

Malignant intracerebral nerve sheath tumor in a patient with Noonan syndrome: illustrative case

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BACKGROUND Malignant peripheral nerve sheath tumors (MPNSTs) within the neuroaxis are rare, usually arising from peripheral and cranial nerves. Even more scarce are cranial subclassifications of MPNSTs termed “malignant intracerebral nerve sheath tumors” (MINSTs). These tumors are aggressive, with a strong tendency for metastasis. With this presentation, alongside resistance to adjunctive therapy, complete excision is the mainstay of treatment, although it is often insufficient, resulting in a high rate of mortality.

OBSERVATIONS The authors report the case of an adult patient with a history of Noonan syndrome (NS) presenting with slowly progressive right-sided hemiparesis and right-sided focal motor seizures. Despite initial imaging and histology suggesting a left frontal lobe high-grade intrinsic tumor typical of a glioblastoma, subsequent molecular analysis confirmed a diagnosis of MINST. The patient’s neurological condition improved after gross-total resection and adjuvant chemo-radiation; he remains on follow-up.

LESSONS MINSTs are rare neoplasms with a poor prognosis; management options are limited, with surgery being the cornerstone of treatment. Reports on rare tumors such as this will increase awareness of this particular pathology and disclose clinical experience. In this case, the authors were unable to establish a definite cause-and-effect relation between NS and MINST. Nevertheless, it remains the first reported case in the literature.

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KEYWORDS malignant peripheral nerve sheath tumor; malignant intracerebral nerve sheath tumor; malignant schwannoma; frontal lobe; intraparenchymal

Malignant peripheral nerve sheath tumors (MPNSTs) occurring within the neuroaxis are rare, usually arising from peripheral and cranial nerves. Even more unusual are subclassifications evolving from the brain parenchyma, widely known as malignant intracerebral nerve sheath tumors (MINSTs). Because of a propensity for fierce regional infiltration and complex metastatic capability, these tumors are considered aggressive.¹ With this presentation, and a resistance to adjunctive treatment, gross-total resection (GTR) is

the mainstay of management; however, recurrence is often considered inevitable, particularly in cases in which GTR is not achieved.² Therefore, MINSTs are associated with a high rate of mortality and poor overall survival; notwithstanding these outcomes, survival figures differ in the literature, particularly at 1 and 5 years after diagnosis, highlighting the need for a review of the literature. Here, we present a rare case of an MINST in the frontal lobe that was initially suspected to be a high-grade glioma, was initially managed with GTR,

ABBREVIATIONS CNS = central nervous system; GFAP = glial fibrillary acidic protein; GTR = gross-total resection; LL = lower limb; MAPK = mitogen-activated protein kinase; MINST = malignant intracerebral nerve sheath tumor; MPNST = malignant peripheral nerve sheath tumor; MRI = magnetic resonance imaging; NF-1 = neurofibromatosis-1; NS = Noonan syndrome; TERT = telomerase reverse transcriptase; UL = upper limb; WHO = World Health Organization.

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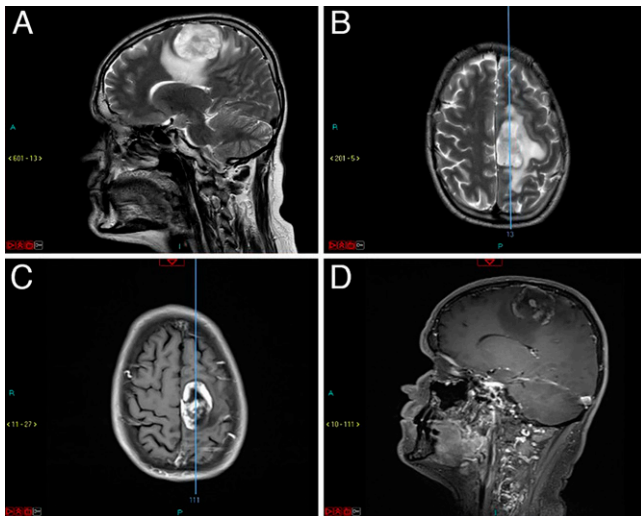


FIG. 1. Preoperative T1- and T2-weighted MRI demonstrating a contrast-enhancing left posterior frontal lesion with necrotic features and surrounding edema. **A:** Preoperative sagittal T2-weighted MRI. **B:** Preoperative transverse T2-weighted MRI. **C:** Postoperative transverse T1-weighted MRI. **D:** Postoperative sagittal T1-weighted MRI.

and has since completed adjuvant treatment. We have also undertaken a review of the medical literature with the aim to present the latest diagnostic and therapeutic developments on this rare entity.

