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Acute non-compressive nucleus pulposus extrusion in cats: clinical features, diagnostic imaging findings, treatment and outcome

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Cat; disc; extrusion; non-compressive; nucleus pulposus; outcome

Abstract

Objectives: The aim of the study was to describe the clinical features, diagnostic imaging findings, treatment and outcome in cats diagnosed with presumptive acute non-compressive nucleus pulposus extrusion.
Methods: Medical records and imaging studies of cats diagnosed with presumptive acute non-compressive nucleus pulposus extrusion were retrospectively reviewed. Long-term follow-up information was acquired from patient records and from either owners or referring veterinary surgeons via a telephone questionnaire.

Results: Eleven cats met the inclusion criteria. All cats had a peracute onset of clinical signs, with eight cats experiencing witnessed (n = 6) or suspected (n = 2) external trauma. Neurological examination findings ranged from ambulatory paresis to plegia with loss of deep nociception. Neuroanatomical localisation included C1-C5 (n = 1), T3-L3 (n = 7) and L4-S3 (n = 3) spinal cord segments. Ten cats were discharged with a median hospitalisation time of 10 days (range 3 days to 26 days). One cat was euthanised during hospitalisation due to complications unrelated to neurological disease. Cats that presented with paraplegia regained voluntary movement within a median of 4 days (range 2 to 7 days). For those cats that presented non-ambulatory, all cats regained an ambulatory status with the median time to ambulation of 17 days (range 6 to 21 days). Five cats had absent voluntary urination at presentation; this resolved in all but one cat that had long-term urinary incontinence. Overall the outcome for cats diagnosed with acute non-compressive nucleus pulposus extrusion was good with almost 90% returning to ambulation with urinary and faecal continence.

Conclusions and relevance: The majority of cats diagnosed with acute non-compressive nucleus pulposus extrusion had characteristic clinical presentations and good outcomes. Acute non-compressive nucleus pulposus extrusion should be considered as a differential diagnosis for cats presenting with peracute onset of spinal cord dysfunction, particularly if there is a clinical history or evidence of trauma.
Introduction

Acute non-compressive nucleus pulposus extrusion (ANNPE), previously referred to as a traumatic intervertebral disc extrusion (IVDE), high-velocity/low-volume IVDE and type III intervertebral disc extrusion occurs when a healthy and hydrated IVDE is exposed to sudden and excessive force and is typically seen following vigorous exercise or trauma\textsuperscript{1-4}. This type of intervertebral disc extrusion results in spinal cord contusion with minimal or no spinal cord compression\textsuperscript{1-3}.

ANNPE has been frequently reported in dogs\textsuperscript{1-3}, however there are only single case reports describing ANNPE in cats\textsuperscript{4,5}. Dogs with ANNPE typically present with a peracute onset of spinal cord dysfunction that is non-progressive after 24 hours \textsuperscript{1,2}. Clinical signs are often strongly lateralised and mild to moderate spinal hyperesthesia may be seen in approximately half of affected cases\textsuperscript{2,6}.

Definitive diagnosis of ANNPE can only be confirmed by histopathology\textsuperscript{1}. However, magnetic resonance imaging (MRI) can be used to make a presumptive diagnosis with with specific characteristics identified to reach a presumptive ante-mortem diagnosis of ANNPE\textsuperscript{2,3}.

Typical treatment involves physiotherapy and supportive care with the use of analgesics as required\textsuperscript{6}. The outcome is considered good in dogs with only a minority failing to regain normal neurological function\textsuperscript{2}.

Despite this disorder being well characterised in dogs, little is known about the clinical presentation, imaging findings and outcome in cats. The aims of this study were therefore to describe the clinical features, diagnostic imaging findings, treatment and outcome in a larger number of cats diagnosed with presumptive ANNPE. We hypothesised that cats diagnosed with presumptive ANNPE would have a characteristic presentation and a good long-term outcome.
Material and Methods

Ethics Statement

Ethics approval was granted by the Royal Veterinary College (RVC) Ethics and Welfare Committee (reference number 2015 1324).

