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A subset of epithelioid and spindle cell rhabdomyosarcomas is associated with *TFCP2* fusions and common *ALK* upregulation

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Abstract

Rhabdomyosarcomas with *TFCP2* fusions represent an emerging subtype of tumors, initially discovered by RNA-sequencing. We report herein the clinicopathological, transcriptional, and genomic features of a series of 14 cases. Cases were retrospectively and prospectively recruited and studied by immunohistochemistry (MYF4, MYOD1, S100, AE1/E3, ALK), fluorescence in situ hybridization with *TFCP2* break-apart probe ($n = 10/14$), array-comparative genomic hybridization (Agilent), whole RNA-sequencing (Truseq Exome, Illumina), or anchored multiplex PCR-based targeted next-generation sequencing (Archer® FusionPlex® Sarcoma kit). Patient's age ranged between 11 and 86 years, including 5 pediatric cases. Tumors were located in the bone ($n = 12/14$) and soft tissue ($n = 2/14$). Most bone tumors invaded surrounding soft tissue. Craniofacial bones were over-represented ($n = 8/12$). Median survival was 8 months and five patients are currently alive with a median follow-up of 20 months. Most tumors displayed a mixed spindle cell and epithelioid pattern with frequent vesicular nuclei. All tumors expressed keratins and showed a rhabdomyogenic phenotype (defined as expression of MYF4 and/or MYOD1). *ALK* was overexpressed in all but three cases without underlying *ALK* fusion on break-apart FISH ($n = 5$) nor next-generation sequencing ($n = 14$). *ALK* upregulation was frequently associated with an internal deletion at genomic level. *TFCP2* was fused in 5' either to *EWSR1* ($n = 6$) or *FUS* ($n = 8$). *EWSR1* was involved in both soft tissue cases. FISH with *TFCP2* break-apart probe was positive in all tested cases ($n = 8$), including one case with unbalanced signal. On array-CGH, all tested tumors displayed complex genetic profiles with genomic indexes ranging from 13 to 107.55 and recurrent *CDKN2A* deletions. *FET-TFCP2* rhabdomyosarcomas clustered together and distinctly from other rhabdomyosarcomas subgroups. Altogether, our data confirm and expand the spectrum of the new family of *FET-TFCP2* rhabdomyosarcomas, which are associated with a predilection for the craniofacial bones, an aggressive course, and recurrent pathological features. Their association with *ALK* overexpression might represent a therapeutic vulnerability.

Introduction

Rhabdomyosarcomas belong to a clinically and biologically heterogeneous family of tumors sharing re-expression of immature skeletal muscle differentiation factors, including MYF4/MYOGENIN and/or MYOD1 proteins. They are currently classified into four main subtypes: embryonal, alveolar, pleomorphic, and spindle cell/sclerosing [1]. Nonetheless, recent studies have deciphered the spindle cell/sclerosing family into molecularly distinct subtypes including *NCOA2*-rearranged infantile rhabdomyosarcomas [2], *VGLL2*-

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rearranged rhabdomyosarcomas [3], and spindle cell/sclerosing rhabdomyosarcomas with *MYOD1* exon 1 mutations [4, 5].

More recently, a new subtype of rhabdomyosarcomas has been identified through an RNA-sequencing screening, defined by *Transcription Factor Cellular Promoter 2 (TFCP2)* rearrangements [6]. These tumors were initially reported as a “new variant of epithelioid rhabdomyosarcomas” and have a striking predilection for bones, especially for the craniofacial skeleton, which is a very uncommon feature for rhabdomyosarcomas [6]. Their morphological spectrum has been subsequently extended with the description of cases with spindle cell morphology in the small case series reported so far, with a total of ten cases now described in the literature [7–10]. The emerging evidence highlight their predilection for bones, female patients, a frequent spindle cell morphology, and common ALK over-expression but substantial clinical follow-up is available for only two cases.

We report herein the clinicopathological and molecular features of a series of 14 *FET-TFCP2* rhabdomyosarcomas, highlighting their recurrent morphological and molecular features, paving the way to their diagnosis and recognition in routine practice. Our data confirm that they represent a distinct type of rhabdomyosarcoma.

Material and methods

Sample selection

Three index cases (*TFCP2* 1–3) were identified through a retrospective RNA-sequencing screening of unclassified sarcomas and were previously reported without detailed clinicopathological data [6]. Seven cases were identified prospectively: six cases were diagnosed on morphological grounds and one case (*TFCP2*-8) diagnosed on molecular grounds. Control cases were retrieved from the archives of the sarcoma pathology review network (RRePS and ResOs) and Institut Bergonie, including malignant peripheral nerve sheath tumors with rhabdomyosarcomatous differentiation (Triton tumor), and epithelioid/pleomorphic/embryonal and alveolar rhabdomyosarcomas. All sarcoma cases are recorded in the national sarcoma pathology RRePS and ResOs databases, approved by the National Committee for Protection of Personal Data (number 910390), in compliance with ethics principles of the chart of Helsinki. Cases 11–14 were retrieved from the files of AHGC and JVMGB, including two retrospective and two prospective cases.

Clinical review

Clinical follow-up was obtained from the medical records of patients provided by participating institutions or through corresponding clinicians. Survival was calculated from the

date of diagnosis to the date of death or last date of follow-up. The clinical onset of disease was defined by the first symptom of cancer. Follow-up duration was calculated from the date of the first clinical consultation. Clinical management data are provided in Table 2. Survival analysis was performed with the Kaplan–Meier method using the dates of initial diagnosis and the dates of death from any cause.

Histopathological analysis

All material (biopsy and surgical specimen if surgery was performed) was centrally reviewed by specialized soft tissue and bone pathologists (FLL, JMC, AHGC, JVMGB).

Immunohistochemistry

The tissue slides were deparaffinized in xylene, hydrated in alcohol, and baked in a microwave (30 min in Tris buffer pH 9). Endogenous peroxidase was blocked. Staining was performed on the Benchmark ultra-automated stainer (Ventana) using diamino-benzidine as chromogen (Dako, Glostrup, Denmark). The following antibodies were used: AE1/E3 (clone PCK26, Ventana), KL1 (clone KL1, HISTOLS reagent), EMA (clone E29, Ventana), P63 (4A4, Ventana), S100 protein (clone poly Z311, Dako), SOX10 (EP268, BioSB), Desmin (clone DE-R11, Ventana), Myogenin (clone LO26, Leica), MYOD1 (clone EP212, Cell Marque), ALK (clones D5F3, Cell Signaling Technology and clone 1A4, Novocastra), CD99/MIC2 (12E7, Dako), ETV4 (clone 16, Santa Cruz), CD30 (BerH2, Dako), CD4 (4B12, Novocastra), and CD7 (SP94, Ventana).