Illustrative Case

A 49-year-old, left-handed man with a background of Noonan syndrome (NS) and who had undergone surgery early in childhood for congenital pulmonary stenosis presented to our neurosurgical department with a 7-month history of progressive right-sided hemiparesis and, more acutely, daily focal motor seizures of the right arm and leg.

Neurological examination elicited normal cranial nerve function, including visual fields with no neglect and normal fundoscopy. Motor examination demonstrated normal power and tone in the left upper limb (UL) and lower limb (LL), with a pyramidal catch in the right UL and obvious pyramidal weakness in the right LL (hip flexion 3/5, extension 4/5, knee flexion 4/5, extension 4/5, dorsiflexion 2/5, plantarflexion 4/5) with symmetrical, pathologically brisk reflexes. There was reduced sensation down the right arm, leg, and torso to all modalities. His performance status was 2 (according to the World Health Organization [WHO] classification).

Initial magnetic resonance imaging (MRI) in September 2020 demonstrated a heterogeneously enhancing lesion with necrosis, suggestive of a high-grade glioma (Fig. 1). Thus, the patient was listed for a neuronavigated craniotomy for debulking of the lesion after multidisciplinary team review.

Intraoperatively, a linear incision was made and a posterior frontal craniotomy was performed using neuronavigation. An encapsulated tumor was subsequently found and macroscopically excised, achieving GTR. Postoperatively, the patient had a Glasgow Coma Scale score of 15, with an improving right-sided weakness (UL 2/5, LL 3/5). Of note, the postoperative MRI and planning MRI for radiotherapy subsequently demonstrated contrast enhancement within the caudal aspect of the surgical bed (Fig. 2), deemed later to be postsurgical inflammatory changes.

Initial histology reports noted an unusual high-grade neoplasm with a predominantly spindle cell appearance. Further immunostaining revealed

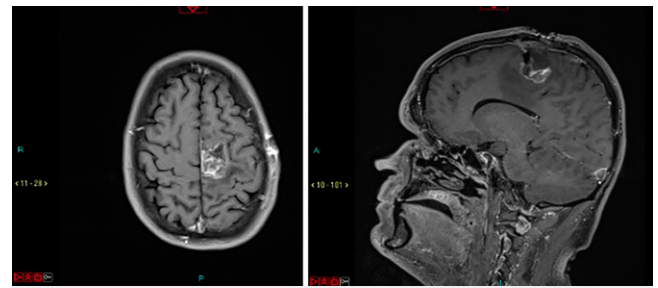


FIG. 2. Postoperative transverse (left) and sagittal (right) T1-weighted MRI demonstrating complete resection of the left paramedian posterior frontal tumor.

a tumor negative for *IDH1* (R132H) and *BRAF* (V600E) mutation with *ATRX* retention and a Ki-67 labeling index of 20% to 30% (complete histomolecular analysis is shown in Table 1). Histological appearances revealed an unusually high-grade, intrinsic central nervous system (CNS) tumor, with a differential diagnosis including gliosarcoma (with a desmoplastic-appearing glial component that also included abnormal ganglion/neuronal cells) and anaplastic pleomorphic xanthoastrocytoma (Fig. 3). Sanger sequencing further showed the tissue to be *IDH1*-R132 and *IDH2*-R172 wild type, with no mutations in the telomerase reverse transcriptase (*TERT*) promoter or histone H3F3A genes. The DKFZ (German Cancer Research Center, Heidelberg) sarcoma methylation classifier

