# Primary Spinal Malignant Melanoma Mimicking a Cervical Nerve Root Schwannoma: Case Report and Literature Review

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# Abstract

Primary spinal malignant melanoma (PSMM) is a rare cancer of the central nervous system (CNS), and PSMM of the spinal nerve root is even more extraordinary. PSMM of a nerve root can mimic the radiographic appearance of benign nerve sheath tumors, thus resulting in misdiagnosis until tissue diagnosis can be made. A 53-year-old African American woman presented with pain primarily involving the left aspect of her neck and shoulder for 2 years. Magnetic resonance imaging (MRI) of the cervical spine demonstrated a T1-hyperintense, T2-hypointense, homogenously enhancing, dumbbell-shaped, intradural extramedullary mass extending out through the left C2–3 foramen. A midline incision was used to perform a C2 and C3 laminectomy, and the mass was removed from the cavity. The histopathologic profile was consistent with the diagnosis of malignant melanoma. The present case report adds to the 110 cases of PSMM and the 20 cases of PSMM of the spinal nerve root in the existing body of literature. Radiographic and clinical features resemble that of the much more common schwannoma or neurofibroma requiring immunohistochemical analysis for definitive diagnosis. The optimal treatment for PSMM has not yet been defined due to its rarity and it is therefore important to report such cases in order to share our clinical experiences and provide data to other clinicians treating this uncommon disease.

#### **Keywords**

- spine surgery
- malignant melanoma
- nerve root
- case report
- schwannoma

# Introduction

Melanoma is the sixth most common cancer in the United States with an increasing incidence.<sup>1,2</sup> While lung and breast cancers are more prevalent than malignant melanoma, metastasis to the central nervous system (CNS) is more likely with

article published online May 27, 2024 DOI https://doi.org/ 10.1055/s-0044-1787081. ISSN 2248-9614. melanoma than with breast or lung cancers—40 to 60% of patients with malignant melanoma will be diagnosed with CNS metastases during the disease course. Autopsy studies indicate the incidence of CNS involvement may be even greater, with up to 80% of metastatic melanoma cases

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Fig. 1 Preoperative magnetic resonance imaging (MRI) of the cervical spine with and without contrast demonstrated a T1-hyperintense, T2-hypointense, homogenously enhancing, dumbbellshaped, intradural extramedullary mass extending out through the left C2-C3 foramen.

involving the CNS.<sup>3</sup> While CNS involvement in metastatic melanoma is common, primary malignant melanoma (PMM) of the CNS is quite rare and accounts for approximately 1% of all melanoma cases.<sup>4</sup> Primary spinal malignant melanoma (PSMM) is even less common and PSMM arising from a spinal nerve root is exceedingly rare. PSMM of a nerve root can mimic the radiographic appearance of benign nerve sheath tumors, thus resulting in misdiagnosis until tissue diagnosis can be made. In this case report, we present a case of PSMM arising from a cervical nerve root mimicking a nerve sheath tumor.

spinal cord

## **Case Report**

#### Presentation

A 53-year-old African American woman presented with pain primarily involving the left aspect of her neck and shoulder for 2 years. The pain also occasionally involved her left mastoid region and posterior aspect of her head. Over the past 2 months, the pain had been progressively worsening. On physical examination, she was noted to be full strength in all major muscle groups. She did not have any signs of myelopathy and her gait was normal. Her sensation to light touch was preserved. Magnetic resonance imaging (MRI) of the cervical spine with and without contrast demonstrated a T1-hyperintense, T2-hypointense, homogenously enhancing, dumbbell-shaped, intradural extramedullary mass extending out through the left C2-3 foramen. The intradural portion of the mass compressed the spinal cord with displacement of the cord to the right. The foraminal portion of the mass resulted in widening of the neural foramen (Fig. 1). Based upon the imaging, the differential diagnosis included schwannoma and neurofibroma, and the patient was counseled on surgical resection of the lesion for which she consented.