Fluorescence in situ hybridization (FISH)

Fluorescence in situ hybridization (FISH) analyses were performed with 4 μ m sections of formalin-fixed paraffin-embedded (FFPE) tissue with the use of Histology FISH Accessory Kit (DAKO, reference: K5799) using commercial break-apart probes targeting *EWSR1* (ZytoLight SPEC *EWSR1* Dual Color Break Apart Probe, Zytovision, reference Z-2096–200), *FUS* (ZytoLight SPEC *FUS* Dual Color Break Apart Probe, Zytovision, reference Z-2130-50), *ALK* (ZytoLight SPEC *ALK* Dual Color Break Apart Probe, Zytovision, reference Z-2124-200), *TFCP2* (*TFCP2* Break Apart probe, Empire Genomics), and reference EG-*TFCP2*BA-20-GROR. Nuclei were scored for non-rearranged patterns (red and green fusion signals), rearranged, and unbalanced patterns (split of red and green signals or extra single red signals) using a Nikon Eclipse 80i fluorescent microscope with appropriate filters. The positive cutoff was $\geq 20\%$. Pictures were captured using a Hamamatsu C4742-95 CCD camera and analyzed with the Genikon software (Alphelys).

Array-comparative genomic hybridization

Genomic DNA was extracted from formalin-fixed paraffin-embedded tissue using QIAamp, DNA micro kit (Qiagen, Hilden, Germany). Genomic DNA and human reference DNA (Promega) were labeled with cyanin 5 and cyanin 3, respectively, using the Genomic DNA *ULS* Labeling Kit (Agilent Technologies, Santa Clara, California) and co-hybridized onto a Sureprint G3 Human CGH microarray 4 × 180 K (Agilent) following the manufacturer's recommendations. Data were analyzed with Agilent Genomic Workbench software (v7.0, Agilent) or by Cytogenomics software (v2.9.2.4, Agilent). Large-size alterations were defined as quantitative anomalies of whole chromosomes or of segments of chromosomes involving many probes of the array, easily detectable without a specific query on a given genomic region. Small-size alterations were quantitative anomalies involving only few probes, not detected by the automatic algorithm but after manual analysis of the genomic regions.

Sanger sequencing of *MYOD1*

Fifty nanograms of FFPE DNA was amplified with forward (5'-AAGCGCAAGACCACCAAC-3') and reverse primers (5'-GTTGCGCAGGATCTCCAC-3') targeting exon 1 of *MYOD1*. The PCR products were analyzed by 2% agarose gel electrophoresis with an expected product size of 153 bp. The PCR products were sequenced using a Big Dye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA) on a 3130XL Genetic Analyzer (Applied Biosystems).

Whole RNA-sequencing

RNA-sequencing was performed on formalin-fixed paraffin-embedded material in tencases (notably, cases TFCP2 1–3, the three previously reported cases studied by RNA-sequencing on frozen material [6], were re-sequenced with FFPE samples to ensure data comparability).

Total RNA was extracted from formalin-fixed paraffin-embedded tissue section using the Formapure RNA kit (Beckman Coulter, Brea, California) following the manufacturer's recommendations. Quantity and quality of total RNA were evaluated using NanoDrop (Thermo Fisher Scientific) and Tape Station with Hs RNA Screen Tape (Agilent) using a cutoff of DV₂₀₀ (defined as the percentage of RNA fragments above 200 nucleotides) above 30%. All samples passed quality criteria. Libraries were prepared with 100 ng of total RNA using TruSeq RNA Access Library Prep Kit (Illumina, San Diego, USA). Libraries were pooled by a group of 12 samples. Paired-end sequencing was performed using the NextSeq

500/550 High Output V2 kit (150 cycles) on Illumina NextSeq 500 platform (Illumina, San Diego, CA). To perform the clustering analysis, gene expression values were extracted using Kallisto v0.42.5 tool [11] with GENCODE release 23 genome annotation based on GRCh38 genome reference. Kallisto Transcripts Per Kilobase Million (TPM) expression values were transformed in $\log_2(\text{TPM} + 2)$ and all samples were normalized together using the quantile method from the R limma package within R (version 3.1.1) environment. Clustering was performed with the R package Cluster v2.0.3 ConsensusClusterPlus v 1.46 [12] using 1000 permutations of 80% of both samples and genes. Agnes was used as the clustering algorithm with Pearson's correlation distance and Ward's clustering method.

Anchored multiplex PCR

Cases 11–14 were analyzed only by targeted RNA-sequencing. For the detection of translocations, total nucleic acid was isolated from five FFPE slides of 10 μm using the Tissue Preparation System (Siemens) after microdissection. After measurement of RNA quantity for FFPE material with Qubit fluorometric quantification system (Life Technologies), the target-enriched cDNA library was prepared with the Archer® FusionPlex® Sarcoma kit as per the manufacturer's description. As described previously [13], reverse transcription of RNA was followed by end repair, adenylation, and universal half-functional adapter ligation of double-stranded cDNA fragments. This was followed by two rounds of low-cycle PCR with universal primers and gene-specific primers, covering 26 target genes that rendered the library fully functional for clonal amplification and sequencing using the Ion Proton™ or the Ion S5™ system. With the Archer analysis software (version 5.0) the produced libraries were analyzed for the presence of relevant fusions. Sequence quality was assessed by the following criteria: Quality control (QC) score of <30, a minimal total read number of 1.5 million with >7% unique fragments and >40% RNA reads.

Results

Clinical, radiological, and macroscopic findings

The age at diagnosis ranged from 11 to 86 years with a mean of 31 years (Table 1). There were five pediatric patients (defined as <18 years of age) and a slight female predominance was seen (eight female vs. six male patients).

Tumor size ranged from 20 to 150 mm with a median size of 60 mm. Tumors were most often located in the bones ($n = 12/14$) or soft tissue ($n = 2/14$). Bone tumors presented

Table 1 *FET-TFCP2* rhabdomyosarcomas cases

Case ID	Age (y)/gender	Location	Survival	Cytomorphology	FISH <i>TFCP2</i>	Fusion transcript ^a	ALK expression
TFCP2-1	16/F	Sphenoid bone (left greater wing)	DOD (15 mo)	Spindled and epithelioid	Break	FUS(ex6)-TFCP2(ex2)	50%
TFCP2-2	26/F	Sacrum	DOD (4 mo)	Epithelioid	NI	FUS(ex6)-TFCP2(ex2)	50%
TFCP2-3	38/F	Peritoneum	DOD (2 mo)	Spindled and epithelioid	Break	EWSR1(ex5)-TFCP2(ex2)	50%
TFCP2-4	32/M	Hard palate and upper lip	DOD (8 mo)	Spindled and epithelioid	Break	EWSR1(ex5)-TFCP2(ex2)	-
TFCP2-5	20/M	Orbito-temporo-sphenoid mass	DOD (6 mo)	Spindled and epithelioid	Break	FUS(ex6)-TFCP2(ex2)	<5%
TFCP2-6	86/M	Inguinal	DOD (6mo)	Spindled and epithelioid	Unbalanced break	EWSR1(ex5)-TFCP2(ex2)	100%
TFCP2-7	18/F	Right femur (bifocal: neck and inter-trochanteric area)	DOD (8mo)	Spindled and epithelioid	Break	EWSR1(ex5)-TFCP2(ex2)	100%
TFCP2-8	17/F	Cervico-occipital junction	AWD (15mo)	Rounded	Break	FUS(ex6)-TFCP2(ex2)	100%
TFCP2-9	31/M	Left occipital bone	DOD (6mo)	Epithelioid and spindled	Break	FUS(ex7)-TFCP2(ex2)	100%
TFCP2-10	32/M	Mandible	AWD (14 mo)	Spindled	Break	FUS(ex6)-TFCP2(ex2)	70%
TFCP2-11	58/F	Mandible	ANED (21 mo)	Spindled and epithelioid	NA	FUS(ex5)-TFCP2(ex2)	70%
TFCP2-12	12/F	Mandible	ANED (21mo)	Spindled and epithelioid	NA	FUS(ex6)-TFCP2(ex2)	50%
TFCP2-13	11/F	Maxilla	DOD (unknown)	Epithelioid	NA	EWSR1(ex6)-TFCP2(ex2)	<5%
TFCP2-14	25/M	Mandible	ANED (20 mo)	Epithelioid	NA	EWSR1(ex5)-TFCP2(ex2)	80%