TABLE 1. Initial immunohistochemistry and molecular pathology analysis results

Molecular/Immunohistochemical Analysis	Result
IDH (R132H) IHC	Negative
<i>IDH1/IDH2</i> sequencing	No mutation
FISH for 1p/19q codeletion	Not performed
<i>ATRX</i>	Retained in neoplastic cell population
MGMT promoter status	Unmethylated
<i>TERT</i> (228, 250)	No mutation
Histone H3F3A (K27, G34)	No mutation
<i>BRAF</i> (V600E) (IHC/sequencing)	Negative
H3K27me3	Loss of nuclear expression
NFP	Highlights abnormal neuronal population w/ entrapped axons in keeping w/ an infiltrative growth pattern
GFAP	Patchy positive staining
P53	A small population of scattered weakly positive cells
STAT6	Negative
Ki-67 proliferation	20%–30%

ATRX = *ATRX* gene; *BRAF* = B-Raf proto-oncogene; FISH = fluorescent in-situ hybridization; H3K27me3 = histone 3 lysine 27 trimethylation; *IDH* = isocitrate dehydrogenase; IHC = immunohistochemistry; MGMT = O6-methylguanine DNA methyltransferase; NFP = neurofilament protein; STAT6 = signal transducer and activator of transcription 6.

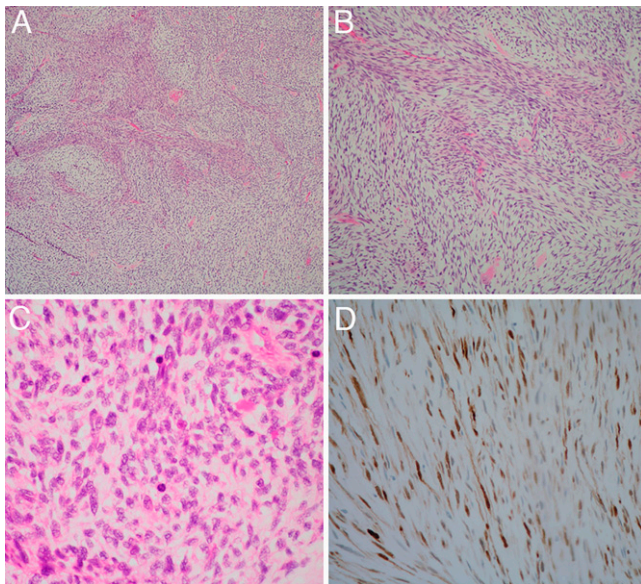


FIG. 3. Light microscopy images showing tumor appearance. **A:** Original magnification $\times 40$. Hematoxylin and eosin (H&E) stain demonstrating the alternating compact and loose growth pattern. **B:** Original magnification $\times 100$. H&E stain demonstrating a higher-power view of panel A. **C:** Original magnification $\times 400$. H&E stain showing four mitotic figures (dark structures scattered among the tumor cell nuclei). **D:** Original magnification $\times 200$. S100 stain demonstrating S100-positive immunohistochemistry labeling in some foci among the tumor cell nuclei and cytoplasm.

categorized this tumor as an MPNST with a high calibrated score. Further support of this diagnosis was provided by the loss of nuclear H3K27me3 expression in the tumor cells. This diagnosis also accounts for the immunohistochemical profile obtained before array analysis.

After bi-institutional multidisciplinary board assessment and review of current data in the medical literature, we offered the patient 6 weeks of adjunctive treatment with concurrent conventional (linear accelerator-based) radiotherapy (60 Gy in 30 fractions) and temozolomide chemotherapy, to which the patient consented; treatment was completed in February 2021. He is currently seizure-free and remains cognitively intact; his right-sided hemiparesis has improved, although he still struggles with his LL (LL power 3–4/5, UL power 4/5) while undergoing physiotherapy. His performance status currently is 1, mainly because of his current neurofunction. His first follow-up MRI (April 2021) showed no evidence of recurrence; the patient is to remain on close surveillance with clinical and MRI follow-ups every 3 months. The interim genetic analysis for neurofibromatosis-1 (NF-1) showed negative results.