## Surgery

A midline incision was used to perform a C2 and C3 laminectomy. Upon opening the dura, a dark pigmented mass was identified to the left of and ventral to the spinal cord arising from a spinal nerve root (**Fig. 2A** and **B**). The mass was gently separated from the spinal cord. The tumor capsule was coagulated using bipolar electrocautery and incised using microscissors. An ultrasonic aspirator was then used to debulk the mass internally to allow greater manipulation of the mass without pressure on the adjacent cervical spinal cord. The rostral attachment to the nerve root was identified, coagulated, and incised (Fig. 2C). We then followed the



amputated as it entered the neural foramen caudally, and delivered from the cavity.

Fig. 2 (A-C) Intraoperative imaging when opening the dura with visualization of a dark pigmented mass to the left of and ventral to the spinal cord arising from a spinal nerve root.

coagulated, and incised.



Fig. 3 Postoperative imaging showing removal of the tumor.

mass caudally to its point of exit into the C3 neural foramen and amputated it as it exited the spinal canal. The mass was then delivered from the cavity. The intradural space was irrigated and inspected for any remaining tumor. Residual tumor was intentionally left in the foraminal and extraforaminal spaces due to the much higher surgical morbidity of removing this portion and the persistent belief that this represented a benign mass. The dura was closed primarily using 4–0 silk suture and fibrin sealant (**- Fig. 3**).

#### **Hospital Course**

The patient was admitted to the neurosurgical floor postoperatively and recovered without complication. She remained at her neurologic baseline without any neurologic deficit and did report some interval improvement in her baseline left shoulder pain. She was discharged home on postoperative day 2. At her 2-week follow-up visit, the preoperative neck and shoulder pain improved by more than 50% and she had expected incisional soreness.

## Pathology

The histopathologic slides demonstrated infiltrating atypical melanocytes with prominent melanin pigmentation, nuclear pseudoinclusions, pleomorphic nuclei, and binucleation. The tumor stained positive for HMB45, Mart-1, SOX-10, and S-100 (**~Fig. 4**) and negative for epithelial membrane antigen (EMA), synaptophysin, and glial fibrillary acidic protein (GFAP). This histopathologic profile was consistent with the diagnosis of malignant melanoma.

#### Follow-up

A thorough skin examination did not reveal any lesions concerning for melanoma of the skin. MRI of the brain with and without contrast likewise did not demonstrate any lesions or enhancement of the leptomeninges. The patient underwent a whole-body positron emission tomography (PET)/computed tomography (CT) scan that demonstrated prominent fluorodeoxyglucose (FDG) uptake in the laminectomy bed consistent



**Fig. 4** The tumor is composed of infiltrating atypical melanocytes with prominent melanin pigmentation, abundant eosinophilic and finely granular cytoplasm, nuclear pseudoinclusions, pleomorphic nuclei, large eosinophilic nucleoli, and binucleation. (A) Hematoxylin and eosin (H&E), ×400. (B) HMB-45, ×400. (C) Mart-1, ×400. (D) SOX-10, ×400.

with postoperative changes but did not demonstrate any other sites of FDG uptake to indicate additional lesions. Based upon these radiographic findings and the histopathologic profile, a diagnosis of PSMM was made. The patient was treated with adjuvant fractionated radiation therapy (RT) with 35 Gy in five fractions to the resection bed and residual tumor left in the foraminal/extraforaminal space. At 8 months since surgery, the patient is doing well with continued improvement of her left neck and shoulder pain. MRI completed at 5 months after surgery demonstrated no progression of the residual tumor and no intradural recurrence.

# **Literature Review**

We identified in the existing body of literature a total of 112 reported cases of PSMM involving the neural elements, excluding 16 cases of PSMM isolated to the vertebral body (>Table 1).5-27 These 112 cases involved the extradural, intradural extramedullary, and/or intramedullary compartments (**-Table 1**). The thoracic spine was the most common location for PSMM (n = 45, 40.2%) followed by cervical (n = 39, 34.8%), lumbosacral (n = 16, 14.3%), and the thoracolumbar junction or conus medullaris (n = 12, 10.7%; **- Table 1**). PSMM of the spinal nerve root is even more uncommon; including the presently reported case, we identified only 21 cases reported in the literature that identified the tumor arising from the nerve root (>Table 1). Among PSMM of the nerve root, the most common location was the cervical spine (n = 10, 47.6%) followed by the lumbosacral spine (n = 7, 33.3%), and an even distribution between the thoracic (n = 2, 9.5%)and thoracolumbar junction or conus medullaris (n=2,9.5%).

