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Source: Journal of Feline Medicine and Surgery Open Reports, 8(2)

Published By: SAGE Publishing

URL: https://doi.org/10.1177/20551169221137536

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Case Report





CTSK variant implicated in suspected pyknodysostosis in a domestic cat

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Journal of Feline Medicine and Surgery Open Reports

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This paper was handled and processed by the European Editorial Office (ISFM) for publication in JFMS Open Reports



Abstract

Case summary A 9-month-old entire male domestic longhair cat presented with a history of pathological fractures, chronic musculoskeletal pain and poor growth. Multiple facial and skeletal abnormalities were identified on physical examination and advanced imaging (CT and radiographs). A variant in CTSK was identified in the affected cat following whole-exome sequencing (WES). The cat was managed symptomatically with diet, environmental modifications and analgesia.

Relevance and novel information This is the first report of a cat with a similar clinical presentation and genetic variant to the hereditary human genetic disorder pyknodysostosis. In this case, WES was performed, which often facilitates the diagnosis of various hereditary disorders (ie, a conceptual framework for practicing feline genomic medicine). Despite the severe skeletal and appendicular abnormalities described, the cat was alive more than 2 years after its initial presentation.

Keywords: Whole-exome sequencing; hereditary; pycnodysostosis; pyknodysostosis; cathepsin K

Accepted: 20 October 2022

Case description

A 9-month-old entire male domestic longhair cat was referred with a lifelong history of poor growth, dysphagia, lameness and pathological fractures, and a 3-month history of stertor and bilateral serous ocular discharge. The cat had three episodes of acute pain and lameness after jumping from furniture. A left radioulnar fracture was suspected clinically following the first episode, while a complete right radioulnar fracture was confirmed radiographically following the second episode. The third episode occurred 2 weeks prior to presentation but was not investigated. All episodes were managed conservatively with cage rest and analgesia with meloxicam (0.05 mg/kg PO q24h [Metacam: Boehringer Ingelheim]) and buprenorphine hydrochloride (0.02 mg/kg transmucosally q12h [Temgesic; Indivior UK]). The cat was up to date with vaccinations, including feline leukaemia virus (FeLV), was indoor-only, had no travel history outside of the UK and had been fed a home-cooked, fishbased liquid diet due to the dysphagia reported. Prior

to referral, measurement of serum thyroxine (50.8 mmol/l; reference interval [RI] 10-60) and insulin-like growth factor 1 (IGF-1; 747 ng/ml; RI >50) excluded congenital hypothyroidism and juvenile hyposomatotropism, respectively.

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Figure 1 Cat with suspected pyknodysostosis and cathepsin K mutation. (a) Front and (b) side views show the face is asymmetrical with distortion of the nasal bone and brachycephaly, hypertelorism and shallow orbits

On presentation, the cat was bright and alert but was lean (body condition score [BCS] 3/9),¹ with poor muscle development, a kyphotic posture, lameness affecting all four limbs and an asymmetrical face with distortion of the nasal bone and hypertelorism (wide spaced eyes) and shallow orbits (Figure 1). Pain was elicitable in the thoracolumbar area and on hindlimb manipulation. Patellar luxation was present in the right hindlimb (grade IV).2 Oral examination revealed multiple persistent deciduous teeth and a cleft palate affecting only the soft palate. Complete ophthalmological examination revealed megalocornea and excessively deep anterior chambers bilaterally but was otherwise unremarkable. Further investigations were performed under general anaesthesia to limit restraint of the cat and the risk of further pathological fractures.

A complete blood count at Langford Vets Diagnostic Laboratories (LVDL) documented abnormalities likely associated with the anaesthesia, including a normocytic, normochromic non-regenerative anaemia (24.4%; RI 27.7–46.8), an absolute reticulocyte count of $0 \times 10^{12}/1$ (RI <50)³ and a mature neutropenia ($2.32 \times 10^9/1$; RI 3–13.4). Serum biochemistry (LVDL) revealed increased alanine aminotransferase (ALT) activity (66 U/1; RI 15–45), alkaline phosphatase (ALP) activity (273 U/1; RI 15–60) and hyperphosphataemia (2.16mmol/1; RI 0.95–1.55). Ionised calcium (1.22mmol/1; RI 1.1–1.5) and serum total calcium (2.36 mmol/1; RI 2.3–2.5) were within