Discussion

Observations

General Aspects Relevant to This Case

MPNSTs are highly aggressive soft tissue sarcomas. Originally termed “malignant schwannomas,” albeit incorrectly because of schwannomas not regularly undergoing malignant transformation, MPNSTs usually arise from the nerves in the limbs or trunk/axillae or form sporadically from existing neurofibromas.^{3–5} Forty percent to 50% of all MPNSTs occur in the presence of NF-1; however, spontaneous occurrences also have been reported (50%).^{6,7} Extraaxial sites

of occurrence predominantly involve peripheral nerves, specifically in the limbs and trunk, although cranial nerves can also be sites of origin.^{7,8} Spontaneous MPNSTs are scarce, with an incidence of 0.1/100,000 persons per year.⁹ MPNSTs can be either extra- or intracranial, with the latter type being subclassified into extraaxial and intraparenchymal tumors. Serious consideration was given to this case because intracranial, intraparenchymal MPNSTs are extraordinarily rare, and reports in the international literature remain scarce. Because of the complex anatomic-topographic deployment inherent to their character, these neoplasms are generally known as MINSTs.¹⁰ Unfortunately, this subclassification is noted to have a penchant for a moderate to poor response to radiation and chemotherapy, with GTR with wide margins considered the cornerstone of management.^{2,11} However, postoperative recurrence and distant metastasis remain likely as a result of the aggressive nature of these tumors;¹ indeed, the 1-year overall survival rate stands at 33%.¹² Therefore, adjunctive radical radiotherapy is often integrated in the portfolio of mainstay therapy.

In the case presented here, there is a background of NS, an autosomal dominant condition with a variable phenotype, in which 50% of cases are due to a germline “gain of function” mutation of the *PTPN11* gene; this particular gene is responsible for encoding the nonreceptor protein tyrosine phosphatase SHP2, positively controlling the RAS function within the RAS–mitogen-activated protein kinase (MAPK) signaling pathway.^{13,14}

Considering the nature of the RAS-MAPK pathway and its role in oncogenesis, patients with certain mutations (so-called RASopathies), and thereby NS, are at an increased risk of certain cancers. Somatic mutations of *PTPN11* have been reported as being present in 35% of persons with juvenile myelomonocytic leukemia, alongside other hematological malignancies and solid organ tumors, such as lung and colon cancer and neuroblastoma.^{15,16} Overall, individuals with NS have an estimated cancer risk of 4% by 20 years old.^{13,17,18}

Currently, limited evidence details the relationship between CNS tumors and NS, with most articles providing case reports of pediatric glial tumors. A case report and literature review of these tumors occurring alongside NS by Lodi and colleagues in 2020 demonstrated that most cases occurred in the pediatric population and were dysembryoplastic neuroepithelial tumors.¹³ Adult cases and primary brain tumor reports remain extremely scarce.

The significance of NS preexisting a diagnosis of MINST is unclear in the literature; indeed, should the relationship prove to be causal, our case would provide the first instance of MINST reported in an adult with NS. However, because of the lack of evidence in the medical literature, we tend to remain cautious on this subject.

Diagnosis

As illustrated by this case, preoperative diagnosis can be difficult, and the lesion’s appearance can be indistinguishable from high-grade glioma on imaging.⁵ MR spectroscopy showing a high choline peak without creatine and *N*-acetyl aspartate resonance has been suggested to differentiate between a glial and nonglial tumor, but awareness and a degree of suspicion are required;¹⁹ the latter study was not made available for our case because local institutional guidelines are not yet set in that direction. From a histological standpoint, the cellular origin of these neoplasms is still unknown; however, some groups have suggested Schwann cells of perivascular nerves or pluripotent mesenchymal cells as plausible sources.²⁰

Important diagnostic markers include the S100 protein, useful to demarcate nerve sheath tumors from tertiary soft tissue neoplasms;