Treatment data were available for 76 cases reported in the literature. All 76 patients underwent surgical resection. Gross total resection (GTR) was achieved in 33 (43.4%) patients, subtotal resection (STR) in 41 (53.9%) patients, and the extent of resection was not reported in the remaining 2 (2.6%) patients. Thirty-five (46.1%) patients underwent adjuvant treatment, which consisted of fractionated RT or chemotherapy or both. The majority of patients (n = 41,53.9%) did not receive any adjuvant treatment. Of the 35 patients who received adjuvant treatment, the most common treatment modality was fractionated RT alone in 23 patients (65.7%). Eight (22.9%) patients received both RT and systemic therapy and four (11.4%) patients received systemic therapy alone. The median fractionated RT dose was 45 Gy (range: 30-60 Gy). Agents used for systemic therapy included dacarbazine, levamisole, temozolomide, cisplatin, carmustine, interferon alpha, interferon beta, and anti-PD1 antibody.

We identified 64 studies that reported survival data for 84 patients. The overall survival (OS) of PSMM varied widely in the literature, ranging from less than 1 month to 25 years from diagnosis with a median survival after diagnosis of 17 months. At the last follow-up, 72.3% of patients were alive. One-year and 3-year OS were 83 and 55%, respectively.

## Discussion

The first description of PSMM was reported by Hirschberg in 1906.<sup>28</sup> Since then, only 110 cases of PSMM involving the neural elements have been reported. While PSMM represents an extremely rare cancer, PSMM of the spinal nerve root is even more uncommon; including the presently reported case, we identified only 21 such cases reported in the literature. Whereas PSMM in general was most commonly found in the thoracic spine, PSMM of the spinal nerve root was most commonly located in the cervical spine. The first description of PSMM arising from a spinal nerve root was described by Kiel et al in 1961 in a 33-year-old woman with a melanoma of the left C5 nerve root.<sup>29</sup> The lesion localized to the intradural extramedullary space and extended laterally into the left C5-6 neural foramen. The authors performed a laminectomy for tumor resection and noted a darkly pigmented tumor that involved the left C5 nerve root. In the present case report, we describe a case of a PSMM arising from the C3 nerve root, mimicking a cervical schwannoma, treated with STR and fractionated radiotherapy.

## Diagnosis

The diagnosis of PSMM of a spinal nerve root is challenging because the radiographic features resemble that of schwannoma or neurofibroma, and melanoma of the nerve root, as described above, is exceedingly rare. Melanoma within the spinal column is characterized on MRI by T1-weighted hyperintensity, T2-weighted iso- or hypointensity, and mild homogenous enhancement after gadolinium administration, mimicking the radiographic findings of benign nerve sheath tumors.<sup>30,31</sup> Definitive diagnosis, therefore, requires immunohistochemical analysis. Malignant melanoma demonstrates melanocytes with melanin pigmentation on hematoxylin and eosin (H&E) staining and positive staining for HMB-45, S-100, Mart-1, and SOX-10. Schwannoma, on the other hand, is characterized by alternating areas of compact spindle cells with nuclear palisading (Antoni A) and hypocellular areas with myxoid stroma (Antoni B) on H&E staining, and positive staining for S-100 and SOX-10. Importantly, schwannoma demonstrates low-grade cytologic features, whereas malignant melanoma demonstrates cytologically malignant cells. Once a diagnosis of spinal melanoma is confirmed, a thorough workup is required to identify whether the tumor represents a primary tumor without metastases versus metastatic melanoma. This workup should include a thorough skin examination and whole-body PET/CT to identify additional melanotic lesions. MRI of the neuraxis is also indicated to rule out other lesions within the CNS. The ultimate diagnosis of PSMM can then be made according to the criteria described by Hayward: (1) absence of melanoma outside of the CNS, (2) absence of melanoma in another area of the CNS, and (3) histologic confirmation of malignant melanoma.<sup>32</sup> PSMM portends a better prognosis than metastatic melanomas that involve the CNS, making this differentiation between PSMM and metastatic malignant melanoma critical for patient counseling and treatment.33-35