normal limits. Serology for FeLV antigen, feline immunodeficiency virus antibodies in blood and feline herpesvirus-1 and feline calicivirus PCR from oropharyngeal samples were all negative. Serum 1,25-dihydroxyvitamin D (248 pmol/l; RI 90–342) and 25-hydroxycholecalciferol (155 nmol/l; RI 127–335) were normal, ruling out hypovitaminosis D. Radiographs of all four limbs revealed multiple severe skeletal abnormalities (Figures 2 and 3), including curvature of the long bones, short diaphyses and angular epiphyses of the metacarpal bones, delayed growth plate closure, right lateral patellar luxation and healed historical fracture sites. CT of the head and spine revealed persistently open skull suture lines and nasal cavity deviation (Figures 4 and 5), as well as retention of multiple deciduous teeth.

The cat was discharged with a plan to gradually transition it to a complete, highly digestible commercial cat food (Feline EN Gastrointestinal; Purina Pro Plan Veterinary Diets) mixed with water to facilitate mastication. Environmental modifications were applied to minimise injury risk, such as keeping the cat indoors and using low-seated furniture. Furthermore, lifelong eye lubrication was prescribed to reduce the risk of corneal injury (q12h [VitaPOS; Ursapharm]), alongside an analgesia plan, titrated based on the Feline Grimace Scale,⁴ consisting of meloxicam (0.05 mg/kg PO q24h [Metacam; Boehringer Ingelheim]) and gabapentin (5 mg/kg PO q12h [Gabapentin; Summit]). A dentistry referral for



Figure 2 Radiographs of the forelimbs of a cat with suspected pyknodysostosis and cathepsin K mutation. (a) Left and (b) right forelimb antebrachial images show mild cranial and moderate lateral curvature and thickening of the mid-diaphysis of the radius and ulna with a reduction in the normal medullary cavity. The distal radius and ulna physes appear open. The metacarpal bones and phalanges appear abnormal with very short diaphyses, more angular and wide flared epiphyses. (c,d) The humeri are markedly abnormal in shape with severe caudal curvature of the entire diaphysis, with increased medullary opacity similar to the cortical opacity

management of the deciduous teeth was offered but declined by the owners.

The cat was re-presented 1 month later for a scheduled recheck and was reported to have been well in the interim. No further episodes of pain or lameness were reported, and transition to the commercial cat food had been successful. The cat had gained 150 g in weight, had an improved BCS (4/9) and appeared comfortable on limb manipulation. The remainder of the physical examination was unchanged. Repeat complete blood count



Figure 3 Hindlimb radiographs of a cat with suspected pyknodysostosis and cathepsin K mutation. (a) The left and (b) right hindlimbs have severe curvature in the distal diaphysis of the left tibia and fibula and moderate curvature in the distal diaphysis of the right tibia and fibula. The femoral diaphyses are abnormally thin. The right patella is positioned distally and is superimposed over the distal femur compatible with grade IV patella luxation

(LVDL) documented borderline leukocytosis $(20 \times 10^9/l;$ RI 6.3–19.6) and associated lymphocytosis $(9.6 \times 10^9/l;$ RI 2–7.2). Serum biochemistry (LVDL) was similar, with increased ALT activity (60U/l; RI 15–45), ALP activity (240 U/l; RI 15–60), hyperphosphataemia (2.29 mmol/; RI 0.95–1.55) and normal total calcium (2.51 mmol/l; RI 2.3–2.5). On the most recent communication with the owners, 2 years after the initial presentation, the cat was reported to be doing well, with no further episodes of fractures.

