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## ACVIM consensus statement on diagnosis and management of acute canine thoracolumbar intervertebral disc extrusion

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### Abstract

**Background:** Thoracolumbar intervertebral disc extrusion (TL-IVDE) is the most common cause of acute paraparesis and paraplegia in dogs; however, guidelines on management of the condition are lacking.

**Objectives:** To summarize the current literature as it relates to diagnosis and management of acute TL-IVDE in dogs, and to formulate clinically relevant evidence-based recommendations.

**Animals:** None.

**Methods:** A panel of 8 experts was convened to assess and summarize evidence from the peer-reviewed literature in order to develop consensus clinical recommendations. Level of evidence available to support each recommendation was assessed and reported.

**Results:** The majority of available literature described observational studies. Most recommendations made by the panel were supported by a low or moderate level of evidence, and several areas of high need for further study were identified. These include better understanding of the ideal timing for surgical decompression, expected

**Abbreviations:** 3D FSE, 3-dimensional fast spin-echo; ACVIM, American College of Veterinary Internal Medicine; CSF, cerebrospinal fluid; CT, computed tomography; CTR, cutaneous trunci reflex; CUSA, cavitron ultrasonic surgical aspirator; DMSO, dimethyl sulfoxide; DPN, deep pain negative; DPP, deep pain positive; DTI, diffusion tensor imaging; EHLD, extensive hemilaminectomy and durotomy; FGF4, fibroblast growth factor 4; GI, gastrointestinal; HASTE, half-Fourier acquisition single-shot turbo spin-echo; IVDE, intervertebral disc extrusion; MMP, matrix metalloproteinase; MPSS, methylprednisolone sodium succinate; MRI, magnetic resonance imaging; MT, magnetization transfer; NMES, neuromuscular electrical stimulation; NP, nucleus pulposus; NSAID, nonsteroidal anti-inflammatory drug; PEMF, pulsed electromagnetic field; PLDA, percutaneous laser disc ablation; PMM, progressive myelomalacia; PROM, passive range of motion; RCT, randomized controlled trial; SCI, spinal cord injury; STIR, short Tau inversion recovery; T1W, T1-weighted; T2W, T2-weighted; TENS, transcutaneous electrical nerve stimulation; TL, thoracolumbar; UTI, urinary tract infection; UWMT, underwater treadmill.

Sarah A. Moore and Natasha J. Olby are Co-Chairs. The rest of the authors are panelists; panel members are listed in alphabetical order.

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surgical vs medical outcomes for more mildly affected dogs, impact of durotomy on locomotor outcome and development of progressive myelomalacia, and refining of postoperative care, and genetic and preventative care studies.

**Conclusions and Clinical Importance:** Future efforts should build on current recommendations by conducting prospective studies and randomized controlled trials, where possible, to address identified gaps in knowledge and to develop cost effectiveness and number needed to treat studies supporting various aspects of diagnosis and treatment of TL-IVDE.

**KEY WORDS**

dog, intervertebral disc herniation, paralysis, spinal cord injury

## 1 | INTRODUCTION

In chondrodystrophic dog breeds, expression of an fibroblast growth factor 4 (FGF4) retrogene on chromosome 12 is associated with dramatically accelerated intervertebral disc degeneration.<sup>1</sup> In these dogs, early chondroid metaplasia, degeneration, and mineralization of the nucleus pulposus (NP) occur with ultimate failure of the intervertebral disc unit and extrusion of mineralized material into the vertebral canal.<sup>2,3</sup> This process, called Hansen type I intervertebral disc extrusion (IVDE), also occurs in non-chondrodystrophic breeds at a much lower frequency, and unassociated with the FGF4 retrogene. As a consequence of contusion and compression of the spinal cord and nerve roots, thoracolumbar (TL-) IVDE causes pain, paresis, or paralysis and retention incontinence. Affected dogs are managed medically or surgically and, although outcomes are usually successful, substantial nursing challenges remain. Thoracolumbar IVDE has been recognized since the 1880s and is the most common cause of acute paraparesis and paraplegia in dogs because of the popularity of chondrodystrophic dog breeds.<sup>4,5</sup> As a result, there is much literature available, the majority describing observational studies. Notably, guidelines on management of the condition are lacking. In this consensus statement on TL-IVDE, we sought to review the available literature, weigh evidence, and generate recommendations for management and highlight areas in need of study. The paucity of high-level evidence for many aspects of managing this condition limits the power of our recommendations, which should be viewed as guidelines and not as standards of care.

## 2 | METHODS

A Qualtrics survey of the American College of Veterinary Internal Medicine (ACVIM)-Neurology membership was distributed in October 2020 to gather priorities related to a consensus statement on diagnosis and management of acute TL-IVDE in dogs. Eleven topics were identified, and subtopics were developed within them (Table 1). The consensus panel formed subgroups for systematic review of the literature in each area, and recommendations were formulated. A modified Delphi method was applied to the recommendations, where a

combination of face-to-face videoconferencing and online anonymous voting was used to modify the statements toward consensus. Complete consensus was achieved for all statements.

PubMed, Google Scholar, Web of Science, and CAB Direct were used along with the following search terms to identify relevant articles: Dog OR canine, intervertebral disc OR intervertebral disk, extrusion OR herniation OR disease OR Hansen type 1, TL. The group also added additional subtopic-relevant terminology. Only peer-reviewed studies containing original data were reviewed by the group. Although not included in the literature review, review articles were used to identify additional relevant papers not captured in the original searches. Papers were excluded if they were published only in abstract form, were not available in English, did not address TL-IVDE, or did not report initial severity of signs or outcome in an understandable way. Level of evidence provided by individual studies was defined by a standard approach ([https://www.elsevier.com/\\_data/promis\\_misc/Levels\\_of\\_Evidence.pdf](https://www.elsevier.com/_data/promis_misc/Levels_of_Evidence.pdf)). For each recommendation developed, the overall body of evidence supporting it was categorized as low, medium, or high level according to panel vote (Table 2). Recommendations are displayed in gray boxes throughout the manuscript, along with their level of evidence.

## 3 | DIAGNOSTIC APPROACHES

Diagnosis of TL-IVDE in dogs is made using magnetic resonance imaging (MRI), computed tomography (CT), CT-myelography, or myelography. These imaging modalities also facilitate surgical planning and prognostication. Each modality differs with respect to diagnostic accuracy, cost, availability, and adverse effects.

### 3.1 | Magnetic resonance imaging

Magnetic resonance imaging has a diagnostic sensitivity >98.5%,<sup>6-9</sup> with enhanced diagnostic performance over CT in dogs with peracute signs, and when differentiating disc extrusion from protrusion.<sup>8,10-12</sup> Residual compression after surgery also can be detected.<sup>13,14</sup>

**TABLE 1** Topics addressed by the ACVIM Neurology consensus panel on canine thoracolumbar intervertebral disc extrusion

Topic	Associated subtopics	No. papers included
Decision making in medical vs surgical management	Expected surgical vs medical outcomes based on severity, influence of level of pain on decision making, influence of recurrence on decision making	114
Components of medical management	Duration of exercise restriction, use of anti-inflammatory medications, analgesia in medical management, adjunctive treatments such as rehabilitation and acupuncture	26
Diagnostic approaches	Imaging modality of choice, standard MRI sequences, prognostic information and CT/myelogram/ MRI sequences, novel or specialized MRI applications	103
Surgical management	Evidence to support 1 approach over another, reported complications by approach, influence of timing on outcome, cut-off time point beyond which recovery is unlikely	88
Fenestration	Fenestration of an affected disc intra-op, fenestration of adjacent discs intra-op, risk vs benefit, impact of approach and technique on success, percutaneous laser disc ablation	38
Neuroprotective strategies	Impact of published treatments on locomotor outcome, adverse effects	46
Postoperative pain management	Current knowledge about postoperative pain, efficacy for published treatments, adverse effects	84
Urination	Current knowledge about recovery of urination, features of optimal management, expected outcomes, prevention and treatment of UTI	33
Rehabilitation therapy	Inclusion as standard therapy, timing of initiation and duration, exercises and modalities to include, cage rest and the balance with rehabilitation, use of mobility aids	21
Progressive myelomalacia	Clinical and histopathologic features, predisposing and protective factors, ante mortem diagnostic approaches	33

Note: Based on ACVIM membership responses to an initial Qualtrics survey, the consensus panel identified 10 topic areas and framed associated subtopics and questions around those to guide a systematic review of the veterinary literature.

Abbreviations: CT, computed tomography; MRI, magnetic resonance imaging; No, number; UTI, urinary tract infection.

**TABLE 2** Definitions of levels of evidence used by the consensus panel

Level of evidence	Definition applied
High	Multiple randomized controlled trials with concordant findings that find the same thing. The evidence strongly supports the conclusions.
Medium	Multiple retrospective studies with concordant findings, controlled trials, or single, small placebo-controlled trials that provide good evidence for a specific defined population, but not the wider population. The evidence suggests findings are likely to be real.
Low	Isolated or small retrospective studies, single non-controlled trials. The evidence suggests findings might be real.

Note: These definitions were used to qualify the body of literature available to support individual recommendations. Once the literature was synthesized to develop a summary document for each topic, and the subtopics that fell within it, the group was asked to vote on the strength of the evidence available to support each developed recommendation.

An additional benefit of MRI is prognostication. Presence and extent of intramedullary T2 hyperintensity, T2 hypointensity, and attenuation of the cerebrospinal fluid (CSF) signal on HASTE/T2\* sequences have been variably associated with worse locomotor outcome<sup>15-22</sup> and development of progressive myelomalacia (PMM).<sup>23-25</sup> However, magnet strength, image plane, and observer might influence reliability of T2 hyperintensity as a prognostic indicator.<sup>8,18</sup> Bladder and urinary outcomes are not associated with cross-sectional severity of spinal cord

compression, length of spinal cord compression or focal vs extensive pattern of disc extrusion on MRI studies.<sup>21,26-28</sup>

Limitations of MRI include prolonged acquisition time,<sup>29,30</sup> availability, and cost.<sup>31</sup> Standard protocols always include T2-weighted (T2W) images and can be abbreviated to focus on sagittal sequences,<sup>32-34</sup> but diagnostic accuracy suffers without transverse images.<sup>32</sup> The HASTE/T2\* or short Tau inversion recovery (STIR) sequences might complement T2W images,<sup>34,35</sup> whereas inclusion of post-contrast imaging does not improve diagnostic accuracy or treatment planning in most cases.<sup>36,37</sup> It remains to be determined if half-Fourier acquisition single-shot turbo spin-echo (HASTE) is complementary or additive to T2W images or only worthwhile if the T2W images do not show evidence of hyperintensity. Novel 3-dimensional fast spin-echo (3D FSE) MRI protocols might offer decreased scan time while preserving accuracy.<sup>38</sup> Although there is emerging evidence in dogs with acute TL-IVDE that specialized MRI applications, including diffusion tensor imaging (DTI) and magnetization transfer (MT), can provide information about injury severity and prognosis, they prolong scan time and require additional expertise, software and post-acquisition processing.

