed for 1h at rt. Staining was visualized with a diaminobenzidine (SK4105, Vector Laboratories, Burlingame, USA) solution under a light microscope
(Eclipse 80i, Nikon Tokyo, Japan).

129 Sampling material for q-PCR

Three 10 µm thick sections from the selected samples were cut with a microtome, place onto slides and left unfixed. These slides were observed under an optic microscope. Tissue that was not relevant for the study was scraped off and the remaining tissue directly collected in a DNase free 1,5 ml tube. To prevent carryover of contaminating DNA the microtome overlay was covered with a new piece of adhesive tape and a new blade was used for each sample.

137 DNA extraction

DNA extraction was performed using DNeasy Blood and Tissue kit (Qiagen, Hilden, Germany) following manufacturer's instruction and applying a preliminary removal of paraffin by extraction with xylene. DNAs were eluted in Multiple and each sample concentration was quantified using the Qubit

spectrophotometer (ThermoFisher Scientific, Waltham, USA), DNA samples
were stored at -20°C until analysis.

144 *q-PCR*

The extracted DNAs were amplified using four specific set of primers amplifying 145 a portion of the four types of FcaPVs L1 gene so far identified in the feline 146 species (Table 2). To normalize the amount of DNA used for each sample so to 147 148 achieve a correct quantification of viral copy number, a q-PCR under the same condition as for FcaPV but with a specific set of primers was run in parallel for 149 the reference gene albumin (ALB).²⁵ The number of copy of the target viral 150 151 gene, measured as Cq value, generates a Δ Cq value when compared to the corresponding Cq value of the reference gene. Moreover, a $\Delta\Delta$ Cq value was 152 calculated comparing the ΔCq value of the sample of interest to the mean ΔCq 153 value obtained from the SCC and AK group, considered as the negative control 154 155 group. The $\Delta\Delta$ Cg value represents a normalized measure of DNA viral quantity and was calculated using REST software.²⁶ In this work, a RQ using the $2^{-\Delta\Delta Cq}$ 156 method ²³ was adapted to estimate in each sample the fold change of FcaPV-1, 157 2, 3 and 4 target viral gene copies relative to the albumin reference gene. 158 Melting curve analysis was performed in conjunction with each four specific 159 FcaPV amplification protocol to determine if non specific products were 160

161 amplified during reaction. The specificity of the melting curve was compared to melting curves values obtained from a plasmid (pFcaPV) containing amplicon of 162 the four FcaPVs spanning the real-time products. The plasmid was generated 163 by inserting the four FcaPVs gene segments into pMA-T vector using GeneArt 164 technology (Thermo Fisher Scientific, Waltham, USA). Serial dilutions of 165 pFcaPV plasmid ranging from 10⁶ to 10² copies/5µl were used to calculate the 166 efficiency of q-PCR for each FcaPV set of primers and compared to the 167 efficiency calculated on five points of a 2-fold serial dilution of a quantified DNA 168 template for ALB gene. The efficiency of each g-PCR assay was similar and 169 ranging from 91% to 98% (average 94,6%). The lesions identified by 170 histological classification as SCC-AK were presumed to serve as the negative 171 control group and each sample belonging to this group was also individually 172 tested by $\Delta\Delta$ Cq analysis for each FcaPV type. Samples that had eventually 173 scored a positive $\Delta\Delta Cq$ value were excluded. In addition, the viral copy 174 number/µl of input DNA for each FcaPV type in each sample tested was 175 calculated (AQ). The assays were performed in Rotorgene thermocycler 176 (Corbett Research, Sydney, Australia) using SSCO SYBR Green master mix 177 (Biorad, Hercules, USA) and 5 µl of extracted DNA. All samples were tested in 178 duplicate and the results were calculated using the mean Cq values. Samples 179 180 positive for only one replica were considered as negative. All samples with a positive $\Delta\Delta$ Cq values were submitted to sequence analysis (BMR genomics, Padova, Italy).

183 **Results**

Histologically all VPs were recognized for areas of focal epidermal 184 dysplasia with koilocytes and keratinocytes with cytoplasm enlarged by blue-185 grevish material. Regarding BISC diagnosis, the presence of koilocytes or 186 koilocyte-like cells pointed to the viral origin in 17/20 cases while in the 187 remaining 3 cases diagnosis was formulated by other morphological details 188 listed in Table 1. SCC and AK showed the morphological alterations listed in 189 Table 1. Among the 20 BISCs cases 5 VPs were also detected in the 190 adjacent skin by histology and in other 5, sites of penetration of keratinocytes 191 192 through the basement membrane into the dermis were observed indicative of progression toward SCC. 193

194 Clinical reports for each of the 29 cases fulfilling the established 195 histological criteria were recorded. Breed, age, gender and anatomical 196 distribution are summarised in Table 3. Twenty-five subjects were shorthair 197 domestic cats, two persian and two main coons. Mean age at presentation 198 varied between 6 and 20 years (median 11,6), 14 out of 29 were female and 199 15 were male. A total of 4 VPs, 20 BISCs, 4 SCCs and 1 AK were detected.

http://mc.manuscriptcentral.com/jfms

200 Lesions were observed in densely haired skin regions in 9 cats (dorsum, flank, and shoulder), in areas non-exposed to sunlight in 1 cat (groin) and in 201 hypotrichotic and solar exposed skin in 18 cats (ears, nose, eyelids, temporal 202 region). All cases of SCC were observed in sun-exposed areas. Clinically 203 VPs were few millimeters wide, alone or grouped, slightly raised, pigmented 204 205 and non-pruritic lesions; BISCs were clinically larger than VPs, usually more 206 than 1 cm of diameter, multifocally coalescing, raised and often verrucous, 207 crusted and hyperpigmented. In SCC, erosive and ulcerative crusted lesions were clinically detected. In the only case in which AK was diagnosed, 208 erythema, scales and crust were seen (Figure 1a, b, c). 209

P16 was immunolocalized in all VPs epidermis and in the epidermis and follicular wall of the BISCs. No signal or only faint staining was detected in the SCC and AK lesion. Immunoreactivity was either nuclear and cytoplasmic; in VPs and BISCs the signal was strong while in the SCCs immunoreactivity was faint and therefore considered as no significant (Figure 1d).

²¹⁶ By q-PCR we were able to detect the albumin reference gene in 28 out of 29 ²¹⁷ DNA samples. Only one sample classified as BISC was not suitable to ²¹⁸ molecular analysis and therefore it was excluded from further analysis.

12

http://mc.manuscriptcentral.com/jfms

By AQ, the presence of FcaPVs has been detected in low copies in almost all 219 cases (Table 4). By RQ ($\Delta\Delta$ Cq method) a reduced number of samples was 220 linked to viral lesions. In details, FcaPVs positive $\Delta\Delta$ Cg values were detected 221 in all VPs (4/4) and in 8/19 BISCs (Table 3). VPs were positive to FcaPV-2 in 222 three samples and to FcaPV-1 in one sample. BISCs were positive to FcaPV-223 2 in three samples, FcaPV-4 and FcaPV-1 in one sample each. Interestingly, 224 as never reported before, in BISCs lesion we found three cases of double 225 infection. In two of those, FcaPV-1 and 2 were detected simultaneously and 226 in one case FcaPV-2 was present along with FcaPV-4 (Table 3). 227

In all cases with a positive $\Delta\Delta$ Cq value, the viral load was always greater than 10² copies /µg DNA (FcaVP-1: min 167 – max 310 Standard Error (SE): 33,0; FcaPV-2: min 100 – max 1150 SE: 163,9; FcaPV-4 min 391 – max 3700 SE: 1654,5) (Table 4). Sequence analysis confirmed the FcaPV types as indicated by q-PCR $\Delta\Delta$ Cq analysis.