Criteria for inclusion

Medical records of cats that had presumptively been diagnosed with ANNPE at the RVC between 2008 and 2014 were reviewed. In order to be included, cats needed to have had an MRI of the affected spinal cord segments within 48 h of the onset of clinical signs, MRI findings consistent with the diagnosis of presumptive ANNPE and have follow-up information for a minimum of 3 months. Recorded information included immediate history preceding onset of clinical signs, treatment prior to referral, signalment, general physical examination findings, neurological examination findings, duration of time from detecting neurological signs to MRI, treatment administered following diagnosis, duration of hospitalisation and presence of complications. In relevant cases the time to recover nociception, voluntary motor activity and unassisted ambulation was also recorded.

Diagnostic imaging

MRI was performed using a 1.5 Tesla scanner (Intera, Philips Medical Systems) and included a minimum of T2- and T1-weighted sagittal and transverse images. All imaging studies were reviewed for diagnostic accuracy by a board certified neurologist (SDD) blinded to the clinical signs and neuroanatomical localisation, and only those cases with imaging features consistent with presumptive ANNPE diagnosis were included in the study. MRI findings compatible with ANNPE included (1) a reduction in volume of the T2-weighted hyperintensity of the nucleus pulposus signal, (2) a focal T2-weighted hyperintensity within the spinal cord overlying an intervertebral disc space, (3) mild narrowing of the intervertebral disc space, and (4) extraneous
material or signal change within the vertebral canal with absent or minimal spinal cord compression\(^2\,^3\,^5\) (Figure 1 a, b).

**Assessment of outcome**

Short-term outcome was defined as the period between the onset of clinical signs up to 6 weeks following presumptive diagnosis of ANNPE, and information was retrieved from medical records. Long-term outcome was defined as a minimum follow-up period of 3 months\(^7\). This information was initially obtained via telephone interview with the referring veterinary surgeons. For cats that were deceased, date and cause of death as well as the last documented neurologic status were recorded. Conforming to local ethics and welfare committee guidelines, only owners of cats that were still alive at the time of data collection were subsequently contacted. Owners were mailed a letter with study details and a standardized questionnaire that had been reviewed and approved by a local ethics and welfare committee. Telephone interviews were conducted using the questionnaire, which included questions covering specific aspects of the disease, such as amount of activity, lameness, paresis and incontinence, type of medical and supportive treatment received, response to treatment and quality of life (supplementary material). A successful outcome was defined as resolution or improvement of clinical signs with the cat being able to ambulate independently with control of urination and defaecation, while an unsuccessful outcome was defined as a cat that required support to ambulate or had persistent urinary or faecal incontinence.

**Results**

Of 14 potential cats identified, 11 were included in the study (Table 1). The cats had a median age of 7 years (range 2 years 9 months to 13 years) at presentation. Eight of the cats were male
neutered and three were female neutered. Breeds comprised the domestic shorthair (n = 6), domestic longhair (n = 3), Egyptian Mau (n = 1) and British Shorthair (n = 1).

**Historical findings**

All cats had an acute or peracute onset of clinical signs. The median time to presentation was 14 h (range 2–48 h) following the onset of neurological signs. Prior to presentation six of the cats had been involved in a witnessed traumatic event (road traffic accident [n = 3] or fall from a height [n = 3]). The remaining five cats were found either in the home or nearby the house and the onset of clinical signs was not witnessed.

**Clinical findings**

The majority of cats (n = 10) had clinical signs referable to the paraparesis or paraplegia (Table 1). Neuroanatomical localisation included the C1–C5 (n = 1), T3–L3 (n = 7) and L4–S3 (n = 3) spinal cord segments. The clinical signs were non-progressive in all cats following presentation. Five of the cats had signs consistent with external trauma, including head trauma, pulmonary contusions and scuffed nails.