A minimum of T2W sagittal and transverse images should be acquired. HASTE and short Tau inversion recovery (STIR), T1-weighted (T1W) and T1W post-contrast sequences might be considered but do not serve as replacements for standard T2W imaging sequences. *Supported by moderate-level evidence.*

### 3.2 | Computed tomography

Computed tomography provides the benefit of rapid acquisition, lower cost and a diagnostic sensitivity of 81% to 100% in specific subgroups, namely chondrodystrophic dogs with mineralized discs.<sup>9,12,29,31,39-43</sup> It also can distinguish acute from chronically extruded mineralized disc material,<sup>41,44</sup> where evidence for this feature with MRI is limited.<sup>37</sup> It is less accurate in older (>5 years) and smaller dogs (<7 kg).<sup>8,10-12</sup> Computed tomography does not provide insight into the severity of parenchymal injury, and prognostic utility is therefore limited, but multilevel IVDE seen on CT had a worse prognosis than focal IVDE in 1 study.<sup>45</sup> Although MRI might predict a greater craniocaudal extent of extradural material compared to CT, both modalities provide similar ability to identify the length of compressive material.<sup>6,8,9,27,45</sup>

decisions (e.g., how long should the decompression be extended, whether to perform durotomy in deep pain-negative (DPN) dogs). *Supported by moderate-level evidence.*

### 3.3 | Myelography/CT-myelography

Diagnostic sensitivity of myelography or CT-myelography is 53% to 97%, and myelography is less accurate than other modalities for determining correct lateralization of extrusion for surgical planning.<sup>6,7,29,40,42,46-52</sup> Myelography also carries a risk of seizures, especially in large dogs or those receiving larger intrathecal contrast volumes.<sup>53-55</sup> Extensive<sup>56,57</sup> spinal cord swelling and infiltration of contrast medium within the spinal cord<sup>58</sup> are poor prognostic factors on myelographic studies.

Magnetic resonance imaging (over CT or myelography) can provide prognostic information in paraplegic dogs, especially paraplegic DPN dogs. Standard sequences, notably T2 sagittal and transverse, might be sufficient but inclusion of the HASTE can also provide prognostic information. *Supported by moderate-level evidence.*

## 4 | MEDICAL VS SURGICAL MANAGEMENT

Treatment of TL-IVDE can be medical (conservative) or surgical. Medical management involves restricted activity and analgesia, whereas surgical management most commonly involves spinal cord decompression by hemilaminectomy, with or without disc fenestration. Decision-making regarding selecting medical or surgical management depends on many factors. Conclusions related to this topic were drawn from a systematic review of cases published after 1983,<sup>59</sup> with data published before 1983 also considered, particularly in the context of medical management.

Magnetic resonance imaging, CT, CT-myelography or myelography are reasonable modalities for diagnosing TL-IVDE. When extruded material is mineralized, especially in young to middle-aged adult, chondrodystrophic dogs, CT is sensitive in diagnosis and treatment planning. It therefore can be recommended as a first-line advanced imaging modality when acute TL-IVDE is suspected, with a low likelihood of missing a compressive surgical lesion and the added benefit of shorter scan time and lower cost (compared to MRI). Magnetic resonance imaging provides definition of intramedullary lesions which can inform surgical approach and prognostication for dogs with severe injuries and has superior ability to make diagnoses other than IVDE. When considering cases outside this typical clinical presentation, there is evidence to support the highest diagnostic sensitivity for high field MRI and higher risk of adverse events with myelography or CT-myelography.

There is insufficient evidence to make recommendations regarding the role of imaging in dictating the type of decompressive surgery that should be performed (i.e., hemilaminectomy vs other) and if 1 imaging modality should be pursued over another for dogs with an inappropriate recovery shortly after decompressive surgery. Magnetic resonance imaging might be useful in guiding some treatment

#### 4.1 | Outcomes for medically vs surgically managed dogs

Expected outcomes for dogs with TL-IVDE managed medically or surgically are summarized in Table 3. Recurrence rates for medical management range from 15% to 66% compared with much lower rates for dogs managed by both hemilaminectomy and fenestration (covered in more detail under fenestration).<sup>60-66,72</sup> There is a group of dogs that present with severe spinal pain and lameness because of lateral extrusion of disc material into the intervertebral foramen for which little is currently published.<sup>65</sup> These dogs represent a unique group that might not respond to medical management.

Ambulatory dogs can be managed successfully medically; however, consideration should be given to risk of recurrence. Surgical management might be considered in a young, active dog with multiple mineralized discs, particularly with recurrent events. Surgical decompression can be considered when neurologic signs are progressive, unimproved or pain is persistent despite appropriate medical management. *Supported by moderate-level evidence.*

**TABLE 3** Outcome of dogs managed medically or surgically, based on severity of presenting signs

Injury severity	Medical outcome	Surgical outcome	Comments	References
Spinal pain only and ambulatory PP	80% (115 dogs)	98.5% (336 dogs)	Lateral extrusion of disc material may lead to reduced response to medical management.	<a href="#">60-70</a>
Non-ambulatory paraparesis	81% (131 dogs)	93% (341 dogs)	Level of recovery of non-ambulatory dogs was less complete with conservative management.	<a href="#">59,60,64,69</a>
Paraplegia DPP	60% (67 dogs)	93% (548 dogs)	Recovery with medical management is prolonged and less complete compared to surgery	<a href="#">59,64</a>
Paraplegia DPN	21% (48 dogs)	61% (502 dogs)	None	<a href="#">59,64</a>

Note: Literature prior to 1983 reports large numbers of dogs that were managed medically but frequently fails to separate paraplegia based on presence of deep pain perception. Rather, paraplegic dogs were grouped according to the presence or absence of “tonus”; these data were not included in this table. However, it is noted that 102 (65%) of 156 dogs that were paraplegic with tonus recovered with medical management, while 0 of 88 paraplegic without tonus dogs recovered.<sup>[60,67,69,71](#)</sup>

Abbreviations: DPN, deep pain negative; DPP, deep pain positive; PP, paraparesis.

Although non-ambulatory paraparetic or paraplegic deep pain-positive (DPP) dogs can be managed successfully medically, success rates, rate of recovery and chance of recurrence are more likely to be improved with surgery. Surgical management for these dogs is recommended. *Supported by moderate-level evidence.*

In paraplegic DPN dogs, success with medical management is largely poor with an increase in the frequency of PMM. Surgical management is recommended. *Supported by moderate to high-level evidence.*

Short-term hospitalization for 1-2 days should be considered in dogs with progressive signs managed medically, to monitor for deterioration in neurological status. *Supported by low-level evidence.*

When dogs cannot be treated surgically, a medical treatment attempt is reasonable for all grades of injury severity unless there is clinical evidence of PMM. *Supported by low to moderate-level evidence.*

## 4.2 | Components of medical management

Core components of medical management include activity restriction, pain management, treatment of retention incontinence, and

prevention and treatment of skin damage and decubital ulcers as needed. Few studies evaluate these components critically, making it impossible to generate evidence-based recommendations. Although treatment failures often are blamed on inadequate restriction, no published literature directly addresses this issue, and published recommendations for confinement range from absolute cage rest to “room rest” or “restricted activity” without details. Frequency and timing of re-evaluation are important considerations, but the literature here also is lacking. Pain management strategies have been studied more rigorously in the postoperative setting, the results of which might also be relevant for medical management.

### 4.3 | Duration of exercise restriction

Because of a lack of critical literature, many clinicians advise a period of decreased activity based on anecdote or institutional convention. Among studies that clearly state the duration of rest, several report a minimum but not total or average duration.<sup>[61,73,74](#)</sup> Others state a defined period of restriction, such as 4 weeks, but report variable degrees such as “absolute cage rest” and “room rest.”<sup>[64](#)</sup> In a retrospective case series in 223 dogs, duration of restricted activity ranging from none to >4 weeks was not associated with outcome.<sup>[63](#)</sup>

A least 4 weeks of restricted activity is recommended, putatively to promote healing of the annulus fibrosus. This period should include confinement to a restricted area (crate ideally, or a small room without furniture) except for when performing rehabilitation exercises or outdoor toileting. There should be no off-leash walking, no jumping on or off furniture and no access to stairs during this time. *Supported by low-level evidence.*

#### 4.4 | Use of anti-inflammatory medications

Several retrospective studies include a subgroup of dogs treated with anti-inflammatory doses of corticosteroids but do not report the effect of this treatment on outcome.<sup>61,64,67,73-77</sup> However, there is limited evidence that corticosteroid use is associated with poorer outcome and decreased quality of life<sup>63</sup> as well as a higher rate of recurrence compared to nonsteroidal anti-inflammatory drugs (NSAIDs) in ambulatory dogs managed medically.<sup>62</sup> Despite insufficient evidence to support corticosteroid use for neuroprotective purposes in dogs with TL-IVDE, anti-inflammatory doses of corticosteroids may be of benefit in some cases with ongoing spinal pain potentially related to epidural inflammation.<sup>66</sup>

Corticosteroids are not recommended for routine use in medical management of the acute phase of presumptive TL-IVDE. In the chronic phase, a short course of anti-inflammatory doses of corticosteroids may be of benefit for some dogs. *Supported by moderate-level evidence.*

No studies specifically evaluate the influence of NSAIDs on outcome of medical management of TL-IVDE, but several studies include dogs treated with NSAIDs as part of their management plan.<sup>60,62-64,77</sup> In a retrospective study, dogs receiving NSAIDs had higher quality of life scores than those receiving corticosteroids.<sup>63</sup> Additional detail on potential adverse effects associated with both corticosteroids and NSAIDs is available in the context of postoperative pain management.

The use of NSAIDs for at least 5-7 days is recommended in dogs managed medically for TL-IVDE, provided there is no specific contraindication. Requirement for analgesia beyond this time period, despite appropriate activity restriction, should be considered as a possible indication for further investigation and potentially surgical management. The use of concurrent corticosteroids or multiple NSAID formulations in combination should be avoided. *Supported by low to moderate-level evidence.*

#### 4.5 | Recommendations regarding analgesia

Most published information focuses on pain management in the post-operative setting.