233 Discussion

In this retrospective study, we detected the concurrent presence of different types of PVs, namely of FcaPV-1, 2 and 4 in feline VPs and BISCs by $\Delta\Delta$ Cq method using specific primers. To the best of our knowledge only the concurrent presence of FcaPV-2 and 3 have been reported from one BISC

and one VP.^{3,22} However differently from previous reports, here we report the
 presence of FcaPV-1 and FcaPV-4, alone or in association to FcaPV-2.

A limitation when performing PCR-based studies on PVs infection is 240 that the viral DNA is often detected and cannot be directly linked to an 241 effective role of virus in pathogenesis and therefore it is not possible to fully 242 establish whether the presence of a viral genome is uneventful (subclinical 243 lesion). In particular, FcaPV-2 has been recently detected in a number of skin 244 swabs from healthy cats, making difficult to discern whether PVs are causing 245 cancer or are merely an "innocent bystander".⁷ The hypothesis that the 246 presence of high viral loads likely represents an infectious state of PV is 247 supported by Thomson et al.⁷ who recently showed that the finding of high 248 copy numbers of FcaPV-2 DNA within a lesion suggests that the detected 249 250 virus may be responsible for it. In fact, while high viral copy numbers were 251 associated with E6/E7 gene expression, no gene expression was detected in association of low copy numbers, an indication of an incidental finding. 252

For the above-mentioned reasons, for this study we used a q-PCR protocol applying a $\Delta\Delta$ Cq method to investigate if and which FcaPVs have induced the lesion in cats. In this sense the obtained results are partially in line with the scientific literature which indicates so far that FcaPV-2 is the

major PV type implicated in skin preneoplastic and neoplastic lesions in cats.⁷ Since the genome organization and the role of viral proteins within the replication cycle are considered similar even between different genera belonging to the *Papillomaviridae* family we assumed that a similar link is maintained for feline PVs other than FcaPV-2. Importantly we reported positive $\Delta\Delta$ Cq values for FcaPV-1 and 4; two FcaPV types rarely reported.^{9–} 11,18

A comparison between our data with those obtained from other studies is difficult for two reasons: i) type of PCR primers used (specific *vs* consensus); ii) type of PCR used (conventional *vs* quantitative). A general prevalence ranging from 24 to 100% of FcaPV-2 both BISCs and VPs has been reported. ^{7,8,27,28} In our study, by using specific primers in a q-PCR analysis applying RQ method, we obtained a prevalence for FcaPV-2 in BISCs of 31,6% (6/19) and for all FcaPV types of 42,10% (8/19).

Failure to demonstrate PV DNA in every BISC have been previously explained through a carcinogenesis model in which papillomaviruses cause transformation but are only for a short period present within the lesion. ²⁹ Surprisingly, cases in which koilocytes were detected did not show any $\Delta\Delta$ Cq positive value; this result is likely due to: i) the detection of koilocytes in very focal lesions that

http://mc.manuscriptcentral.com/jfms

could have been no more present when additional serial paraffin sections were

277 prepared for DNA extraction and PCR analysis; ii) the presence of a PCR

278 undetectable PV variant.

Low viral copies for all PV types have been detected in almost all cases, 279 however the $\Delta\Delta Cq$ analysis was necessary to associate the FcaPVs as 280 biological agent potentially causing the lesions. $\Delta\Delta Cq$ positive values for the 281 different PV types were detected in all VPs and in 42% of BISCs. 282 Noteworthy, mean $\Delta\Delta Cq$ of positive samples differed among PV types being 283 62.5, 189.1 and 3083.5 for FcaPV-1, FcaPV-2 and FcaPV-4 respectively. 284 Studies considering a larger panel of cases might be necessary to establish a 285 286 further $\Delta\Delta$ Cq cut off value.

Here we document the presence of FcaPV-2 with high prevalence in both VPs and BISCs while we rescued FcaPV-1 and FcaPV-4 less frequently; these last two PVs are rarely described as playing an active role in skin and oral mucosa lesions in cats.^{9,10,18,30} None of the lesions was associated to FcaPV-3 when adopting $\Delta\Delta$ Cq method, confirming the rare occurrence of this FcaPV type also in Italy.

293 Either cytoplasmic or nuclear p16 signals were detected in all cases where 294 cytopathic effects were found (VP, BISC) while p16 immunolocalisation was

present with faint and mainly cytoplasmic signal in cases of SCCs and no p16-295 immunoreactivity was found in the case of AK. Despite the difficulties of 296 interpretation when the signal was faint, these results are in line with what 297 already documented previously. About half of the SCC studied by Thomson et 298 al.⁷ were negative to p16 immunostaining. A strong association between 299 FcaPV-2 E6/E7 gene expression and p16 immunostaining have been recently 300 found in feline SCC with 18 of 20 (90%) E6/E7-positive SCCs being also 301 positive for p16 compared to 13 of 40 (33%) E6/E7-negative SCCs.⁷ E6/7 gene 302 expression investigation has not been carried out in our study but our results 303 might reflect this correlation. 304

305 **Conclusion**

 $\Delta\Delta$ Cq analysis has proved to be necessary to associate the FcaPVs as biological agent potentially causing the lesions. Based on this method the presence of FcaPV-2 is confirmed to be the most representative FcaPV in feline skin lesions referable to diagnosis of VP and BISC in Italy. To a lesser extent also FcaPV-1 and FcaPV-4 has been detected in our examined samples.

313 Acknowledgement: The authors thank all the colleagues who referred the

314 **Cases**.

- 315 **Conflict of interest**: The authors declared no potential conflicts of interest with
- respect to the research, authorship, and/or publication of this article
- **Source of funding**: This study was financially supported by Fondi di Ateneo
- 318 University of Pisa.

319 Bibliografy

- Sundberg JP, Van Ranst M, Montali R, et al. Feline papillomas and
 papillomaviruses. Vet Pathol 2000; 37: 1–10.
- Gil da Costa RM, Peleteiro MC, Pires MA, et al. An Update on Canine,
 Feline and Bovine Papillomaviruses. Transboundary and Emerging
 Diseases, 11 September 2016. Epub ahead of print 11 September 2016.
 DOI: 10.1111/tbed.12555.
- Munday JS, Dunowska M, Hills SF, et al. Genomic characterization of
 Felis catus papillomavirus-3: a novel papillomavirus detected in a feline
 Bowenoid in situ carcinoma. Vet Microbiol 2013; 165: 319–25.
- Munday JS, French A, Thomson N. Detection of DNA sequences from a
 novel papillomavirus in a feline basal cell carcinoma. Vet Dermatol 2017;
 28: 236-e60.
- Munday JS, Gwyther S, Thomson NA, et al. Bilateral pre-auricular
 papillary squamous cell carcinomas associated with papillomavirus
 infection in a domestic cat. Vet Dermatol 2017; 28: 232-e58.
- 335 6. Scott DW, Miller WH, Griffin CE, et al. Muller & Kirk's Small Animal 336 Dermatology. Epub ahead of print 2001. DOI: 10.1016/B978-0-7216-

337 **7618-0.50002-X**.

