

CSH Molecular Case Studies

# Genomic profiling of a metastatic anaplastic melanocytic neuroectodermal tumor arising from a mature thymic teratoma as part of a mediastinal germ cell tumor

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Abstract Following chemotherapy, a mediastinal germ cell tumor can lead to a mature teratoma that is composed of tissues derived from all three germ layers. Although teratoma is usually curable, in rare cases it can give rise to various somatic tumors and exceptionally it undergoes melanocytic neuroectodermal tumor (MNT) transformation, a process that is not well-described. We report a patient with a postchemotherapy thymic teratoma associated with an MNT component who, 10 years later, additionally presented a vertebral metastasis corresponding to an anaplastic MNT. Using exome sequencing of the mature teratoma, the MNT and its metastatic vertebral anaplastic MNT components, we identified 19 somatic mutations shared by at least two components. Six mutations were common to all three components, and three of them were located in the known cancer-related genes KRAS (p.E63K), TP53 (p.P222X), and POLQ (p.S447P). Gene set enrichment analysis revealed that the melanoma tumorigenesis pathway was enriched in mutated genes including the four major driver genes KRAS, TP53, ERBB4, and KDR, indicating that these genes may be involved in the development of the anaplastic MNT transformation of the teratoma. To our knowledge, this is the first molecular study realized on MNT. Understanding the clinicopathological and molecular characteristics of these tumors is essential to better understand their development and to improve therapeutics.

[Supplemental material is available for this article.]

# INTRODUCTION

Mediastinal germ cell tumors (GCTs) represent a heterogeneous group of neoplasms originating from primitive germs left in the mediastinum during embryogenesis. They can be

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**Ontology term:** neoplasm of the thymus

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classified as seminomatous or nonseminomatous tumors (Marx et al. 2022). Their prognosis depends on tumor histology, stage, and serum markers. Nonseminomatous neoplasms represent up to 85% of mediastinal GCTs including teratoma (Pini and Colecchia 2022). Their treatment requires chemotherapy followed by surgical resection of the residual tumor (Albany and Einhorn 2013). This systemic treatment can select the teratomatous component present in the mediastinal GCT, leading to a residual postchemotherapy mature teratoma component at the initial and/or metastatic locations of the GCT (Green et al. 2021).

Teratoma is a GCT defined as a neoplasm originating from pluripotent cells that form differentiated somatic-type tissues, which may be either mature, immature, or both (Marx et al. 2022). The majority of teratomas carry mature tissues and have a favorable prognosis. Immature teratomas represent only 4% of mediastinal teratomas but have a malignant potential that correlates with the proportion of immature tissues (Dulmet et al. 1993; Moran and Suster 1997; De Backer et al. 2008). In rare cases, a part of the mature teratoma may undergo a somatic transformation into a malignant nongerminal cell tumor that is equivalent to a somatic malignancy (Motzer et al. 1998). A wide variety of somatic-malignancy transformations have been described but the most frequent ones are adenocarcinoma, primitive neuroectodermal tumors, and rhabdomyosarcoma (Motzer et al. 1998; Ryu et al. 2013; Marx et al. 2022). In addition, exceptional cases of melanocytic neuroectodermal transformation (MNT) have been reported (Misugi et al. 1965; Hameed and Burslem 1970; Tobo et al. 1981; King et al. 1985; Anagnostaki et al. 1992; Vajtai et al. 2000). MNTs are extremely rare, yet rapidly growing, benign tumors consisting of a biphasic population of small neuroblast-like and larger melanin-producing epithelioid cells. Children are mostly affected by MNTs that occur in the head and neck region (Almomani and Rentea 2022), but extracranial locations have also been reported (Styczewska et al. 2021). Nearly 500 cases have already been reported but only six were linked to a teratoma (Misugi et al. 1965; Hameed and Burslem 1970; Tobo et al. 1981; King et al. 1985; Anagnostaki et al. 1992; Vajtai et al. 2000).

The presence of somatic mutations induced by genomic instability of a precursor cell has been proposed to be the trigger of the development of GCTs and in particular teratoma (Feldman et al. 2014; Taylor-Weiner et al. 2016). Similar molecular events might contribute to the development of the malignant component. Although studying these cases is an opportunity to better understand the mechanisms that are involved in the malignant transformation, such studies are sparse in the literature, mainly descriptive and lacking molecular data. This caveat is mainly due to the rarity but also to the technical challenges of selecting and reliably analyzing the various neoplastic components.

