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Total prostatectomy as a treatment for prostatic carcinoma in 25 dogs

This is a pre print version of the following article:

Original Citation:	
Availability:	
This version is available http://hdl.handle.net/2318/1661178	since 2018-09-02T11:14:02Z
Published version:	
DOI:10.1111/vsu.12768	
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Total Prostatectomy as a Treatment for Prostatic Carcinoma in 25 Dogs

Journal:	Veterinary Surgery
Manuscript ID	VSU-17-032
Manuscript Type:	Original Article - Clinical
Keywords:	prostate, dog, neoplasia, urogenital surgery

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3 ABSTRACT

- 4 **Objective:** To describe the complications and outcome following total prostatectomy in
- 5 dogs with histologically-confirmed prostatic carcinoma.
- 6 **Study Design:** Multi-institutional retrospective case series
- 7 **Animals:** Twenty-five client-owned dogs
- 8 **Methods:** Medical records of dogs undergoing total prostatectomy were reviewed from
- 9 20014-2016. Data retrieved included signalment, presenting signs, preoperative clinical
- findings, including laboratory data, diagnostic imaging, surgical technique, histologic
- diagnosis, postoperative complications, occurrence of postoperative metastasis, and
- 12 survival.
- 13 **Results:** Twenty-five dogs underwent total prostatectomy for prostatic carcinoma.
- 14 Urinary anastomotic techniques included urethrourethral anastomosis in 14 dogs,
- cystourethral anastomosis in 9 dogs, ureterocolonic anastomosis in 1 dog, and
- anastomosis between the bladder neck and penile urethra in 1 dog. All dogs survived to
- discharge. Fifteen dogs were diagnosed with transitional cell carcinoma, 8 dogs with
- prostatic adenocarcinoma, 1 dog with prostatic cystadenocarcinoma, and 1 dog with an
- 19 undifferentiated carcinoma. Permanent postoperative urinary incontinence was present in
- 20 8 dogs. The median survival time was significantly shorter in dogs with extracapsular
- 21 tumor extension compared to those with intracapsular tumors. The overall median
- survival time was 231 days (range, 24-1255 days) and the 1- and 2-year survival rates
- were 32% and 12%, respectively.

24	Conclusions: Based on information from this study, the median survival time for dogs
25	with prostatic carcinoma undergoing total prostatectomy is longer and complication rates
26	lower than previously reported.
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Prostatic neoplasia is relatively rare in dogs ¹ making evaluation of various therapeutic
interventions difficult. Despite the uncommon occurrence in dogs, they are one of the few
domestic species known to develop spontaneous prostatic neoplasia, with carcinomas
being the most common histologic diagnosis. ^{2, 3} Prostatic carcinomas include transitional
cell carcinoma, adenocarcinoma, and squamous cell carcinoma ⁴ . Attempts have been
made to develop more objective methods than traditional light microscopy for
differentiating prostatic adenocarcinoma and transitional cell carcinoma ⁴ . To the authors'
knowledge there is not yet a well accepted objective method of differentiation and the
distinction between adenocarcinoma and transitional cell carcinoma of the prostate
remains controversial.
Prostatic neoplasia carries a poor prognosis in dogs because of aggressive local invasion
and a high rate of regional and distant metastasis. ⁵ Hematuria, stranguria and tenesmus
are common clinical signs in dogs with prostatic disease, 1,6 but a diagnosis of prostatic
neoplasia is often delayed because these clinical signs are not pathognomonic for
prostatic tumors. A diagnosis of prostatic neoplasia is made based on physical
examination findings, diagnostic imaging, cytology, and histology. Metastatic disease is
often present at the time of initial diagnosis. ^{1,6}
Various treatments have been described for prostatic neoplasia including non-steroidal
anti-inflammatory drugs (NSAIDs), chemotherapy, radiation therapy, photodynamic
therapy, and surgery. Surgical options include curative-intent total prostatectomy and
palliative-intent procedures such as partial prostatectomy, transurethral resection,