^aANED alive no evidence of disease, AWD alive with disease, DOD died of disease, FISH fluorescence in situ hybridization, NA not available, NI non-interprettable, y years
^bExons number are respective to ENST00000332050 (EWSR1), ENST00000254108 (FUS), and ENST00000257915 (TFCP2)

at locally advanced stage (Figs. 1, 3, 4 and Supplementary Fig. S1), invading surrounding tissues in all but one case ($n = 11/12$). Two cases involved two contiguous bones ($n = 2/12$). In one case, the tumor was bifocal within femoral bone (case TFCP2-7). Involved bones included the cranial bones ($n = 3$), bones of the orofacial complex ($n = 6$, mandible being the most commonly affected bone), vertebra, and long bone and sacrum (one case each). When performed, gross examination confirmed, in case of bone origin, destruction of cortical bone and extension in surrounding soft tissue (Supplementary Fig. S1). Regarding soft tissue tumors, locations included the peritoneum and inguinal area (the origin and precise level of extension of this case were not available, TFCP2-6).

Overall, all patients received chemotherapy with varied regimens, including two patients with neoadjuvant chemotherapy and four patients with chemotherapy as the sole therapeutic modality (Table 2). No consistent therapeutic response was achieved with conventional chemotherapy. Other treatment modalities included surgery ($n = 7$). Local control was achieved with surgery in six patients, including two R1 micro surgery (i.e., defined as the presence of microscopic tumor residue) and four R0 surgery but local relapse occurred in all patients (Table 2). Seven patients had metastatic disease, either at presentation ($n = 3/14$) or at follow-up ($n = 4/14$). Metastases involved lungs ($n = 6/7$), bones ($n = 2/7$), and lymph nodes ($n = 1/7$).

The mean survival was 8 months. The cohort included eight patients dead of disease and five patients alive, of whom two had evidence of clinical progression (Tables 1 and 2). The probability of survival was estimated to 46% at 12 months with Kaplan–Meier analysis (Supplementary Fig. S2). Among alive patients in this series ($n = 5$), all but one had undergone radical surgery and the mean follow-up was 6 months, with two patients currently in clinical progression. The sole survivor without surgery had unresectable tumor, which was progressive despite combined VDC-IE

(vincristine–doxorubicin–cyclophosphamide–ifosfamide–etoposide) chemotherapy and radiation therapy, and was subsequently switched to targeted therapy (Alectinib), which halted progression (slight tumor regression from 53×27 to 32×29 mm) at 7 months of therapy (Table 2, TFCP2-8).

Microscopic findings

Tumors had been initially heterogeneously classified with diagnosis of spindle cell rhabdomyosarcoma ($n = 3$), epithelioid rhabdomyosarcoma ($n = 2$), embryonal and unclassified rhabdomyosarcomas not otherwise specified (NOS) ($n = 2$ and 2, respectively), malignant Triton tumor ($n = 2$), desmoplastic small round cell tumor ($n = 1$),

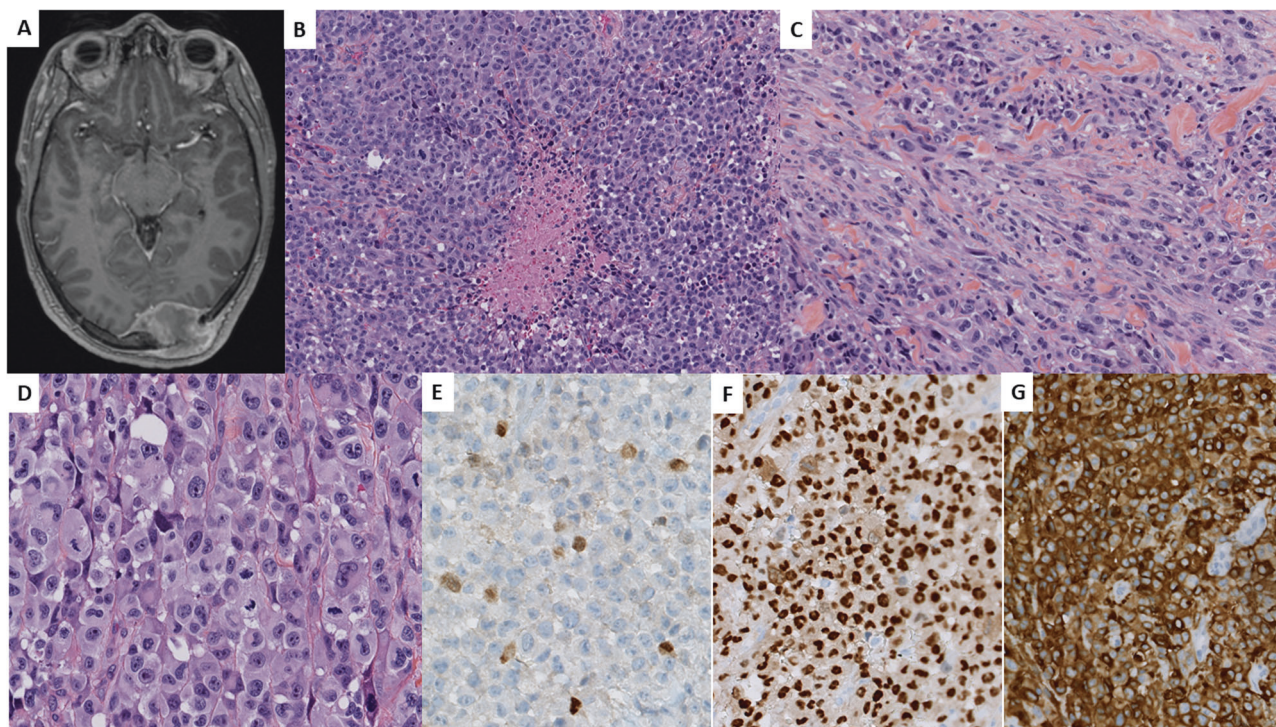


Fig. 1 Typical case of *FET-TFCP2* rhabdomyosarcoma. MR image demonstrating an expansive and destructive mass of the occipital bone in a 31yo male (a, *TFCP2-9*). The proliferation was arranged in solid sheets (b) and fascicles (c). Tumor cells displayed middle-sized nuclei with distinct nucleoli and mild anisocaryosis along with abundant

eosinophilic cytoplasm (d). Cells stained focally for Myogenin (e), diffusely for MYOD1 (f), and diffusely for ALK (g). Cytokeratins AE1/3 and Desmin were heterogeneously and diffusely expressed, respectively (not shown)

unclassified sarcoma with a spindle cell phenotype ($n = 1$), and anaplastic lymphoma ($n = 1$) (Table 3).