TABLE 2. Summarized cases reported in the literature to date

Case No.	Authors & Yr	Age (yrs) at Diagnosis	Gender	Laterality	Location	Surgical Procedure	Postop Therapy	Recurrence (mos)	FU (mos)	Survival at Last FU
1	Current study	49	M	Lt	Frontal	GTR	RT & CT	—	Ongoing (6 at time of writing)	Alive
2	Le Fèvre et al., 2016 ²⁴	68	F	Lt	Fronto-temporal	STR	RT	7	15	Dead
3	Le Fèvre et al., 2016 ²⁴	47	F	Rt	Frontal	GTR	RT	6, 10, 13	20	Alive
4	Smith et al., 2014 ²⁰	26	M	—	Bifrontal	STR	—	1	12	Dead
5	Lee et al., 2013 ²²	13	M	Rt	Frontal	Resection (not specified)	RT	50, 54, 60	77	Alive
6	Shweikeh et al., 2013 ⁵	18	M	Rt	Fronto-parietal	GTR	RT	44	52	Dead
7	Gong et al., 2012 ²⁵	55	F	Lt	Cerebellopontine angle	Resection (not specified)	—	No	5	Alive
8	van den Munckhof et al., 2011 ²⁶	6	F	Rt	Fronto-parietal	GTR	RT & CT	15	48	Alive
9	Ellis et al., 2011 ²⁷	9	F	Rt	Fronto-temporal	STR	CT & RT	No	6	Alive
10	Bamard et al., 2011 ²⁸	75	F	Lt	Frontal	GTR	RT	No	12	Alive
11	Oztanir et al., 2009 ²⁹	1	F	Rt	Fronto-temporo-parietal	STR	—	—	1.5	Dead
12	Scheithauer et al., 2009 ⁸	69	M	Rt	Frontal	No treatment	—	—	4	Dead
13	Scheithauer et al., 2009 ⁸	26	M	—	Posterior fossa	Not specified	—	—	Lost to FU	—
14	Scheithauer et al., 2009 ⁸	41	M	—	Posterior fossa	Resection (not specified)	RT	—	5	Dead
15	Kozić et al., 2008 ¹⁹	39	M	Lt	Pontine	Biopsy	—	—	—	Not Specified
16	De Cauwer et al., 2007 ³⁰	68	F	Lt	Parieto-frontal (rolandic area)	GTR	RT	5	5	Dead
17	Maiuri et al., 2004 ³¹	36	M	—	Cerebellar vermis	GTR	RT	6	8	Dead
18	Beauchesne et al., 2004 ³	35	M	Rt	Cerebral peduncle	Biopsy	RT & CT	17	29	Dead
19	Bornstein-Quevedo et al., 2003 ³²	3	M	Rt	Parieto-occipital	STR	—	No	0.33	Dead
20	Takahashi et al., 2000 ³³	57	M	Rt	Lateral ventricle	GTR	RT & CT	—	4	Dead
21	Tanaka et al., 2000 ³⁴	4	F	Rt	Parieto-occipital	GTR	—	No	19	Alive
22	Sharma et al., 1998 ³⁵	8	F	Rt	Temporal	GTR	—	No	17	Alive
23	Jung et al., 1995 ³⁶	40	M	Rt	Lateral ventricle	GTR	RT	8	8	Dead
24	Singh et al., 1993 ³⁷	61	F	Rt	Cerebellar	GTR	RT	10	18	Dead
25	Stefanko et al., 1986 ³⁸	15	M	Lt	Parieto-occipital	GTR	RT & CT	5, 8	9	Dead
26	Bruner & Armstrong, 1984 ³⁹	18	M	—	Bifrontal	GTR	—	24, 48, 66	66	Alive

CT = chemotherapy; FU = follow-up; RT = radiotherapy; STR = subtotal resection.

loss of nuclear H3K27me3 expression, seen in most MPNSTs;^{7,21} negative glial fibrillary acidic protein (GFAP) expression, which helps exclude desmoplastic astrocytoma, gliofibroma, and gliosarcoma; and negative synaptophysin and neurofilament protein, which reliably distinguish MINSTs from desmoplastic ganglioglioma.²² Other differential diagnoses include other tumors such as rhabdomyosarcoma, gastrointestinal stromal tumor, and meningioma.²⁰