The cat's clinical picture, appearance and imaging abnormalities were clinically similar to pyknodysostosis, a rare human disease caused by variations in CTSK. Cat DNA samples were donated by the owner and archived in accordance with LVDL protocol and the University of Missouri Institutional Animal Care and Use Committee protocol (study protocols 9056, 9178 and 9642). DNA was isolated from the whole blood sample by organic extraction, and approximately 2.9µg was submitted to whole-exome sequencing (WES), which was performed as previously described.⁵ Felis catus 9.0 reference alignment⁶ and Ensembl 94 gene annotation were used, as previously reported.5 The variant call file was combined with variants determined from 39 additional cats with WES data. Variant data were analysed using VarSeq software (GoldenHelix) by assuming the affected 4

Figure 4 CT of a cat with suspected pyknodysostosis and cathepsin K mutation. (a) Dorsal plane of the cat's skull shows marked deviation of the nasal septum. (b) Transverse plane of the cat's skull shows a large defect at the level of the fontanelle, between the frontal and parietal bones

cat was homozygous for a causal variant and no other cat in the data set had the variant present. The cat WES data set contained over 1 million variants; 109,688 were homozygous in the case, of which 2344 were private variants. Thirty-six high-impact variants altering protein coding, including a stop gain (termination codon) variant in a candidate gene (CTSK), were identified. The CTSK variant at position C1:105945826 in the Felis catus cat genome assembly Version 9.06 is a c.724C>T in transcript ENSFCAT00000003643, causing a p.Arg242Ter (see Figure 1 in the supplementary material). The depth of read for the variant was high (n = 158). The candidate variant was validated by direct Sanger sequencing conducted at the University of Missouri DNA Core Facility using an Applied Biosystems 3730xl DNA Analyzer with a BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems). DNA sequences were visualised with Sequencer (GeneCodes).

The identified variant was also absent in the 99 Lives data set that includes whole-genome sequencing (WGS)

data from 195 cats.⁶ All of the sequence variants unique to the affected cat are included in the supplementary file, as well as all variants identified in CTSK in the data set.⁷ All CTSK variants following the Human Genome Variation Society nomenclature are summarised in Table 1. Exome sequencing data are available at the Sequence Read Archive under accession number PRJNA627536. The sequence for the 195 cat analysis of the 99 Lives Cat Genome Sequencing Project was submitted to the NCBI short-read archive under BioProject PRJNA627536; SAMN14675162; SRS6521458; with the accession for this case as SRX8159688.8 Protein-coding alignment of CTSK was produced using Protein Variation Effect Analyzer (PROVEAN)9,10 and cat CTSK has 93% identity to the human CTSK transcript (see Figure 2 in the supplementary material).

Discussion

In humans, pyknodysostosis is a rare autosomal recessive bone disorder (OMIM: 265800) caused by a variation in *CTSK*.¹¹ The disease is characterised by multiple musculoskeletal abnormalities, including short stature, osteosclerosis with increased bone fragility, facial dysmorphism, persistent deciduous teeth and large fontanels with delayed closure of sutures.¹² To our knowledge, this is the first report of a cat with very similar skeletal abnormalities and concurrent associated *CTSK* variation identified in genome sequencing.

Pyknodysostosis is a severely debilitating disease in humans, causing a variety of complications, including pathological fractures, craniosynostosis and respiratory, sleep and dental problems.¹³ Patients with a more severe phenotype may die at a very young age.¹⁴ The cat in this case report suffered from similar complications, including pathological fractures, chronic musculoskeletal and oral pain. The severe conformational abnormalities reported likely contributed to the additional ocular and upper respiratory signs seen in this cat. In addition, the serum biochemistry revealed increases in ALP and phosphate that may have been age related or due to the increased osteoblastic activity associated with healing fractures and the unbalanced home-cooked diet.

Based on the human literature, no specific treatment is available for pyknodysostosis and affected patients are managed symptomatically.¹³ In this case, the cat was managed with long-term analgesia and environmental modifications, to prevent further injury, and diet modifications, to combat dysphagia. Regular dental care and deciduous tooth extractions were recommended, but the owner declined these due to the risk of mandibular fracture and the post-extraction osteomyelitis reported in humans with the disease.¹⁵ Regular recheck appointments were scheduled to ensure that the cat's quality of life was maintained and to adjust treatment as necessary. Euthanasia should be considered in cases where chronic pain cannot be addressed effectively.