Most tumors ($n = 9/14$) showed a hybrid spindle cell and epithelioid cytology (Fig. 1b–d), whereas the other cases had essentially epithelioid ($n = 3$) (Fig. 2a, b), round cell ($n = 1$) (Fig. 2c, d), or spindle cell morphology (Fig. 3b, c).

The lesional cells had abundant eosinophilic cytoplasm and oval nuclei dotted with distinct nucleoli in most cases ($n = 9/14$) (Fig. 4e). Nuclear hyperchromasia and pleomorphism (defined as variations in size and shape of the nuclei) were focally seen in four cases (Fig. 2f). Scattered rhabdomyoblasts were seen in two cases ($n = 2/14$) (Supplementary Fig. S1f). There was no significant stroma in half of the cases but focal fibrous changes were seen in six cases (Fig. 2e). A single case displayed a diffuse hyalinized stroma that encased the tumor cells, accounting for a trabecular growth pattern (Fig. 2c, d). Tumor necrosis was present in all but two tumors (Table 3).

Immunohistochemical findings

All but one case expressed desmin either diffusely ($n = 8/14$), heterogeneously ($n = 3/14$), or only focally ($n = 2/14$). All tumors expressed either MYOD1 or myogenin.

MYOD1 was strongly expressed (i.e., >50%) in all but three cases ($n = 11/14$), whereas myogenin staining was only focally (<35%) positive in most cases ($n = 9/14$), negative in three cases ($n = 3/14$), and diffusely positive in two cases (Table 3 and Fig. 1). Notably, MYOD1 was diffusely positive in all tumors negative for myogenin. Keratins AE1/E3 were positive in all cases ($n = 14$), with focal staining in four cases (Figs. 1 and 4). ALK was overexpressed in all but three cases, with diffuse cytoplasmic overexpression (>50%) in all positive cases (Figs. 1, 4 and Supplementary Fig. S1). In one case, CD30 and CD4 were focally positive, accounting for its initial misdiagnosis as anaplastic lymphoma but no significant staining was seen in other tested cases ($n = 8$). S100 protein was focally positive in six cases and diffusely positive in one case, without concomitant expression of SOX10 in tested cases ($n = 0/10$).

Fluorescence in situ hybridization

EWSR1 break-apart FISH was positive in all but one *EWSR1-TFCP2* tumors ($n = 2/3$), including one case with 3' loss, consistent with unbalanced translocation (Supplementary Table S1). Break-apart FISH for *FUS* was positive in all tested cases ($n = 5/5$).

Table 2 Clinical features of *FET-TFCP2* rhabdomyosarcomas

Case #	Clinical symptoms	Location	Tumor size (mm)	Locoregional extension	Metis	First-line therapy	Evolution	Second-line therapy	Evolution	Third-line therapy	Evolution	Survival
TFCP2-1	Headache and left exophthalmos	Sphenoid bone (left greater wing)	90	Dura, left orbit, and infratemporal fossa	NxM0	4 Cycles of etoposide, ifosfamide neoadjuvant (100%)	Progression	Resection of left orbit, sphenoid, ethmoid, and temporal bones with patch of dura (R1 micro)	Local relapse at 2 mo post surgery, metastasis of right femoral bone	1 Cycle (cyclophosphamide, doxorubicin) and cerebral radiotherapy	Respiratory failure due to pleural effusion	DOD (15 mo)
TFCP2-2	Right hip pain	Sacrum	100	Right sacro-iliac joint and medullary involvement	NxM1 (lung, mediastinal lymphadenopathies)	2 Cycles of IVADO (100%)	Locoregional progression and pleural effusion	1 Cycle of IVADO (100%)	Local progression	1 Cycle VAI (100%)	-	DOD (4 mo)
TFCP2-3	Peritoneal effusion	Peritoneum	Multiple nodules	Pleural effusion (no mass on CT nor PET-CT)	NxM0	1 Cycle of carboplatin, paclitaxel (100%)	Local progression and fistule post-celoscopy	1 Cycle of adriamycin, holoxan (100%)	Lymphangitic carcinomatosis	-	-	DOD (2 mo)
TFCP2-4	Nodule of gingiva and hard palate of 3 cm	Mass of hard palate and upper lip	Multiple confluent nodules	Hard palate, gingiva	NxM1 (vertebra, ribs, pelvis)	3 Cycles of VAI (100%)	Local progression	1 Cycle of etoposide-carboplatin (100%)	Local progression	1 Cycle of actinomycin-cyclophosphamide	Local progression and bacterial infection	DOD (8 mo)
TFCP2-5	Soft tissue mass and left exophthalmos	Orbito-temporo-sphenoid mass	150 × 145	Left orbit, sphenoid and temporal bones, dura, left infratemporal fossa, soft tissue, and dermis of left temporal region	NxM0	1 Cycle doxorubicin–ifosfamide (100%)	Local progression	3 Cycles of VAC (100%)	Local progression	Radiation therapy and pazopanib	Local progression	DOD (6 mo)
TFCP2-6	Inguinal mass	Inguinal	65 × 40	NA	NxMx	Resection (hemia surgery, R1)	Local relapse at 4 mo	Palliative care	Local progression	-	-	DOD (6 mo)
TFCP2-7	Right thigh pain	Right femur (bifocal: neck and inter-trochanteric area)	51	-	NxM1 (lungs and bones, L5 vertebra, left ischio-pubic branch, ribs)	2 Cycles of VDC-IE (100%)	Local progression (left iliac bone, sacrum) with soft tissue extension in pelvis and right hip	Pazopanib	Local progression	-	-	DOD (8 mo)
TFCP2-8	Cervical pain	Cervico-occipital junction	53 × 37 × 44	Lysis of left occipital condyle and vertebra C1, and left clivus; extension into left jugular, hypoglossar foramen, and paravertebral soft tissue	NxM0	2 Cycles of VDC-IE (100%) and adjuvant RT (60 Gray)	Local progression with extension to the odontoid process and into the surrounding soft tissue measuring 20 × 30 × 34 mm	Crizotinib 500 mg/d for 1 month switched to alectinib 1200 mg/d	Stable disease for 9 mo since initiation of anti-ALK therapy (32 × 29 × 42 mm)	-	-	AWD (15 mo)
TFCP2-9	Headache, clinical hypotthesis of meningioma	Left occipital bone	118	Extracranial component (65 × 17 mm); soft tissue of scalp; intracranial component (53 × 16 mm): meninges, left superior sagittal sinus	NxM0	Fragmented resection (R1 micro)	Local relapse at 1 mo post surgery	Adjuvant CT; 3 cycles of AI	Local progression and metastasis (lung, mediastinum)	-	Progression with pleural effusion and infiltration of the superior sagittal sinus	DOD (6 mo)

Table 2 (continued)