69	radiation therapy, photodynamic therapy, urethral stenting, and urinary diversion
70	procedures. ⁷⁻¹⁶
71	Improved survival in dogs with urogenital carcinoma has been shown with the use of
72	non-steroidal anti-inflammatory drugs (NSAIDs) alone and in combination with
73	chemotherapeutic agents such as mitoxantrone and carboplatin. 17-20 Reported survival
74	times in dogs with prostatic malignancies vary widely, depending on the stage at
75	diagnosis and treatment pursued. 6,7,9-11,13,14, 20-22 Furthermore current therapeutic strategies
76	have been associated with poor response and high complication rates.
77	Total prostatectomy involves removal of the entire prostate gland and prostatic urethra
78	with subsequent reconstruction of the lower urinary tract. Criteria for appropriate case
79	selection for total prostatectomy in dogs have been previously suggested to include small,
80	intracapsular primary lesions, without evidence of metastatic disease. ²³ There have been
81	few reports evaluating total prostatectomy with most concluding that complication rates
82	are too high and survival times are too short to routinely recommend this technique for
83	treatment of dogs with prostatic neoplasia. 21,22,24-26 The most common complication
84	reported following total prostatectomy is urinary incontinence, which has been reported
85	in 33-100% of cases. ^{24,27}
86	The purpose of this retrospective multi-institutional study was to report the signalment,
87	presenting signs, intraoperative and postoperative complications, histologic diagnosis,
88	and outcome in dogs treated with total prostatectomy for prostatic neoplasia. We
89	hypothesized total prostatectomy would be associated with a complication rate and
90	survival time similar to other currently available therapeutic interventions.
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Total Prostatectomy for Prostatic Carcinoma In Dogs

MATERIALS AND METHODS

This investigation was a multi-institutional retrospective case series approved by the
Veterinary Society of Surgical Oncology Research Committee. The study period ranged
from October 2004 to August 2016. Medical records from contributing institutions were
searched to identify dogs which had undergone total prostatectomy for prostatic
neoplasia. Dogs were included if they had undergone total prostatectomy for confirmed
prostatic carcinoma. Dogs were excluded where the diagnosis was not confirmed to be
carcinoma. Data retrieved included neuter status, breed, age, body weight, presenting
clinical signs, dates of presentation and surgery, results of preoperative staging and
diagnostic testing, surgical technique, use and duration of postoperative indwelling
urethral catheterization, histologic criteria (histologic diagnosis, surgical margin
evaluation, and presence of lymphatic and/or vascular invasion), postoperative
complications (incidence and severity), use of adjunctive therapy, date and method of
detection of recurrent disease, and date and cause of death. Postoperative complications
were classified as minor or major. Minor complications were defined as self-limiting or
those managed with medical intervention. Major complications were defined as any
complication that was expected to cause death without rapid intervention or those
requiring a second surgical procedure. Postoperative urinary incontinence was graded
from 0-4 using a scheme modified from that reported by Byron et al. (Table 1) ²⁸ with
information recorded from veterinarian assessment and owner reporting in the medical
history.
Disease-free interval (DFI) was defined as the time between total prostatectomy and
detection of confirmed or suspected metastasis or local recurrence of neoplasia.

Metastasis was suspected if there were consistent imaging or clinical examination
findings and was confirmed with cytology or histology. Tumor recurrence was suspected
if there was a recurrence of clinical signs or if imaging findings were consistent with a
recurrent mass in the region of the previous surgical site; and tumor recurrence was
confirmed with cytologic or histologic evidence of neoplasia.
Survival time was defined as the time between total prostatectomy and death. Cause of
death was classified as either tumor-related or unrelated. Dogs for which the cause of
death was unknown were presumed to have died or been euthanized as a result of tumor-
related causes. Dogs that died from unrelated causes or were still alive at the time of
writing were censored from the survival analysis.
Statistical Analysis
Descriptive statistics for signalment, historical, preoperative, and postoperative data were
generated and reported as the arithmetic mean and range. Disease-free intervals and
survival times were reported as medians with 95% confidence intervals. Median survival
times (MST) were estimated from Kaplan-Meier survival analysis. A log-rank test was
used to compare survival curves of dogs with transitional cell carcinoma and prostatic
adenocarcinoma. $P \le 0.05$ was considered significant. Statistical software (Medcalc
version 16.8.4 for Windows, Medcalc Software, Ostend, Belgium, www.medcalc.org)
was used for descriptive statistical modelling and Kaplan-Meier survival analysis.