	_		_															_								_		_
Systemic therapy																												
Radiation																												
Extent of resection																											STR	STR
Location	IM	Extradural	IDEM	IDEM	IDEM	M	IM	IM	IDEM	IDEM	M	Extradural	IDEM	IM				IDEM	IM			IDEM	IM	IDEM	IDEM		Extradural/IDEM	IM exophytic
Level	Thoracolumbar	T8-11	T1-2	Cervical	C1–3	C7	Thoracic	T7–8	Thoracic	C7-T1	T7–8	T6	T9-10	T6	T5	11	12	Cauda equina	Cervicomedullary	19 1	L2-4	Cauda equina	Thoracic	Cervical	Cauda equina	T5–6	Cauda equina	Cauda equina
Alive at last follow-up	Dead														Alive		Alive				Alive						Dead	Dead
Survival (mo)													6		120	2	300			9	228		2	11	11	8	4	9
Recurrence or metastases (mo)																												
Sex	ц	Σ	Σ	ш	ш	Σ	ш	Σ	Σ	щ	щ	Σ	щ	ш	ш	Σ	ш	Σ	ш	ш	Μ	ц	Μ	ш	Μ		Σ	Μ
Age (y)	67	51	32	26	45	25	61	71	29	48	71	38	49	55	55	49	29	52	32	62	45	52	57	25	26	34	53	47
Year	1906	1907	1907	1910	1912	1916	1926	1926	1929	1930	1930	1933	1938	1939	1940		1940	1941	1942	1944	1950	1950	1950	1950	1951	1952	1952	
Author	Hirschberg	Boit	Esser	Kawashima	Lindborn	Koelichen	Ringertz	Schmid	Bau-Prussak	Bell	de Blasi	Van Bogaert	Schnitker	Da Costa	Moersch		Ray and Foot	Garcin	Mackay	Woods	Bakody	Castaner Vendrell	Forbes	Kissel	De Assis	Declich	King	

													rbazine,													(pointing
Systemic therapy								No			No		Carmustine, daca and levamisole		No	No	No	No	No	No		No	No		No	
Radiation								6,000 rads (cobalt)			No		No		50 Gy	60 Gy	45 Gy	50 Gy	No	Yes		No	50 Gy		No	
Extent of resection				No surgery			GTR		STR		GTR	GTR			STR	STR	STR	STR	STR	GTR	STR	GTR	STR	GTR	GTR	
Location	IDEM	IDEM		M			Σ	Σ	IDEM	Extradural	IDEM	MI	Extradural	IDEM	MI	IM	IM	IM	IM	Extradural	MI	IDEM	IM	IM	MI	
Level	T10-12	Lumbosacral	T12-1	Thoracolumbar	T6	T9-10	Cervical	T8-10	C4–6	S2	C4	C2-5	T7-T10	C1–6	T6-8	19	T9-11	C1–3	T9-10	L3-4	C7-T1	C5–6	C3-5	T8	T8-9	
Alive at last follow-up					alive		Dead	Dead	Dead	Dead	Dead		Alive	Alive	Alive	Dead	Alive	Dead	Dead	Alive	Alive	Alive	Alive	Alive	Alive	
Survival (mo)					42	4	surgical mortality	6.5	25	12	24		16	18	84	156	1	30	45	10	17	36	14	18	28	
Recurrence or metastases (mo)								No	10	6	18, metastasis		No	No	No	36	No	No	No	No		No			No	
Sex	Μ	щ		ц	Σ	Σ	Σ	Σ	щ	щ	Σ	ш	ц	ш	Σ	Μ	ч	ш	Ŀ	щ	щ	ц	Δ	Μ	Σ	
Age (y)	40	50		51	62	42		42	33	20	64	62	30	15	73	63	67	57	69	68	20	20	41	64	62	
Year	1953	1954	1956	1957	1957	1958	1959	1960	1961	1968	1968	1974	1984		1987					1987	1987	1994	1996	1996	1998	
Author	Perino	Roca de Vinals	Gros and Rotgen	Gibson	Leger	Zimmerman and Adams	Lang and Bridge	Hirano	Kiel <sup>a</sup>	Holaday <sup>a</sup>	Clifford	Jung	Ozden		Larson					Schneider <sup>a</sup>	Yoo	Skarli <sup>a</sup>	Bae	Magni	Francois	