Case #	Clinical symptoms	Location	Tumor size (mm)	Locoregional extension	Mets	First-line therapy	Evolution	Second-line therapy	Evolution	Third-line therapy	Evolution	Survival
TFCP2-10	toothache	Mandible (symphysis and from teeth 37 to 44)	45 × 32	Vestibular gingiva, mylohyoid, and hyoglossus muscles	NOM0	Partial mandibulectomy (RO)	-	Adjuvant CT: 5 cycles: doxorubicin, ifosfamide (100%)	-	Adjuvant CT: 3 cycles of gemcitabine-doxetaxel (100%)	-	AWD (14 mo)
TFCP2-11	-	Mandible	16 × 15 × 13	Surrounding soft tissue (muscle)	NOM0	Surgery (RO)	-	Adjuvant CT and RT	-	No recurrence or metastatic disease	-	ANED (21 mo)
TFCP2-12	Local painful swelling for 4 months	Mandible	55	Surrounding soft tissue (muscle)	NOM0	Neoadjuvant CT	-	Surgery (RO), RT, adjuvant CT	-	No recurrence or metastatic disease	-	ANED (21 mo)
TFCP2-13	-	Maxilla	60	Surrounding soft tissue (muscle)	NOM0	CT	-	Radiotherapy	-	Progression	-	DOD (unknown)
TFCP2-14	Local painful swelling for 1 month	Mandible	34	Surrounding soft tissue (muscle)	NOM0	Surgery (RO)	-	CT	-	No recurrence or metastatic disease	-	ANED (20 mo)

At doxorubicin–ifosfamide, ANED alive no evidence of disease, AWD alive with disease, BEP bleomycin–etoposide–platin; CT chemotherapy, IVADO ifosfamide–vincristine–actinomycin–D–doxorubicin, IVE ifosfamide–vincristine–etoposide, Mets metastasis, VAC vincristine–actinomycin D–cyclophosphamide, VAI vincristin–actinomycin–ifosfamide, VDC-IE vincristine–doxorubicin–cyclophosphamide–ifosfamide–etoposide

No rearrangement of *ALK* was seen nor by a FISH break part assay for *ALK* ($n = 0/5$) nor by RNA-sequencing ($n = 0/14$) (Supplementary Table S1).

FISH with *TFCP2* break-apart probe was positive in all tested case ($n = 9$) (Figs. 3 and 4), including one case with an unbalanced split signal. To test the specificity of *TFCP2* rearrangements in the setting of rhabdomyosarcomas, we tested 31 cases of rhabdomyosarcomas spanning the different subtypes by break-apart FISH with a break-apart probe for *TFCP2* (Supplementary Table S2). All control cases were negative for rearrangement, including four sporadic malignant triton tumors, one neurofibromatosis type 1-associated malignant triton tumor, ten pleomorphic rhabdomyosarcomas, one case each of embryonal rhabdomyosarcoma and alveolar rhabdomyosarcoma, threespindle cell rhabdomyosarcomas, nine cases of epithelioid rhabdomyosarcomas, one case of carcinoma with rhabdomyosarcomatous component, and two cases of unclassified sarcomas with epithelioid phenotype (Supplementary Table S2).

Array-comparative genomic hybridization and *MYOD1* mutational analysis

Mutations in the exon 1 of *MYOD1* were absent at DNA level ($n = 0/8$) as well as at RNA level ($n = 0/10$) (Supplementary Table S1).

Genomic profiling was performed in ten cases, showing overall rearranged profiles with high genomic index ranging from 13.5 to 107.55. Two *EWSR1-TFCP2* tumors ($n = 2/4$) and one *FUS-TFCP2* tumor ($n = 1/5$) displayed genomic breakpoints within *EWSR1* and *FUS* loci, respectively. Two tumors had a genomic breakpoint within *TFCP2* locus. All cases displayed homozygous deletion of the tumor suppressor gene *CDKN2A* ($n = 9/9$) (Supplementary Table S3 and Fig. 4). Two cases displayed additional homozygous deletions spanning in the first case: *PRKAR2A* in 3p21.31, *AUTS2* in 7q11.22, *PTPN12* in 7q11.23-q21.11, *PDE3A* in 12p12.2, *ST8SIA1* in 12p21.1, *LARGE1* in 22q12.3, and in the second case *MAP3K15*, *GPR63*, and *PHKA2* in Xp22.13-p22.12 and the locus 16p11.2-q12.1. Genomic amplifications were seen in one case spanning most notably *MDM2* in 12q15, *CCNE1* in 19q12-q13.11, and *MAP3K1* in 5q11.2-12.1. Focusing on *ALK* locus, five cases displayed alterations ($n = 5/9$), including an internal hemizygous deletion in three cases, a hemizygous deletion, and a homozygous internal deletion in one case each.

RNA-sequencing findings

Whole RNA-sequencing was performed on FFPE material in ten cases and four cases were studied by Anchored multiplex PCR (Supplementary Table S1). Four cases of

Table 3 Pathological features of *FET-TFCP2* rhabdomyosarcomas

Case #	Material reviewed	Microscopy				Immunohistochemistry														
		Initial diagnosis	Patterns	Stroma	Cytology	Mitotic index	Necrosis	Post-therapy changes	AE1/ E3/ KL1	EMA	p63/ CK5/ 6	ALK 1A4/ ALK D5F3	H- caldesmon	Desmin	Myogenin/ MYOD1	S100P/ SOX10	CD99/ ETV4	CD30	CD4/ CD7	
TFCP2- 1	Biopsy and resection specimen	Malignant Triton tumor	Spindled and solid	Focal fibrous	Fine chromatin, focal hyperchromatism	12	15%	Necrosis 50%, fibrosis 10%, focal pleomorphism	+	-	-	50%/50% (cytop.)	-	+	Diffuse	10%/75%	-/-	NA	NA	NA
TFCP2- 2	Biopsy	Embryonal rhabdomyosarcoma	Solid	Absent	Epithelioid, vesicular nuclei	18	60%	NA	+	(65%)	-	50%/- (cytop.)	-	70%	Diffuse	10%/65%	-	NA	-	-
TFCP2- 3	Surgical specimen	Desmoplastic small round cell tumor	Spindled and solid	Focal fibrous	Epithelioid vesicular nuclei	12	15%	NA	+	Focal	-	50%/10% (cytop.)	-	70%	Diffuse	35%/65%	-	-	-	-
TFCP2- 4	Biopsy	Unclassified rhabdomyosarcoma, NOS	Spindled and solid	Absent	Epithelioid, vesicular nuclei, focal hyperchromatism	21	10%	NA	+	-	-	-	-	+	Diffuse	-75%	+	Focal/-	-	-
TFCP2- 5	Surgical biopsy	Epithelioid rhabdomyosarcoma	Spindled and solid	Absent	Epithelioid, vesicular nuclei, focal hyperchromatism	32	30%	NA	+	-	-	-<5%	-	+	Diffuse	15%/65%	NA	-	-	-
TFCP2- 6	Resection specimen	Spindle cell sarcoma, grade 3	Spindled and solid	Focal fibrous and myxoid	Epithelioid, vesicular nuclei, focal hyperchromatism	10	30%	NA	+	(20%)-	-	+5% (cytop.)	-	+	Diffuse	15%/35%	+	Focal/-	NA	NA
TFCP2- 7	Biopsy	Epithelioid rhabdomyosarcoma	Spindled and solid	Absent	Epithelioid, clear chromatin	8	10%	NA	+	-	NA	+	(cytop.)	+	Diffuse	+/+focal	-	-	-	-
TFCP2- 8	Biopsy	Anaplastic lymphoma	Solid	Focal fibrous and myxoid	Rounded, epithelioid and plasmacytoid, clear chromatin	16	10%	NA	NA	Focal	NA	+	(cytop.)	NA	+	20%	-75%	-/NA	+ Focal	10% + Focal
TFCP2- 9	Biopsy	Embryonal rhabdomyosarcoma	Solid and spindled	Fibrous septa	Epithelioid, vesicular nuclei	66	15%	NA	+	(25%)	-	NA	+	(cytop.)	+	Diffuse	15%/ + diffuse	NA	+	Focal
TFCP2- 10	Biopsy	Spindle cell rhabdomyosarcoma	Spindled	Fibrous	Focal pleomorphism, vesicular nuclei	21	Absent	NA	+	(20%)	-	-	+	(cytop.) 65%	+	20%	15%/ + diffuse	NA	-	-
TFCP2- 11	Resection specimen	Unclassified rhabdomyosarcoma, NOS	Spindled and solid	Absent	Vesicular nuclei, scattered rhabdomyoblasts	11	Absent	NA	+	-	NA	NA/70% (cytop.)	-	+	Diffuse	15%/70%	+	Focal/-	NA	NA
TFCP2- 12	Biopsy and resection specimen	Malignant Triton tumor	Spindled and solid	Absent	Vesicular nuclei, scattered rhabdomyoblasts	4	30%	NA	+	Focal	NA	NA/50% (cytop.)	-	+	Diffuse	50%/40%	+/N-	NA	NA	NA
TFCP2- 13	Biopsy	Spindle cell rhabdomyosarcoma	Solid	Absent	Fine chromatin	26	30%	NA	+	Weak	+	Focal	NA	+	Diffuse	30%/80%	+	Focal/-	NA	NA
TFCP2- 14	Biopsy and resection specimen	Spindle cell rhabdomyosarcoma	Solid	Fibrous	Fine chromatin	3	30%	NA	+	-	NA	NA/80% (cytop.)	-	-	Diffuse	-80%	+	Focal/-	NA	NA