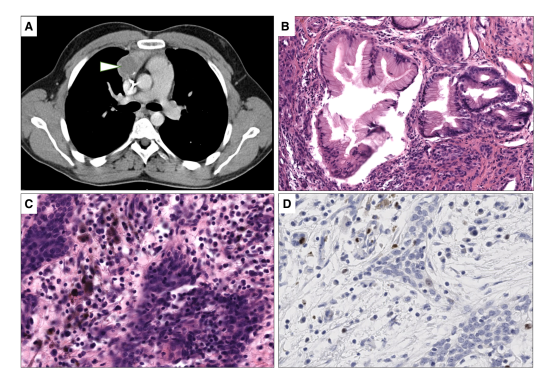
Here, we report an extremely rare case of a patient who presented with a mediastinal GCT presenting as a mature teratoma after chemotherapy with an MNT component and secondly developed vertebral metastatic localization with an anaplastic transformation of the MNT that was histologically close to a malignant melanoma. Besides clinical and pathological evaluation, we aimed to characterize the molecular events associated with the teratoma and its melanocytic transformation using both targeted gene-panel and exome sequencing.

#### RESULTS

#### **Clinical Presentation and Family History**

A Caucasian male born with anal imperforation and bilateral cryptorchidism was referred at 26 yr of age for dyspnea. No pathogenic germline variant in known genes of cryptorchidism, congenital anorectal malformations, and cancer susceptibility genes were identified. Physical and radiological examination showed a mediastinal tumor measuring  $9 \text{ cm} \times 8 \text{ cm} \times 7 \text{ cm}$  (Fig. 1A) metastatic to the pleura, liver, and an isolated brain metastasis. Initial assessment showed  $\alpha$ -fetoprotein at 40  $\mu$ g/L,  $\beta$ -human chorionic gonadotropin at

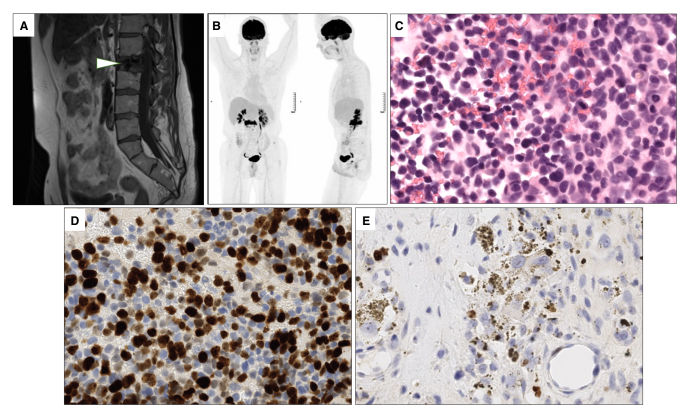




**Figure 1.** Radiological and histological features of mediastinal lesion. (*A*) Computed tomography scan imaging showing presence of hypodense lesion of the right anterior mediastinum (arrowhead). (*B*) Mature teratoma with mature fundic differentiation without cytologic atypia and representing 70% of the viable tumor (10×, hematoxylin and eosin [H&E] stain). (*C*) Melanocytic neuroectodermal transformation (MNT) component with one pleiomorphic cell population containing intracytoplasmic melanin pigments and a second small basophilic cell population (40×, H&E stain). (*D*) Ki67 immunohistochemistry showing a low proliferation index of the MNT component. Brown nuclear stain highlights a Ki67-positive tumor cell and granular cytoplasmic brown corresponds to melanin pigments.

61114 U/L, and lactate dehydrogenase at 1100 U/L, diagnosing nonseminomatous germcell cancer. He received several lines of chemotherapy including BEP (bleomycin, etoposide, cisplatin), T (paclitaxel)-BEP/oxaliplatin, and cisplatin-ifosfamide because of insufficient marker response. Five months later, he underwent brain surgery for the brain metastasis, which was partially removed, showing teratoma. He was treated with six cycles of doxorubicin, cyclophosphamide, and methotrexate followed by whole-brain radiation therapy and was considered as in complete remission. A recurrence of the occipital metastasis was removed at 27 yr of age and revealed reactional and benign fibromatous gliosis corresponding to post-treatment remodeling. Five months later, radical thymic compartment resection was performed. Microscopic examination revealed a tumoral mass composed of 50% fibro-necrotic remodeling. The viable territory was composed of 70% fundic and intestinal glandular mature teratoma (Fig. 1B), 25% MNT (Fig. 1C), and 5% angiosarcomatous component. MNT tumoral fraction was composed of pleiomorphic large cells with vesicular nuclei and intracytoplasmic melanin pigment inclusions and small round cells with little basophilic cytoplasm and small dark nuclei. Immunohistochemistry on MNT showed a strong expression of PS-100 as well as the melanocytic markers SOX10, Melan-A, and HMB45. MNT was negative for glypican 3, bHCG, synaptophysin, panKeratin AE1/AE3, and SALL4. The Ki67 proliferative index was low (<1%) (Fig. 1D)