- Thomson NA, Munday JS, Dittmer KE. Frequent detection of
 transcriptionally active Felis catus papillomavirus 2 in feline cutaneous
 squamous cell carcinomas. J Gen Virol 2016; 97: 1189–97.
- Munday JS, Kiupel M, French AF, et al. Detection of papillomaviral
 sequences in feline Bowenoid in situ carcinoma using consensus primers.
 Vet Dermatol 2007; 18: 241–5.
- 344 9. Tachezy R, Duson G, Rector A, et al. Cloning and genomic
 345 characterization of Felis domesticus papillomavirus type 1. Virology 2002;
 346 301: 313–21.
- Dunowska M, Munday JS, Laurie RE, et al. Genomic characterisation of
 Felis catus papillomavirus 4, a novel papillomavirus detected in the oral
 cavity of a domestic cat. Virus Genes 2014; 48: 111–119.
- Terai M, Burk RD. Felis domesticus papillomavirus, isolated from a skin
 lesion, is related to canine oral papillomavirus and contains a 1.3 kb non coding region between the E2 and L2 open reading frames. J Gen Virol
 2002; 83: 2303–7.
- Lange CE, Tobler K, Markau T, et al. Sequence and classification of
 FdPV2, a papillomavirus isolated from feline Bowenoid in situ carcinomas.
 Vet Microbiol 2009; 137: 60–65.
- Ravens PA, Vogelnest LJ, Tong LJ, et al. Papillomavirus-associated
 multicentric squamous cell carcinoma in situ in a cat: An unusually
 extensive and progressive case with subsequent metastasis. Vet
 Dermatol 2013; 24: 642-e162.
- 14. Munday JS. Papillomaviruses in felids. Vet J 2014; 199: 340–347.
- Altamura G, Corteggio A, Pacini L, et al. Transforming properties of Felis
 catus papillomavirus type 2 E6 and E7 putative oncogenes in vitro and
 their transcriptional activity in feline squamous cell carcinoma in vivo.
 Virology 2016; 496: 1–8.

Munday JS, Gibson I, French AF. Papillomaviral DNA and increased
 p16CDKN2A protein are frequently present within feline cutaneous
 squamous cell carcinomas in ultraviolet-protected skin. Vet Dermatol
 2011; 22: 360–366.

- Geisseler M, Lange CE, Favrot C, et al. Geno- and seroprevalence of
 Felis domesticus Papillomavirus type 2 (FdPV2) in dermatologically
 healthy cats. BMC Vet Res 2016; 12: 147.
- Munday JS, Fairley RA, Mills H, et al. Oral Papillomas Associated With
 Felis catus Papillomavirus Type 1 in 2 Domestic Cats. Vet Pathol 2015;
 52: 1187–1190.
- Munday JS, Fairley R, Atkinson K. The detection of Felis catus
 papillomavirus 3 DNA in a feline bowenoid in situ carcinoma with novel
 histologic features and benign clinical behavior. J Vet Diagnostic Investig
 2016; 28: 612–615.
- Wilhelm S, Degorce-Rubiales F, Godson D, et al. Clinical, histological and
 immunohistochemical study of feline viral plaques and bowenoid in situ
 carcinomas. Vet Dermatol 2006; 17: 424–431.
- Egberink HF, Berrocal A, Bax HA, et al. Papillomavirus associated skin
 lesions in a cat seropositive for feline immunodeficiency virus. Vet
 Microbiol 1992; 31: 117–25.
- Alberti A, Tore G, Scagliarini A, et al. Regressing Multiple Viral Plaques
 and Skin Fragility Syndrome in a Cat Coinfected with FcaPV2 and
 FcaPV3. Case Reports Vet Med 2015; 2015: 1–5.
- Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the
 comparative C T method. 2008; 3: 1101–1108.
- Levison DA, Parratt D. Skin Diseases of the Dog and Cat. Oxford, UK:
 Blackwell Science Ltd. Epub ahead of print 20 July 2005. DOI:
 10.1002/9780470752487.
- Helfer-Hungerbuehler AK, Widmer S, Hofmann-Lehmann R. GAPDH
 Pseudogenes and the Quantification of Feline Genomic DNA Equivalents.

396 Mol Biol Int 2013; 2013: 587680.

Pfaffl MW, Horgan GW, Dempfle L. Relative expression software tool
 (REST) for group-wise comparison and statistical analysis of relative
 expression results in real-time PCR. Nucleic Acids Res 2002; 30: e36.

AND 27. Nespeca G, Grest P, Rosenkrantz WS, et al. Detection of novel papillomaviruslike sequences in paraffin-embedded specimens of invasive and in situ squamous cell carcinomas from cats. Am J Vet Res 2006; 67: 2036–2041.

- 404 28. Munday JS, Peters-Kennedy J. Consistent detection of Felis domesticus
 405 papillomavirus 2 DNA sequences within feline viral plaques. J Vet Diagn
 406 Invest 2010; 22: 946–9.
- Smith KT, Campq MS. 'Hit and run' transformation of mouse C127 cells
 by bovine papillomavirus type 4: the viral DNA is required for the initiation
 but not for maintenance of the transformed phenotype. Virology 1988;
 164: 39–47.
- Munday JS, French AF, Gibson IR, et al. The presence of p16 CDKN2A
 protein immunostaining within feline nasal planum squamous cell
 carcinomas is associated with an increased survival time and the
 presence of papillomaviral DNA. Vet Pathol 2013; 50: 269–73.

415

416

417 LEGEND

Figure 1. Multiple symmetrical raised, hyperpigmented and crusted plaques on the face of a cat with Bowenoid in situ carcinoma (a). Histopathology from a viral plaque in a cat. Focal epidermal hyperplasia, absence of follicular wall involvement, evident viral cytopathic effects (koilocytes) (bar=50µm) (b);

Histopathology from a Bowenoid in situ carcinoma in a cat. Epidermal and 422 follicular dysplasia with upward in situ keratinocyte proliferation and a few 423 koilocytes (bar=50µm) (c); p16 immunohistochemistry, groups of 424 . a 425 keratinocytes show nuclear and cytoplasmic positivity (ABC system, bar=100 426 µm) (d).

1 Papillomavirus induced skin lesions in cats from Italy.

- 2 A study of multiple Felis catus Papillomavirus types (FcaPV-1, 2, 3, 4) in
- 3 cat skin lesion in Italy by q-PCR.
- 4
- 5 Mazzei M.*, Forzan M.*, Carlucci V.*, Anfossi A.G.§, Alberti A.§, Albanese F.°,
- 6 Binanti D.^, Millanta F., Baroncini L.*, Pirone A.*, Abramo F.*
- * Department of Veterinary Sciences, University of Pisa, viale delle Piagge 2, 56124 Pisa,
 Italy;
- [§] Department of Veterinary Medicine, University of Sassari, via Vienna 2, 07100 Sassari,
 10 Italy;
- [°] Private Veterinary Laboratory "LaVallonea", via G. Verdi 39, 73031 Alessano (LE), Italy;
- Private Veterinary Laboratory "AbLab", via Privata Massa Neri 13, 19038 Sarzana (SP),
 Italy.
- 14
- 15 **Corresponding author:** Francesca Abramo, associate professor.
- 16 Department of Veterinary Sciences, University of Pisa, Viale delle Piagge 2, 56124 Pisa, Italy
- 17 Phone number +39 0502216988 Mobile +39 3287119710; e-mail: francesca.abramo@unipi.it

18

Key words: feline papillomavirus, skin, viral plaque, *in situ* Bowenoid
 carcinoma, squamous cell carcinoma, actinic keratosis, q-PCR

21 Abstract

Objective: Aim of the study is to investigate by q-PCR the presence of PV
Papillomavirus in feline viral plaques (VP), Bowenoid *in situ* carcinoma (BISC),
squamous cell carcinoma (SCC), and actinic keratosis (AK).