NA not available

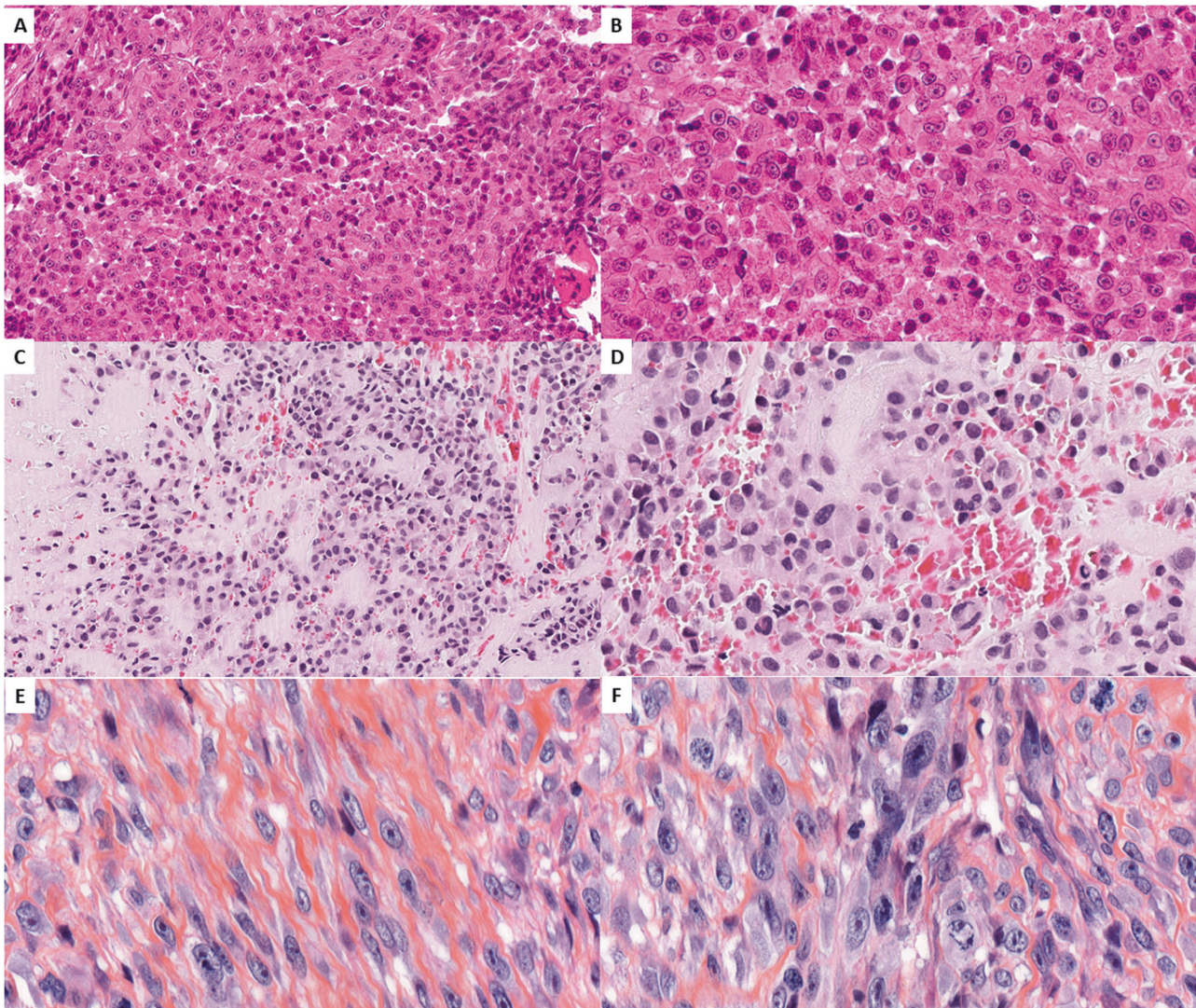


Fig. 2 Morphological variants seen in *FET-TCF2* rhabdomyosarcomas. Three *FET-TCF2* cases displayed pure epithelioid cytomorphology where tumor cells are arranged in solid sheets (**a**, case TFCP2-2) and harbored abundant globoid eosinophilic cytoplasm (**b**). One case of *FET-TCF2* rhabdomyosarcoma (TFCP2-8) displayed more abundant fibrous and myxoid stromal changes than seen in other cases (**c**) and tumor cells displayed small to medium rounded nuclei

with fine chromatin without nucleoli (**d**). This tumor was associated with a *FUS-TCF2* fusion and displayed otherwise similar immunophenotypic and RNA expression features to other *FET-TCF2* tumors. *FET-TCF2* rhabdomyosarcomas may focally display fibrous stromal changes (**e**). Tumor nuclei were mostly monomorphic while a subset of cases harbored marked anisokaryosis with variations in size and shape of the nuclei (**f**)

this series, including the three index cases of *FET-TCF2* rhabdomyosarcomas, were diagnosed based on RNA-sequencing findings without prior pathological suspicion of the diagnosis. *TCF2* was either fused to *FUS* ($n = 8$) or to *EWSR1* ($n = 6$). Breakpoints involved constantly exon 2 of *TCF2*, exons 5 or 6 of *FUS* and *EWSR1*. The transcriptomes of the seven cases sequenced on the same whole RNA-sequencing platform were compared with embryonal rhabdomyosarcomas ($n = 6$), alveolar rhabdomyosarcoma ($n = 8$), *MYOD1*-mutated rhabdomyosarcomas ($n = 4$), and *VGLL2*-fused rhabdomyosarcomas ($n = 5$). Unsupervised consensus clustering analyses highlighted that these seven *FET-TCF2* samples clustered apart from the other

rhabdomyosarcomas (Supplementary Fig. S3). Expression analysis evidenced *ALK* RNA upregulation in all but two samples available for the analysis ($n = 10$). These two cases were notably negative for *ALK* in immunohistochemistry. *ALK* transcripts upregulation in *FET-TCF2* rhabdomyosarcomas was at least at the level of *ALK*-rearranged inflammatory myofibroblastic tumors or *ALK*-rearranged Spitz nevi (Supplementary Fig. S3). No fusion involving *ALK* was identified by targeted or whole transcriptome sequencing. At the transcriptional level, *ALK* upregulation was stronger in the subset of tumors associated with genomic deletion of *ALK* and was uneven along the transcript. The upregulation predominated on the sequences