Appropriate options for management of pain in dogs with TL-IVDE managed medically include an NSAID,

gabapentin or pregabalin for neuropathic pain and potentially muscle relaxants such as diazepam or methocarbamol. For dogs with pain severe enough to merit treatment with opioids, hospitalization should be recommended until that pain is controlled adequately. *Supported by low-level evidence.*

#### 4.6 | Adjunctive treatments such as rehabilitation and acupuncture

Several studies investigate acupuncture for treatment of TL-IVDE, with substantially variability in methodology, as well as lack of details on outcome measures and control group management protocol. These studies suggest that electroacupuncture could be associated with improved outcomes compared to medical management alone.<sup>73,74,76,77</sup> One study on electroacupuncture reported vomiting or diarrhea in 5 (16%) of 43 dogs, although dogs also received anti-inflammatory doses of prednisolone for 7 to 12 days<sup>73</sup>; most other studies do not report adverse event monitoring.<sup>74,76,77</sup>

A study including 82 dogs of variable injury severity identified a tendency toward improved outcome in conservatively treated dogs undergoing physical rehabilitation.<sup>75</sup> Most recent research into rehabilitation in dogs with TL-IVDE pertains to postoperative management.<sup>78-80</sup> Although frequency of adverse events associated with physical rehabilitation is low,<sup>80</sup> injury prevention is mandatory when considering a rehabilitation program.

Although there is a low level evidence to support the use of acupuncture as a component of medical management of TL-IVDE, this treatment option currently is not recommended as an alternative to surgical management. Despite limited evidence to support the use of physical rehabilitation in medically treated dogs, basic rehabilitation exercises are recommended as an additional treatment (e.g., passive range of motion exercises and massage), with an emphasis on restricted activity for at least 4 weeks, followed by increased levels of physical activity. *Supported by low-level evidence.*

#### 5 | SURGERY

Several surgical approaches for dogs with TL-IVDE have been published. These include hemilaminectomy, mini-hemilaminectomy/pediculectomy, dorsal laminectomy, partial corpectomy, fenestration of the

intervertebral disc with or without concurrent laminectomy for removal of extruded disc material, and laminectomy with concurrent durotomy. Few studies provide direct comparisons of 1 surgical technique to another.<sup>13,81-84</sup> Descriptions of the various techniques and their associated benefits and complications have been summarized previously.<sup>85</sup>

In addition, several authors recently have reported minimally invasive techniques to address intervertebral disc herniation in dogs. To date, most published veterinary studies are cadaver studies describing specific approaches, and only a few report outcomes in the clinical setting without employing a control group<sup>86-92</sup>; however, there is a growing body of literature supporting their potential utility, which might gain future popularity.

The rationale for the addition of durotomy at the time of surgical decompression is a topic of re-emerging interest and stems initially from experimental literature suggesting that durotomy could improve spinal cord blood flow, that the procedure was safe and that functional recovery in dogs with spinal cord injury (SCI) might be better with the addition of durotomy as compared to laminectomy alone.<sup>93-95</sup> Several veterinary clinical studies have evaluated the impact of durotomy on spinal cord perfusion or on outcome for dogs with TL-IVDE that underwent laminectomy with durotomy, with mixed results.<sup>96-99</sup> One recent study retrospectively reported the outcomes of 51 paraplegic DPN dogs treated by hemilaminectomy with durotomy compared to 65 dogs treated by hemilaminectomy alone.<sup>98</sup> In that study, dogs that underwent durotomy were 3.32 times more likely to regain ambulation compared those treated by hemilaminectomy alone. No cases of PMM were observed in the durotomy group, but 21.5% of dogs treated with hemilaminectomy alone died of presumed PMM. An additional recent study prospectively described a cohort of 26 consecutive paraplegic dogs presented with loss of deep pain sensation treated by laminectomy and durotomy, for which 22 had postoperative follow-up. Sixteen (72%) of 22 dogs recovered independent ambulation and only 1 (5%) dog developed PMM.<sup>99</sup> These more recent veterinary studies, along with the ongoing DISCUS trial (<https://discus.octru.ox.ac.uk>) in humans highlight renewed interest in the role of durotomy to improve outcome after SCI.

Each disc extrusion is different and therefore requires unique consideration for the best approach. Hemilaminectomy and mini-hemilaminectomy (with or without concurrent fenestration) typically are considered the surgical approaches of choice because of increased ability to access and remove compressive disc material. Durotomy for dogs with severe neurologic signs may improve outcome and lessen risk of PMM, although further evaluation in a large cohort of dogs is needed. Lateral corpectomy has utility for more chronic or very ventrally located disc extrusions. Fenestration alone, without decompression, is not recommended for dogs with severe (non-ambulatory paraparesis or worse) IVDE. Minimally invasive approaches appear safe and feasible but require further evaluation before they can be routinely adopted. *Supported by moderate-level evidence.*

## 5.1 | Timing of surgical decompression

Whether urgent surgical decompression improves neurologic recovery in dogs with TL-IVDE has been controversial for decades. There are several challenges with the currently available literature addressing this question. First, a positive outcome usually is defined dichotomously and simply as the ability to ambulate without support (or not), rather than assessing residual neurologic deficits using finer locomotor scoring systems. Second, dogs with the most severe disc-associated SCI (paraplegic DPN) represent a small number in any given study and frequently are grouped with less severe injuries. Third, the extent of time delay before presentation often is not well reported by owners, making it difficult to draw firm conclusions on timing of onset of signs, particularly loss of deep pain perception. Interpretation of the current literature is complicated further by the fact that outcome also might be influenced by the rate of onset and features of the injury itself, independent of timing of decompression. Finally, study design and definition of successful outcome differ across studies, making it difficult to make comparisons.

Several retrospective and prospective case series have reported neurologic outcome and its association with surgical timing.<sup>15,100-110</sup> Taken individually, each study represents relatively low-level evidence to inform clinical practice, and overall, the literature provides mixed evidence with respect to the influence of timing of surgical decompression on outcome in dogs with less severe injury, where most larger and more recent studies suggest a lack of association (Table 4). The suggestion that surgical decompression must be performed within 24 hours for paraplegic DPN dogs (a time-point sometimes treated as a “cut-off” by clinicians), in order to achieve a possibility of return of ambulation has been addressed by several studies. Although 1 study reported that no dogs in a very small group that had been DPN for >24 hours recovered ambulation,<sup>105</sup> a number of dogs reported elsewhere to have been paraplegic DPN for as much as a week or more before surgery went on to recover ambulation.<sup>15,104,108</sup> Publications addressing this topic generally are limited by their retrospective or prospective observational nature, because a prospective randomized controlled trials (RCTs) evaluating timing of surgical decompression, particularly in paraplegic DPN dogs, might be viewed as unethical.

The pathogenesis of IVDE-induced SCI supports that surgical decompression ought to be performed as early as possible for dogs with substantial neurologic deficits. However, taken as a whole, the current literature does not generally support the use of a specific timeline for urgent surgical decompression, even for DPN dogs that are paraplegic at the time of presentation. Some evidence suggests that delayed decompression may result in a longer time to achieve postoperative ambulation, which requires further investigation. The influence of surgical timing on locomotor outcome should be considered independently of its influence on development of PMM. *Supported by moderate-level evidence.*

**TABLE 4** Studies presenting data regarding the influence of surgical timing on outcome in dogs with thoracolumbar intervertebral disc extrusion

Studies suggesting no influence			Studies suggesting possible influence		
Ref	N	Study design/outcome	Ref	N	Study design/outcome
111	98	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• No difference between surgery &lt;12 h and 12- 36 h after losing ability to walk</li> </ul>	112	22	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Suggests better outcome if operated within 4 d</li> <li>• Statistics not performed</li> </ul>
101	71	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• No association between duration of clinical signs prior to surgery and outcome</li> <li>• Clinical signs of ≥6 d had longer recovery</li> </ul>	113	99	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Trend for better outcome if operated within 4 d</li> <li>• Statistics not performed</li> </ul>
102	70	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Duration of signs or of DPN were not associated with recovery</li> </ul>	114	187	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Outcome better if surgery ≤48 h of onset of paresis</li> </ul>
103	112	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Duration of non-ambulatory paraparesis was not associated with recovery</li> </ul>	57	46	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Better outcome at hospital discharge for dogs DPN for &lt;12 h prior to surgery</li> <li>• Statistics not performed</li> </ul>
104	30	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Duration of signs was not associated with recovery</li> </ul>	115	32	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Paradoxically, the success rate was higher for cases with longer duration of signs</li> </ul>
15	77	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Presumptive duration of DPN status not associated locomotor outcome</li> </ul>	105	46	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Better outcome in paraplegic DPN dogs if surgery &lt;24 h after losing deep pain</li> </ul>
116	36	<ul style="list-style-type: none"> <li>• Prospective</li> <li>• Duration of non-ambulatory status before surgery did not influence recovery</li> </ul>	106	28	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Rate of recovery of ambulation was higher if surgery &lt;24 h after onset of clinical signs</li> </ul>
108	78	<ul style="list-style-type: none"> <li>• Prospective</li> <li>• Time between onset of signs and referral evaluation was not associated with locomotor recovery</li> </ul>	107	197	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• No influence on locomotor recovery</li> <li>• Risk of development of PMM 3.1X higher with delayed surgery</li> </ul>
107	197	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Delayed surgery in paraplegic DPN dogs &gt;12 h after loss of ambulation not associated outcome at 6 m</li> </ul>	109	273	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Number of dogs who lost deep pain sensation was higher with delayed decompression than same-day surgery</li> <li>• Likelihood of regaining pain sensation at 3 wk was higher with early vs delayed surgery</li> </ul>
117	131	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Timing of decompression not associated with outcome</li> </ul>	110	1501	<ul style="list-style-type: none"> <li>• Retrospective</li> <li>• Time between onset of signs and surgery was significantly associated with overall outcome using a multivariable regression model, but not in bivariate analysis</li> <li>• May suggest timing itself did not influence outcome, but rather combination of timing with other factors</li> </ul>

Note: The literature provides mixed evidence with respect to the influence of timing of surgical decompression on outcome in dogs with less severe injuries, where the predominance of larger and more recent studies suggest a lack of association.

Abbreviations: DPN, deep pain negative; DPP, deep pain positive; N, number; PMM, progressive myelomalacia.

Surgical treatment should not be declined simply because the dog has been paralyzed for an extended period, because the literature documents that recovery of ambulation may occur for dogs that present with DPN status and have been paralyzed > 1 week before surgery. *Supported by moderate-level evidence.*

## 6 | FENESTRATION

Prophylactic fenestration refers to removing disc material *in situ* with the goal of preventing future extrusion. When first described as a surgical technique, fenestration was used as a means of treating disc extrusion without concurrent laminectomy; however, fenestration is now most commonly performed at the site of extrusion, along with

decompression, to lessen further extrusion through the ruptured annulus in the early postoperative period. Fenestration also may be performed at adjacent disc spaces, typically between T11 and L4, as a prophylactic measure. Fenestration is an advanced procedure that requires excellent skill and knowledge of local anatomy to perform effectively and safely. Several surgical approaches and techniques have been described but direct comparisons are challenging because methods for reporting recurrence vary among studies and recurrences are not always confirmed by imaging or surgery. Finally, surgeon experience is likely to influence outcome, and RCTs in this area are difficult to design. The evidence for fenestration can be considered in the context of minimizing further extrusion of an already extruded disc, and in preventing future extrusion of non-ruptured discs.