Methods: Twenty-nine cases with previously established diagnosis of feline VP, 25 BISC, invasive SCC, and AK were selected from a dermatopathological 26 database. A critical re-evaluation of diagnosis was performed by defining clear 27 criteria toward carcinomatous vs non carcinomatous, in situ vs invasive (if 28 carcinomatous) and viral vs actinic. Cases were evaluated for p16 29 immunolocalisation. The presence of the target viral genes for FcaPV-1, 2, 3 30 and 4 was determined by q-PCR. Data generated $\Delta\Delta Cq$ values, which 31 represent a normalized measure of DNA viral quantity. Samples with a positive 32 $\Delta\Delta$ Cg values were submitted to sequence analysis. 33

Results: 4 VPs, 19 BISCs 4 SCCs and 1 AK were included. By $\Delta\Delta$ Cq analysis we found that all VPs were positive for FcaPV-1 or FcaPV-2; and 8 BISCs for

36 FcaPV-1, 2, 4. The AACq detected 3 out of 4 types of FcaPVs. AACq detected

- 37 FcaPV-1 in 1 VP and 3 BISCs, FcaPV-2 in 3 VPs and 6 BISCs, FcaPV-4 in 2
- ³⁸ BISCs. FcaPV-2 was the most prevalent among the group of VPs and BISCs.
- 39 **Conclusion and relevance:** Using $\Delta\Delta$ Cq method we report the first evidence of
- 40 FcaPV-1, 2 and 4 in Italy, FcaPV-2 was the most frequently detected. To a
- 41 lesser extent also FcaPV-1 and FcaPV-4 has been detected in our examined
- samples while FcaPV-3 was never associated to viral induced lesions by $\Delta\Delta Cq$
- 43 investigation. Compared to conventional PCR the $\Delta\Delta$ Cq method has the
- 44 advantage to allow establish a possible role of the virus in the outcome of
- 45 infection.
- 46

47 Introduction

Detection of Feline papillomaviruses (FcaPVs) from feline cutaneous 48 lesions has been reported worldwide^{1,2} and these viruses are considered liable 49 of causing several feline skin conditions.^{1–5} These include a wide spectrum of 50 proliferative skin lesions ranging from non-neoplastic viral plaques (VP) to pre-51 neoplastic Bowenoid in situ carcinoma (BISC), and to sarcoid and invasive 52 squamous cell carcinoma (SSC).⁶ A progression from non neoplastic viral 53 induced lesions to overt neoplasia has also been documented to occur.⁷ Four 54 FcaPV types (FcaPV-1 to FcaPV-4) have been so far described in cats, having 55 different anatomical distribution and presentation.^{1,8–11} The most frequent 56 FcaPV (FcaPV-2, belonging to Dyothetapapillomavirus genus) was isolated 57 from a cutaneous pigmented plaque,¹² and later related to cutaneous SCC.^{13–15} 58 FcaPV-2 seems to play a role in a high proportion of nasal planum squamous 59 cell carcinomas and, animals with tumours associated to papillomaviral 60 aetiology with p16 upregulation, show increased survival compared with those 61 attributable to UV radiation.¹⁶ The relevance of FcaPV-2 in the pathogenesis of 62 both premalignant and malignant lesions has been recently investigated by 63 concurrently assessing number of DNA viral copies and viral gene expression.^{5–} 64 ⁷ However, the effective role of FcaPV-2 in the development of malignant 65 lesions is still a debated question, especially in view of recent results indicating 66

a high FcaPV-2 prevalence in healthy domestic cats.^{7,17} The role of other feline 67 PV types, namely FcaPV-1, 3, and 4, remains obscure. FcaPV-1, belongs to the 68 Lambdapapillomavirus genus and has been rescued from a hyperkeratotic 69 cutaneous lesion in a Persian cat by Tachezy et al.⁹ In the same year another 70 FcaPV was also identified in a shorthair domestic cat with papillomatosis.¹¹ 71 More recently the virus was detected in multiple small sessile raised lesions on 72 the ventral surface of the tongue in two 13-year-old domestic cats.¹⁸ FcaPV-3 is 73 a Taupapillomavirus, isolated for the first time from a cutaneous in situ 74 carcinoma.³ Recently FcaPV-3 has been detected from a BISC with a novel 75 behaviour.¹⁹ same histological feature and a benign clinical The 76 histopathological cell changes were observed in a feline basal cell carcinoma, 77 from which DNA sequences of a novel PV closely related to FcaPV-3 were 78 detected by the authors.⁴ FcaPV-4 belongs to genus *Taupapillomavirus*, and 79 has been detected in oral cavity following a severe gengivitis.¹⁰ Pathogenicity of 80 FcaPV-1 and FcaPV-4 is still unclear and these viruses are rarely detected in 81 oral inflammatory as well as neoplastic lesions. Therefore, an active role of 82 FcaPV-1 and FcaPV-4 in carcinogenesis is still debated. 83

Based on literature, it emerges that most cases of FcaPVs infection have been documented in domestic felids from New Zealand, North and South America and, to a lesser extent from Europe. This might just reflect the active research

http://mc.manuscriptcentral.com/jfms

87	done in these countries without any epidemiological meaning. In Europe PVs
88	infections have been reported in Switzerland in only nine cases, ^{12,20} in one case
89	in Germany ²¹ and in another one case in Italy. ²² Here we assess by a
90	combination of histology, immunohistochemistry and q-PCR the concurrent
91	presence of FcaPV-1, 2, 3 and 4 viral DNA and p16 immunostaining in 29 feline
92	lesions collected in Italy including non neoplastic and preneoplastic, viral and
93	non viral induced skin lesions namely: VP, BISC, invasive SSC and actinic
94	keratosis (AK). Two methods were applied to analyse the viral load data:
95	absolute quantification (AQ) and relative quantification (RQ). The AQ was
96	applied to determine the viral copy number per μ g of DNA; however, by this
97	method wide range of values can be obtained without a cut-off value that would
98	allow to discriminate FcaPVs presence as innocent bystander or as responsible
99	of viral lesion. To address this problem we approached the RQ by $\Delta\Delta$ Cq
100	calculation. ²³ This method measures viral genome presence by relating the
101	obtained values of each of the 4 FcaPVs in each lesion classified as VP or
102	BISC to those of SCC and AK.

103 Materials and methods

- 104 Biopsies
- 105 Feline skin biopsies have been identified from dermatopathological databases

and 29 cases with a diagnosis of VP, BISC, invasive SCC and AK were 106 selected. The retrieved formalin fixed cases were routinely processed for 107 histopathology and 4 µm thick paraffin embedded tissues sections were stained 108 with haematoxylin and eosin (HE). Before inclusion in the study, the selected 109 samples were subjected to critical re-evaluation of diagnosis by defining clear 110 111 distinctive criteria, such as carcinomatous vs non carcinomatous, in situ vs 112 invasive (if carcinomatous), and viral vs actinic. The established criteria were 113 defined as described in the scientific literature and are summarized in table 1.^{14,20,24} Data from signalment and lesion distribution, when available, were 114 included. 115

116 *P16 immunohistochemistry*

Cases were evaluated for the detection of p16 by immunohistochemistry 117 employing an ABC system (Vector Laboratories, Burlingame, USA). Antigen 118 unmasking was carried out at 120°C for 3 min in a pressure cooker with a TRIS-119 EDTA buffer (1,2 g/L Tris; 0,36 g/L EDTA; pH 9.0). Sections were pre-treated 120 for 10 min with 1% H₂O₂ in 0.1 M phosphate buffered saline (PBS), pH 7.4 40 121 min to quench endogenous peroxidase activity, and blocked for 30 min at room 122 123 temperature (rt) in PBS with a 2% normal horse serum (PK-7200, Vector 124 Laboratories, Burlingame, USA) and 0.05% TritonX-100. Sections were then 125 incubated overnight at 4°C with a 1:100 diluted monoclonal primary antibody

126	p16-INK4a (mouse monoclonal IgG, BiorByt, San Francisco, USA). After
127	washing in PBS, an incubation with a secondary universal biotinylated anti-
128	mouse/rabbit antibody (PK-7200, Vector Laboratories, Burlingame, USA) was
129	performed for 1h at rt. Staining was visualized with a diaminobenzidine (SK-
130	4105, Vector Laboratories, Burlingame, USA) solution under a light microscope
131	(Eclipse 80i, Nikon Tokyo, Japan).