138	RESULTS
139	Signalment
140	Twenty-five dogs met the inclusion criteria. The median age was 9.3 years (range 4.9-
141	13.0 years). The median weight was 25.0 kg (range 6.1-47.4 kg). All dogs were neutered
142	males. Breeds were Labrador retriever (n=5), mixed breed (5), German shepherd (2), and
143	1 each of West Highland White Terrier, Lhasa Apso, Boxer, Dachshund, Jack Russell
144	Terrier, Australian Cattle Dog, Siberian Husky, American Staffordshire Bull Terrier,
145	Wheaten Terrier, Rhodesian Ridgeback, Shetland Sheepdog, Boston Terrier, and Golden
146	Retriever (Table 2).
147	
148	Clinical Findings
149	The most common clinical signs on presentation were dysuria (n=12), dyschezia (6),
150	gross hematuria (6), pollakiuria (6), hyporexia (4), and lethargy (4). Prostatic
151	enlargement was detected incidentally on routine digital rectal examination in 4 dogs.
152	Two dogs presented with pre-existing urinary incontinence (grade 2, n=1; grade 4, 1).
153	An enlarged prostate palpated on rectal examination was the most common physical
154	examination finding (n=16). A caudal abdominal mass was detected on abdominal
155	palpation in 2 dogs.
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157	Preoperative Diagnostic Tests
158	Preoperative serum biochemistry, hematology and urinalysis results were available for
159	24, 23, and 24 dogs, respectively. Serum biochemistry abnormalities included increased
160	alkaline phosphatase (n=4; 227-634 U/L [reference range, 23-212 U/L]), increased

161	alanine transferase (2; 189-657 U/L [reference range, 10-125 U/L]), and
162	hypertriglyceridemia (1; 539 mg/dL [reference range, 20-112 mg/dL]). Hematologic
163	abnormalities included anemia (1; 4.88 x10 ¹² cells/L [reference range, 5.65-8.87
164	$x10^{12}/L$]), neutrophilia (4; 12.5-24.6 x $10^{9}/L$ [reference range, 2.95-11.64 x $10^{9}/L$]) and
165	monocytosis (1; 3.01x10 ⁹ /L [reference range, 0.16-1.12 x10 ⁹ /L]). Urinalysis
166	abnormalities included hematuria (13) and neoplastic epithelial cells on sediment
167	examination (1). Urine was cultured preoperatively in 6 dogs and 3 of these were
168	positive, including Streptococcus canis (1), Gram negative rods and Gram positive rods
169	and cocci (1), and a positive culture without further information available (1).
170	Preoperative imaging for clinical staging included three-projection thoracic radiographs
171	(n=20), orthogonal abdominal radiography (3), abdominal ultrasonography (19), thoracic
172	and abdominal computed tomography (CT) (8), and abdominal magnetic resonance
173	imaging (MRI) (1). No dog had evidence of pulmonary metastatic disease.
174	Prostatomegaly was detected in all but one dog. The prostate was intrapelvic and not
175	detected on abdominal ultrasonography in this dog. Additional imaging findings included
176	mild internal iliac lymphadenomegaly (3), pyelectasia (1), and ureteral dilation (1).
177	Preoperative cytology (20) and histopathology (3), reports were available for 23 dogs.
178	Cytology results included carcinoma, not further classified in (17), epithelial dysplasia
179	(2) and squamous metaplasia (1). Histology results were in agreement with the final post-
180	operative diagnosis in all three cases with transitional cell carcinoma in 2 dogs and
181	prostatic adenocarcinoma in 1 dog.
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Total Prostatectomy for Prostatic Carcinoma In Dogs

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The prostate was approached via a caudal ventral midline celiotomy in all dogs. Pubic
and ischial osteotomies (n=3) or pubic symphysiotomy (2) were required for further
exposure in 5 dogs with an intrapelvic prostate. Enlarged medial iliac lymph nodes were
detected intra-operatively and removed for histologic evaluation in 2 dogs. A retrograde
urinary catheter was placed in all dogs prior to prostatectomy. The deferent ducts were
ligated and transected. The periprostatic fat was dissected from the prostate, with
dissection as close as possible to the prostatic capsule, especially dorsally, to minimize
the risk of iatrogenic damage to the neurovascular supply to the urinary bladder and
urethra. The prostatic vascular supply was ligated or cauterized as close to the prostate as
possible. The urinary catheter was then partially withdrawn to allow for pre- and post-
prostatic urethral transection before advancing the catheter back into the bladder
following completion of the total prostatectomy. Urethrourethral anastomosis was
performed in 14 dogs and cystourethral anastomosis in 9 dogs. One dog had gross disease
extending into the bladder and post-prostatic urethra. A total cystoprostectomy was
performed with bilateral ureterocolonic anastomosis in this dog. Another dog had gross
disease involving a large section of the post-prostatic urethra and an anastomosis between
the bladder neck and penile urethra was performed. The suture materials and patterns
used for anastomosis were recorded for 19 dogs. Anastomosis was performed with a
monofilament absorbable suture in all dogs using either a simple interrupted (11) or a
simple interrupted and simple continuous pattern (8). Closure of the celiotomy incision
was routine.