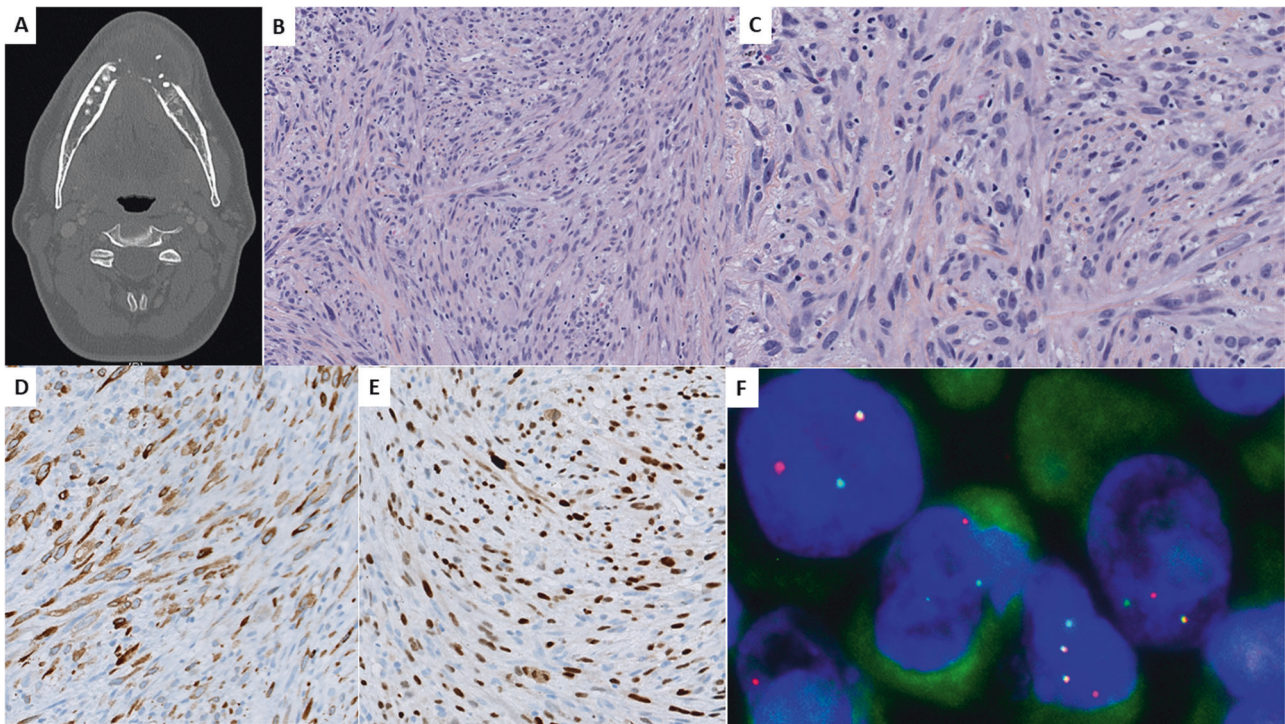


Fig. 3 Case of *FET-TFCP2* rhabdomyosarcoma with pure spindle cell morphology. CT of an osteolytic lesion of the mandible with cortical destruction and soft-tissue extension (**a**, TFCP2-10). Tumor cells were arranged in intersecting fascicles with abundant eosinophilic

cytoplasm (**b**, **c**). Tumor cells expressed heterogeneously desmin (**d**) and diffusely MYOD1 (**e**). Tumors cells displayed a split signal by FISH with the *TFCP2* break-apart probe (**f**)

beyond exon 17, a finding which has been found in case of alternative transcription initiation [14] (Supplementary Fig. S4).

Fusions were characterized by anchored multiplex PCR in four cases, showing two cases each of *FUS-TFCP2* and *EWSR1-TFCP2* fusions, involving the same exons than those seen in the cohort explored by RNA-sequencing (Table 1).

Discussion

Rhabdomyosarcomas represent a clinically and biologically heterogeneous group of tumors that share by definition the re-expression of the immature myogenic markers Myogenin and MYOD1 [15]. However, rhabdomyosarcomatous differentiation can be seen in many tumor types, both epithelial with the example of carcinosarcomas [16] and mesenchymal malignancies, including malignant Triton tumors, biphenotypic sinonasal sarcomas [17], and mesenchymal chondrosarcomas [18]. The most well-defined categories of rhabdomyosarcomas are those associated with recurrent molecular alterations, which include alveolar rhabdomyosarcoma underlined by *FOXO1* fusions [19], *MYOD1*-mutated

spindle cell rhabdomyosarcomas [4, 5], and *NCOA2*- and *VGLL2*-fused spindle cell rhabdomyosarcomas [2, 3]. The identification of new recurrent molecular features may be helpful to break down the heterogeneous category of spindle cell rhabdomyosarcomas into homogeneous and clinically more accurate subtypes. Our initial description of *FET-TFCP2* rhabdomyosarcomas [6] has hence been followed by the description of seven additional cases to date [6–8, 10, 20] (Table 4). Our series report 11 additional cases and provides a comprehensive clinical, pathological, genomic, and transcriptional characterization of this emerging entity.

Clinically, *FET-TFCP2* rhabdomyosarcomas display a striking bone predilection with only two cases exclusively located in soft tissue in this series (Table 1), whereas no other soft tissue case is reported in the literature. Most bone cases destroyed the cortex and invaded surrounding soft tissue (Table 2). Most intriguingly, craniofacial bones are the commonest bones affected ($n = 14/18$). This is in keeping with available literature of intra-osseous rhabdomyosarcomas, as despite their extreme rarity, most published series are available in craniofacial bones, although without molecular characterization of the cases [21, 22]. Our data confirm that *FET-TFCP2* tumors mostly affect young adult patients with a median age at diagnosis of 31