132 Sampling material for q-PCR

Three 10 µm thick sections from the selected samples were cut with a microtome, place onto slides and left unfixed. These slides were observed under an optic microscope. Tissue that was not relevant for the study was scraped off and the remaining tissue directly collected in a DNase free 1,5 ml tube. To prevent carryover of contaminating DNA the microtome overlay was covered with a new piece of adhesive tape and a new blade was used for each sample. and the microtome overlay covered with a new piece of adhesive tape.

140 DNA extraction

DNA extraction was performed using DNeasy Blood and Tissue kit (Qiagen,
Hilden, Germany) following manufacturer's instruction and applying a
preliminary removal of paraffin by extraction with xylene. DNAs were eluted in
30 µl, and each sample concentration was quantified using the Qubit Nanodrop

spectrophotometer (ThermoFisher Scientific, Waltham, USA), DNA samples
 were stored at -20°C until analysis.

147 *q-PCR*

The extracted DNAs were amplified using four specific set of primers amplifying 148 a portion of the four types of FcaPVs L1 gene so far identified in the feline 149 species (Table 2). To normalize the amount of DNA used for each sample so to 150 151 achieve a correct quantification of viral copy number, a q-PCR under the same condition as for FcaPV but with a specific set of primers was run in parallel for 152 the reference gene albumin (ALB).²⁵ The number of copy of the target viral 153 gene, measured as Cq value, generates a Δ Cq value when compared to the 154 corresponding Cq value of the reference gene. Moreover, a $\Delta\Delta$ Cq value was 155 calculated comparing the ΔCq value of the sample of interest to the mean ΔCq 156 value obtained from the SCC and AK group, considered as the negative control 157 158 group. The $\Delta\Delta$ Cg value represents a normalized measure of DNA viral quantity and was calculated using REST software.²⁶ In this work, a RQ using the $2^{-\Delta\Delta Cq}$ 159 method ²³ was adapted to estimate in each sample the fold change of FcaPV-1, 160 2, 3 and 4 target viral gene copies relative to the albumin reference gene. 161 Melting curve analysis was performed in conjunction with each four specific 162 FcaPV amplification protocol to determine if non specific products were 163

amplified during reaction. The specificity of the melting curve was compared to 164 melting curves values obtained from a plasmid (pFcaPV) containing amplicon of 165 the four FcaPVs spanning the real-time products. The plasmid was generated 166 by inserting the four FcaPVs gene segments into pMA-T vector using GeneArt 167 technology (Thermo Fisher Scientific, Waltham, USA). Serial dilutions of 168 pFcaPV plasmid ranging from 10⁶ to 10² copies/5µl were used to calculate the 169 efficiency of q-PCR for each FcaPV set of primers and compared to the 170 efficiency calculated on five points of a 2-fold serial dilution of a quantified DNA 171 template for ALB gene. The efficiency of each g-PCR assay was similar and 172 ranging from 91% to 98% (average 94,6%). The lesions identified by 173 histological classification as SCC-AK were presumed to serve as the negative 174 control group and each sample belonging to this group was also individually 175 tested by $\Delta\Delta$ Cq analysis for each FcaPV type. Samples that had eventually 176 scored a positive $\Delta\Delta Cq$ value were excluded. In addition, the viral copy 177 number/µl of input DNA for each FcaPV type in each sample tested was 178 calculated (AQ). The assays were performed in Rotorgene thermocycler 179 (Corbett Research, Sydney, Australia) using SSCO sybr green master mix 180 181 (Biorad, Hercules, USA) and 5 µl of extracted DNA. All samples were tested in duplicate and the results were calculated using the mean Cq values. Samples 182 183 positive for only one replica were considered as negative. All samples with a

positive $\Delta\Delta$ Cq values were submitted to sequence analysis (BMR genomics, Padova, Italy).

186 **Results**

Histologically all VPs were recognized for areas of focal epidermal 187 dysplasia with koilocytes and keratinocytes with cytoplasm enlarged by blue-188 grevish material. Regarding BISC diagnosis, the presence of koilocytes or 189 koilocyte-like cells pointed to the viral origin in 17/20 cases while in the 190 remaining 3 cases diagnosis was formulated by other morphological details 191 listed in Table 1., were used to formulate the diagnosis. SCC and AK showed 192 the morphological alterations listed in Table 1. Among the 20 BISCs cases 5 193 VPs were also detected in the adjacent skin by histology and in other 5, sites 194 195 of penetration of keratinocytes through the basement membrane into the dermis were observed indicative of progression toward SCC. 196

¹⁹⁷Clinical reports for each of the 29 cases fulfilling the established ¹⁹⁸histological criteria were recorded. Breed, age, gender and anatomical ¹⁹⁹distribution are summarised in Table 3. Twenty-five subjects were shorthair ²⁰⁰domestic cats, two persian and two main coons. Mean age at presentation ²⁰¹varied between 6 and 20 years (median 11,6), 14 out of 29 were female and ²⁰²15 were male. A total of 4 VPs, 20 BISCs, 4 SCCs and 1 AK were detected.

http://mc.manuscriptcentral.com/jfms

203 Lesions were observed in densely haired skin regions in 9 cats (dorsum, flank, and shoulder), in areas non-exposed to sunlight in 1 cat (groin) and in 204 hypotrichotic and solar exposed skin in 18 cats (ears, nose, eyelids, temporal 205 region). All cases of SCC were observed in sun-exposed areas. Clinically 206 VPs were few millimeters wide, unique alone or grouped, slightly raised, 207 208 pigmented and non-pruritic lesions; BISCs were clinically larger than VPs, 209 usually more than 1 cm of diameter, multifocally coalescing, raised and often 210 verrucous, crusted and hyperpigmented. In SCC, erosive and ulcerative crusted lesions were clinically detected. In the only case in which AK was 211 diagnosed, erythema, scales and crust were seen (Figure 1a, b, c). 212

P16 was immunolocalized in all VPs epidermis and in the epidermis and follicular wall of the BISCs. No signal or only faint staining was detected in the SCC and AK lesion. Immunoreactivity was either nuclear and cytoplasmic; in VPs and BISCs the signal was strong while in the SCCs immunoreactivity was faint and therefore considered as no significant (Figure 1d).

²¹⁹ By q-PCR we were able to detect the albumin reference gene in 28 out of 29 ²²⁰ DNA samples. Only one sample classified as BISC was not suitable to ²²¹ molecular analysis and therefore it was excluded from further analysis.