Systemic therapy	No	No		No	ON	No	NO	NO	ON	No		Interferon alpha		No		Yes	Interferon beta, intrathecal dacarbazine		Interferon	No	Yes	Interferon alpha	
Radiation	30 Gy, 8 fractions	44 Gy, 22 fractions		40 Gy	ON	54 Gy (details NS)	NO	NO	ON	60 Gy		30 Gy over 2 wk		30 Gy, 10 fractions		No	50 Gy		30 Gy, 10 fractions	No	No	45 Gy, 25 fractions	
Extent of resection	STR	STR	GTR	STR	STR	STR	STR	STR	GTR	GTR	GTR	STR	GTR	STR	STR	GTR	STR	STR	STR	GTR	GTR	GTR	STR
Location	M	MI						IM	IDEM	Extradural	IDEM	Extradural	IDEM	IDEM		Extradural	IM	Extradural	Extradural	IM	MI	IDEM	IDEM
Level	T9-10	ΰ	T10	CI	C4	C4	Lumbar	T12-L1	L3	C6-7	L1–2	12	C2-4	C6–7	T6–8	17	T6	C6-7	T7–8	T4	T10	C1–6	C6-7
Alive at last follow-up	Alive	Dead	Alive	Alive	Alive	Dead	Alive	Alive	Alive	Alive	Alive	Alive	Alive	Alive	Alive	Alive	Alive		Dead	Alive	Alive	Alive	
Survival (mo)	21	14	22	16	20	8	2	6	24	8	3		3	6	3	9	216		6.5	36	12	17	
Recurrence or metastases (mo)	No	Brain metastasis	No	16	20	8	2	No	No	No	3, metastasis		No	No		No	216		No	No	6	No	
Sex	щ	Σ	ш	Σ	ш	щ	Σ	ш	Σ	ш	ш	ш	ш	ц	Σ	ш	Μ	Σ	щ	ш	ш	Μ	ш
Age (y)	76	62	71	52	20	57	53	80	26	45	57	42	41	76	34	37	31	65	68	34	57	39	71
Year	1998	1998	1999	_	_	_	_	2001	2003	2004	2004	2004	2005	2007	2007	2007	2009	2009	2010	2010	2010	2010	2010
Author	Salame	Salpietro	Brat					Farrokh	Sanz-Trelles <sup>a</sup>	Kwon <sup>a</sup>	Montinaro <sup>a</sup>	Naing <sup>a</sup>	Kounin	Kanatas <sup>a</sup>	Mekni	Unal	Nishihara	Roh <sup>a</sup>	oſ	Kim	Kolasa	Lee	Lee <sup>a</sup>

Table 1 (Continued)

Systemic therapy	No			No	No	Temozolomide		No	No		No		No	No	No	No	No	Temozolomide	No	No	No	No	No	No	No	No
Radiation	30 Gy, 10 fractions			No	Yes	Yes		No	No		No		No	No	No	No	No	30 Gy, 10 fractions	Yes	40 Gy, 16 fractions	No	No	45 Gy, 25 fractions	45 Gy, 25 fractions	No	45 Gy, 25 fractions
Extent of resection	STR	GTR	GTR	GTR	STR	GTR	GTR	STR	STR	GTR	GTR	GTR	GTR	GTR	STR	GTR	GTR		STR	STR	GTR	STR	STR	STR	GTR	STR
Location	IM	IDEM	IDEM	Extradural		Extradural	IDEM	IDEM	IDEM	Extradural	M	IDEM	IDEM	IDEM/ Extradural	IDEM	IDEM	IDEM	IDEM	MI	IDEM	IDEM	IDEM	MI	M	IDEM	IDEM
Level	C2-3	C1-2	C1-5	17	12	L3	L2-4	C2-6	12	L4	T9-L1	T4-T5	C2-C3	C5	T9-10	C4-5	L2-3	T8-9	T7-8	T12	C2-6	T12-L1	C1-C2	T6–7	T4–6	C5-6
Alive at last follow-up	Alive	Alive	Alive	Dead	Dead	Alive		Dead	Alive	Alive	Alive		Alive	Dead	Alive	Alive	Alive	Alive	Alive	Alive	Dead	Dead	Dead	Alive	Alive	Alive
Survival (mo)	11	4	4	204	2	48		2	22	38	6		24	-	7	76	67	24	84	24	25	10	14	72	96	38
Recurrence or metastases (mo)	No	No	No	Yes, metastasis		Metastasis		No	22	No	No		No	No	No	No	No	24	No	No	24	8	12	No	No	No
Sex	Μ	Σ	Σ	Σ	щ	щ	ш	Σ	Σ	Σ	ш	щ	щ	Σ	Σ	Σ	Σ	Σ	щ	ш	Σ	Μ	Σ	ц	ш	ш
Age (y)	62	40	16	30	82	49	44	48	42	55	47	57	54	54	39	47	76	28	51	57	47	47	51	23	39	57
Year	2011	2011		2011	2012	2012	2012	2012	2013	2013	2014	2014	2014	2015	2015			2015	2016	2016	2017					
Author	Fuld	Jaiswal		Katalinic <sup>a</sup>	Cicuendez	Ganiusmen <sup>a</sup>	Yan <sup>a</sup>	λu	Jeong	Sinha <sup>a</sup>	Cetinalp	Li	Marx	Beculic <sup>a</sup>	Liu <sup>a</sup>			Mallick	Agarwalla	Hering	Mu					