**Figure 2.** Radiological and histological features of the vertebral metastases. (A) T1-weighted magnetic resonance image of the hypointense L2 spinal metastatic lesion (arrowhead). (B) 18F-fluorodeoxyglucose-positron emission tomography-computed tomography (18F-FDG-PET/CT) scan imaging revealing a hypermetabolism of the L2 spinal metastasis. (C) Microscopic examination of the pretreatment vertebral metastasis corresponding to an anaplastic melanocytic neuroectodermal transformation (MNT). Cells are pleiomorphic and numerous mitoses are visible (40×, hematoxylin and eosin [H&E] stain). (D) Ki67 immunohistochemistry in the same pretreatment vertebral biopsy showing an important proliferative index of the tumoral cells. Brown nuclear staining highlights Ki67-positive tumor cells and granular cytoplasmic brown corresponds to melanin pigments. (E) Ki67 immunohistochemistry of the post-therapy vertebral cell highlighting the presence of a residual MNT component with a lower proliferative index.

At age 28, standard follow-up showed an osteolytic lesion of L2 vertebra (Fig. 2A). Multiple biopsy samples from this lesion showed the presence of mesenchymatous tissue corresponding immature teratoma remnants with a low (1%) Ki67 index.

The patient remained in complete remission until age 38, when a fluorodeoxyglucosepositron emission tomography (FDG-PET) scan showed an increased L2 glucose uptake (Fig. 2B). Biopsy revealed a proliferation consisting of small-sized cells with chromatic nuclei and faint cytoplasm. Numerous mitoses were found with a Ki67 proliferative index of 50% (Fig. 2C,D). The lesion was histologically close to a malignant melanoma, expressed the melanocytic markers HMB45 and Melan-A and was classified as an anaplastic transformation of the MNT. The patient received another chemotherapy line with carboplatin and etoposide targeting the immature pathological features of the tumor were done and complete surgical L2 resection was performed. Microscopical analysis showed an important fibro-necrotic post-treatment remodeling with presence of 10% neoplastic proliferation close to the original MNT described above. The Ki67 proliferative index was 1% but reached 10% focally (Fig. 2E).



Radiological evaluation performed at age 39 showed locoregional retroperitoneal lymph node evolution leading to an immunotherapy initiation (ipilimumab–nivolumab). To date, the patient is still alive with poor therapeutic response.

## **Genomic Analyses**

An oncogenetic assessment including 26 target genes was carried out only on the post-treatment proliferative MNT component of the vertebral metastasis. This analysis revealed the presence of a p.E63K mutation in KRAS (variant allele frequency [VAF] 37.5% and 38% in targeted panel and exome, respectively). Then, we performed exome sequencing in order to identify somatic mutations specific of each neoplastic component and to further analyze their similarities. We analyzed the germline thymic sample, the mature teratoma, and the MNT components that were derived from the radical thymic compartment resection (at age 27 yr) and the metastatic anaplastic MNT that was derived from the vertebral metastasis (at age 37 yr). The sequencing of the mature teratoma, MNT, metastatic MNT, and matched germline thymic samples resulted in an average depth of coverage of 317.5, 405, 194.4, and 172, respectively. The three components shared six mutations, and 13 mutations were present in at least two components. Among the variants shared by the three components, three are modifying amino acids in proteins known to be involved in oncogenesis. The p.E63K mutation in KRAS is one of the mutations that had previously been described in the targeted gene-panel sequencing. The two others are p.P222X in TP53 and p.S447P in POLO (Table 1; Fig. 3A; Supplemental Table S1). Other variants in genes potentially involved in oncogenesis were also identified in each component (Fig. 3A). However, none of them are localized in codons that are already known to impact oncogenesis. Using gene set enrichment analysis (GSEA), we observed specific mutational profiles in genes involved in the "Melanoma," "Epithelial to mesenchymal transition in colorectal cancer," "Regulation of actin cytoskeleton," "Chemokine signaling pathway," and "Folate metabolism" pathways (Fig. 3B; Supplemental Table S2). Interestingly, the "Melanoma" pathway is the most enriched one and contains four genes that are all implicated in oncogenesis (KRAS, TP53, ERBB4, and KDR).

# DISCUSSION

We report a rare case of a postchemotherapy mediastinum teratoma leading to a somatictype malignant transformation toward MNT and anaplastic MNT that was histologically close to a malignant melanoma (Supplemental Fig. S1).