## 6.1 | Fenestration of an already extruded disc at the time of decompressive surgery

Early recurrence has been reported within the first month after decompression and is almost exclusively caused by extrusion of additional disc material at the site of previous decompression.<sup>71,118-120</sup> Although some studies report few early recurrences without fenestration,<sup>100,121</sup> most published studies support that fenestration of the extruded disc at the time of decompression prevents early recurrence.<sup>72,118,119,122-125</sup> The overall low rate of recurrence at the site of decompression regardless of fenestration status may be related to much of the disc having extruded and been removed by decompression, differences in length and quality of follow-up, how the definition of recurrence is applied and confirmed, and what endpoints are used.

Fenestration of the herniated disc space at the time of surgical decompression is recommended to minimize risk of recurrence at the site of herniation. *Supported by moderate-level evidence.*

## 6.2 | Prophylactic fenestration of adjacent or distant, non-extruded discs

A separate consideration is the impact of fenestration on future extrusion at new disc spaces, which is reported in up to 19% of all dogs undergoing decompressive surgery.<sup>72,100,118,121,123-125</sup> Most (87.5%) of these late recurrences develop within 1 to 2 disc spaces of the original extruded disc,<sup>121,124</sup> with rates of recurrence being higher for dachshunds (25%<sup>121</sup>) and French bulldogs (44%<sup>126</sup>). Although more extensive prophylactic fenestration remains controversial, several studies have identified a protective effect.<sup>72,123,124,127</sup> For instance, recurrence rates have been reported to be 17.89% with single site fenestration and 7.45% with multiple site fenestration,<sup>124</sup> where dogs with single site fenestration were 2.7 times as likely to develop

confirmed recurrence than those with multiple fenestration.<sup>124</sup> The odds of recurrence at a non-fenestrated disc space were 5.86 times that of a fenestrated site,<sup>123</sup> and the recurrence rate at a non-fenestrated site was 11 times that of a fenestrated site.<sup>124</sup> Similarly, other authors reported that the prevalence of recurrence at non-fenestrated discs was 26.2 times that of fenestrated discs.<sup>127</sup> Additional considerations for prophylactic fenestration include breed, where dachshunds<sup>121,123</sup> and French bulldogs<sup>126</sup> are more prone to recurrences than other breeds. The presence and number of mineralized discs at the time of decompression also are positively associated with rate of recurrence.<sup>121,124,128</sup>

How many, and which, additional discs to fenestrate is difficult to address based on the currently available literature. Several studies support fenestrating adjacent mineralized discs accessible through the same incision because most recurrences develop within 1 to 2 disc spaces of the original extrusion.<sup>121,124</sup> This consideration may be particularly relevant for at-risk breeds. Mineralized discs cannot be specifically identified on MRI, but discs that appear degenerate on MRI have increased risk of recurrence if not fenestrated.<sup>128</sup>

Fenestration of adjacent, degenerated but non-ruptured disc spaces, should be considered. In breeds predisposed to IVDE such as dachshunds and French bulldogs, fenestration is recommended even if the discs are not mineralized. The decision to fenestrate must take into account the status of the dog, surgical time, and other relevant factors. In situations where decompression is not performed, prophylactic fenestration could be considered. *Supported by moderate-level evidence.*

## 6.3 | Adverse events, approach, and technique

Potential arguments against fenestration include the possibility of additional surgical trauma, increased anesthetic time, and associated costs. Risks or complications associated with fenestration that have been reported in the veterinary literature include extrusion of additional disc into the vertebral canal at the site of extrusion,<sup>127,129</sup> the potential to induce vertebral instability,<sup>130,131</sup> increased morbidity when performed at L5-6 & L6-7,<sup>124,132</sup> pneumothorax<sup>68,123,133,134</sup> or hemothorax (chemonucleolysis<sup>123</sup>), neuromuscular complications secondary to trauma to the peripheral nerve or nerve root,<sup>124,134</sup> hemorrhage from the sinus or vertebral artery,<sup>68,124</sup> and development of spondylosis deformans.<sup>135</sup> Additionally, and specific to laser disc ablation, abscess and discospondylitis have been reported.<sup>134</sup> The overall reported complication rate associated with various fenestration techniques is quite low, and a recent survey of the literature reported a published complication rate of 0.01%, or only 15 instances in over 1100 published cases.<sup>85</sup> Cost benefit analysis of the additional anesthetic and surgical time required for fenestration and the clinical relevance of any additional tissue trauma is not addressed by the currently available literature.

Multi-site fenestration is associated with an overall very low complication rate, rarely leading to clinically relevant morbidity when performed by experienced surgeons. Reported complications associated with fenestration cranial to the L4-5 disc are limited and are not a threat to life or mobility. Caudal to L3-4, fenestration carries increased risk; therefore, routine fenestration of L4-5 and more caudal sites is not recommended. Fenestration at T10-11 and above usually is not recommended due to the low rate of disc extrusion at these sites. *Supported by moderate-level evidence.*

Percutaneous laser disc ablation (PLDA) appears to be safe and, based on a low-level of evidence in a large number of dogs, may limit recurrent disc extrusion. *Supported by low to moderate-level evidence.*

Some evidence supports that surgical approach influences retrieval of disc material, with a ventral or lateral approach providing better success than a dorsolateral approach.<sup>135-137</sup> However, most of these studies involved fenestration alone. Thus, the practicality of a dorsolateral approach for fenestration must be considered when a decompressive laminectomy also is being performed.

Technique has been shown to influence yield of disc material during fenestration, where power-assisted fenestration resulted in more complete removal of the NP compared to manual fenestration.<sup>119,122</sup> However, these studies involved a single surgeon and therefore the results potentially are user dependent. Cavitronic ultrasonic surgical aspirator (CUSA) and vacuum aspiration also are described as options in the literature.<sup>132,138</sup> Importantly, the literature suggests that complete removal of the NP is not achieved by any technique or approach, but that the effectiveness of fenestration is governed by the amount of NP removed at surgery.<sup>139</sup> Additionally, an inflammatory reaction that might be hypothesized to dissolve the remaining nucleus does not occur.<sup>139</sup> The void created by fenestration and the annulus defect heals relatively quickly (4-16 weeks) by invasion of fibrocartilage after fenestration.<sup>140</sup> Although some fenestration techniques may result in more removed NP, the clinical relevance of such differences with respect to recurrence has yet to be evaluated.

Insufficient evidence exists to support a single surgical approach to fenestration over another, and surgical approach tends to be dictated by the decompressive procedure and surgeon experience. Creating a fenestration without subsequently curetting NP from the disc space does not lead to removal of disc material by an inflammatory response. Although power fenestration has been shown to improve yield over manual (blade) fenestration, complete fenestration is not achieved by any technique, and the relationship between increased NP removal and recurrence after fenestration has not been explored. *Supported by low-level evidence.*

With respect to percutaneous laser disc ablation (PLDA) as a unique approach, the group made the following recommendation:<sup>134,141,142</sup>

## 7 | NEUROPROTECTIVE STRATEGIES

Evaluation of cell-based treatments was considered beyond the scope of this discussion. Many neuroprotective strategies have been proposed to mitigate secondary injury or promote recovery of neuronal function or functional connectivity. A subset of these strategies evaluated in the veterinary clinical setting have evidence suggesting a lack of effectiveness to improve locomotor outcome (Table 5); their routine use therefore is not currently advocated.

Despite conflicting findings, in a double-blind RCT in paraplegic DPN dogs undergoing concurrent surgical decompression, dogs treated with the matrix metalloproteinase (MMP) inhibitor GM6001 with a dimethyl sulfoxide (DMSO) carrier or DMSO alone showed significantly improved locomotor outcome compared to those treated with a saline control. Although these findings suggest that DMSO might be the primary influence on improved outcome, group sizes were small and no dogs in the saline-treated group recovered locomotor function.<sup>157,166</sup>

When used in combination with decompressive surgery for dogs that are paraplegic DPN at the time of presentation, DMSO might improve locomotor outcome. Routine administration of DMSO, although currently not recommended as standard adjunctive neuroprotective treatment for acute TL-IVDE, deserves further evaluation in a larger cohort. *Supported by moderate-level evidence.*

## 8 | PAIN MANAGEMENT

It is accepted that both TL-IVDE and its surgical treatment cause pain. Nociceptive pain (perception of a noxious stimulus processed by a normally functioning somatosensory system) is caused by both the disc extrusion and decompressive surgery and is the primary target of many treatment strategies. Neuropathic pain, caused by a lesion that leads to damage or dysfunction of the somatosensory system,<sup>167</sup> also is known to be a

**TABLE 5** Neuroprotective strategies not currently recommended for routine treatment of TL-IVDE

Intervention	References	Comments
Pulsed electromagnetic field (PEMF) therapy	<a href="#">143,144</a>	<ul style="list-style-type: none"> <li>Possible positive effects on wound healing and pain control</li> </ul>
Electroacupuncture (EA)	<a href="#">73,145-151</a>	<ul style="list-style-type: none"> <li>Might facilitate functional improvement</li> <li>Validity of results is influenced by study design and reporting</li> <li>Might improve perioperative pain control</li> </ul>
Photobiomodulation/Laser	<a href="#">78,116,152-154</a>	<ul style="list-style-type: none"> <li>There is conflicting evidence from 2 unblinded studies.<a href="#">116,152</a></li> <li>Cadaveric study reported lower irradiance at the level of the spinal cord than has been proposed as necessary to modulate the area after SCI.</li> </ul>
Polyethylene glycol (PEG); intravenous	<a href="#">155,156</a>	<ul style="list-style-type: none"> <li>There is continued interest from an experimental perspective, using different routes of administration</li> </ul>
GM6001 (MMP inhibitor)	<a href="#">157,158</a>	<ul style="list-style-type: none"> <li>Improved bladder compliance compared to placebo</li> <li>Potential for reduced incidence of urinary tract infection</li> </ul>
Dexamethasone	<a href="#">159-161</a>	<ul style="list-style-type: none"> <li>Increased risk of urinary tract infection and gastrointestinal disease</li> <li>Reduced quality of life in dogs managed medically for IVDE</li> </ul>
Methylprednisolone sodium succinate (MPSS)	<a href="#">155,159,162-164</a>	<ul style="list-style-type: none"> <li>No adverse events were described in 1 study (pretreatment with steroids or NSAIDs excluded)</li> <li>Experimental administration has resulted in gastric hemorrhage or suppressed neutrophil function in healthy dogs</li> </ul>
Other steroid (eg, prednisone)	<a href="#">82,107,108,159,165</a>	<ul style="list-style-type: none"> <li>References <a href="#">82</a> and <a href="#">165</a> cite potential benefit but missing comparative aspects hamper conclusions.</li> <li>Possible decreased risk of PMM for dogs receiving corticosteroids</li> </ul>

Note: Existing veterinary clinical evidence suggests these are not effective at improving locomotor outcome. Some may have evidence supporting their utility in other aspects of thoracolumbar intervertebral disc extrusion (noted where applicable).

