

A case of spongiform polioencephalomyelopathy in a cat with a history of behavioural problems

Authors: Camps, Tomàs, de la Fuente, Cristian, Pumarola, Martí, Amat, Marta, Le Brech, Susana, et al.

Source: Journal of Feline Medicine and Surgery Open Reports, 1(2)

Published By: SAGE Publishing

URL: https://doi.org/10.1177/2055116915599172

The BioOne Digital Library (<u>https://bioone.org/</u>) provides worldwide distribution for more than 580 journals and eBooks from BioOne's community of over 150 nonprofit societies, research institutions, and university presses in the biological, ecological, and environmental sciences. The BioOne Digital Library encompasses the flagship aggregation BioOne Complete (<u>https://bioone.org/subscribe</u>), the BioOne Complete Archive (<u>https://bioone.org/archive</u>), and the BioOne eBooks program offerings ESA eBook Collection (<u>https://bioone.org/esa-ebooks</u>) and CSIRO Publishing BioSelect Collection (<u>https://bioone.org/csiro-ebooks</u>).

Your use of this PDF, the BioOne Digital Library, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <u>www.bioone.org/terms-of-use</u>.

Usage of BioOne Digital Library content is strictly limited to personal, educational, and non-commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne is an innovative nonprofit that sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Case Report





A case of spongiform polioencephalomyelopathy in a cat with a history of behavioural problems

Journal of Feline Medicine and Surgery Open Reports 1–5 © The Author(s) 2015 Reprints and permissions: sagepub.co.uk/journalsPermissions.nav DOI: 10.1177/2055116915599172 jfmsopenreports.com



Tomàs Camps¹, Cristian de la Fuente², Martí Pumarola², Marta Amat¹, Susana Le Brech¹ and Xavier Manteca¹

Abstract

A 7-month-old, entire female, domestic shorthair cat was referred to our behavioural service owing to soiling in the house and a play-related problem. The owners' complaints were that the cat had never used the litter tray, and it did not know how to play. After reviewing the behavioural history, a problem of substrate preferences acquisition was suspected with regard to the elimination problem. During the consultation, the physical examination was unremarkable, but the neurological examination revealed a moderate and hypermetric ataxic gait, and a bilateral lack of menace response. Some degree of visual impairment was suspected. The problem was located in the central nervous system (CNS); specifically, an intracranial and multifocal problem was diagnosed. After a complete work-up (complete ophthalmological examination, complete blood count and a complete biochemistry panel, feline immunodeficiency virus/feline leukaemia virus test, thorax radiographs, abdominal ultrasound, brain magnetic resonance imaging [0.2 T], cerebrospinal fluid analysis and a urinary metabolic screen test), a degenerative CNS problem was suspected. No treatment was prescribed for the neurological problem. Regarding the problem of soiling in the house, reward-based training with a clicker was used, and the cat partially improved in a few weeks. Three months later, the cat was referred to the neurology service in status epilepticus. A symptomatic treatment was prescribed, with a mild response. After 2 years of treatment and a progressive worsening, the cat was euthanased. Necropsy revealed spongiform polioencephalomyelopathy. In order to rule out prion aetiology a PrPsc inmunohistochemistry assay was performed, and the results were negative. Congenital spongiform polioencephalomyelopathy (CSP) was diagnosed. We strongly suggest that the cat's behavioural clinical signs were caused by the CSP, causing learning impairment. To the best of our knowledge, this would be the first case in which a congenital degenerative disease affected a cat's capability to learn, leading to behavioural signs as the main complaint of the owners, even before neurological signs are detected by the owners.

Accepted: 14 July 2015

A 7-month-old, entire female, domestic shorthair cat was referred to our behavioural service owing to house soiling and a play-related problem. The owners' complaints were that the cat had never used the litter tray, and it did not know how to play. The environment consisted of two young adult humans with no children. They lived in a flat of 85 m², with two terraces of 5 m² each. There were three separated litter boxes at home, all of which were non-covered with low sides. The owners had used clumping, non-clumping, silica-based and soil-based litter during the months between the adoption (when the

¹Department of Animal and Food Science, School of Veterinary Science, Autonomous University of Barcelona, Barcelona, Spain ²Department of Animal Medicine and Surgery, School of Veterinary Science, Autonomous University of Barcelona, Barcelona, Spain

Corresponding author:

Tomàs Camps DVM, MsC, Dip ECAWBM-BM, Department of Animal and Food Science, School of Veterinary Science, Autonomous University of Barcelona, Despatx V0-135, Edifici V, Campus UAB, Bellaterra, Barcelona 08193, Spain Email: tomas.camps@uab.cat

Creative Commons CC-BY-NC: This article is distributed under the terms of the Creative Commons Attribution-NonCommercial 3.0 License (http://www.creativecommons.org/licenses/by-nc/3.0/) which permits non-commercial use, reproduction and distribution of the work without further permission provided the original work is attributed as specified on the SAGE and Open Access page (https://us.sagepub.com/en-us/nam/open-access-at-sage).