12

http://mc.manuscriptcentral.com/jfms

222	By AQ, the presence of FcaPVs has been detected in low copies in almost all
223	cases (Table 4). By RQ ($\Delta\Delta$ Cq method) a reduced number of samples was
224	linked to viral lesions. In details, FcaPVs positive $\Delta\Delta$ Cq values were detected
225	in all VPs (4/4) and in 8/19 BISCs (Table 3). VPs were positive to FcaPV-2 in
226	three samples and to FcaPV-1 in one sample. BISCs were positive to FcaPV-
227	2 in three samples, FcaPV-4 and FcaPV-1 in one sample each. Interestingly,
228	as never reported before, in BISCs lesion we found three cases of double
229	infection. In two of those, FcaPV-1 and 2 were detected simultaneously and
230	in one case FcaPV-2 was present along with FcaPV-4 (table 3).
231	The presence of FcaPVs has been detected in low copies in almost all cases
232	(Table 4). By ∆∆Cq analysis a reduced number of cases was identified:
233	FcaPVs positive AACq value were detected in all VPs (4/4) and in 8/19
234	BISCs (table 3). The PV type most frequently detected was FcaPV-2 that
235	was the unique PV in 3 out of 4 VPs and in 3 BISCs; in 2 BISCs FcaPV-2
236	was associated to FcaPV-1, and in 1 case FcaPV-2 was associated to
237	FcaPV-4. FcaPV-1 was the other detected unique PV type in 1 VP and 1
238	BISC, while FcaPV-4 was the unique PV type in only 1 BISC (Table 3).

In all cases with a positive $\Delta\Delta Cq$ value, the viral load was always greater 239 than 10² copies /µg DNA (FcaVP-1: min 167 – max 310 Standard Error (SE): 240

33,0; FcaPV-2: min 100 – max 1150 SE: 163,9; FcaPV-4 min 391 – max 241 3700 SE: 1654,5) (Table 4). Sequence analysis confirmed the FcaPV types 242 as indicated by q-PCR AACq analysis. Number of AACq FcaPV1-4 positive 243 samples for each type of lesion are summarized and showed in 244 supplementary material. 245 246 Discussion In this retrospective study, we detected the concurrent presence, we 247 investigated the presence of different types of PVs, namely of FcaPV-1, 2, 248 and 4 in feline VPs and BISCs by $\Delta\Delta$ Cg method using specific primers. To 249 the best of our knowledge only the concurrent presence of FcaPV-2 and 3 250 have been reported from one BISC and one VP.^{3,22} 251 . The data obtained on concurrent presence of more than one PV type 252 253 is not unexpected since multiple PVs infection has been previously detected

²⁵⁴ in single skin samples from cats; FcaPV-2 and FcaPV-3 have been ²⁵⁵ concurrently detected in a feline BISC by Munday at al and from a viral ²⁵⁶ plaque by Alberti et al.</sup> However differently from previous reports, here we ²⁵⁷ report the presence of FcaPV-1 and FcaPV-4, alone or in association to ²⁵⁸ FcaPV-2.

A limitation when performing PCR-based studies on PVs infection is 259 that the viral DNA is often detected and cannot be directly linked to an 260 effective role of virus in pathogenesis and therefore it is not possible to fully 261 establish whether the presence of a viral genome is uneventful (subclinical 262 lesion). In particular, FcaPV-2 has been recently detected in a number of skin 263 264 swabs from healthy cats, making difficult to discern whether PVs are causing cancer or are merely an "innocent bystander".⁷ The hypothesis that the 265 presence of high viral loads likely represents an infectious state of PV is 266 supported by Thomson et al.⁷ who recently showed that the finding of high 267 copy numbers of FcaPV-2 DNA within a lesion suggests that the detected 268 virus may be responsible for it. In fact, while high viral copy numbers were 269 associated with E6/E7 gene expression, no gene expression was detected in 270 association of low copy numbers, an indication of an incidental finding. 271

For the above-mentioned reasons, for this study we used a q-PCR protocol applying a $\Delta\Delta$ Cq method to investigate if and which FcaPVs have induced the lesion in cats. In this sense the obtained results are partially in line with the scientific literature which indicates so far that FcaPV-2 is the major PV type implicated in skin preneoplastic and neoplastic lesions in cats.⁷ Since the genome organization and the role of viral proteins within the replication cycle are considered similar even between different genera

15

279	belonging to the Papillomaviridae family we assumed that a similar link is
280	maintained for feline PVs other than FcaPV-2. Importantly we reported
281	positive $\Delta\Delta$ Cq values for FcaPV-1 and 4; two FcaPV types rarely reported. ^{9–}
282	^{11,18} It is noteworthy the observation that also FcaPV-1 and FcaPV-4 showed
283	positive ∆∆Cq values however no data are available on these two types for
284	comparison.
285	A comparison between our data with those obtained from other studies
286	is difficult for two reasons: i) type of PCR primers used (specific vs
287	consensus); ii) type of PCR used (conventional vs quantitative). A general
288	prevalence ranging from 24 to 100% of FcaPV-2 both BISCs and VPs has
289	been reported. ^{7,8,27,28} The rate of PV detection by ∆∆Cq analysis was in
290	agreement with previous studies which reported a prevalence ranging from
291	<mark>24 to 61% . Moreover, using ⊿∆Cq might have lowered the prevalence. In</mark>
292	our study, by using specific primers in a q-PCR analysis applying RQ
293	method, we obtained a prevalence for FcaPV-2 in BISCs of 31,6% (6/19) and
294	for all FcaPV types of 42,10% (8/19).
295	Failure to demonstrate PV DNA in every BISC have been previously explained

- through a carcinogenesis model in which papillomaviruses cause transformation
- ²⁹⁷ but are only for a short period present within the lesion. ²⁹ Surprisingly, cases in

which koilocytes were detected did not show any $\Delta\Delta$ Cq positive value; this

result is likely due to: i) the detection of koilocytes in very focal lesions that

300 could have been no more present when additional serial paraffin sections were

- ³⁰¹ prepared for DNA extraction and PCR analysis; ii) the presence of a PCR
- 302 undetectable PV variant.

Low viral copies for all PV types have been detected in almost all cases, 303 however the $\Delta\Delta$ Cq analysis was necessary to associate the FcaPVs as 304 305 biological agent potentially causing the lesions. $\Delta\Delta Cq$ positive values for the different PV types were detected in all VPs and in 42% of BISCs. Low viral 306 307 copies number was also detected from the SCC-AK group resulting in a not detectable $\Delta \Delta Cq$. Noteworthy, mean $\Delta \Delta Cq$ of positive samples differed 308 among PV types being 62.5, 189.1 and 3083.5 for FcaPV-1, FcaPV-2 and 309 FcaPV-4 respectively. Studies considering a larger panel of cases might be 310 necessary to establish a further $\Delta\Delta$ Cg cut off value. 311

Here we document the presence of FcaPV-2 with high prevalence in both VPs and BISCs while we rescued FcaPV-1 and FcaPV-4 less frequently; these last two PVs are rarely described as playing an active role in skin and oral mucosa lesions in cats.^{9,10,18,30} None of the lesions was associated to FcaPV-3 when adopting $\Delta\Delta$ Cq method, confirming the rare occurrence of this

317 FcaPV type also in Italy.

Either cytoplasmic or nuclear p16 signals were detected in all cases where 318 cytopathic effects were found (VP, BISC) while p16 immunolocalisation was 319 present with faint and mainly cytoplasmic signal in cases of SCCs and no p16-320 immunoreactivity was found in the case of AK. Despite the difficulties of 321 322 interpretation when the signal was faint, these results are in line with what already documented previously. About half of the SCC studied by Thomson et 323 al.⁷ were negative to p16 immunostaining. A strong association between 324 FcaPV-2 E6/E7 gene expression and p16 immunostaining have been recently 325 found in feline SCC with 18 of 20 (90%) E6/E7-positive SCCs being also 326 positive for p16 compared to 13 of 40 (33%) E6/E7-negative SCCs.⁷ E6/7 gene 327 expression investigation has not been carried out in our study but our results 328 might reflect this correlation. 329

330 Conclusion

 $\Delta\Delta$ Cq analysis has proved to be necessary to associate the FcaPVs as biological agent potentially causing the lesions. Based on this method the presence of FcaPV-2 is confirmed to be the most representative FcaPV in feline skin lesions referable to diagnosis of VP and BISC in Italy. To a lesser 335 extent also FcaPV-1 and FcaPV-4 has been detected in our examined

336 samples. while FcaPV3 was never detected.