Figure 5 Three-dimensional CT reconstruction of the skull of a cat with suspected pyknodysostosis and cathepsin K mutation. (a) Dorsal plane of the cat's skull shows a large defect at the level of the fontanelle between the frontal and parietal bones (arrow). (b) Dorsal plane of the cat's skull shows a large defect at the level of the fontanelle (arrow) and marked deviation of the nasal septum (star)

IGF-1 was found to be low in eight paediatric patients with pyknodysostosis and short stature vs normal agematched children with a constitutional short stature, likely owing to concurrent pituitary hypoplasia.¹⁶ In the same study, IGF-1 levels and the children's growth clinically improved significantly after treatment with growth hormone. In contrast, this cat had normal IGF-1 levels and therefore juvenile hyposomatotropism was considered unlikely, although a growth hormone stimulation test and MRI of the brain were not performed.

Cathepsin K (OMIM: 601105) is a member of the papain family of cysteine proteinases and has an important role in osteoclast function.¹⁷ Approximately 70 CTSK variants described in ClinVar are implicated in human pyknodysostosis, of which 41 are considered pathogenic, or likely pathogenic (Table 1).¹⁸ A knockout model for CTSK in mice has many of the phenotypic features of pyknodysostosis, including increased bone density and bone deformity.¹⁹ In an American Hispanic patient with pyknodysostosis and non-consanguineous parents, heteroallelism for the G146R mutation (NM_000396.4[CTSK]:c.436G>C [p.Gly146Arg]) and a C-to-T transition of a CpG dinucleotide at nucleotide 826 of the cDNA sequence was found (NM_000396.4[CTSK]:c.721C>T[p.Arg241Ter]), predicting an arginine at protein position 241 change to a termination codon (R241X) nonsense mutation, which is likely the same position as identified in the cat (see Figure 2 in the supplementary material).¹⁷

The long-read assembly of the domestic cat genome⁶ has been used to identify DNA sequence variants that might be associated with pyknodysostosis in the cat in this report.^{5,20} As with genome approaches in human medicine, the success of the WES approach for identifying disease-associated sequence variants is highly dependent on the heritability and frequency of the disease. In this case, WES was conducted on a solitary cat. The collaborative efforts of the cat genetics community have produced the WGS 99 Lives data set that complements exome data evaluations and has also facilitated identification of sequence variants associated with other feline hereditary disorders, supporting precision/ genome medicine healthcare for domestic cats.^{21,22} At the time of writing this case report, both genome approaches are commercially available. Currently, WGS costs around US\$800-\$1000, while WES costs substantially less (around US\$200-\$300).

A few limitations apply to this case report. First, other congenital bone disorders reported in humans were not excluded in this case, including cleidocranial dysostosis, acro-osteolysis (also known as Hajdu–Cheney syndrome), osteogenesis imperfecta and osteopetrosis.²³ Nevertheless, these were considered unlikely based on the clinical findings. In cleidocranial dysostosis, for example, the clavicle is predominantly involved, while this bone is rarely affected in pyknodysostosis.²⁴ Furthermore, acro-osteolysis is not expected to involve facial bones,²⁵ and osteopetrosis is usually associated

(HGVS) nomenclature
Variation Society
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Table 1