207	Postoperative Management
208	A urinary catheter was maintained postoperatively in 20 dogs for a median of 4 days
209	(range, 1-7).
210	All dogs were treated with postoperative analgesia and protocols were variable including
211	NSAIDs (n=22), opioids (20), tramadol (9), ketamine (1), and acetaminophen (1).
212	
213	Surgical Complications and Outcome
214	All dogs survived to discharge and no intraoperative or perioperative deaths were
215	recorded. There were 4 major complications in 4 dogs and 16 minor complications in 15
216	dogs.
217	Major complications included minor incisional dehiscence (n=2), uroabdomen (1), and
218	prepubic herniation (1). Revision surgery was performed in all dogs with major
219	complications. Uroabdomen was detected 1 day post-operatively in 1 dog and a 15 mm
220	laceration was found at the bladder neck. This was presumed to be iatrogenic. Prepubic
221	herniation occurred 20 days post-operatively in another dog and was repaired with
222	polypropylene mesh.
223	Minor complications included permanent urinary incontinence (n=8), urinary tract
224	infection (6), and superficial surgical site infection (2).
225	Postoperative urinary incontinence was recorded in 23 dogs overall. Eleven dogs had
226	grade 0 urinary incontinence. Urinary incontinence resolved completely in 3 additional
227	dogs within 1-4 weeks postoperatively. In 1 dog, continence was maintained when treated
228	with phenylpropanolamine. Eight dogs exhibited some degree of permanent urinary
229	incontinence: grade 1 (n=2), grade 2 (3), grade 3 (1) and grade 4 (2). One of the dogs

with grade 4 urinary incontinence had pre-existing grade 4 urinary incontinence. Fifteen dogs returned to complete urinary continence within 4 weeks; although recurrent urinary incontinence, secondary to suspected local tumor recurrence, was recorded in 1 dog 148 days after surgery (Table 2). Some degree of post-operative urinary incontinence was seen in 4 of 9 dogs with cystourethral anastomosis (one of which was pre-exisiting) and 4 of 11 dogs with urethrourethral anastomosis.

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Histologic Diagnosis

Histologic examination of the excised prostate was performed in all dogs, with a report available in 20 dogs. Diagnoses included transitional cell carcinoma (n=15), adenocarcinoma (8), undifferentiated carcinoma (1), and papillary cystadenocarcinoma (1). For the two dogs with sublumbar lymph node excision, one dog had evidence of nodal metastasis and the other dog had a reactive lymph node. Of the 20 dogs with histologic margin evaluation, 8 dogs had complete excision and 12 dogs had incomplete excision. Local recurrence was either suspected or confirmed in 3 dogs with complete histologic margins and in 4 dogs with incomplete margins. There was no significant difference in the rate of local recurrence between dogs with complete histologic margins compared to those with incomplete margins (P = 0.84). There was histologically diagnosed extracapsular extension in 11 dogs and no evidence of extracapsular extension in 9 dogs. There was no significant difference in the rate of local recurrence between dogs with extracapsular extension and those without extracapsular extension (P = 0.44). There was histologic evidence of lymphatic and/or vascular invasion in 13 dogs and no evidence of lymphatic and/or vascular invasion in 7 dogs.

253	Metastatic disease was either confirmed or suspected in 7 dogs with lymphatic and/or
254	vascular invasion and in 2 dogs without lymphatic and/or vascular invasion. There was
255	no significant difference in the rate of metastasis between dogs with lymphatic and/or
256	vascular invasion compared to those without lymphatic and/or vascular invasion (P =
257	0.29).
258	There was no significant difference in MSTs between dogs with and without complete
259	histologic margins or dogs with and without lymphatic and/or vascular invasion ($P = 0.23$
260	and 0.11, respectively). The MST was significantly shorter in dogs with extracapsular
261	extension compared to those with intracapsular tumors ($P = 0.02$) (Table 3).
262	
263	Adjuvant Therapy
264	Twenty-one dogs received adjunctive therapy, including mitoxantrone and NSAIDs
265	(n=14); NSAIDs alone (3); metronomic thalidomide, cyclophosphamide, and piroxicam
266	(3); and carboplatin and deracoxib (1). Dosing and protocols were variable, but of the 15
267	dogs treated with curative-intent chemotherapy protocols, 10 dogs completed their
268	targeted chemotherapy protocols. Of the five dogs that did not complete their protocols
269	reasons for termination of the protocol were available for 3 dogs and were all due to the
270	development of metastatic disease. Adverse effects were recorded for 2 dogs both of
271	which were episodes of neutropenia during treatment with mitoxantrone and piroxicam
272	which resolved with dose reduction of mitoxantrone.
273	One dog was treated with adjunctive radiation therapy to the local surgical site (27 Gy
274	divided into 10 fractions of 2.7 Gy daily, Monday through Friday), starting 20 days
275	following total prostatectomy and bilateral medial iliac lymphadenectomy.