Downloaded From: https://complete.bioone.org/journals/Journal-of-Feline-Medicine-and-Surgery-Open-Reports on 15 Jul 2025 Terms of Use: https://complete.bioone.org/terms-of-use cat was 4 months old) and the first visit. One of the latrines always had clumping substrate. There were three food and three water troughs, all of them far from the latrines.

The impression of the owners was that the cat eliminated where it was at any given moment. It eliminated many more times in front of the owner (90%) than when it was alone (10%). The cat never tried to cover its faeces or urine after depositions. Occasionally, the owners punished the cat verbally and physically, but only when it eliminated in front of them. The substrates used by the cat were ceramic tiles, the sofa and beds. It always adopted an emptying-body posture. Spots were always located on horizontal surfaces. The owners used bleachbased products in order to clean the spots, and just water in the case of the sofa and beds.

Regarding the play-related problem, the owners said that the cat did not understand the body language of other cats and commonly crashed into other cats or people. It also 'tried to bite, catch and scratch the air' when playing. It did not find balls or other toys when the owners threw them to the cat to play.

The rest of the cat's behavioural history was unremarkable.

The differential diagnoses of the house-soiling problem included problems with the litter trays, including insufficient number, incorrect type, competition with other cats for the latrines, incorrect location, acquired aversion, or inappropriate substrate; a preference for another location or substrate; a problem with preference acquisition (ie, because of a cognitive impairment, or a sensory impairment or an unavailability of appropriate latrines or substrate during the first few weeks of life); and marking behaviours. Finally, a medical illness can contribute to all of these problems or to be the main cause.^{1,2}

We could rule out most of these problems after interviewing the owners. Firstly, it was unlikely that there was a problem with the litter tray because the number, type and location were correct. The locations were correct because the animal eliminated near the litter tray if it was there. Many different substrates had been used. Secondly, it was not a problem of preference because the cat eliminated in different locations and surfaces. The age of the animal, the distribution of the spots and the body posture during elimination ruled out marking behaviours.

Alterations in play behaviours described by the owners could have been due to a cognitive impairment, and/ or a sensory impairment (ie, blindness). Play behaviours depend on learning capability and sensory systems.³ Additionally, an enriched environment is necessary to learn and display play behaviours in a proper manner.⁴ In that case, the social and instrumental environment was good. Regarding elimination, a problem of substrate preferences acquisition was diagnosed. A cognitive impairment, a medical condition or both could have been the cause of the problem during the elimination habits acquisition. Moreover, cognitive impairment and some medical conditions also could explain the play-related behavioural problems.

During the consultation, the physical examination was unremarkable; however, the neurological examination revealed a moderate and hypermetric ataxic gait, and a bilateral lack of menace response. A complete ophthalmological examination was performed by the ophthalmological service in order to rule out ocular diseases. No ophthalmological abnormalities were detected. Additionally, based on the behaviours at home described by the owners (the inability to find some toys, and the behaviour of 'scratching and biting the air'), some degree of visual impairment was suspected but not confirmed with the neurological examination. The cat did not crash with objects either at home or at the consultation room. The problem was located in the central nervous system (CNS); specifically, an intracranial and multifocal problem was diagnosed. A complete blood count and a complete biochemistry panel were performed, and all of the results were within normal limits. The feline immunodeficiency virus/feline leukaemia virus test was negative. A thorax radiograph, abdominal ultrasound, brain magnetic resonance imaging (MRI; 0.2 T) and cerebrospinal fluid analysis showed no abnormalities. Although some small lesions could be missed with low-field MRI, we had to assume the absence of lesions obtained in the work-up. Thus, a degenerative condition such as a lysosomal storage disease, organic aciduria or mitochondrial encephalopathy was suspected. Samples of blood and urine were sent to the University of Pennsylvania School of Veterinary Medicine for metabolic screen tests. Amino acids, organic acids, carbohydrates, nitroprusside, ketone and mucopolysaccharide concentrations were analysed, as was α-mannosidase, β-mannosidase, fucosidase and hexosaminidase A and B activity. All of these were within normal limits.