Abbreviations: NSAID, nonsteroidal anti-inflammatory; PMM, progressive myelomalacia; SCI, spinal cord injury.

common phenomenon in people after SCI; however, limited data are available on its occurrence in dogs. Pain after TL-IVDE has been quantified in dogs using ordinal pain scores, frequency of rescue medication use, and mechanical sensory thresholds (MST).[143-146,168-175](#)

reluctance to do certain activities, flinching when touched) for up to 6 weeks. *Supported by low-level evidence.*

## 8.1 | Postoperative pain

The prevalence and time course for resolution of paraspinal pain in dogs after surgical decompression treated using a standard postoperative analgesic protocol (opioids, anti-inflammatory drugs and gabapentin) have been quantified prospectively.[176](#) This analgesic protocol controlled immediate postoperative pain based on ordinal pain scales. However, MSTs were decreased for up to 6 weeks in the vicinity of the surgical wound. These values normalized by 6 months in 85% of dogs, but 15% had persistent hyperesthesia suggesting chronic neuropathic pain.[176](#)

Dogs can experience surgical site discomfort for up to 6 weeks after hemilaminectomy and a small percentage might develop chronic neuropathic pain. The need for prolonged treatment of pain is unclear, but veterinarians should evaluate dogs for pain at the time of first re-evaluation and advise owners to observe for signs of pain (e.g., vocalizing,

## 8.2 | Management of postoperative pain

No standardized protocol for postoperative pain management is available, but based on studies that report specific protocol details, combinations of opioids and anti-inflammatory drugs are used frequently.[28,91,97,102,119,144,176-178](#) The impact of adding various additional treatments to an existing analgesic protocol has been examined in RCTs and case cohort retrospective studies (Table 6). No studies compare NSAIDs, corticosteroids or different opioids. Numerous other treatments commonly are used, many of which have no published data on efficacy in the context of TL-IVDE. (Table 7).

Given the breadth of analgesic options published for use in dogs, coupled with variability in study designs and outcome assessments, it is difficult to compare findings across studies. However, the following postoperative analgesic protocol is proposed: IV or SC opioids for 24 to 48 hours postoperatively (longer if needed), fentanyl patch for 3 to 5 days

**TABLE 6** Details of studies investigating different intra- and postoperative pain therapies

Paper	Design	Treatment	Findings
171	RCT, n = 26	Epidural morphine	Better postoperative (48 h) pain control
168	RCT, n = 12		
172	RCT, n = 30	Epidural morphine & dexmedetomidine & hydromorphone	
174	Retrospective case cohort study n = 114	Erector spinae block	Better intra- and postoperative (48 h) pain control
175	RCT, n = 30		Better intra- and postoperative (24 h) pain control
169	Prospective, n = 10	Fentanyl patch	Plasma levels therapeutic, pain control adequate (72 h)
173	RCT, n = 46	Adjunctive Pregabalin 4 mg/kg q8h	Better pain control for 5-d postop
170	RCT, n = 63	Adjunctive Gabapentin 10 mg/kg q12h	No benefit over placebo for 5-d postop
145	RCT, n = 15	Postoperative electroacupuncture	No benefit for 3-d postop
146	RCT, n = 24	Preoperative acupuncture	Reduced intraoperative need for fentanyl, reduced pain on recovery from anesthesia
144	RCT, n = 16	Pulsed electromagnetic fields	Reduced postoperative pain from 2 to 6 wk
143	RCT, n = 53	Pulsed electromagnetic fields	Reduced postoperative pain for 6 wk
178	RCT, n = 20	Harmonic blade for surgical approach	Reduced postoperative pain for 30 d
91,92	a: Controlled trial (normal dogs) n = 6, b: case report, n = 1	Minimally invasive surgery	Reduced need for opioids postoperatively Reduced postoperative pain for 7 d

Note: Various analgesic protocols have been examined in RCT and case cohort retrospective studies.

Abbreviation: RCT, randomized controlled trial.

**TABLE 7** Therapies for postoperative pain reported but not investigated for efficacy

Therapeutic	Paper
Tramadol	14,99,179
Amantadine	180,181
Ketamine and lidocaine	77
Bee venom	77
Cold laser therapy	78
Cryotherapy	176
Massage	176
Therapeutic ultrasound	116
Methocarbamol/Diazepam	176

decrease intra- and postoperative pain and their use can be considered. *Supported by low-level evidence.*

### 8.3 | Adverse effects of pain medications

Medications used for pain control are known to have potential adverse effects. However, it is difficult to determine the relative contribution of pain medications to the development of complications in the setting of disc-associated SCI, hospitalization, anesthesia, and surgery.<sup>182</sup>

Many dogs with IVDE treated with corticosteroids alone or in combination with NSAIDs develop subclinical or clinical adverse gastrointestinal (GI) effects such as diarrhea, gastritis, GI ulceration, regurgitation and pancreatitis.<sup>159,170,183-188</sup> They occur at comparable rates in dogs treated with NSAIDs or corticosteroids alone, but at higher rates if these drug classes are combined.<sup>182</sup> However, fatal colonic perforations have been described in dogs treated with high doses of dexamethasone,<sup>189-191</sup> and dogs treated with dexamethasone are more likely to develop urinary tract infections (UTIs) and diarrhea than those treated with other or no glucocorticoids.<sup>159</sup> Although high doses of methylprednisolone sodium succinate (MPSS) cause GI bleeding on endoscopy,<sup>162</sup> 1 RCT did not encounter a higher rate or severity of GI signs in dogs receiving high dose MPSS when

postoperatively, NSAIDs for 7 days postoperatively in addition to or instead of a fentanyl patch. A medication for neuropathic pain such as pregabalin at a dosing interval of q8h might be added to the suggested protocol to improve pain control. A similar positive effect could not be shown for gabapentin administered q12h. Gabapentin might be beneficial at a dosing interval of q8h, although evidence to support that assumption currently is lacking. In addition, pre- and intraoperative interventions such as erector spinae block, epidural morphine as well as postoperative pulsed electromagnetic field therapy (PEMF) have been proven to

compared with 2 other study arms, although dogs that already had received an NSAID or other corticosteroid were excluded from this trial.<sup>155</sup> Different GI protectants have been used to decrease or prevent GI signs, but studies suggest that these are not effective.<sup>182,185</sup>

Opioids can increase the risk of gastro-esophageal reflux and regurgitation,<sup>192,193</sup> as can medications that cause sedation such as gabapentin and pregabalin. Vomiting and regurgitation are risk factors for aspiration pneumonia, which has been reported in up to 6.8% of dogs treated surgically for IVDE.<sup>20,194</sup>

Gastrointestinal mucosal lesions are common in dogs with IVDE, but often are subclinical. It is not known if these lesions are caused by the disease itself, the medications used or a combination of both. Life-threatening GI lesions have been reported in dogs treated with high doses of dexamethasone or with combinations of NSAIDs and glucocorticoids. If GI signs are present, NSAIDs or glucocorticoids should be discontinued. The use of opioids should be considered as a potential contributing factor to regurgitation.

*Supported by low-level evidence.*

## 9 | MANAGEMENT OF URINATION

One of the most challenging consequences of TL-IVDE is loss of voluntary urination and the resulting retention incontinence. This complication negatively impacts overall recovery and the quality of life of both dogs and owners.<sup>195,196</sup>

### 9.1 | Recovery of voluntary urination

Data that summarize the rate and success of recovery of voluntary urination are provided in Table 8. Ultrasonography and cystometrography indicate that recovery of normal bladder function lags behind the initial appearance of voiding and still can be suboptimal 6 weeks after motor recovery.<sup>158,198</sup>

Paraplegic dogs cannot urinate voluntarily, whereas dogs with motor function and pain perception can. As voiding recovers, it is commonly incomplete for a period of weeks post-injury, resulting in urine retention. This complication places dogs at higher risk than usual for development of

**TABLE 8** Studies describing timing and level of recovery of voluntary urination

Paper	Design	Outcomes	Findings																				
<i>Timing and quality of recovery of voiding</i>																							
198	Prospective (n = 15)	Voiding efficiency via US	<ul style="list-style-type: none"> <li>Non-ambulatory PP: normal within 4 d</li> <li>DPP: urinating by 6 d, increased residual volume</li> <li>DPN: large residual volumes at 25 d</li> </ul>																				
197	Prospective (n = 26)	Onset of urination	<ul style="list-style-type: none"> <li>Non-ambulatory PP, DPP, DPN dogs: 31% urinating within 24 h, 92% by 1 mo</li> </ul>																				
158	Prospective case (n = 20), cohort (n = 10)	Cystometry	<ul style="list-style-type: none"> <li>Bladder parameters in DPP and non-ambulatory PP are abnormal at 7 d</li> <li>Max bladder P and p void (leakage point) are abnormal at 42 d</li> </ul>																				
199	Retrospective (n = 48 T3-L3, 48 L4-S3)	Urination recovery	<ul style="list-style-type: none"> <li>Rate of urination recovery at 3 wk was lower with L4-S3 lesions than T3-L3 lesions</li> </ul>																				
28	Retrospective (n = 57)	Urination recovery	<ul style="list-style-type: none"> <li>Ambulatory PP: 1.9 d to urination; Non ambulatory PP: 2.9 d to urination</li> <li>Paraplegic DPP: 6 d to urination; DPN: 15.5 d to urination</li> </ul>																				
144	PEMF RCT (n = 16)	DPN dogs, urination recovery	<ul style="list-style-type: none"> <li>6 of 11 dogs that recovered DP urinating at 14 d</li> </ul>																				
<i>Rate of long-term urinary and fecal incontinence</i>																							
102	Retrospective (n = 64)	Long-term incontinence, DPN dogs	<p>37 dogs recovered deep pain perception: 32%—UI &amp; FI; 41%—FI 18 dogs persistent DPN: 100%—UI &amp; FI</p> <table border="1"> <thead> <tr> <th></th> <th>UI</th> <th>FI</th> <th>Both</th> </tr> </thead> <tbody> <tr> <td>Amb: 270</td> <td>15/5.6%</td> <td>9/3.3%</td> <td>6/2.2%</td> </tr> <tr> <td>Non-amb PP: 171</td> <td>9/5.3%</td> <td>5/2.9%</td> <td>3/1.8%</td> </tr> <tr> <td>DPP: 158</td> <td>26/16.5%</td> <td>14/9%</td> <td>10/6%</td> </tr> <tr> <td>DPN: 110</td> <td>42/38.2%</td> <td>20/18.2%</td> <td>16/14.5%</td> </tr> </tbody> </table>		UI	FI	Both	Amb: 270	15/5.6%	9/3.3%	6/2.2%	Non-amb PP: 171	9/5.3%	5/2.9%	3/1.8%	DPP: 158	26/16.5%	14/9%	10/6%	DPN: 110	42/38.2%	20/18.2%	16/14.5%
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125	Retrospective (n = 709)	Presence of long-term incontinence	<table border="1"> <tbody> <tr> <td>Amb: 270</td> <td>15/5.6%</td> <td>9/3.3%</td> <td>6/2.2%</td> </tr> <tr> <td>Non-amb PP: 171</td> <td>9/5.3%</td> <td>5/2.9%</td> <td>3/1.8%</td> </tr> <tr> <td>DPP: 158</td> <td>26/16.5%</td> <td>14/9%</td> <td>10/6%</td> </tr> <tr> <td>DPN: 110</td> <td>42/38.2%</td> <td>20/18.2%</td> <td>16/14.5%</td> </tr> </tbody> </table>	Amb: 270	15/5.6%	9/3.3%	6/2.2%	Non-amb PP: 171	9/5.3%	5/2.9%	3/1.8%	DPP: 158	26/16.5%	14/9%	10/6%	DPN: 110	42/38.2%	20/18.2%	16/14.5%				
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Abbreviations: DP, deep pain; DPN, deep pain negative; DPP, deep pain positive; FI, fecal incontinence; n, number; PP, paraparetic; UI, urinary incontinence; US, ultrasound.