MNT is a rare tumor particularly affecting children of <1 yr of age and localized in the craniofacial region in 90% of cases (Barrett et al. 2002). To our knowledge, only six cases of MNT have been described in a context of teratoma. Three of them were related to ovarian teratoma (Hameed and Burslem 1970; King et al. 1985; Vajtai et al. 2000), two arose from mediastinal teratoma (Misugi et al. 1965; Anagnostaki et al. 1992), and one from the sellar region (Tobo et al. 1981). These patients are often older than those with classical MNT. Although MNTs are generally benign, three of the six previously described cases were very aggressive and metastatic (Hameed and Burslem 1970; King et al. 1985; Anagnostaki et al. 1992). Altogether, these data suggest that in the context of teratoma, the mechanism of tumor development is different from classical MNT.

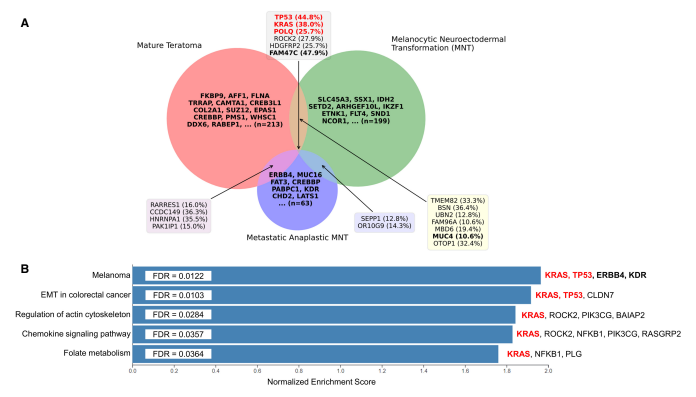
The absence of anatomopathological examination of the primitive tumor before neoadjuvant therapy did not allow us to clearly conclude about the origin of the teratoma component. It may have been initially present in the GCT and selected by the chemotherapy or it may result from the chemotherapy-induced maturation of immature cells of the GCT. Although the etiopathogenesis of teratoma is not fully understood, it has been suggested

Table	<b>3 1.</b> List o	f codi	ing sc	Table 1. List of coding somatic variants in census genes according	i census ger	ies according		to the COSMIC database	C database						
										Cosr in	Cosmic database information	Cancer type observation	Tera-		Ana- nlastic
Chr.	Pos.	Ref.	. Alt.	Type of variant	Gene symbol	HGVSp short	SIFT	FT PolyPhen	rsID/Cosmic ID	Census gene	Role in cancer	in COSMIC Database	()	MNT (VAF %)	MNT (VAF %)
~	7723485	U	4	Missense	CAMTA1	p.R293O	F	Prob. dam.	rs753954585	Tier 1	Tumor suppresor, fusion	Large intestine adeinocarcinoma, malignant melanoma	9.26	1	1
-	17965150	U	⊢	Missense	ARHGEF10L p.R416C	p.R416C		Prob. dam.	rs1304791222	Tier 2	Tumor suppresor	I	I	8.77	I
~	205628612		⊢	Missense	SLC45A3	p.R471H	⊢	Benign	rs765881137/ COSM6887586	Tier 1	Fusion	I	I	8.47	I
2	46605084	υ	⊢	Missense	EPAS1	p.P434L	Ω	Prob. dam.	rs745357489	Tier 1	Oncogene, tumor suppressor	I	9.62	I	I
5	212286781	U	∢	Missense	ERBB4	p.A972V	⊢	Benign	1	Tier 1	Oncogene, tumor suppressor	I	I	I	12.2
т	47165718	υ	∢	Missense and NMD transcript	SETD2	p.R8S		Prob. dam.	I	Tier 1	Tumor suppresor	I	I	12.82	I
т	121238847	A	G	Missense	POLQ	p.S447P		Poss. dam.	I	Tier 1	Oncogene, tumor suppressor	I	25.68	8.57	23.53
4	55948198	U	∢	Missense	KDR	p.T1258M	Ω	Prob. dam.	rs147627339	Tier 1	Oncogene	Large intestine adenocarcinoma, pancreatic ductal carcinoma	I	I	20.71
4	88035579	υ	⊢	Missense	AFF1	p.P525S	⊢	Prob. dam.	I	Tier 1	Fusion	I	13.89	I	I
5	180058713	U m	⊢	Missense	FLT4	p.D42N	⊢	Benign	rs777334601	Tier 1	Oncogene	1	I	10.14	I
9	150001575	0	∢	Missense	LATS1	p.A677S	⊢	Benign	I	Tier 1	Tumor suppresor	I	I	I	11.36
7	50468251	U	۹	Missense	IKZF1	p.G496S		Prob. dam.	rs548117828	Tier 1	Tumor suppresor, fusion	I	I	13.64	I
7	98579527	υ	⊢	Missense	TRRAP	p.R2899C		Prob. dam.	rs1792941314	Tier 1	Oncogene	1	10.58	I	I
7	127326707		۹	Missense	SND1	p.R40H	⊢	Benign	rs773076395/ COSM6786017	Tier 1	Oncogene, Fusion	Large intestine adenocarcinoma	I	11.76	I
ω	101730335	0	4	Missense	PABPC1	p.G123C	Ω	Prob. dam.	rs755674364	Tier 2	Oncogene, tumor suppressor	Astrocytoma, acute myeloid leukemia, colon adenocarcinoma, squamous cell carcinoma	I	I	11.67
11	46329469	υ	⊢	Missense	CREB3L1	p.S145L	⊢	Prob. dam.	rs746245139	Tier 1	Tumor suppresor, fusion	I	6.33	I	I
11	118629536	U v	⊢	Missense	DDX6	p.A314T	⊢	Prob. dam.	I	Tier 1	Oncogene, fusion	I	6.09	I	I
12	22797153	U	⊢	Missense	ETNK1	p.S231L		Benign	rs764046781	Tier 1	Tumor suppresor	I	I	10.91	I
12	25380271	υ	F	Missense	KRAS	p.E63K	۵	Prob. dam.	COSM27159	Tier 1	Oncogene	Malignant melanoma, colon adenocarcinoma, bladder carcinoma, carcinoid endocrine tumour, seminoma, acute lymphoblastic leukaemia	11.32	10.07	38.04
12	48373317	U	۷	Missense	COL2A1	p.V835I		Prob. dam.	rs121912882	Tier 1	Fusion	I	10.71	I	I
15	90628058	U	⊢	Missense	IDH2	p.G421S	Dlc	Prob. dam.	rs1023592505	Tier 1	Oncogene	Rectum adenocarcinoma, ovary mixed adenosquamous carcinoma	I	10.42	I
													(Con	(Continued on next page.)	iext page.)