337

- Acknowledgement: The authors thank all the colleagues who referred the cases.
- 340 **Conflict of interest**: The authors declared no potential conflicts of interest with
- respect to the research, authorship, and/or publication of this article
- 342 **Source of funding**: This study was financially supported by Fondi di Ateneo
- 343 University of Pisa.

344 **Bibliografy**

- Sundberg JP, Van Ranst M, Montali R, et al. Feline papillomas and
 papillomaviruses. Vet Pathol 2000; 37: 1–10.
- Gil da Costa RM, Peleteiro MC, Pires MA, et al. An Update on Canine,
 Feline and Bovine Papillomaviruses. Transboundary and Emerging
 Diseases, 11 September 2016. Epub ahead of print 11 September 2016.
 DOI: 10.1111/tbed.12555.
- Munday JS, Dunowska M, Hills SF, et al. Genomic characterization of
 Felis catus papillomavirus-3: a novel papillomavirus detected in a feline
 Bowenoid in situ carcinoma. Vet Microbiol 2013; 165: 319–25.
- Munday JS, French A, Thomson N. Detection of DNA sequences from a novel papillomavirus in a feline basal cell carcinoma. Vet Dermatol 2017; 28: 236-e60.

- 5. Munday JS, Gwyther S, Thomson NA, et al. Bilateral pre-auricular papillary squamous cell carcinomas associated with papillomavirus infection in a domestic cat. Vet Dermatol 2017; 28: 232-e58.
- Scott DW, Miller WH, Griffin CE, et al. Muller & Kirk's Small Animal
 Dermatology. Epub ahead of print 2001. DOI: 10.1016/B978-0-72167618-0.50002-X.
- Thomson NA, Munday JS, Dittmer KE. Frequent detection of
 transcriptionally active Felis catus papillomavirus 2 in feline cutaneous
 squamous cell carcinomas. J Gen Virol 2016; 97: 1189–97.
- Munday JS, Kiupel M, French AF, et al. Detection of papillomaviral sequences in feline Bowenoid in situ carcinoma using consensus primers. Vet Dermatol 2007; 18: 241–5.
- 369 9. Tachezy R, Duson G, Rector A, et al. Cloning and genomic
 370 characterization of Felis domesticus papillomavirus type 1. Virology 2002;
 371 301: 313–21.
- Dunowska M, Munday JS, Laurie RE, et al. Genomic characterisation of
 Felis catus papillomavirus 4, a novel papillomavirus detected in the oral
 cavity of a domestic cat. Virus Genes 2014; 48: 111–119.
- Terai M, Burk RD. Felis domesticus papillomavirus, isolated from a skin
 lesion, is related to canine oral papillomavirus and contains a 1.3 kb noncoding region between the E2 and L2 open reading frames. J Gen Virol
 2002; 83: 2303–7.
- Lange CE, Tobler K, Markau T, et al. Sequence and classification of
 FdPV2, a papillomavirus isolated from feline Bowenoid in situ carcinomas.
 Vet Microbiol 2009; 137: 60–65.
- Ravens PA, Vogelnest LJ, Tong LJ, et al. Papillomavirus-associated
 multicentric squamous cell carcinoma in situ in a cat: An unusually
 extensive and progressive case with subsequent metastasis. Vet
 Dermatol 2013; 24: 642-e162.
- 14. Munday JS. Papillomaviruses in felids. Vet J 2014; 199: 340–347.

- Altamura G, Corteggio A, Pacini L, et al. Transforming properties of Felis
 catus papillomavirus type 2 E6 and E7 putative oncogenes in vitro and
 their transcriptional activity in feline squamous cell carcinoma in vivo.
 Virology 2016; 496: 1–8.
- Munday JS, Gibson I, French AF. Papillomaviral DNA and increased
 p16CDKN2A protein are frequently present within feline cutaneous
 squamous cell carcinomas in ultraviolet-protected skin. Vet Dermatol
 2011; 22: 360–366.
- 395 17. Geisseler M, Lange CE, Favrot C, et al. Geno- and seroprevalence of
 396 Felis domesticus Papillomavirus type 2 (FdPV2) in dermatologically
 397 healthy cats. BMC Vet Res 2016; 12: 147.
- Munday JS, Fairley RA, Mills H, et al. Oral Papillomas Associated With
 Felis catus Papillomavirus Type 1 in 2 Domestic Cats. Vet Pathol 2015;
 52: 1187–1190.
- 401 19. Munday JS, Fairley R, Atkinson K. The detection of Felis catus
 402 papillomavirus 3 DNA in a feline bowenoid in situ carcinoma with novel
 403 histologic features and benign clinical behavior. J Vet Diagnostic Investig
 404 2016; 28: 612–615.
- Wilhelm S, Degorce-Rubiales F, Godson D, et al. Clinical, histological and
 immunohistochemical study of feline viral plaques and bowenoid in situ
 carcinomas. Vet Dermatol 2006; 17: 424–431.
- Egberink HF, Berrocal A, Bax HA, et al. Papillomavirus associated skin
 lesions in a cat seropositive for feline immunodeficiency virus. Vet
 Microbiol 1992; 31: 117–25.
- Alberti A, Tore G, Scagliarini A, et al. Regressing Multiple Viral Plaques
 and Skin Fragility Syndrome in a Cat Coinfected with FcaPV2 and
 FcaPV3. Case Reports Vet Med 2015; 2015: 1–5.
- 414 23. Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the 415 comparative C T method. 2008; 3: 1101–1108.
- 416 24. Levison DA, Parratt D. Skin Diseases of the Dog and Cat. Oxford, UK:

417 Blackwell Science Ltd. Epub ahead of print 20 July 2005. DOI: 418 10.1002/9780470752487.

- 419 25. Helfer-Hungerbuehler AK, Widmer S, Hofmann-Lehmann R. GAPDH
 420 Pseudogenes and the Quantification of Feline Genomic DNA Equivalents.
 421 Mol Biol Int 2013; 2013: 587680.
- 422 26. Pfaffl MW, Horgan GW, Dempfle L. Relative expression software tool
 423 (REST) for group-wise comparison and statistical analysis of relative
 424 expression results in real-time PCR. Nucleic Acids Res 2002; 30: e36.
- A25 27. Nespeca G, Grest P, Rosenkrantz WS, et al. Detection of novel papillomaviruslike sequences in paraffin-embedded specimens of invasive and in situ squamous cell carcinomas from cats. Am J Vet Res 2006; 67: 2036–2041.
- 429 28. Munday JS, Peters-Kennedy J. Consistent detection of Felis domesticus
 430 papillomavirus 2 DNA sequences within feline viral plaques. J Vet Diagn
 431 Invest 2010; 22: 946–9.
- Smith KT, Campq MS. 'Hit and run' transformation of mouse C127 cells
 by bovine papillomavirus type 4: the viral DNA is required for the initiation
 but not for maintenance of the transformed phenotype. Virology 1988;
 164: 39–47.
- Munday JS, French AF, Gibson IR, et al. The presence of p16 CDKN2A
 protein immunostaining within feline nasal planum squamous cell
 carcinomas is associated with an increased survival time and the
 presence of papillomaviral DNA. Vet Pathol 2013; 50: 269–73.
- 440
- 441
- 442 LEGEND
- 443 Figure 1. Multiple symmetrical raised, hyperpigmented and crusted plaques
- on the face of a cat with Bowenoid in situ carcinoma (a). Histopathology from

a viral plaque in a cat. Focal epidermal hyperplasia, absence of follicular wall 445 involvement, evident viral cytopathic effects (koilocytes) (bar=50µm) (b); 446 Histopathology from a Bowenoid in situ carcinoma in a cat. Epidermal and 447 follicular dysplasia with upward in situ keratinocyte proliferation and a few 448 (bar=50µm) (c); koilocytes p16 immunohistochemistry, groups of 449 ι cytor keratinocytes show nuclear and cytoplasmic positivity (ABC system, bar=100 450 451 µm) (d).