UTI. Dogs with severe injury, even if they recover pain perception and motor function, might have suboptimal continence. *Supported by moderate-level evidence.*

## 9.2 | Bladder evacuation

Techniques used for bladder evacuation include manual expression and intermittent and indwelling catheterization. Efficacy of manual expression varies widely with a mean of 50% bladder emptying.<sup>200</sup> Although the method of bladder evacuation used does not influence the frequency of UTI, prolonged placement of indwelling catheters might delay recovery of voluntary urination and increase the risk of UTI.<sup>201,202</sup> No information is available on the impact of technique on factors such as pain or distress or regarding complications such as urethral or bladder damage.

Bladder expression technique chosen should be tailored to the dog, the clinician and the technicians caring for the dog. Placement of an indwelling urinary catheter is an effective and low risk management method in the short term; however, duration of indwelling catheterization should be minimized whenever possible because of increased risk of UTI. As voluntary urination returns, the bladder should be palpated, or ultrasonography performed, to ensure adequate emptying has been achieved. *Supported by moderate-level evidence.*

## 9.3 | Pharmacologic intervention

Use of drugs to relax urethral sphincters is common practice, but literature on efficacy is extremely limited. Details provided in clinical trials and retrospective studies were used to summarize standard protocols for dogs undergoing manual expression.<sup>144,158,197,199,203</sup> These protocols aim to relax the internal and external urethral sphincters using 1 or both of an alpha-adrenergic antagonist (eg, prazosin, phenoxybenzamine, alfuzosin, tamsulosin) and a centrally acting muscle relaxant (eg, diazepam). Experimental studies characterize the effects of alpha-adrenergic antagonists on urethral tone, but clinical relevance in dogs with alterations in urethral tone is difficult to establish.<sup>204</sup> Apart from a single retrospective study,<sup>205</sup> no data validate clinical efficacy. Although hypotension is a possible adverse effect of alpha-adrenergic antagonists, no reports of clinically relevant hypotension in this population have been published. Based on the literature, bethanechol is not used to enhance detrusor activity routinely in dogs recovering from acute TL-IVDE. Some panel members noted anecdotally that

unwanted adverse cholinergic effects can occur at doses that induce detrusor contraction.

Although evidence is lacking to guide use of medications to relax the urethral sphincters in dogs with TL-IVDE, use of an alpha-adrenergic antagonist to relax the internal urethral sphincter with or without a centrally acting muscle relaxant such as diazepam to relax the external sphincter is recommended in dogs in which the bladder cannot be manually expressed easily. These drugs should be continued until voluntary urination has been re-established. Long-term use of these medications usually is not necessary in chronically paralyzed dogs. *Supported by low-level evidence.*

## 9.4 | Complications of retention incontinence

Urinary tract infections are the most common complication reported in the literature along with hematuria and bacteriuria.<sup>197</sup> No reports of bladder rupture in this population of dogs have been published, but because of the gravity of this complication, the panel felt it should be taken into consideration when managing retention incontinence. The literature on UTIs is challenging to summarize because of different exclusion criteria, definitions of UTI, and sample timing. Most studies define UTI as bacterial growth of  $>1000/\text{mL}$ , but distinguishing active infection from bacteriuria requires pyuria and documentation of clinical signs, which can be challenging if the dog is paralyzed.<sup>206</sup> Frequency of UTI ranges from 0% to 42% in the first postoperative week, 17% to 36% in the first 6 weeks and 15% at 3 months.<sup>197,202,203</sup> *Escherichia coli* is the most common isolate and frequently is associated with antibiotic resistance.<sup>197,202,203,207</sup>

Risk factors for UTI include the severity of neurologic deficits, duration of retention incontinence and duration of indwelling catheterization.<sup>197,202,203</sup> Conflicting evidence exists on the effect of perioperative antibiotics on subsequent UTI frequency,<sup>202,203,208</sup> but good antibiotic stewardship guides veterinarians to avoid prophylactic use of antibiotics postoperatively.<sup>209</sup> Cranberry extract does not decrease the risk of UTI.<sup>210</sup>

Antibiotics should not be used prophylactically to decrease the frequency of UTIs in dogs with TL-IVDE and it is important to establish a specific diagnosis of UTI versus clinically insignificant bacteriuria. The risk of UTI can be minimized by limiting duration of indwelling catheterization, monitoring voiding efficiency, and using medications to relax urethral sphincters as needed. *Supported by low-level evidence.*

## 10 | PHYSICAL REHABILITATION

Physical rehabilitation is increasingly utilized as a component of the management plan for dogs diagnosed with TL-IVDE, especially postoperatively.

Questions remain regarding the value of incorporating rehabilitation protocols, which populations benefit, as well as when to initiate protocols, how long to continue therapy and the optimal treatment regimens.

### 10.1 | Efficacy of physical rehabilitation

In non-ambulatory dogs with TL-IVDE managed surgically, 2 RCTs provide evidence that in-hospital, staged, intensive physical rehabilitation for 10-14 days postoperatively is well-tolerated but does not improve the recovery of ambulation compared to a control population receiving only basic rehabilitation in the form of passive range of motion (PROM) and assisted walking.<sup>78,80</sup> In dogs with incomplete injuries, rehabilitation also did not impact the rate of recovery of walking, walking coordination, proprioceptive placing or muscle mass.<sup>80</sup> One RCT excluded DPN dogs (80), and the other was not adequately powered to evaluate them; evidence is conflicting regarding the role for rehabilitation in these severely affected dogs.<sup>78,79,152,211-216</sup> Study populations and rehabilitation protocols vary widely, and control groups frequently are lacking. Intensive rehabilitation might enhance the recovery of ambulation in paraplegic DPN dogs.<sup>212</sup>

Although not related to locomotor outcome, rehabilitation has been associated with weight loss and muscle mass gain in dachshunds during a 3-month postoperative study period.<sup>217</sup> Therefore, rehabilitation could have positive implications for recovery of general mobility and overall well-being, but such effects cannot be definitively attributed to the exercises performed.

In dogs with incomplete injuries, rehabilitation performed postoperatively is safe but fails to demonstrate benefit on the rate or extent of recovery of walking compared to dogs receiving only basic exercises (e.g., PROM, assisted walking). Although these findings suggest that more intensive, tailored rehabilitation protocols are not needed for all dogs with incomplete injuries to achieve a successful outcome (i.e., independent, coordinated ambulation), available evidence does support inclusion of basic exercises as standard components of routine postoperative care in all dogs regardless of severity. *Supported by moderate-level evidence.*

### 10.2 | Rehabilitation protocols

Postoperative rehabilitation can be initiated safely as soon as 24 hours postoperatively without adverse consequences such as

increased pain or neurologic deterioration,<sup>78,80,216</sup> but delayed initiation up to 2 weeks postoperatively still might benefit recovery.<sup>79</sup> When specified, hydrotherapy or underwater treadmill (UWMT) therapy is safely initiated anywhere from 3 (with incision protection) to 14 days postoperatively.<sup>78,80,152,212,213,216,218</sup> Among persistently paraplegic DPN dogs, earlier initiation of rehabilitation (at approximately 1 vs 3 weeks) might improve the likelihood of recovery of ambulation.<sup>212</sup>

Reported duration of physical rehabilitation varies from initial hospitalization only to many months in some dogs.<sup>75,78-80,152,212,217,218</sup> Longer duration of rehabilitation (more days participating in rehabilitation, more sessions, more UWMT sessions) is associated with better improvement in neurologic status and shorter time to recovery of weight-bearing stepping.<sup>218</sup>

Rehabilitation protocols vary widely among studies, but exercises included as components of rehabilitation protocols for dogs recovering from TL-IVDE are outlined in Table 9. Mechanistic or physiologic evidence to support inclusion of specific exercises generally is lacking, but a single cadaver-based study proposed a neuroanatomic basis for performing specific mobilization stretching exercises targeting the TL spinal cord and nerve roots.<sup>219</sup> Reported frequency of exercises ranges from 1 to 3 times per day while hospitalized, 1 to 3 times per week for outpatient sessions, and 1 to 5 times per day for at-home exercises, but frequency commonly is not specified.

Several additional modalities have been included under the umbrella of rehabilitation and are listed in Table 10. No specific evidence is available regarding the benefit of inclusion of neuromuscular electrical stimulation (NMES), transcutaneous electrical nerve stimulation (TENS) or infrared therapy, but the utility of photobiomodulation, PEMF and electroacupuncture is considered in the neuroprotective strategies section.