Table	Table 1. (Continued)	tinuec	(r												
										Cosr in	Cosmic database information	Cancer type observation	Tera-		Ana- plastic
Chr.	Pos.		Ref. Alt.	Type of . variant	Gene symbol	HGVSp short	SIFT	SIFT PolyPhen	rsID/Cosmic Census ID gene	Census gene	Role in cancer	in COSMIC Database	toma (VAF %)	toma MNT (VAF %) (VAF %)	MNT (VAF %)
15	93524121	U	A	Missense	CHD2	p.G985R	۵	Prob. dam.		Tier 2	Tumor suppresor	I	I	I	11.49
16	3778563	υ	⊢	Missense	CREBBP	p.G2162E	DIc	Poss. dam. rs1401593486	rs1401593486	Tier 1	Oncogene, tumor suppressor, fusion	I	11.63	I	I
16	3808047	G	⊢	Missense and splice region	CREBBP	p.D1124E		Poss. dam.	I	Tier 1	Oncogene, tumor suppressor, fusion	I	I	I	17.95
17	5268516	U	۷	Missense	RABEP1	p.E590K		Benign	rs757825581	Tier 1	Fusion	Endometrioid carcinoma	8.93	I	I
17	7578183	CG	U	Frameshift	TP53	p.P222X	I	I	rs1567551073	Tier 1	Oncogene, tumor suppressor, fusion	I	44.83	32	42.86
17	30325720	U	A	Missense	SUZ12	p.V6401	⊢	Poss. dam.	I	Tier 1	Oncogene, Tumor suppressor, Fusion	I	17.65	I	I
×	48123289	U	A	Missense	SSX1	p.D135N	⊢	Poss. dam.	rs782680032	Tier 1	Oncogene, fusion	Astrocytoma, papillary renal cell carcinoma	I	15.79	I
Gene	svmbols an	e hase	no be	the HUGO Gene	Nomenclature	- Committee (	(DNDH)	. All variants	were validated by	v visualizatio	on with the Integrativ	Gene symbols are based on the HUGO Gene Nomenclature Committee (HGNC). All variants were validated by visualization with the Integrative Genomics Viewer (IGV).			