No treatment was prescribed for the neurological problem. The owners were given advice on correcting the soiling problem in the house using reward-based training with a clicker. During the first week, the clicker was conditioned by a food reward, and the entire floor was covered with newspaper. Each time that the cat eliminated, it was rewarded with a clicker and food. Newspaper was removed progressively. Three months later, the cat used a small newspaper-covered area to eliminate. This partial improvement suggests that there was a learning impairment during the acquisition of habits but not a total lack of learning capability.

After 3 months, the cat was referred to the neurology service again, in status epilepticus. Neurological

findings, after the postictal phase, included a lack of bilateral menace response and cerebellar ataxic gait. A bilateral carpal valgus that had already been found in the first visit and a visible suture line in the posterior capsule of both crystalline lenses were also detected. The owners reported progressive gait deficits over the previous month and compulsive running episodes with a partially impaired mental status (probably seizure activity). A symptomatic treatment with diazepam (1 mg/kg intrarectally only if seizures) and phenobarbital (2 mg/kg PO q24h) was started, with a very poor response. After 15 days of treatment, levetiracetam was added (10 mg/kg PO q8h) owing to an increase in seizure activity, with an initially good response. However, seizures reappeared after 2 months with 1-2 episodes every 15 days each lasting <2 mins. After 2 years of treatment and a progressive worsening, the cat was euthanased. A complete necropsy was immediately performed. No gross lesions were found, and based on the CNS histological lesions shown in Figures 1-4, a diagnosis of spongiform polioencephalomyelopathy was made. The spongiform degeneration of the grey matter was extensively distributed in the whole CNS. In order to rule out prion aetiology, a PrPsc inmunohistochemistry assay was performed, and the results were negative. Thus, congenital spongiform polioencephalomyelopathy (CSP) was diagnosed postmortem.

Spongy vacuolation seen by light microscopy in the neural tissue is defined as spongy degeneration, and may take the form of vacuoles within processes of the neuropil, vesiculation of myelin sheaths, or swelling of astrocyte or oligodendrocyte cytoplasm.⁵ A congenital problem, retrovirus infection and prion disease have been suggested as possible aetiologies.^{6–10} Congenital



Figure 1 Mesencephalon. Intraneuronal vacuolisation of the red nucleus. Prominent vacuoles of different sizes are located in the perikaryon of some neuronal bodies (arrows) occupying most of the perikaryon and displacing the neuronal nucleus to the periphery. Small vacuoles are also present in the neuropil together with a moderate gliosis (microgliosis)



Figure 2 Lumbosacral spinal cord. (a) A general view of the lumbosacral spinal cord shows a generalised spongiform appearance of the grey matter in the dorsal and ventral spinal horns. (b) Detail of the ventral spinal horn showing a moderate spongiosis of the neuropil and the presence of vacuoles of different sizes with an irregular greyish content in most of the perikaryon of the neuronal bodies (arrows)



Figure 3 Cerebellar cortex. High-power field view of the cerebellar cortex showing a generalised mild spongiosis of the molecular layer with round, empty spaces in the neuropil



Figure 4 Cerebral cortex. Prominent and empty vacuoles are multifocally distributed throughout the neuropil of the parietal cortex (arrow). The lesion is located in the deeper layer of parietal cortex while no histopathological changes are observed in the subcortical white matter (left side)

spongiform degeneration of the grey matter has been previously described in a few cases of cats;^{6–8} however, the cause remains still unknown. Common clinical signs include gait alteration, seizures, blindness, bilateral cataracts, behavioural changes and cranial nerve alterations. These signs appear during very early stages of development (just after birth), with a progressive fatal outcome in a few days or months. Behavioural signs are poorly described in animal science literature.^{6–8} Although the degeneration usually affects diffusely all the grey matter, the behavioural alterations and the evolution of the clinical signs depend on the affected area of the brain in each

case. Nevertheless, the clinical signs do not always correlate with the degree of the histological lesions. A spongy degenerative problem of grey matter has also been described in humans, and occurs in isolated cases and in sibs.^{11–13} In all of human cases reported, the problem appears early in infancy and the outcome is always fatal. All the affected children show learning disabilities (ie, retarded speech development) during the early periods of infancy. Additionally, they rapidly develop neurological signs, especially seizures. The clinical findings and neuropathological changes are very similar in humans and the present case. There are no studies regarding degeneration of grey matter and its effect on learning ability in animals. However, other neurodegenerative problems (ie, lysosomal storage diseases) and problems that lead to structural abnormalities of the forebrain (ie, hydrocephaly) may be correlated with learning disabilities in animals and humans.14-17

The acquisition of the elimination habits occurs during the first weeks of a kitten's life. Most kittens naturally seek out sand-like materials for elimination purposes. However, the preference for a substrate needs to be learnt during those first weeks of age. Learning disabilities and/or sensory impairments could modify the acquisition of these habits.