Although some studies did not use standardized exercise protocols, the RCTs described phased exercise combinations and indicated that pre-defined recovery benchmarks can be used to establish standardized yet dynamic protocols that evolve over time as specific functions are regained. This approach includes information regarding which specific exercises and additional treatment modalities are indicated at what time points or functional status (eg, NMES is poorly tolerated once motor function returns) as part of an overall multimodal approach to recovery from TL-IVDE.<sup>75,78,80,217</sup>

Reasonable timing for postoperative rehabilitation in dogs consists of initiation within a 24-hour to 14-day window postoperatively and continuing for at least 2 to 6 weeks. However, it is possible that optimal timing might fall outside of this timeframe or might vary among patient subsets (e.g., based on severity of neurologic status). *Supported by moderate-level evidence.*

**TABLE 9** Rehabilitation exercises with timeline of implementation described in the literature

Rehabilitation Exercise	Timing of implementation and therapy duration	References
Cryotherapy (cold or warm packing)	Initiation: 24 to 48 h postop Duration: 48 h (cold packing) ± 1 to 4 wk (warm packing)	<a href="#">75,78-80,143,144</a>
Range of motion (passive and active stretching) and massage	Initiation: 24 to 48 h postop Duration: 10 d to 6 wk or until ambulation or normal mobility	<a href="#">75,78-80,116,144,152,211,212,214,216-218</a>
Sensory stimulation (eg, toe pinching, hair brushing, different flooring surfaces)	Initiation: 48-h to 3-wk postop Duration: 10 d to 6 wk or until ambulation	<a href="#">78,116,212,215,218</a>
Deep tendon reflex stimulation	Initiation and duration not specified	<a href="#">213</a>
Assisted standing, weight shifting, sit-to-stand	Initiation: 24 to 48 h postop, once able to bear some weight Duration: 5 d to 6 wk or until normal body weight support, ambulation or normal mobility	<a href="#">75,78-80,116,144,212,213,215,217,218</a>
Assisted walking (over ground)	Initiation: 24 to 48 h postop Duration: 4 to 6 wk or until ambulation	<a href="#">75,79,80,144,212,214,215,217,218</a>
Land treadmill walking	Initiation: 3 d to 3 wk postop Duration: 6 wk to 3 mo or normal mobility	<a href="#">79,214,215</a>
Hydrotherapy (swimming or UWWTM walking)	Initiation: 2 (swimming) or 3 (UWWTM) to 14 d postop, typically once able to bear weight or motor is present (for UWWTM) Duration: 7 d to 3 mo or until ambulation or normal mobility	<a href="#">78-80,152,211-218</a>
Manual gait patterning (using land treadmill, UWWTM or cart/lift-assisted)	Initiation: 3 d to 3 wk postop Duration: 6 wk to 3 mo or until ambulation or normal mobility	<a href="#">79,214,215,217,218</a>
Balance exercises (eg, balance boards)	Initiation: 3 d to 14 d, once able to stand Duration: 14 d to 3 mo or until ambulation or normal mobility	<a href="#">79,80,152,213,214,218</a>
Advanced gait & proprioception exercises (eg, cavaletti rails, variable terrain or inclines)	Initiation: 7 to 14 d, not before ambulatory and added progressively Duration: 6 wk to 3 mo or until strong ambulation	<a href="#">79,152,214,215,218</a>

Abbreviation: UWWTM, underwater treadmill.

The specific exercises and adjunctive modalities that should be included in an optimal postoperative rehabilitation regimen remain to be determined. At a minimum, a basic rehabilitation protocol can be recommended to include cryotherapy, PROM, massage, assisted standing and walking, which can be performed with no specialized equipment or training. Although more data are needed to evaluate precisely how such treatment protocols should be adapted over time, a stepwise approach with increasing intensity and incorporation of additional exercises can be tailored to the individual patient as neurologic status changes. *Supported by low-level evidence.*

commonly poorly articulated, dogs strictly rested in a hospital setting for 2 weeks after surgery regained ambulation more quickly compared to historical controls recovering at home.<sup>80</sup> However, activity restriction also was recommended for dogs at home and several other factors could explain or have contributed to this increased rate of improvement.

Despite limited evidence, there is consensus agreement that confinement and activity restriction for a period of at least 4 weeks be recommended as a component of standard postoperative care for dogs with TL-IVDE. Note that activity restriction does not mean avoidance of rehabilitation exercises. *Supported by low-level evidence.*

### 10.3 | Activity restriction

Activity restriction, as described in the medical management section, also is important in postoperative care to allow healing of both the surgical site and the annulus fibrosus, and presumably to minimize the risk of early re-extrusion at the surgical site. Although details are

### 11 | PROGRESSIVE MYELOMALACIA

Progressive myelomalacia is a clinical syndrome characterized by progressive necrosis, ischemia and hemorrhage of the spinal cord that

**TABLE 10** Rehabilitation treatment modalities with timeline of implementation described in the literature

Rehabilitation treatment modality	Timing of implementation and duration of therapy	References
Photobiomodulation (ie, laser therapy)	Initiation: <24 h to 5 d postop Duration: 5 d to 6 wk	78,79,116,152,153
Transcutaneous electrical nerve stimulation (TENS) (including inferential)	Initiation: <24 h to 5 d postop Duration: up to 4 wk	75,212-216
Neuromuscular electrical stimulation (NMES)	Initiation: 1 to 7 d postop, prior to any motor function Duration: 5 to 10 d or until motor or ambulation present	78,80,215,218
Functional electrical stimulation (FES)	Initiated within 7 d postop. Duration: 2 wk	215
Infrared radiation treatment	Initiated within 5 d post-injury Duration: up to 4 wk	75,213
Ultrasound therapy	Initiated within 5 d post-injury. Duration: up to 4 wk	75
Pulsed electromagnetic field (PEMF) therapy	Initiation: preop to 24 h postop Duration: 7 to 14 d (q2-12hr treatment frequency) and 14 d to 4 wk (q12h treatment frequency)	80,143
Acupuncture or electroacupuncture	Initiation: preop to 3 d post-presentation or surgery Duration: up to 72 h (pain control) or 1 wk to 6 mo (functional recovery)	73,74,76,77,145,146,149,150,213,220-223

expands cranially and caudally from an initial site of insult.<sup>224</sup> Clinical signs reflect progressive tissue destruction, and include loss of pelvic limb reflexes, and pelvic limb, trunk, and abdominal muscle tone; cranial progression of the cutaneous trunci reflex (CTR) caudal border; and ultimately thoracic limb involvement and ventilatory failure.<sup>24,225-227</sup> Progressive myelomalacia usually develops within days (24 hours up to 14 days) of an acute IVDE-associated injury to the TL spinal cord and is not always apparent at time of presentation.<sup>227</sup>

The pathophysiology of PMM, although not completely understood, involves primary mechanical damage to and compression of the spinal cord, followed by secondary damage.<sup>228</sup> Increased pressure within the neuraxis might play a role in longitudinal propagation of damage.<sup>226</sup> Studies focused on paraplegic DPN dogs report prevalence of PMM of 10% to 33%<sup>102,229</sup>; however, development of PMM has been reported in a small number of dogs that presented with less severe injury and subsequently deteriorated.<sup>23,227</sup>

## 11.1 | Diagnosis of progressive myelomalacia

Although histopathologic examination presently is the gold standard for diagnosis of PMM, a presumptive antemortem diagnosis often is needed in the clinical setting. A combination of clinical findings and their evolution over time is reported in cases with histopathologic confirmation of PMM.<sup>24,225-228</sup> These include ascending paralysis, hypventilation, loss of segmental spinal reflexes, cranial migration of the

CTR caudal border, decreased abdominal tone, Horner syndrome, diffuse pain, thermoregulation, and malaise.<sup>24,225-228</sup> Although the diagnostic accuracy of these clinical findings requires further study, additional literature describes clinical signs in DPN dogs with presumptive PMM (Table 11). Clinical suspicion can be further heightened by findings on MRI and by serum biomarker measurements (Table 12).

A combination of clinical findings can support a high level of concern for PMM and, when further coupled with imaging findings and longitudinal monitoring for specific changes in the neurologic examination and serum biomarkers, can be even more highly suggestive of the condition. *Supported by moderate-level evidence.*

## 11.2 | Risk factors for PMM

Risk factors for PMM have been evaluated in retrospective studies.<sup>23,107,229</sup> Injury severity is the most important risk factor, with paraplegic DPN dogs at highest risk for developing PMM.<sup>23</sup> Lesion location also is important, with lumbar intumescence disc extrusions more likely to result in PMM than others.<sup>23,107</sup> A retrospective comparison between

**TABLE 11** Clinical features suggestive of an ante mortem diagnosis of progressive myelomalacia in paraplegic DPN dogs with thoracolumbar intervertebral disc extrusion

Clinical signs in DPN dogs	Reference	Predictive/diagnostic utility—for presumptive PMM
CTR cut-off $\geq$ 1–2 spinal cord segments cranial to the site of IVDE	155,225 107	<ul style="list-style-type: none"> <li>All 9 dogs with this sign developed PMM</li> <li>11 of 51 PMM dogs had this sign at presentation</li> </ul>
Progressive CTR cut-off advancement	230	<ul style="list-style-type: none"> <li>145X more likely to develop presumptive PMM in 6 of 36 dogs</li> </ul>
Weak to absent patellar reflexes with a disc extrusion located cranial to the lumbar intumescence	24 155 227	<ul style="list-style-type: none"> <li>Present in 6 of 12 dogs at presentation, all 12 developed it</li> <li>All 8 dogs with this sign developed PMM</li> <li>8 of 51 dogs had this sign at presentation and 23 developed it</li> </ul>
Weak to absent anal tone and perineal reflex with a disc extrusion located cranial to the lumbar intumescence	24 227	<ul style="list-style-type: none"> <li>6 of 12 dogs had this sign at presentation, all 12 developed it</li> <li>Present in 9 of 51 dogs at presentation, and developed in 21</li> </ul>
Loss of abdominal tone	155 227	<ul style="list-style-type: none"> <li>All 11 dogs with this sign developed PMM</li> <li>4 of 51 dogs had this sign at presentation, and 20 developed it</li> </ul>
Difficulty retaining sternal recumbency	227	<ul style="list-style-type: none"> <li>Present in 2 of 51 dogs at presentation, developed in 9 dogs</li> </ul>
Thoracic limbs paresis or proprioceptive deficits in the absence of an explanatory lesion	24 25 227	<ul style="list-style-type: none"> <li>Notes cranial migration of paralysis in all 12 dogs</li> <li>Progression to involve thoracic limbs in 5 dogs</li> <li>Present in 2 of 51 dogs at presentation, developed in 23 of 51 dogs</li> </ul>

Note: Absence of these clinical signs does not preclude later development of PMM and progression of signs in the immediate postoperative period (1–7 days) is key to providing evidence of a clinical diagnosis.

Abbreviations: CTR, cutaneous trunci reflex, IVDE, intervertebral disc extrusion, PMM, progressive myelomalacia.

dachshunds and French bulldogs concluded that French Bulldogs had a higher risk of PMM; however, the study did not control for location of injury and the higher prevalence of lumbar intumescence IVDE in French bulldogs might explain the increased rate of PMM.<sup>229</sup>

and development of PMM in a group of 197 paraplegic DPN dogs.<sup>107</sup> Although NSAIDs were not associated with altered risk, treatment with corticosteroids of any type or dose decreased risk of PMM in this cohort of dogs. Given the retrospective nature of the study, limited information was available related to dosing strategy in individual dogs.