452

Table 1. Histological criteria for the diagnosis of VP VP, BISC, SCC and AK.

Type of lesion	Histopathological criteria
VP	Focal epidermal hyperplasia, koilocytes, clear cells, giant keratoyalin granules, cytoplasm enlarged by blue-greish fibrillar material, hyperpigmentation;
BISC	Epidermal and follicular dysplasia with upward in situ keratinocyte proliferation, loss of nuclear polarity, windblown nuclei, koilocytes, mitosis;
SCC	Epidermal dysplasia and atypia with downword keratinocyte proliferation, keratonocyte invasion to the dermis through the basal lamina, mitosis;
AK	Epidermal dysplasia and atipia with upward in situ keratinocyte proliferation, squamotisation, apoptosis

Legend: VP viral plaque; BISC Bowenoid *in situ* carcinoma; SCC squamous cell carcinoma; AK actinic keratosis

Table 2. Primers nucleotide sequences

Oligonucleotide name	Nucleotide sequence (5'-3')	5' Nucleotide position	Reference sequence accession number	
FcaPV1-F	AGGATGGTGACATGGTGGAT	7143	AE480454	
FcaPV1-R	TTTGCACTGTGTGTCTGCAA	7246	AF480454	
FcaPV2-F	TACACGCGGTACCAATTTCA	7191	EU796884	
FcaPV2-R	AGAGTGACCACGCACACTTG	7331	EU/90004	
FcaPV3-F	AAGATTGGTATGGCGTTTGC	5960	JX972168	
FcaPV3-R	TTTGCCTTTCATCTGCTGTG	6105	JX972100	
FcaPV4-F	ATGCAAATGGCCAGACTTTC	6356	KF147892	
FcaPV4-R	AAAAATGGCGGCAGTACAAC	6453	KF 147892	
Fel Alb F	GATGGCTGATTGCTGTGAGA	3706	NC_018726.2	
Fel Alb-R	CCCAGGAACCTCTGTTCATT	3855	GPC_000001738	

1

Table 3.Signalment, lesion distribution, p16 immunolocalisation and FcaPV type assessed by $\Delta\Delta$ Cq analysis in cats with of VP, BISC, SCC and AK diagnosis								
Case N°	Diagnosis	Breed	Years	Gender	Lesion distribution	p16	FcaPV type	2 ^{–(ΔΔCq)}
1	VP	DSH	10	М	Flank	+	2	119
2	VP	persian	15	F	Groin	+	2	53
3	VP	DSH	9	М	Nose	+	1	72
4	VP	main coon	10	F	Nose	+	2	271
5	BISC	DSH	10	F	Flank	+	-	
6	BISC	main coon	6	М	Shoulder	+	4	5836
7	BISC	DSH	13	М	Nose	+	-	
8	BISC	DSH	13	F	Nose	+	1	49
9	BISC	DSH	9	F	Temporal	+	-	
10	BISC	DSH	20	F	Dorsal	+	ND	
11	BISC	DSH	15	М	Temporal	+	2-4	224-331
12	BISC	DSH	15	М	Nose	+	-	
13	BISC	DSH	16	F	Shoulder	+	-	
14	BISC	DSH	12	М	Ear	+	-	
15	BISC	DSH	12	М	Dorsum	+	2	172
16	BISC	DSH	8	F	Temporal	+	1-2	58-167
17	BISC	DSH	10	F	Temporal	+	-	
18	BISC	persian	15	М	Eyelid	+	2	211
19	BISC	DSH	10	F	Ear	+	-	
20	BISC	DSH	7	F	Flank	+	-	
21	BISC	DSH	10	М	Nose	+	2	159
22	BISC	DSH	15	М	Shoulder	+	-	
23	BISC	DSH	12	М	Nose	+	1-2	71-326
24	BISC	DSH	10	М	Flank	+	-	
25	SCC	DSH	9	F	Ear	+-	-	
26	SCC	DSH	8	М	Nose	+-		
27	SCC	DSH	15	F	Ear	+-	-	
28	SCC	DSH	10	М	Eyelid	+-	-	
29	AK	DSH	11	F	Ear	+-	-	

2

3 Legend: VP viral plaque, BISC Bowenoid *in situ* carcinoma, SCC squamous cell carcinoma, AK actinic

4 keratosis, DSH domestic shorthair, ND non determined.

- 5 $2^{-(\Delta\Delta Cq)}$ indicates the fold expression change of FcaPV normalised to Albumin.
- 6

Table 4. Absolute viral load quantification expressed	as viral copy number/µg of input
DNA	

Cases	Type of lesion	FcaPV-1	FcaPV-2	FcaPV-3	FcaPV-4
1	VP	10	<u>153</u>	10	20
2	VP		<u>100</u>		6
3	VP	<u>167</u>	33	23	28
4	VP		<u>1130</u>		
5	BISC	30	48		
6	BISC	18	55	26	<u>3700</u>
7	BISC	13	2		4
8	BISC	<u>201</u>			
9	BISC	42	12		
10	BISC		15		
11	BISC	87	<u>1150</u>	59	<u>391</u>
12	BISC		10		
13	BISC	33	55	28	120
14	BISC				
15	BISC	14	<u>436</u>		
16	BISC	<u>176</u>	<u>210</u>	13	
17	BISC		18		
18	BISC		<u>1120</u>	79	45
19	BISC	3			
20	BISC	33	10		
21	BISC	6	<u>193</u>		
22	BISC				
23	BISC	<u>310</u>	<u>1140</u>	18	26
24	BISC				
25	SSC		6	2	
26	SSC	12	10		17
27	SSC	20	15	7	23
28	SSC	10	5	14	14
29	AK	12	18		
Mean v	iral load	63	247.7	<mark>2.4</mark>	366.2
	iral load of ΔΔCq samples	213.5	625.8		<mark>2045.5</mark>
Standard error of $\Delta\Delta Cq$ positive samples		33	163.9		1654.5

Bold and underlined numbers indicate $\Delta\Delta$ Cq positive samples.

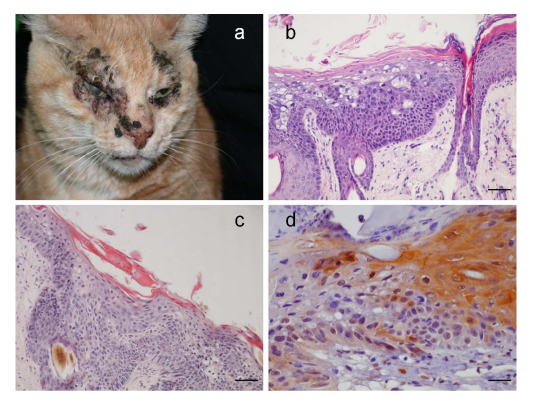


Figure 1

136x103mm (300 x 300 DPI)