In our case, S100 positivity coexisting with patchy GFAP expression proved challenging and certainly delayed diagnosis. Additionally, as mentioned above, this tumor initially appeared similar to a glioblastoma on imaging. Histologically, this was thought not to be the immediate case because of certain morphological clues such as the appearance of a malignant tumor with a mesenchymal appearance, alongside the aforementioned absent GFAP expression. Gliosarcoma was indeed considered; however, it was deemed unlikely because of the lack of biphasic architecture (only malignant mesenchymal with no high-grade glioma component). A tumor falling within the hemangiopericytoma/solitary fibrous tumor spectrum was also excluded based on STAT6 immunohistochemistry and DNA methylation profiling. The latter also helped to rule out a diagnosis of anaplastic pleomorphic xanthoastrocytoma. Further analysis to differentiate between MINST and other high-grade tumors, such as SOX10 and reticulin staining, could have been conducted; however, local guidelines favor the reliability of H3K27me3 and DNA methylation profiling from a diagnostic perspective. Local capabilities of analyzing Olig2 expression were not available to us at the time of diagnosis. To that end, the histological and immunohistochemical profile of this tumor did not fit with any known high-grade glioma or any other specific entity recognized by the current WHO classification of CNS tumors.

Treatment and Follow-Up

Surgical management of intracranial MPNSTs is technically complex because of frequent involvement with a cranial nerve. Indeed, Patankar et al. described the challenges associated with resection of an MPNST in the middle cranial fossa and its close involvement with the facial nerve and geniculate ganglion, resulting in a postoperative facial palsy.¹⁰ Scheithauer and colleagues also reported a similar finding.⁸ In the case discussed within this report, no cranial nerves were involved. Of note, this is not surprising because intraparenchymal MINSTs arise from the brain parenchyma, and as a result cranial nerve involvement remains a less common finding in similar cases.²³ In terms of adjuvant therapy, radiotherapy remains the most favored approach; Lee et al. found 60 Gy in 30 fractions beneficial in cases of recurrence.²² Alternatively, different lines of chemotherapy have been suggested by different authors, including combination vincristine, carboplatin, etoposide, single-agent temozolomide, and ifosfamide-based treatment;³ from our institutional experience, combination procarbazine, lomustine, and vincristine or single-agent lomustine may also be options worth considering. However, the outcome is still poor. Overall, we suggest that chemotherapy be given in chemoradiation settings (as in this case) or when radiotherapy is not indicated; also, as shown in Table 2, lifelong follow-up (including MRI) should be considered for those few long-term survivors. In this context, follow-up should be tailored to clinical evolution and survival timeline.

Lessons

Multiple lessons can be learned from this case, particularly regarding diagnosis and management. (1) There have been few reports of MINSTs in the literature, with only 25 reported since 1984; none of these cases had a background of NS (Table 2). As a result, it is hypothesized that the rarity of malignant intraparenchymal peripheral

nerve sheath tumors brings about a lack of evidence on the best management of these lesions, and although there is insufficient evidence to establish a proper cause-and-effect relation between the two entities, it cannot be fully discarded considering NS's oncogenic capability. (2) MINST can mimic glioblastoma clinically, radiologically, and, to some extent, even histopathologically. This possibility highlights the importance of reliable immunohistochemistry and institutional experience with these rare neoplasms. (3) With surgery being the mainstay treatment, postoperative treatment remains surrogate to the clinical evolution of MINSTs. However, the scarce number of patients makes retrospective analytical work and prospective studies hardly feasible; thus, there is a lack of consensus regarding the adequate postoperative and/or adjuvant management of these tumors, although it seems that radiotherapy has a clearer therapeutic role than chemotherapy.

We believe this case to be the 26th reported case of MINST in the available literature; additionally, to our knowledge, this is the first reported case of MINST in an adult patient with a background of NS. However, as pointed out above, the lack of data in the medical literature makes it difficult to discern between a complex causal relation and a mere incidental finding. Finally, we suggest that an international registry focusing on diagnostics, applied treatments, and therapeutic outcome should be considered with the aim of achieving a wider consensus in the management of these rare and complex entities.

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Disclosures

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions

Conception and design: Sinclair, Allison, Shumon, Surash. Acquisition of data: Shumon, Allison. Analysis and interpretation of data: Sinclair, Shumon, Allison, Joshi, Quaegebeur. Drafting the article: Sinclair, Allison, Shumon, Quaegebeur. Critically revising the article: Sinclair, Allison, Shumon, Quaegebeur, Surash. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Sinclair. Administrative/technical/material support: Sinclair, Shumon, Joshi. Study supervision: Sinclair, Surash.

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