Conclusions

We strongly suggest that the behavioural signs (elimination and the play-related problem) were caused by CSP, causing learning impairment.

This case contributes to scientific knowledge for two reasons. Firstly, it describes a CSP, which is a very rare condition described in cats. Secondly, to the best of our knowledge, this is the first case in which a congenital degenerative disease affects a cat's capability to learn, leading to behavioural signs as the main complaint, even before neurological signs are detected by the owner.

Finally, this case illustrates the importance of considering medical conditions in all behavioural cases and of using an accurate diagnostic protocol.

Acknowledgements This manuscript has been proofread by Mr Chuck Simmons, a native, English-speaking University Instructor of English.

Conflict of interest The authors do not have any potential conflicts of interest to declare.

Funding The authors received no specific grant from any funding agency in the public, commercial or not-for-profit sectors for the preparation of this case report.

References

1 Manteca X, Amat M and Camps T. House soiling in cats. *Feline Focus* 2015; 2: 51–54.

- 2 Bowen J and Heath S. Feline house-soiling and marking problems. In: Bowen J and Heath S (eds). Behaviour problems in small animals: practical advice for the veterinary team. 1st ed. Philadelphia, PA: Elsevier Saunders, 2005, pp 185–203.
- 3 Spinka M, Newberry RC and Bekoff M. Mammalian play: training for the unexpected. *Q Rev Biol* 2001; 76: 141–168.
- 4 Boissy A, Manteuffel G, Jensen MB, et al. Assessment of positive emotions in animals to improve their welfare. *Physiol Behav* 2007; 92: 375–397.
- Summers B, Cummings JF and DeLahunta A. Degenerative diseases of the central nervous system. In: Summers B, Cummings JF and DeLahunta A (eds). Veterinary neuropathology. 1st ed. St Louis, MO: Mosby-Year Book, 1995, pp 208–305.
- 6 Jones BR, Alley MR, Shimada A, et al. An encephalomyelopathy in related Birman kittens. *N Z Vet J* 1992; 40: 160–163.
- 7 Morita T, Shimada A, Ishibashi T, et al. Congenital spongiform change in the brain stem nuclei of a domestic kitten. *J Comp Pathol* 2002; 126: 212–215.
- 8 Vidal E, Montoliu P, Añor S, et al. A novel spongiform degeneration of the grey matter in the brain of a kitten. J Comp Pathol 2004; 131: 98–103.
- 9 Abramo F, Bo S, Canese MG, et al. Regional distribution of lesions in the central nervous system of cats infected with feline immunodeficiency virus. *AIDS Res Hum Retro*viruses 1995; 11: 1247–1253.

- 10 Wyatt JM, Pearson GR, Smerdon TN, et al. Naturally occurring scrapie-like spongiform encephalopathy in five domestic cats. Vet Rec 1991; 129: 233–236.
- 11 Jellinger K and Seitelberger F. Spongy glio-neuronal dystrophy in infancy and childhood. Acta Neuropath 1970; 16: 125–140.
- 12 Janota I. Spongy degeneration of grey matter in 3 children. Arch Dis Child 1974; 49: 571–575.
- 13 Ropper AH and Brown RH. The inherited metabolic diseases of the nervous system. In: Fotlin J, Nogueira I, Edmmonson KG and Sheinis LA (eds). Adams and Victor's principles of neurology. 8th ed. New York: McGraw-Hill, 2005, pp 797–849.
- 14 Blakemore WF. A case of mannosidosis in the cat: clinical and histopathological findings. J Small Anim Pract 1986; 27: 447–455.
- 15 Robinson AJ, Crawley AC, Auclair D, et al. Behavioural characterisation of the α-mannosidosis guinea pig. Behav Brain Res 2008; 186: 176–184.
- 16 Del Bigio MR, Crook CR and Buist R. Magnetic resonance imaging and behavioural analysis of immature rats with kaolin-induced hydrocephalus: pre- and postshunting observations. *Exp Neurol* 1997; 148: 256–264.
- 17 Lindquist B, Carlsson G, Persson EK, et al. Learning disabilities in a population-based group of children with hydrocephalus. Acta Paediatr 2005; 94: 878–883.