Gene symbols are based on the HUGO Gene Nomenclature Committee (HGNC). All variants were validated by visualization with the Integrative Genomics Viewer (IGV). (Chr.) Chromosome, (Pos,) genomic position according to genome version hg19 (GRCh37), (Ref.) reference base, (Alt.) alternative base, (MNT) melanocytic neuroectodermal tumor, (SIFT T) tolerated variant as predicted by the SIFT (Sorting Intolerant From Tolerant) program (https://sift.bii.a-star.edu.sg/), (SIFT D) deleterious variant as predicted by the SIFT program, (Folyphen Prob. dam.) probably damaging variant as predicted by the SIFT program, (Polyphen Prob. dam.) probably damaging variant as predicted by the PolyPhen program (http://genetics.bwh.harvard.edv.phb2/), (Polyphen Poss. dam.) possibly damaging variant as predicted by the PolyPhen program (active base), (SIFT D) deleterious variant as predicted by the SIFT program, (Folyphen Prob. dam.) probably damaging variant as predicted by the PolyPhen program (http://genetics.bwh.harvard.edu/phb2/), (Polyphen Poss. dam.) possibly damaging variant as predicted by the PolyPhen, (rsID) rs identification number, (VAF) variant allele frequency.





**Figure 3.** Somatic mutation profile of the mature teratoma, melanocytic neuroectodermal transformation (MNT), and metastatic anaplastic MNT. (A) Venn diagram analysis of the three different components showing six mutations shared by the three different component and 13 mutations common to at least two components. Variant allele frequencies are indicated in brackets. (*B*) Gene set enrichment analysis (GSEA) using Wikipathway reveals a significant normalized enrichment score in the melanoma pathway. For *A* and *B*, genes involved in oncogenesis according to the COSMIC database are indicated in bold. The oncogenic specific mutations appear in red.

that it originates from a germ cell precursor that erroneously stopped in midline migration during embryogenesis (Ronchi et al. 2019). Because of the high expression of KIT ligands that are implicated in germ cell proliferation, the thymus might be the preferential site for this germ cell precursor arrest (Oosterhuis et al. 2007; Ronchi et al. 2019). In some cases, this precursor may lead to a germ cell neoplasia which is a type II GCT precursor and then lead to the different subtypes including teratoma (Fichtner et al. 2021).

Teratoma seems to share genetic characteristics with other types of GCT. As the other type II GCT, adult teratoma are characterized by gain of Chromosome 12p or an isochromosome 12p (Lee et al. 2019; Fichtner et al. 2021). During the transition from precursor to the invasive component, acquisition of additional genetic alterations in genes involved in germ cell tumorigenesis and proto-oncogenes have been reported (Blanco and Tirado 2018; Fichtner et al. 2021). As the different lines of chemotherapy administered to the patient including cisplatin, etoposide, and paclitaxel have been described in the development of postexposure secondary neoplasms, mainly acute myeloid leukemia (Larson 2009; Smith et al. 2011), it is possible that the malignant transformation of teratoma is chemotherapy-induced.

Exome sequencing on the postchemotherapy mature thymic teratoma, the MNT, and its metastatic vertebral anaplastic MNT revealed six common mutations, among which three are located in known oncogenic factors or tumor suppressor genes. The first one is p.P222X



in the R175–R282 hotspot region of the DNA binding domain of TP53, which is mutated in close to 50% of all human solid cancers (Chiang et al. 2021; Marei et al. 2021). According to a recent genomic study on intracranial teratomas, TP53 is not among the most frequently mutated genes (Zhang et al. 2022). Nevertheless, this study showed that the TP53 signaling pathway is affected in 18% of cases and that TP53 mutations might be associated with poor prognosis. The role of TP53 alterations on the development of teratoma and malignant somatic transformation is, however, not well-understood (Zhang et al. 2022). It has also been shown that TP53 mutations play a crucial role in squamous cell carcinoma arising from ovarian mature cystic teratoma (Cooke et al. 2017; Tamura et al. 2020, 2023). Finally, in mediastinum, GCT with TP53 mutation have a worsening response to platinum-based chemotherapy (Timmerman et al. 2021). The second mutation is the p.E63K mutation of KRAS, which is part of the curated mutation list of the COSMIC database and has been reported in genital tract, pancreas, lymphoid tissue, large intestine, and thyroid neoplasms (Bamford et al. 2004). KRAS mutation seems to be the most common somatic alteration observed in GCT (Taylor-Weiner et al. 2016). Interestingly, the p.E63K mutation has also been described in three cases of extragonadal seminoma, a subtype of GCT, in a case of primitive neuroectodermal tumor-medulloblastoma and in a case of malignant lymphoma (Bamford et al. 2004; Fukushima et al. 2014; Wang et al. 2014; Ichimura et al. 2016). This mutation leads to an amino acid charge alteration at a critical position for the guanosine triphosphate hydrolysis reaction. In addition, NIH3T3 cells overexpressing KRAS p.E63K mutation showed altered morphology and displayed higher proliferative and migratory rates (Angeles et al. 2019). Interestingly, some cases of KRAS mutations have been reported as teratoma with adenocarcinoma transformation. However, this mutation is observed in both benign intestinal-type epithelium and adenocarcinoma component of the teratoma, suggesting that it is an early event in the carcinogenic sequence (Hershkovitz et al. 2013; Li et al. 2014; Kim et al. 2015). The third common mutation is the p.S447P mutation in the ATPase domain of the POLQ DNA double strand repair enzyme known to participate in cancer progression and resistance to therapy (Schrempf et al. 2021). Interestingly, two studies have shown that mutations in this domain may lead to its inactivation and thereby impact the sequence of the repaired products (Wyatt et al. 2016; Zahn and Jensen 2021). Although, the SIFT and PolyPhen prediction tools indicate a deleterious impact of the p.S447P mutation on protein function, additional data are needed to definitively confirm its potential involvement in the mutagenesis process.