Chromosome: position	Ref/alt	Allele counts	Allele frequencies	No. of alleles	No. of hetero	No. of homo	Sequence ontology	Effect	HGVS c (cDNA)	HGVS p (protein)	Exon	Distance of ranscript (%)
C1:105944311	СЛ		0.0125	80	-	0	Intron variant	Other	ENSFCAT0000003643:c.894-49G>A		00	
C1 105945321	C/T	43	0.5375	08	0	10	Intron variant	Other	FNSFCAT000003643.c 893+73G>A		7	
C1 105945347	۲/۵	0	0.025		0	ic	Intron variant	Other	ENSECATOOOOOO3643.c 893+47C>T		. ~	
C1-105945453	T/C	J .	0.0125		J .		Synonymous variant	Other		n Ala278-	. ~	33 8870
C1:105945516	C/T		0.0125	800			Intron variant	Other	ENSFCAT0000003643.c.788-17G>A		. ~	1
C1:105945532	-/9	,	0.0125	80	,	0	Intron variant	Other	ENSFCAT0000003643:c.788-33delC		2	
C1:105945562	C/T		0.0125	80	—	0	Intron variant	Other	ENSFCAT0000003643:c.788-63G>A		2	
C1:105945563	A/G		0.0125	80		0	Intron variant	Other	ENSFCAT0000003643:c.788-64T>C		2	
C1:105945702	G/A		0.0125	80		0	Intron variant	Other	ENSFCAT0000003643:c.787+61C>T		9	
C1:105945725	G/A	25	0.3125	80	15	Q	Intron variant	Other	ENSFCAT0000003643;c.787+38C>T		9	
C1:105945734	C/G		0.0125	80		0	Intron variant	Other	ENSFCAT0000003643;c.787+29G>C		9	
C1:105945746	AC	Ţ	0.0125	80	-	0	Intron variant	Other	ENSFCAT0000003643:c.787+17T>G		9	
C1:105945826	G/A	0	0.025	80	0	-	Stop gained	LoF	ENSFCAT0000003643:c.724C>T	p.Arg242Ter	9	72.8097
C1:105945848	_/T	80	-	80	0	40	Splice acceptor variant	LoF	ENSFCAT0000003643:c.702+1delA)	D.	
C1:105945876	A/G	,	0.0125	80	,	С	Svnonvmous variant	Other	ENSFCAT0000003643:0.675T>C	n.Tvr225=	LC.	37.8751
C1:105949867	G/C	· .	0.0125	80	. . .		Intron variant	Other	ENSFCAT0000003643:c.621+38C>G		0 4	
C1 105949894	5/P	-	0.0125	80			Intron variant	Other	ENSECAT0000003643 c 621+11C>T		4	
C1.105950155	√0		0.0125				Intron variant	Other	ENSECATOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOO		. 4	
01.100000100		- +	0.0100		- +			Othor Othor	ENERC A TOODOODOOD 412:0 400 - 001 - 0		t c	
01.103931023	5 Q 2 F	- 0	0.0120		- 0	ΣL		Ourier			೧ (
C1:105951040	2	28	0.35	0 N N	20	Q	Intron_variant	Other	ENSFCA10000003643:c.402+/3A>G		n	
C1:105951045	G/A		0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.402+68C>T		ო	
C1:105951061	A/C	-	0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.402+52T>G		ო	
C1:105951157	T/C	-	0.0125	80		0	Missense_variant	Missense	ENSFCAT0000003643:c.358A>G	p.lle120Val	с С	35.9517
C1:105951484	G/T	0	0.025	80	0	0	Splice_region_variant	Other	ENSFCAT00000003643:c.124-8C>A		2	
C1:105951516	-/TACA		0.0125	80		0	Intron_variant	Other	ENSFCAT00000003643:c.124-43_124-		2	
									40dupTGTA			
C1:105951529	A-		0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.124-53deIT		0	
C1:105951875	G/A	Ţ	0.0125	80	, - -	0	Intron_variant	Other	ENSFCAT0000003643:c.123+71C>T		, -	
C1:105951898	C/G	-	0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.123+48G>C		, -	
C1:105951908	G/A	-	0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.123+38C>T		, -	
C1:105951921	C/T	-	0.0125	80		0	Intron_variant	Other	ENSFCAT0000003643:c.123+25G>A		, -	
C1:105951923	-/CTT	-	0.0125	80	-	0	Intron_variant	Other	ENSFCAT0000003643:c.123+21_123		. 	
									+23dupAAG			
C1:105951951	T/C		0.0125	80		0	Missense_variant	Missense	ENSFCAT0000003643:c.118A>G	p.Asn40Asp	, -	11.7825
C1:105951964	C/T	-	0.0125	80	, - -	0	Synonymous_variant	Other	ENSFCAT0000003643:c.105G>A	p.Gly35=	,	10.4733
C1:105951982	T/A	-	0.0125	80	-	0	Synonymous_variant	Other	ENSFCAT0000003643:c.87A>T	p.Leu29=	. 	8.66062
C1:105952007	AG	-	0.0125	80		0	Missense_variant	Missense	ENSFCAT0000003643:c.62T>C	p.Val21Ala	—	6.143
C1:105952050	GЛ	Ţ	0.0125	80		0	Missense_variant	Missense	ENSFCAT0000003643:c.19C>A	p.Leu7lle	. 	1.81269
C1:105945848	T/-	388	-	388	0	194	Splice_acceptor_variant	LoF	ENSFCAT0000003643:c.702+1deIA		5	
C1:105952056	T/G	CJ	0.00518135	386	0	-	Missense_variant	Missense	ENSFCAT0000003643:c.13A>C	p.Lys5GIn	. 	1.20846
C1:105951157	T/C	-	0.00259067	386		0	Missense_variant	Missense	ENSFCAT0000003643:c.358A>G	p.lle120Val	e	35.9517
C1:105951210	C/T	Ţ	0.0025641	390	, - -	0	Missense_variant	Missense	ENSFCAT0000003643:c.305G>A	p.Arg102His	e	30.6143
C1:105951951	T/C	-	0.00257732	388		0	Missense_variant	Missense	ENSFCAT0000003643:c.118A>G	p.Asn40Asp	, -	11.7825
C1:105952050	G/T	-	0.00259067	386		0	Missense_variant	Missense	ENSFCAT0000003643:c.19C>A	p.Leu7lle	. 	1.81269
C1:105952007	A/G	2	0.00515464	388	S	0	Missense_variant	Missense	ENSFCAT0000003643:c.62T>C	p.Val21Ala		6.143
C1:105951484	G/T	0	0.0230769	390	2J	0	Splice_region_variant	Other	ENSFCAT0000003643:c.124-8C>A		2	