Dogs with IVDE of the lumbar intumescence, that progress to paraplegic DPN status might be at higher risk for the development of PMM compared to those with lesions at other sites in the thoracic and lumbar spinal cord. *Supported by low-level evidence.*

Currently, there is not sufficient evidence to support the use of corticosteroids as a protective strategy against the development of PMM. *Supported by low-level evidence.*

Although French bulldogs appear to have a higher incidence of PMM, the study that reported breed-specific increased risk did not control for other factors that could have contributed to higher numbers of affected French bulldogs. *Supported by low-level evidence.*

Two retrospective studies have evaluated the influence of timing of surgical intervention on the development of PMM. Of these, 1 study including only paraplegic DPN dogs identified an association between delayed decompression and increased odds ratio for the development of PMM where odds were decreased when surgical decompression occurred within 12 hours of non-ambulatory status.<sup>107</sup> The other study, including dogs of all neurological grades, did not show a relationship.<sup>23</sup>

### 11.3 | Prevention and treatment of progressive myelomalacia

Few studies have evaluated the influence of medical interventions on the development of PMM. One retrospective study evaluated the potential association between treatment with anti-inflammatory drugs

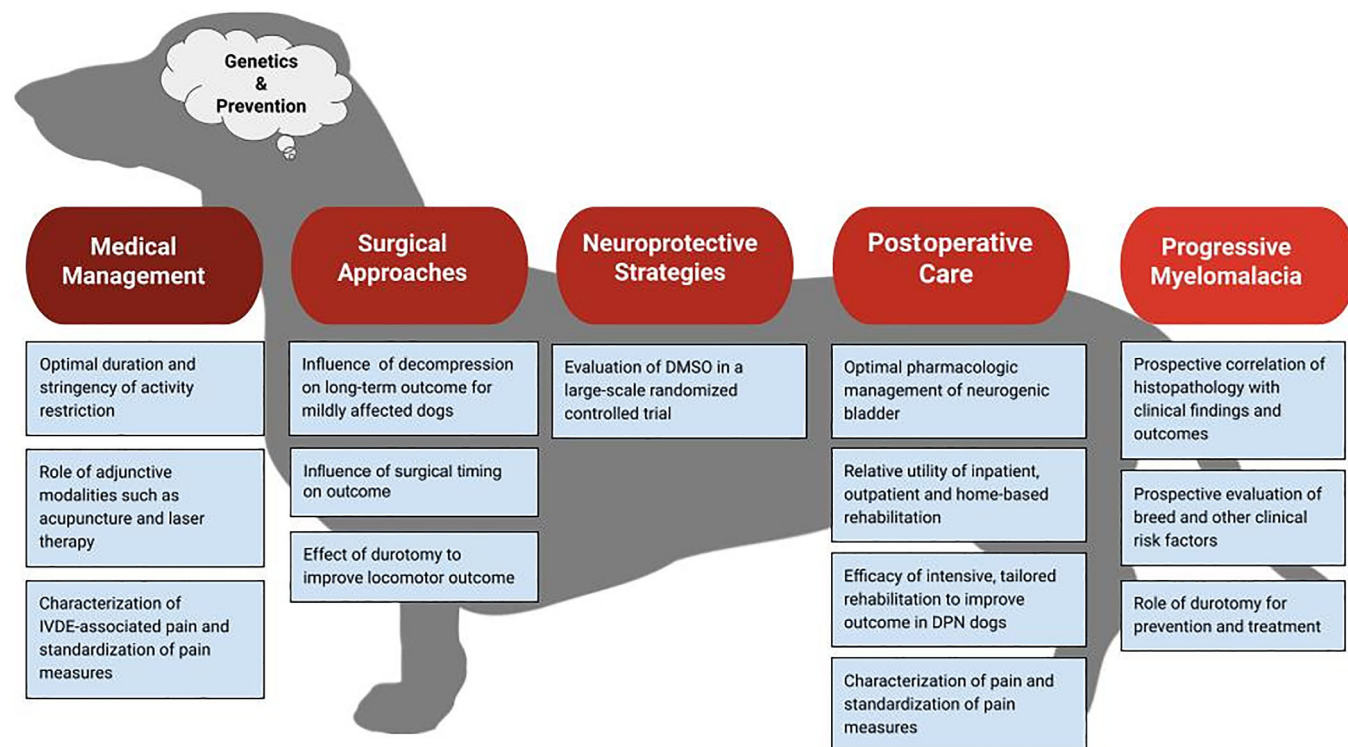
The potential clinical benefit of prompt surgical decompression to minimize likelihood of development of PMM for paraplegic DPN dogs with TL-IVDE should be further explored. This benefit is considered separately from overall improved locomotor outcome, for which the literature may provide more specific evidence. *Supported by low-level evidence.*

**TABLE 12** Imaging and clinical biomarkers evaluated in progressive myelomalacia (PMM)

Feature	References	Summary
Serum [GFAP]	231 232	<ul style="list-style-type: none"> <li>Literature is mixed with respect to utility</li> <li>Reference 231 reports 97.7% specific but only 75% sensitive</li> <li>Reference 232 included only 3 PMM dogs - all with measurable serum GFAP</li> <li>GFAP was noted in the serum of deep pain negative dogs that did not develop PMM.</li> </ul>
Serum (pNFH)	233	<ul style="list-style-type: none"> <li>No difference in serum pNFH concentrations at presentation</li> <li>Dogs that develop PMM have significant elevation at 24 h.</li> </ul>
Myelography: diffuse intraparenchymal contrast	58	<ul style="list-style-type: none"> <li>Presence of contrast within the cord parenchyma was a typical feature of myelomalacia</li> <li>Identified in both focal or progressive myelomalacia</li> </ul>
T2 hyperintensity length ratio > 4.57	23	<ul style="list-style-type: none"> <li>OR for PMM: 17.22</li> </ul>
T2 hyperintensity length > 6 L2	24	<ul style="list-style-type: none"> <li>Observed in only 45% of PMM cases</li> </ul>
SSTSE CSF:L2 ratio > 7.4	25,227	<ul style="list-style-type: none"> <li>Sensitivity 100% and specificity of 75% for diagnosis PMM<sup>25</sup></li> <li>Noted in (17/20) 85% of cases<sup>227</sup></li> </ul>

Note: In addition to clinical findings, *ante mortem* suspicion can be further heightened by findings on MRI and by serum biomarker measurements.

Abbreviations: GFAP, glial fibrillary acidic protein; h, hours; OR, odds ratio; PMM: progressive myelomalacia; pNFH, phosphorylated neurofilament heavy protein.

**FIGURE 1** Areas identified by the consensus group as important opportunities for future study

A few publications have focused on the influence of focal or extensive durotomy on development and progression of PMM. One study retrospectively reported the outcome from a group of paraplegic DPN dogs that underwent hemilaminectomy and spinal decompression with (65 dogs) or without (51 dogs) durotomy performed along the length of the laminectomy window, which varied from single to multiple sites.<sup>98</sup> The rate of PMM was 21.5% for hemilaminectomy alone compared to 0% for hemilaminectomy plus durotomy,

suggesting that addition of durotomy might be associated with decreased risk of PMM. Another case series reported outcomes of dogs with clinical and imaging features suggestive of PMM and treatment with focal hemilaminectomy or more extensive hemilaminectomy and durotomy (EHLD).<sup>238</sup> Ten dogs that had EHLD, some extending as many as 13 vertebral segments based on spinal cord changes on MRI, had a postoperative survival rate of 100%. Postoperative survival rate was 61% for 18 dogs with focal hemilaminectomy alone, and EHLD was identified as the independent variable

associated with increased survival using multivariate analysis. In a retrospectively reported case series of 34 dogs with presumed PMM and treated with EHLD of variable length, postoperative survival rate was 91% with a median follow-up of 82.5 weeks.<sup>234</sup> No dogs in this series regained ambulation, but the authors attributed the high rate of postoperative survival to EHLD and attenuated PMM progression. Finally, a prospective case series of 26 paraplegic DPN dogs treated using EHLD, reported a very low rate of PMM (4%) compared to published rates of the condition.<sup>99</sup>

Conventional wisdom suggests that removal of numerous consecutive articular processes and their soft tissue attachments can result in decreased vertebral column stability, and, a recent retrospective study identified a negative association between prolonged anesthetic time and neurologic outcome in paraplegic DPN dogs.<sup>235</sup> However, a study describing very extensive laminectomy and durotomy for some animals (often >10 vertebral segments) reported that no dogs required spinal stabilization or developed serious complications or radiographic signs of instability.<sup>234</sup> One study did report a single case with short-term severe allodynia that resolved within 10 days after treatment with PO gabapentin.<sup>99</sup>

Durotomy can lead to spinal cord herniation, meningeal scarring and adhesion formation.<sup>236,237</sup> These were not reported in the literature on EHLD; however, a laminectomy membrane can be a very late complication requiring MRI for proper diagnosis. As such, longer-term studies incorporating advanced imaging follow-up might be helpful to assess risk. Given the difficulty in identifying dogs with PMM at the time of original presentation and the propensity for these cases to evolve over time, future studies might focus on the utility of EHLD for all dogs considered clinically at risk for the development of PMM.

Emerging evidence suggests that focal or extensive hemilaminectomy and durotomy might decrease the risk of development of PMM in dogs that are paraplegic DPN secondary to TL-IVDE and may improve survival in dogs with clinical signs suggestive of PMM by halting progression of the condition. This outcome is a separate consideration from whether or not the technique might improve locomotor outcome (discussed elsewhere). Focal or more extensive hemilaminectomy with durotomy can be considered for dogs with imaging and clinical risk or suspicion of PMM; however, specific surgical approaches, including required extent of both hemilaminectomy and durotomy, require further investigation as do potential long-term morbidities associated with these procedures. *Supported by low-level evidence.*

## 12 | CONCLUSIONS

Systematic review of the veterinary clinical literature relating to acute TL-IVDE in dogs has identified some areas of strong

understanding, including the natural history of progression and recovery. Additionally, it has highlighted some areas of opportunity for further exploration (Figure 1), including a need for better understanding of the ideal timing for surgical decompression, expected surgical vs medical outcomes for more mildly affected cases, impact of durotomy on outcome with a particular emphasis on PMM, refining of postoperative care, cost/benefit analysis and number needed to treat studies for certain common interventions, and preventative measures.

The neurology specialty is well-poised to address these knowledge gaps with well-defined clinical outcome measures, strong infrastructure for effective clinical trial designs that can inform veterinary practice, new attention to important features of postoperative care such as bladder and pain management as well as rehabilitation, and a much better understanding of the genetic basis of IVDE. Our consensus panel is hopeful that this statement will be revised in 5-10 years with many new studies directed at these important issues.

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## OFF-LABEL ANTIMICROBIAL DECLARATION

Authors declare no off-label use of antimicrobials.

## INSTITUTIONAL ANIMAL CARE AND USE COMMITTEE (IACUC) OR OTHER APPROVAL DECLARATION

Authors declare no IACUC, or other approval was needed.

## HUMAN ETHICS APPROVAL DECLARATION

Authors declare human ethics approval was not needed for this study.

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## SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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