GSEA showed an enrichment of the mutated genes in the melanoma pathway. The four genes (KRAS, TP53, ERBB4, and KDR) driving this enrichment are recognized for their implication as Cancer Census genes according to the COSMIC database (Bamford et al. 2004). This result is in accordance with the histologic findings showing that the anaplastic transformation of the MNT is histologically close to a malignant melanoma and expresses melanocytic protein markers SOX10, Melan-A, and HMB45. TP53 and KRAS are oncoproteins which have been described as potentially involved in melanoma (Cicenas et al. 2017; Hayward et al. 2017). In addition, mutations of ERBB4 and KDR have also been described in melanoma and these mutations are significantly more present in tumors with any known recurrent mutations in classical driver genes including BRAF, NRAS, KIT, GNAQ, and GNA11 (Xia et al. 2014). ERBB4 is a member of EGFR subfamily of receptor tyrosine kinase containing well-known EGFR and HER2 oncoproteins. In our study, we observed a p.A972V ERBB4 variant present specifically in the anaplastic MNT component. This mutation has not been previously described as a classical hotspot mutation in the protein but it is located in the intracytoplasmic tyrosine kinase domain, which is classically mutated in cancer development (Lucas et al. 2022). We also observed a somatic variant in the intracytoplasmic domain of KDR (p.T1258M), also known as VEGFR2, which is specific to the anaplastic MNT component. Altogether, these results suggest a potential involvement of these four variants in the development of the anaplastic MNT transformation of the teratoma.



## **METHODS**

#### **Patient and Samples**

The different samples were collected at the Oncology department of Strasbourg University Hospitals and analyzed at the Pathology department. The research protocol was approved by the institutional review board of Strasbourg University Hospitals, and a written informed consent was obtained from the patient. Samples were collected and processed under the principles of the Helsinki declaration. Tissue samples for histopathologic analysis were immediately fixed in 4% neutral buffered formalin for 24 h and processed according to standard procedures on slides stained with haematoxylin and eosin (H&E) method. All relevant pathology materials were carefully reviewed by expert pathologists. The tumor cellularity was 40% for the teratoma and 60% for the MNT and the anaplastic MNT.

#### Immunohistochemistry and In Situ Hybridization Studies

Immunohistochemical studies were performed on formalin-fixed paraffin-embedded (FFPE) sections following routine protocols on a Ventana Benchmark XT automated slide stainer (Ventana). An extensive antibody panel—which included the following antibodies—was used for diagnostic purposes: PS-100 (clone EP-32), SOX10 (clone EP268), Melan-A (clone A103), HMB45 (clone MO634), glypican 3 (clone GC33), bHCG (clone CGO4/CGO5), synaptophysin (Clone 27G12), panKeratin AE1/E3 (clone AE1 + AE3), and SALL4 (clone 6E3).