Table 1 (Continued)

Author	Year	Age	Sex	Recurrence or	Survival	Alive at last	Level	Location	Extent of	Radiation	Systemic
		( <u>v</u>		metastases (mo)	(mo)	follow-up			resection		therapy
		44	ц	No	52	Alive	T2-3	IDEM	STR	45 Gy, 25 fractions	No
Martinez <sup>a</sup>	2017	47	Σ	No	76	Alive	C4-5	IDEM	GTR	No	No
lga	2018	39	Σ	No	24	Alive	C2-5	IDEM	STR	No	Anti-PD1 antibody
Wuerdeman	2018	64	ш	No	96	Alive	T8	WI	STR	50.4 Gy	No
Zou <sup>a</sup>	2018	42	ц	No	16	Alive	C8	Extradural	GTR	Yes	Temozolomide and cisplatin
Chatterjee	2019	78	Μ	No	18	Alive	C7	WI	GTR	No	No
Hironaka	2019	39	Σ	n/a	14	Dead	L1–S5	IDEM	No surgery		
Sharma <sup>a</sup>	2019	67	щ	6	6	Alive	L1–2	IDEM	STR	No	No
Yoshizaki	2019	49	Σ	No	60	Alive	T12	Extradural	STR	36 Gy	Dacarbazine
Hanft <sup>a</sup>	2023	53	ч	No	8	Alive	C2-3	IDEM	STR	Yes	No
Abbreviations: GTR <sup>a</sup> Cases of PSMM ari:	, gross to sing from	tal resec the spir	ction; IC nal nerv	JEM, intradural extrame ve root.	dullary; IM, in	tramedullary; STR	, subtotal resection.				

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Table 1 (Continued)