276	Clinical Outcome
277	Local tumor recurrence was confirmed in 3 dogs and suspected in 5 dogs. Metastatic
278	disease was confirmed in 4 dogs and suspected in 9 dogs. Confirmed metastatic sites
279	included lungs (n=1), sublumbar lymph nodes (1), sublumbar lymph nodes and pelvis (1)
280	and lungs, pelvis, vertebrae, adrenal glands and sublumbar lymph nodes (1). Sites of
281	suspected metastasis included lungs (6), skin (1), bone (1) and sublumbar lymph nodes
282	(1).
283	Data to calculate the DFI was available for 14 dogs. The median DFI was 81.5 days (95%)
284	confidence interval (CI) 48.4-263, range 11.0-630 days). Data to calculate the median
285	DFI was available for 5 dogs with suspected or confirmed local recurrence (median DFI
286	85.0 days, 95% CI 27.7-208.7, range 76-247) and 9 dogs with suspected or confirmed
287	metastatic disease (median DFI 76.0 days, 95% CI 31.4-305, range 24.0-630). Two dogs
288	with recurrent disease were still alive at the time of writing 65 and 190 days post-
289	operatively.
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291	Death was attributed to tumor-related causes in 19 dogs: local recurrence in 7 dogs
292	(confirmed in 2 dogs and suspected in 5 dogs) and metastasis in 12 dogs (confirmed in 3
293	dogs and suspected in 9 dogs). Three dogs were euthanized for reasons unrelated to
294	prostatic neoplasia. Two dogs were euthanized for clinical progression of chronic kidney
295	disease; and 1 dog was euthanized for suspected degenerative myelopathy. Three dogs
296	were still alive at the time of writing, ranging from 65-1255 days post-operatively.
297	The MST for all dogs was 231 days (95% CI 138-628, range 24-1255 days). The MST
298	for dogs with prostatic transitional cell carcinoma was 189 days (95% CI 135-628, range

299	34.0-664) and the MST for dogs with prostatic adenocarcinoma was 248 days (95% CI
300	169-789 days, range 24-1255 days). There was no significant difference in MSTs
301	between dogs with transitional cell carcinoma and adenocarcinoma ($P = 0.27$). The 1- and
302	2-year survival rates following total prostatectomy were 32% and 12% of dogs,
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This report describes the surgical technique and outcome of dogs undergoing total
prostatectomy for the treatment of prostatic carcinoma. The signalment and clinical
presentation of dogs in the present study was similar to previous reports of dogs with
prostatic tumors. 6,29
Total prostatectomy has been associated with an unacceptably high complication rate in
previous studies in dogs. ^{21, 23, 24-27} However, the majority of these studies included total
prostatectomy for treatment of various prostatic diseases and only one specifically
investigated total prostatectomy for treatment of prostatic neoplasia. ²¹ In this study,
survival times were short (range, 5-45 days) and the incidence of urinary incontinence
was not described. ²¹ However, urinary incontinence was the most common postoperative
complication in other studies of total prostatectomy for dogs with prostatic diseases, with
an incidence ranging from 33%-100%. ^{24,26, 27} Permanent postoperative urinary
incontinence was recorded in 34.8% of dogs in the present study. The severity of urinary
incontinence was subjectively graded based on owner and veterinary assessment. Of the 8
dogs with permanent urinary incontinence, 5 dogs had grade 1 or 2 urinary incontinence.
A previous study classified the severity of urinary incontinence following total
prostatectomy as minor if it only occurred with excitement or activity, and major if it was
permanent. ²⁷ If this classification were used in the present study then only 17.4% of dogs
(4/23) would have been assessed to have major urinary incontinence. A grading system to
describe the severity of urinary incontinence may assist owners in determining whether
the postoperative outcome will be compatible with their expectations. Permanent urinary
incontinence may lead to secondary complications such as recurrent urinary tract