**MRI findings**

MRI revealed ANNPE located at C3–C4 (n = 1), T12–T13 (n = 1), T13–L1 (n = 1), L1–L2 (n = 1), L3–L4 (n = 3), L4–L5 (n = 1) and L5–L6 intervertebral disc spaces (n = 3). One cat had a dorsal spinous process fracture of the L7 vertebra, which was not associated with the neuroanatomical localisation nor the anatomical localisation of the ANNPE and was therefore considered incidental. There was evidence of ill-defined T2-weighted hyperintensity within the epaxial musculature compared with surrounding muscle suggestive of contusion, haemorrhage or oedema in five cats (Figure 1c). Of these five cats, two cats had no history or examination findings consistent with trauma, while the other three cats were involved in a witnessed trauma.
Treatment and short-term outcome

All cats received physiotherapy performed by a veterinary physiotherapist and/or qualified veterinary nurse consisting of massage, passive range of motion exercises, assisted standing and exercises to develop strength and coordination, as appropriate and tolerated by each cat. Five cats that demonstrated signs of spinal hyperaesthesia received analgesic medication that included opioids (ie, methadone and buprenorphine; n = 3), non-steroidal anti-inflammatory drugs (n = 1) and gabapentin (n = 1).

The median time for cats with absent deep nociception (n = 3) to regain sensation was 2 days (range 1–3 days). Of the cats that presented with paraplegia, including those with absent deep nociception (n = 5) the median time for them to regain voluntary movement (non-ambulatory) was 4 days (range 2–7 days). For those cats that presented non-ambulatory (including paraplegic cats; n = 9) the median time to ambulation was 17 days (range 6–21 days).

Five cats required bladder management during hospitalisation, including indwelling catheter placement (n = 1), intermittent catheterisation (n = 1) and manual bladder expression (n = 3). Two cats received a sympatholytic medication (prazosin) to aid in bladder management. Three cats were discharged with improved motor function but continued to require manual bladder expression.

The cats had a median hospitalisation time of 10 days (range 3–26 days). Four of the 10 cats that survived to discharge were ambulatory at that time. One cat did not survive until discharge, and was euthanased owing to respiratory deterioration as a result of pulmonary contusions.

Short-term outcome (4–6 weeks following diagnosis of presumptive ANNPE) in six cats revealed all cats were ambulatory and had improved neurological function compared with the time of discharge; however, none of the cats were considered to be neurologically normal.

Long-term outcome
Long-term outcome in eight cats (four cats were also included in the assessment of short-term outcome) was obtained from the referring veterinary surgeons (n = 2) or veterinary surgeons and owners (n = 6). The median duration of time between the onset of clinical signs and assessment of outcome was 44 months (range 4–68 months). None of the cats displayed signs of further improvement 6 months after reaching a presumptive diagnosis of ANNPE. Although all cats were ambulatory and did not demonstrate any signs of spinal hyperaesthesia, none were reported to have become neurologically normal. Owners or veterinary surgeons assessed all cats to have regained a good quality of life; however, quality of life was considered decreased compared with before the onset of clinical signs in all of the cats with 3/8 cats now indoor-only cats.

One cat (cat 2) had ongoing urinary incontinence requiring twice daily manual bladder expression, and the same cat had intermittent faecal incontinence (Table 1).

Overall, 7/8 cats (88%) were considered to have a successful long-term outcome, and one cat was considered to have an unsuccessful outcome.

Discussion

The differential diagnosis for cats presenting with an acute or peracute onset of paresis or plegia includes aortic thrombo-embolism, ischemic myelopathy, fibrocartilaginous embolism, intervertebral disk disease, and vertebral fractures and luxations.\(^8,9\) It has previously been reported that trauma accounts for 14% of cases of feline spinal cord injury,\(^10\) and the occurrence of a vertebral fracture or luxation is generally considered the most important differential diagnosis for cats presenting with a peracute onset of spinal cord dysfunction after a witnessed or suspected traumatic event. Of the cats included in this study nearly three-quarters of the cats had experienced a witnessed traumatic event or there was evidence of trauma based on their clinical exam or imaging findings. This highlights the need to include ANNPE as a possible
differential diagnosis for any cat presenting with an acute or peracute onset of spinal cord
dysfunction, particularly if there is any history or evidence of trauma.