## **Targeted Panel Sequencing**

DNA was extracted from FFPE tissue sections using the QIAamp DNA FFPE Tissue Kit (QIAGEN) or the Maxwell 16 FFPE Plus LEV DNA Purification Kit (Promega), respectively, and according to the manufacturers' recommendations. Targeted sequencing was made only on anaplastic NMT FFPE sample within the routine clinical care process in order to identify potential therapeutic targets. Libraries were prepared with the Multiplicom Tumor Hotspot MASTR Plus kit (Agilent) covering hotspot mutations in 26 genes validated by the French Cancer Institute (INCa). The list of analyzed genes and exons is as follows: AKT1 (exon 3), ALK (exons 20–29), BRAF (exons 11–15), CDKN2A (complete coding sequence), CTNNB1 (exon 3), DDR2 (complete coding sequence), EGFR (exons 18-21), ERBB2 (exons 19-21), ERBB4 (exons 10 and 12), FGFR2 (exons 7, 12, and 14), FGFR3 (exons 7, 9, 14, and 16), H3F3A (exon 2), HIST1H3B (exon 1), HRAS (exons 2-4), IDH1 (exon 4), IDH2 (exon 4), KIT (exons 8-11, 13-14, and 17-18), KRAS (exons 2-4), MAP2K1/MEK1 (exons 2-3), MET (exons 2, 10, and 14–20), NRAS (exons 2–4), PDGFRA (exons 12, 14, and 18), PIK3CA (exons 2–3, 10-11, and 21), PIK3R1 (exons 11-13), PTEN (complete coding sequence), and STK11/LKB1 (complete coding sequence). Sequencing libraries were prepared with the SureSelectQXT Target Enrichment System kit (Agilent Technologies) following the manufacturer's instructions and sequenced on an Illumina MiSeq instrument (Illumina) (Okutman et al. 2021). Bioinformatics analysis was performed using STARK (version 0.9b), as previously described (Okutman et al. 2021). Only nonsilencing variants with an allelic frequency of >4% and a sequencing depth of >300 reads were reported.

## **Exome Sequencing**

Exome sequencing was performed on four FFPE samples including mature teratoma, MNT, metastatic anaplastic MNT component, and a matched germline thymic sample serving as a nontumoral control. The purification of genomic DNA was performed with the GeneRead DNA FFPE kit (QIAGEN) according to the manufacturer's instructions. The genomic DNA was quantified using a Qubit instrument and the dsDNA BR Assay kit (Thermo Fisher Scientific). Exome sequencing libraries were prepared with the Twist Human Core Exome

Kit (Twist Bioscience) following the manufacturer's recommendations. Paired-end (2-bp × 75-bp) sequencing was performed on a NextSeq500 sequencer (Illumina) with an average depth of coverage of 317.5, 405, 194.4, and 172 for the mature teratoma, MNT, metastatic MNT, and matched germline thymic samples, respectively. Sequences were mapped to the hg19 reference genome using BWA v0.7.17 (Li and Durbin 2010). Sequence variants were called using the GATK mutect2 v4.1.8.0 and vardict v1.8.2 tools and only those with the "PASS" filter were considered (Cibulskis et al. 2013; Lai et al. 2016). Variants were annotated using VEP (version 94) (McLaren et al. 2016) and filtered as follows: depth of coverage in tumor and corresponding normal samples >10, depth of coverage of alternative bases in tumor  $\geq$ 5 and in normal equal to 0, and VAF > 5%. Variants described in the 1000 Genomes Project and gnomAD database with an allele frequency of >1% were discarded. We focused only on protein-altering variants (missense, nonsense, splice site variants, start and stop gain/loss, and coding indels). Raw exome data (FASTQ files) are available at the National Center for Biotechnology Information's Sequence Read Archive (accession PRJNA825743).

For GSEA, we defined a score for each gene carrying a somatic coding or splice site variant. This score was defined as the weighted sum of conditions in which this variant was found. Weights were chosen as 1, 2, and 3 for mature teratoma, MNT, and anaplastic MNT, respectively, as a gradation of their involvement in anaplastic transformation. This weighted gene list was then analyzed using the online WEB-based Gene Set Analysis toolkit for Human Transcriptome (www.webgestalt.org) and the WikiPathway database. Functional enrichment analyses were based on false discovery rate (FDR) < 0.05 and normalized enrichment score (NES) (Benjamini and Hochberg 1995; Subramanian et al. 2005).

## **ADDITIONAL INFORMATION**

#### **Data Deposition and Access**

Raw exome data (FASTQ files) are available at the National Center for Biotechnology Information's Sequence Read Archive (accession: PRJNA825743). Variants of Table 1 are available in ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/) under the accession numbers VCV000501513.5, VCV000017379.26, VCV000634705.4, and SCV003915799– SCV003915823.

#### **Ethics Statement**

The research protocol was approved by the institutional review board of Strasbourg University Hospitals, and a written informed consent was obtained from the patient.

#### **Authors' Contributions**

S.Ma., B.L., and R.C. designed research, performed research, and wrote the manuscript; J.G., A.M., T.S., S.Me, A.K., and J-E.K. analyzed data and reviewed the manuscript; and S.B. designed research and reviewed the manuscript.

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Competing Interest Statement

The authors have declared no competing interest.

Referees

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