## Management

Management of PSMM, and of malignant melanoma in general, requires a multimodality, multidisciplinary treatment approach that includes surgery, RT, targeted therapy, and immunotherapy. A challenging aspect of the treatment of PSMM is that the diagnosis is unlikely to be known prior to surgical intervention because these rare tumors mimic benign tumors of the nerve sheath or meninges (i.e., schwannoma, meningioma). Additionally, because they are primary lesions, there are no lesions elsewhere in the body as would be observed with metastatic disease. Surgeons, therefore, often go into surgery without high suspicion for PSMM. The tailoring of treatment thus does not begin until malignant melanoma is confirmed on immunohistochemical analysis and further workup does not reveal any other melanotic lesions, confirming the diagnosis of PSMM. Because the number of PSMM cases in the literature is so low, there is no accepted standard treatment. If PSMM is somehow known prior to surgery (e.g., biopsy of an extradural lesion or extradural portion of dumbbell-shaped tumor), one may consider aggressive resection to achieve GTR with spinal column stabilization and reconstruction, as necessary.<sup>35</sup> More likely, however, surgeons will resect as much tumor as possible safely without risk of neurological injury, perhaps leaving residual tumor if it is adherent to the spinal cord or if it extends into and/or out of the neural foramen, as we did in our presented case. The question then becomes what to do with the residual tumor. The literature suggests that surgical resection followed by adjuvant fractionated RT may be a reasonable treatment paradigm. Our literature review revealed that 45% of patients underwent adjuvant treatment following surgical resection and the median RT dose was 45 Gy. However, a greater majority of patients who underwent STR went on to receive adjuvant treatments compared to patients who underwent GTR of the lesion (60 vs. 21.9%). All patients with subtotally resected lesions who did receive adjuvant treatment received RT, with or without systemic therapy, except for one patient who received chemotherapy only. In this case, if we had known that the pathology was melanoma, which the darkly pigmented nature of the tumor did suggest, would we have performed a facetectomy with instrumented fusion in an attempt for a GTR? This is a difficult argument to make as there is significant additional morbidity of this extended operation, including increased operative time, increased blood loss, upfront instrumentation secondary to facet takedown, oft-challenging cerebrospinal fluid (CSF) leak repair from resection of the extradural tumor and widened nerve root sleeve, dissection in proximity to the vertebral artery, and persistent possibility of still leaving tumor behind. In drawing upon the growing body of literature that supports the notion of separation surgery in epidural melanoma metastatic cases (as with other malignant metastases) followed by adjuvant radiation, we believe similar logic applies in this case. In essence, this was an intradural version of a separation surgery operation, and we are hopeful that the patient will experience long-term progression-free and OS with this combination of STR and adjuvant radiotherapy.

#### Survival

The OS of PSMM varied widely in the literature. The majority of publications, however, reported only on a single patient. There exist only three publications that reported on a case series of five or more patients. The earliest, in 1987, reported on a series of five patients with intramedullary PSMM who were treated with subtotal surgical resection followed by adjuvant fractionated radiotherapy (45-60 Gy) in four patients, achieving a mean OS of 6.7 years (median: 45 months).<sup>33</sup> Brat et al reported on their series of primary melanocytic neoplasms of the CNS, including five cases of PSMM.<sup>36</sup> GTR was achieved in one case with no tumor recurrence after 22 months of follow-up. The remaining four tumors underwent STR with two receiving adjuvant fractionated RT. All subtotally resected tumors were found to have recurrence during follow-up. One-year recurrence-free survival (RFS) was 60% (median RFS: 16 months). Finally, Wu et al reported on seven patients with PSMM.<sup>34</sup> Treatment included surgical resection with adjuvant fractionated RT (45 Gy in 25 fractions) for STR. One-year and 3-year RFS were 71.4 and 57.1%, respectively. One-year and 3-year OS were 85.7 and 71.4%, respectively. Of note, all three patients who had tumor recurrence died within 2 months of recurrence diagnosis. Taken altogether, one may conclude from the available data that, although limited due to the small sample size, median survival following a diagnosis of PSMM is around 17 months. Compared to the dismal historical median survival of metastatic melanoma to the CNS of about 4 to 5 months (4.7 months in Davies et al<sup>37</sup> and 5.2 months in Raizer et a<sup>38</sup>) and more recently 6 to 12 months (Kotecha 6 BRAF 9, McHugh 8, Rauschenberg 12)<sup>36,39-41</sup> with combined modern treatment modalities consisting of surgery, radiotherapy, targeted therapy, and/or immunotherapy,<sup>37,38,40,41</sup> PSMM appears to portend a better prognosis. As targeted therapies and immunotherapies continue to evolve, prognosis of both PSMM and metastatic melanoma to the CNS can be expected to improve.

## Conclusion

PSMM is a rare cancer of the CNS, and PSMM of the spinal nerve root is even more extraordinary. The present case report adds to the 110 cases of PSMM and the 20 cases of PSMM of the spinal nerve root in the existing body of literature. Radiographic and clinical features resemble that of the much more common schwannoma or neurofibroma requiring immunohistochemical analysis for definitive diagnosis. PSMM may require adjuvant treatment postoperatively to limit recurrence or metastases, unlike benign nerve sheath tumors, which makes the diagnosis crucial for patient survival. The optimal treatment for PSMM has not yet been defined due to its rarity, and it is therefore important to report such cases in order to share our clinical experiences and provide data to other clinicians treating this uncommon disease.

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