When considering the location of the ANNPE the most frequent sites were the L3-L4 and L5-
6 intervertebral discs. There was also one patient with a cervical ANNPE. This is consistent
with the previous case reports that describe a lumbar and cervical ANNPE. Whilst this
contrasts to the findings in dogs, which predominantly have T12-T13 and T13/L1 ANNPE, it
is more consistent with data looking at the location of IVDE, with previous studies suggesting
that the mid to caudal lumbar region is more commonly affected in cats.

When considering the outcome for patients diagnosed with ANNPE it is overall very good with
almost 90% of the cats being ambulatory with full urinary and faecal continence. None of the
cats were described as returning to ‘normal’ following the onset of clinical signs, and this is
consistent with one of the previous case reports that suggested there was ataxia present six
months following diagnosis. However, 50% of cats had returned to former behaviours
including outside activity and climbing on to furniture. It is currently unclear for those cats that
were no longer allowed outside, if this reflected a concern on part of the owners or an actual
inability to perform activities as before the onset of clinical signs. From the results of this
study, it is difficult to draw any conclusions on potential prognostic indicators for cats with a
presumptive ANNPE.

The incidence of ANNPE in cats is not known, although it appears to be infrequent, however
it is possible that this reflects a decreased awareness of the condition and therefore an under-
diagnosis. The treatment involved in caring for cats following the diagnosis of ANNPE is
primarily supportive, involving the use of analgesics as appropriate, bladder management
where required and intensive physiotherapy. The cost of treatment compared to cats diagnosed
with vertebral fracture/luxation or IVDD is often reduced owing to the fact that there is no need
for surgery. In addition it is often possible for caregivers to be trained to provide physiotherapy and bladder management at home. This combined with the evidence presented in this study that suggests cats with ANNPE appear to have a favourable prognosis highlights the need for ANNPE to be considered as an important differential diagnosis in cats with a peracute onset of spinal cord dysfunction.

Conclusions

This study is obviously limited by its retrospective nature and the small number of included cases. However, the majority of cats diagnosed with presumptive ANNPE presented with paraparesis or paraplegia and had neuroanatomical localisation of T3–L3 and L4–S3 spinal cord segments. Nearly 75% of the cats were involved in a witnessed trauma or had evidence of trauma based on clinical examination or imaging findings. The majority of cats diagnosed with ANNPE had good outcomes. ANNPE should be considered as a differential diagnosis for cats presenting with peracute onset of spinal cord dysfunction, particularly if there is a clinical history or evidence of trauma.

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Conflict of Interest

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

1 Griffiths IR. A syndrome produced by dorso-lateral “explosions” of the cervical intervertebral discs. Vet Rec 1970; 87: 10-11


## Table 1

Signalment, clinical presentation and outcome of 11 cats diagnosed with presumptive ANNPE