infection, pyelonephritis, and urine scalding; and owners should be aware of these
possible sequalae.
Urinary incontinence has been assessed in both normal dogs and dogs with prostatic
disease undergoing total prostatectomy. ^{24,31} For dogs without prostatic disease, total
prostatectomy does not result in urinary incontinence. ³¹ However, in another study by the
same authors, 93% of dogs with prostatic disease had urinary incontinence following
total prostatectomy with 54% of these dogs having permanent incontinence. ²⁴ This
suggests that the disease process itself may play a role in the development of urinary
incontinence, and urinary incontinence may not solely be a consequence of the surgical
technique. This is further supported by a study in which the surviving 3 dogs that
underwent inadvertent prostatectomy during cryptorchidectomy did not have urinary
incontinence following surgical correction. ³² In normal dogs treated with total
prostatectomy, there were minimal functional changes to the urinary tract, but there was a
decrease in the maximal urethral closing pressure in these dogs. ³¹ However, this decrease
in maximal urethral closing pressure was not sufficient for urethral sphincter pressure to
be overcome by intravesicular pressure. ³¹ Dogs with prostatic disease have abnormally
low external urethral sphincter pressures. ²⁴ This reduced external urethral sphincter
pressure, in combination with a decrease in maximal urethral closing pressure following
total prostatectomy, likely predisposes to urinary incontinence in dogs with prostatic
disease.
Surgical technique may also influence the development of urinary incontinence post-total
prostatectomy. The neurovascular supply to the bladder neck and prostatic urethra
courses along the dorsal aspect of the prostate and disruption of this neurovascular supply

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during total prostatectomy may result in urinary incontinence postoperatively. ²⁷ Total
prostatectomy was performed in all dogs in the present study with close attention to the
dorsal dissection technique to minimize the risk of disrupting the innervation to the
bladder neck and proximal urethra.
It is possible that the comparatively low rate of urinary incontinence in the present study
is related to the primary pathology. However, it has been suggested that total
prostatectomy in dogs with prostatic neoplasia may be associated with a higher incidence
postoperative incontinence compared to other causes of prostatic pathology. ²⁷ The
numbers in that study were small with only 3 of 9 dogs being diagnosed with prostatic
neoplasia, two of which were carcinomas. Case selection may also have played in role in
the lower rate of urinary incontinence in the present study as dogs may not have been
deemed appropriate surgical candidates if there was gross disease extending beyond the
prostatic capsule. This may have contributed to less aggressive dissection being required
to excise the prostate thus limiting collateral damage to the surrounding neurovascular
structures.
Adjunctive treatment was used in the majority of the dogs in this study, with 21 dogs
receiving some form of adjunctive therapy following total prostatectomy. While no
comment can be made on the adjunctive treatment protocol of choice, adjunctive
treatment is still recommended for dogs with prostatic carcinoma because of the high risk
of metastatic disease. Additionally, because very few dogs in this study were treated with
total prostatectomy alone, no comparison of outcome for surgical intervention with and
without adjunctive therapy can be made from this population.

There was no significant difference found between completeness of histologic margins or
the presence extracapsular extension and local recurrence of prostatic carcinoma. There
was no significant difference in MST between dogs with local recurrence of prostatic
carcinoma compared to those without local recurrence; however the MST was
significantly shorter for dogs with extracapsular extension compared to those with
intracapsular tumors. The reason for this difference is unclear. Since there was no
association found between extracapsular extension and local recurrence; or local
recurrence and MST, it is unlikely that extracapsular extension contributed to the
significant difference in MST as a consequence of an increased rate of local recurrence.
Nonetheless the significantly shorter survival time of dogs with extracapsular extension
indicates that this information may be of prognostic value.
There was no association found between the presence of lymphatic and/or vascular
invasion at the time of surgery and the occurrence of metastatic disease. Likewise there
was no association found between the presence of lymphatic and/or vascular invasion and
MST. Overall metastatic disease was either suspected or confirmed in 13 of 25 dogs
(52.0%). This is comparable to previous reports where metastasis to sublumbar
lymph nodes, bone, and lungs was reported in 63–89% of dogs at the time of diagnosis. ⁸
There was no significant difference in survival time between dogs with transitional cell
carcinoma or adenocarcinoma. Some authors have referred to prostatic neoplasia with
various morphologic features, such as glandular and urothelial differentiation,
collectively as prostatic carcinoma ⁶ as we have done in the present study. This may be
reasonable given the lack of a proven objective means of differentiation. Given the
similar morphologic features of prostatic adenocarcinoma and transitional cell carcinoma