with severe anaemia and hepatosplenomegaly, which were absent in this case.²⁶ A further limitation is that genome sequencing of the cat's parents was not available to confirm the significance of the CTSK variant identified (the owner did not consent to them being tested). In addition, the remaining private DNA variants of the case could be valid causal mutations. Nevertheless, the relevant variant was not present in any other WGS of the 99 Lives data set and, alongside the compatible clinical signs, was strongly suggestive of the disease. Finally, the home-cooked diet may have also contributed to the clinicopathological abnormalities identified, for example due to phosphate imbalance or nutritional hyperparathyroidism, although the phosphate did not improve after the diet change and the clinical signs predated the start of this diet. Nevertheless, measurement of parathyroid hormone levels would have been useful to exclude nutritional hyperparathyroidism in this case.

Conclusions

To our knowledge, this is the first report of suspected pyknodysostosis in a cat. The diagnosis was reached by WGS identification of a *CTSK* variant, a technique that is able to facilitate the diagnosis of various hereditary disorders and support precision medicine healthcare in domestic cats. Although pyknodysostosis is a severely debilitating disease, a reasonable quality of life may be achieved with appropriate symptomatic management.

Acknowledgements We appreciate the laboratory assistance of Thomas R Juba, Juan A Valencia and the 99 Lives Cat Genome Consortium (felinegenetics.missouri.edu) for sharing domestic cat variant frequency information, and the assistance of Chris Warren Smith (Langford Vets) for the creation of the CT images. Our thanks to the owners of the cat that was the subject of this study for allowing us to collect samples from their pet, and the primary veterinarian for referring the case.

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Conflict of interest The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding Funding for the whole-genome sequencing was provided by the Gilbreath McLorn Endowment and donations to the 99 Lives project, EveryCat Health Foundation (formerly the Winn Feline Foundation), the Miller Trust (MT18-009; MTW18-009) (LAL) and Mars Wisdom Health (RMB).

Ethical approval The work described in this manuscript involved the use of non-experimental (owned or unowned) animals. Established internationally recognised high standards ('best practice') of veterinary clinical care for the individual patient were always followed and/or this work involved the use of cadavers. Ethical approval from a committee was therefore not specifically required for publication in *JFMS Open Reports*. Although not required, where ethical approval was still obtained, it is stated in the manuscript.

Informed consent Informed consent (verbal or written) was obtained from the owner or legal custodian of all animal(s) described in this work (experimental or non-experimental animals, including cadavers) for all procedure(s) undertaken (prospective or retrospective studies). For any animals or people individually identifiable within this publication, informed consent for their use in the publication (verbal or written) was obtained from the people involved.

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Supplementary Figure 1: *CTSK* variants in cat whole-genome sequences.

Supplementary Figure 2: Protein alignment of human and cat CTSK.

Supplementary file: All sequence variants unique to the affected cat, as well as all variants identified in *CTSK* in the data set.

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