<table>
<thead>
<tr>
<th>Cat</th>
<th>Age</th>
<th>Sex</th>
<th>Breed</th>
<th>Neurological Examination Findings</th>
<th>Deep nociception present</th>
<th>Neurolocalisation</th>
<th>Site of ANNPE</th>
<th>Time to Ambulation (days)</th>
<th>Duration of hospitalisation (days)</th>
<th>Follow-up time and outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>12 y 5 mo</td>
<td>MN</td>
<td>DSH</td>
<td>Paraplegic</td>
<td>Yes</td>
<td>T3-L3</td>
<td>L3-L4</td>
<td>15</td>
<td>11</td>
<td>Now indoor cat</td>
</tr>
<tr>
<td>2</td>
<td>2 y 9 mo</td>
<td>MN</td>
<td>Egyptian Mau</td>
<td>Non-ambulatory paraparetic; right pelvic limb plegic</td>
<td>Absent</td>
<td>L4-S3</td>
<td>L5-L6</td>
<td>9</td>
<td>23</td>
<td>11 mo Urinary incontinence and occasional faecal incontinence</td>
</tr>
<tr>
<td>3</td>
<td>10 y 6 mo</td>
<td>FN</td>
<td>DLH</td>
<td>Non-ambulatory paraparetic</td>
<td>Yes</td>
<td>T3-L3</td>
<td>L3-L4</td>
<td>n/a</td>
<td>n/a</td>
<td>Euthanised due to pulmonary contusions</td>
</tr>
<tr>
<td>4</td>
<td>6 y</td>
<td>MN</td>
<td>DSH</td>
<td>Non-ambulatory paraparetic</td>
<td>Yes</td>
<td>T3-L3</td>
<td>T12-T13</td>
<td>16</td>
<td>20</td>
<td>Now indoor cat</td>
</tr>
<tr>
<td>5</td>
<td>4 y</td>
<td>MN</td>
<td>BSH</td>
<td>Non-ambulatory tetraparetic</td>
<td>Yes</td>
<td>C1-C5</td>
<td>C3-C4</td>
<td>18</td>
<td>9</td>
<td>LTF</td>
</tr>
<tr>
<td>6</td>
<td>4 y</td>
<td>FN</td>
<td>DSH</td>
<td>Ambulatory paraparetic</td>
<td>Yes</td>
<td>T3-L3</td>
<td>L4-L5</td>
<td>n/a</td>
<td>3</td>
<td>41 mo Only allowed outside in daylight hours</td>
</tr>
<tr>
<td>7</td>
<td>13 Y</td>
<td>MN</td>
<td>DLH</td>
<td>Non-ambulatory paraparetic; right pelvic limb plegic</td>
<td>Absent</td>
<td>T3-L3</td>
<td>L3-L4</td>
<td>21</td>
<td>7</td>
<td>46 mo Returned to previous lifestyle; now deceased due to</td>
</tr>
<tr>
<td>No.</td>
<td>Age</td>
<td>Gender</td>
<td>Breed</td>
<td>Condition</td>
<td>Ambulatory Status</td>
<td>Lesion Location</td>
<td>Lesion Extent</td>
<td>Follow-Up</td>
<td>Notes</td>
<td></td>
</tr>
<tr>
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<td></td>
</tr>
<tr>
<td>8</td>
<td>5 y</td>
<td>FN</td>
<td>DSH</td>
<td>Ambulatory paraparetic</td>
<td>Yes</td>
<td>L4-S3</td>
<td>L5-L6</td>
<td>n/a</td>
<td>3</td>
<td>67 mo</td>
</tr>
<tr>
<td>9</td>
<td>8 y</td>
<td>MN</td>
<td>DSH</td>
<td>Non-ambulatory paraparetic</td>
<td>Yes</td>
<td>L4-S3</td>
<td>L5-L6</td>
<td>6</td>
<td>7</td>
<td>68m</td>
</tr>
<tr>
<td>10</td>
<td>7 y</td>
<td>MN</td>
<td>DSH</td>
<td>Paraplegic</td>
<td>Yes</td>
<td>T3-L3</td>
<td>L1-L2</td>
<td>18</td>
<td>13</td>
<td>LTF</td>
</tr>
<tr>
<td>11</td>
<td>8 y 6 mo</td>
<td>MN</td>
<td>DLH</td>
<td>Paraplegic</td>
<td>Absent bilaterally in pelvic limbs</td>
<td>T3-L3</td>
<td>T13-L1</td>
<td>21</td>
<td>26</td>
<td>46 mo</td>
</tr>
</tbody>
</table>

ANNPE = acute non-compressive nucleus pulposus extrusion; y = years; mo = months; MN = male neutered; FN = female neutered; DSH = domestic shorthair; DLH = domestic longhair; BSH = British Shorthair; LTF = lost to follow up
Figure 1. (a) Sagittal T2-weighted and (b) transverse T2-weighted images at the level of the L5–L6 intervertebral disc space, and (c) L4 vertebral body of an Egyptian Mau aged 2 years and 9 months (cat 2). (a) A focal intraparenchymal hyperintensity is present at the level of the L5–L6 intervertebral disc space (long arrow). Although the nucleus pulposus has a reduced volume compared with the adjacent discs, it has remained a homogeneous hyperintense signal. (b) A small amount of extraneous material present in the epidural space (arrow). (a,c) A poorly demarcated hyperintensity within the epaxial musculature at the level of the L4 vertebral body, suggestive of epaxial muscle contusion, oedema or haemorrhage, was considered indicative for external trauma (short arrow [a] and arrow [c])