when assessed using light microscopy it may be difficult to distinguish the precise cell
origin. ⁴ Immunohistochemical methods have been investigated for the purpose of
differentiating prostatic adenocarcinoma from transitional cell carcinoma but as yet a
valid method has not been found. ⁴ Accurate classification of prostatic epithelial neoplasia
may be significant from a prognostic perspective though currently this remains
controversial.
The MST time for dogs undergoing total prostatectomy for prostatic carcinoma in this
study was 231 days (Figure 1). All dogs survived to discharge and almost one-third of
dogs survived longer than one year following total prostatectomy. A MST of 17 days was
reported in the only other study to report a MST following total prostatectomy in dogs
with prostatic neoplasia ²¹ Published survival times for prostatic carcinoma including
various other treatments range from 17 to 654 days (Table 4). 7-9,11,14,21,22,30,33 However a
number of these studies included cases with urogenitial carcinomas arising from locations
other than the prostate and did not provide separate analysis of survival for patients with
prostatic carcinoma alone. 11, 18, 33
The MST in this study population is markedly longer than that previously published for
dogs with prostatic carcinoma following total prostatectomy. The study by Vlasin, et al. ²¹
reporting a MST of 17 days in dogs was prospective and randomised. This study design
may have contributed, in part, to the poor overall survival of those dogs as randomization
may have prevented selection of cases most appropriate for total prostatectomy. In
contrast the retrospective nature of our report may have resulted in reporting of cases
deemed more suitable for total prostatectomy by the surgeon. While no reported effort
was made to select cases with primary lesions under a certain size and without

436	extracapular extension or pre-existing metastatic disease, it is reasonable to assume that
437	these factors played a role in case selection. Only one case in this study had documented
438	metastatic disease prior to total prostatectomy.
439	The limitations of this study are shared with other multi-institutional retrospective
440	studies. Medical records can be inaccurate or incomplete. There was no standardization
441	of perioperative or adjunctive treatments, and the surgeons performing the procedure
442	differed. The cause of death was only confirmed by post-mortem evaluation in 4 dogs.
443	Retrospective inference of the degree of post-operative urinary incontinence from owner
444	and veterinarian descriptions may be inaccurate. Also, the power of statistical analyses
445	performed was likely limited by the small sample size.
446	Based on our findings, total prostatectomy may be considered as a viable treatment
447	option in dogs with prostatic neoplasia, particularly if presenting with urethral
448	obstruction. However, we propose that case selection is likely to play an important role in
449	postoperative complications and survival. Our study suggests that the incidence and
450	severity of urinary incontinence in dogs with prostatic carcinoma treated with total
451	prostatectomy may be lower and survival times longer than previously reported. Further
452	prospective evaluation of risk factors for post-operative complications and outcome
453	following total prostatectomy in dogs is necessary to determine appropriate case selection
454	criteria.
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460	DISCLOSURE
461	The authors have no conflicts of interest related to this report.
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Table 1 Urinary Incontinence Scoring System*

- 0 Dog always continent
- 1 Dog urine soils where is has been sleeping more than 50% of the time. Does not dribble urine or have a wet prepuce/ventrum when awake.
- 2 Dog urine soils where it has been sleeping more than 50% of the time. Dribbles urine or has a wet prepuce when awake up to 25% of the time.
- 3 Poorly continent. Dog urine soils where it has been sleeping more than 50% of the time and has a wet prepuce/ventrum 25-75% of the time.
- 4 Dog is never continent. Dribbles urine when awake wet when risi. and when sleeping. Constantly has a wet prepuce/ventrum and leaves urine when rising from a sitting to standing position.

^{*}Modified from Byron, et al.28

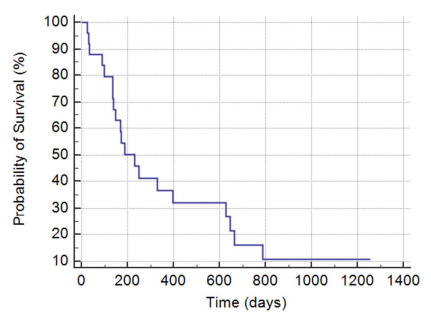


Figure 1 Kaplan-Meier survival curve for 25 dogs with prostatic carcinoma that underwent total prostatectomy. Dogs were censored from analysis if they were still alive at the time of writing or if they had died from causes unrelated to prostatic neoplasia.

Table Signalment, diagnosis, surgery details, incontinence and outcome for dogs undergoing total prostatectomy

1 10.6 Labrador retriever CU 2 PA 3 789 Euthanasia (renal insufficiency)	
2 11.3 West-Highland terrier CU 2 TCC 0 [§] 190 Alive	
3 4.9 Lhasa apsoa UU 1 TCC 0 628 Euthanasia (suspected metastasis)	
4 12.75 mixed breed CU 4 TCC 2 34 Euthanasia (confirmed pulmonary met	rastasis)
5 9.3 boxer UU 4 TCC 1 489 Euthanasia (renal insufficiency)	
6 10.58 dachshund UU 1 TCC 0 231 Euthanasia (confirmed local recurrence	e)
7 9.25 mixed breed CU 1 TCC 2 135 Euthanasia (suspected local recurrence	e)
8 8.5 Jack Russell terrier CU 2 PA 0 172 Euthanasia (suspected local recurrence	e)
9 7.9 Australian cattle dog UU 2 TCC 4 134 Euthanasia (suspected local recurrence	e)
10 8.25 Siberian husky UU 3 PA 0 65 Alive	
11 11 German shepherd UU 5 PA 0 798 Euthanasia (suspected degenerative my	yelopathy)
12 8 mixed breed UU 7 PA 0 1255 Alive	
13 7.66 American Staffordshire bull terrier CU 3 PA 0 ⁸ 169 Euthanasia (suspected metastasis)	
14 6.5 wheaten terrier UU n/a TCC 2 99 Euthanasia (suspected local recurrence	e and metastasis)
15 10 golden retriever UU n/a TCC 0 ⁹ 189 Unknown	
16 10 Labrador retriever CU n/a TCC 0 ⁸ 396 Euthanasia (suspected metastasis)	
17 10 Labrador retriever CU n/a TCC 4 (pre-existing) 647 Euthanasia (suspected metastasis)	
18 9 Rhodesian ridge back UU 5 TCC 0 664 Euthanasia (confirmed local lymph no	de metastasis)
19 10 mixed breed UU 7 PA 0 248 Euthanasia (suspected local recurrence	e)
20 9 Shetland sheepdog UU 5 PA 1 31 Euthanasia (confirmed multifocal meta	astatic disease)
21 12 Labrador retriever UU 5 TCC 0 88 Euthanasia (suspected metastasis)	
Anastomosis between bladder neck and 22 9 Boston terrier penile urethra 7 PA Unknown 24 Euthanasia (suspected metastasis) Ureterocolonic	
23 13 German shepherd anastomosis n/a TCC n/a 138 Euthanasia (melena and lethargy)	
24 9 mixed breed UU 7 TCC 0 149 Euthanasia (confirmed local recurrence	e)
25 9 Labrador retriever CU 5 PCA 0 330 Euthanasia (suspected metastasis)	<i>'</i>

^{*}Urethrourethral anastomosis (UU), cystourethral anastomosis (CC)

[†] Prostatic carcinoma (PA), transitional cell carcinoma (TCC), prostatic cystadenocarcinoma (PCA)

^{*} Modified from Byron et al²⁸ *Had initial post-operative incontinence which resolve

No incontinence when on phenylpropanolamine. Incontinence recurred 148 days post-operatively, suspected secondary to local recurrence

Table 3 Comparison of MST with and without lymphatic/vascular invasion, extracapsular extension and complete histologic margins

MST	With	Without	With Extracapsular	Without	With Complete	Without Complete
	Lymphatic/vascular	Lymphatic/vascular	Extension	Extracapsular	Histologic Margins	Histologic Margins
	Invasion	Invasion		Extension		
MST	149	248	138	248	248.	172
95% CI	99.0-189	135-664	88.0-169	172-628	99.0-628	134-396
P value	0.12		0.02		0.23	



Table 4 Published Survival Times in Dogs with Prostatic Carcinoma

Reference	Treatment	MST (range)
Vlasin et al ²¹	Total prostatectomy	17 days (5-45)
Vlasin et al ²¹	Subtotal intracapsular prostatectomy	130 days (2-220)
Liptak et al ⁸	Transurethral resection	32, 74, and 264 days
L'Epplattenier et al ³⁰	Partial prostatectomy with Nd:YAG laser	103 days (5-239)*
Weisse et al ¹¹	Urethral stenting	20 days (6-105) [†]
L'Epplattenier et al ¹⁴	Photodynamic therapy with 5-aminolevulinic acid	41 days (10-68)
Turrel et al ⁹	Intraoperative radiation therapy	114 days (41-750)
Nolan et al ³³	Intensity modulated and image guided radiation therapy	654 days [‡]
Sorenmo et al ¹⁸	NSAIDS	6.9months
Sorenmo et al ¹⁸	Untreated	21 days

^{*}Excluding the three dogs that died within 16 days MST =183 (91-239)

ays MS, rival for cases v. *Survival time for all 12 dogs in this study (survival for cases with prostatic carcinoma not reported separately).

^{*} Survival time for all 21 dogs in this study (survival for cases with prostatic carcinoma were not reported separately).