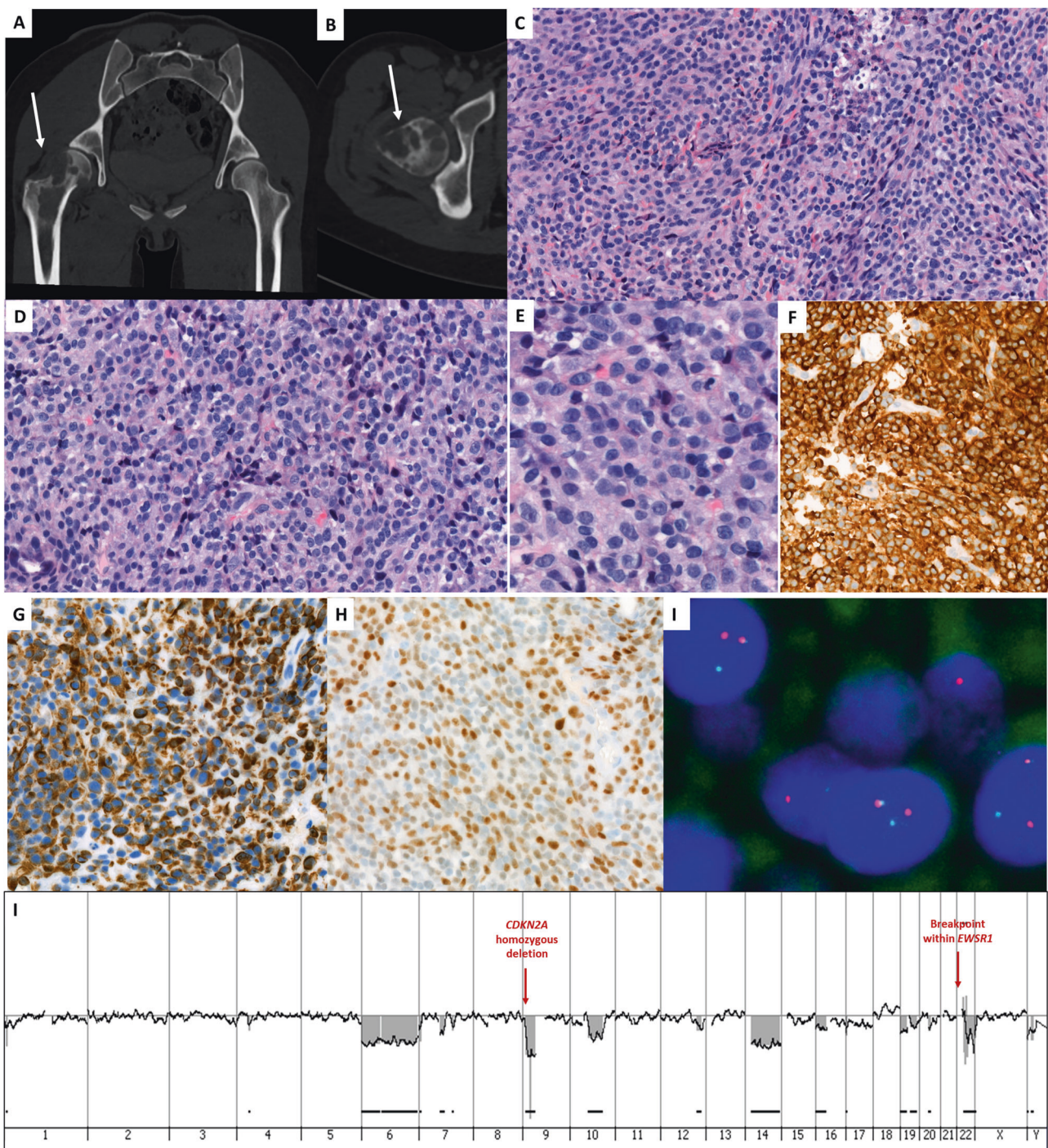


Fig. 4 Case of *FET-TCF2* rhabdomyosarcoma with small nuclear cytomorphology. CT of a multifocal osteolytic lesion of the right femur in the inter-trochanteric area and femoral neck (**a**, **b**, coronal reformatted and axial views, respectively) (case TFCP2-7). Core-needle biopsy evidence a mixed proliferation arranged in short fascicles (**c**) and solid sheets (**d**). Tumor cells harbor ovoid nuclei with fine chromatin without nucleoli, relatively monomorphic (**e**). The

proliferation stained diffusely for ALK (**f**), AE1/E3 (**g**), and MYOD1 (**h**). A split signal is seen with the FISH assay using a break-apart probe spanning *TCF2* (**i**). Array-comparative genomic hybridization evidenced copy number alterations across the genome including homozygous deletion of *CDKN2A* and breakpoint within *EWSR1* locus, in keeping with the presence of an underlying *EWSR1-TCF2* fusion (**j**)

years with a slight female predominance ($n = 13/21$). Regarding outcomes, our series provides the best detailed clinical follow-up so far, with a median survival of

8 months. Among patients still alive, two have evidence of progressive disease ($n = 2/5$) despite the administration of multimodal therapy (Table 2) with available median follow-

Table 4 *FET-TFCP2* rhabdomyosarcomas reported in the literature

Author	Case ID	Age (y)/gender	Location	Tumor size	Morphology	IHC	Molecular	Therapy	Survival
Dashti et al. [7], 1 case	-	72/M	Mandible	4 cm	Monomorphic spindled and epithelioid cells arranged in short fascicles. No stroma.	Des+ MYOD1+ S100 focal+ Keratin+ ALK+	<i>FUS (exon 6)-TFCP2 (exon 2)</i>	Radical mandibulectomy	2 Months after surgery disease free
Zhu et al. [10], 1 case	Case 5	74/F	Maxilla	4 cm	Monomorphic spindle cells with inflammatory background. Frequent mitotic figures and focal necrosis.	Des+ MYOD1+ S100- Keratin- ALK+	<i>FUS (exon 6)-TFCP2 (exon 2)</i>	NA	Regional metastasis to the neck
Agaram et al. [8], 4 cases	Case 3 Case 4	27/F 33/F	Skull Maxilla	NA	Epithelioid Epithelioid	Des+ MYOD1+ Keratin + ALK+	<i>EWSR1-TFCP2</i> <i>EWSR1-TFCP2</i>	NA Surgical resection	NA ANED (108 months FO) NA
	Case 5	20/M	Iliac bone		Pure spindle cell morphology		<i>EWSR1-TFCP2</i>	NA	NA
	Case 6	37/F	Femoral bone		Pure spindle cell morphology		<i>FUS-TFCP2</i>	NA	NA
Tagami et al. [9], 1 case	-	70/F	Lumbar vertebra	NA	Spindled to round cells, scattered rhabdomyoblasts	Des focal + MYOD1 + S100 - Keratin + ALK +	<i>FUS</i> rearrangement	Surgery followed by chemotherapy and radiotherapy	Still alive after 6 month of diagnosis
Wong et al. [20], 1 case	-	23/M	Nasal cavity	8 cm	Monomorphic epithelioid, and rhabdoid and spindled cells. High mitotic count >30 per 10 HPF	Des focal + MYOD1+ MYF4 focal+ S100- Keratin- ALK+	<i>FUS-TFCP2</i>	No response to combination of chemotherapy and radiotherapy, currently treated with Crizotinib (ALK inhibitor)	AWD (2 months FO)

ANED alive no evidence of disease, *AWD* alive with disease, *FO* follow-up, *Des* Desmin

up of 20 months. Altogether, our series highlights the aggressive behavior of this entity.

Microscopically, in addition to our initial report, most *FET-TFCP2* rhabdomyosarcomas actually display a hybrid spindle cell and epithelioid pattern. All tumors displayed high-grade features with vesicular nuclei, high mitotic activity, and tumor necrosis. A rhabdomyosarcomatous immunophenotype was always present, *MYOD1* being more sensitive than Myogenin (Table 3). At transcriptional level, *FET-TFCP2* tumors expressed high levels of skeletal muscle transcripts, especially *sarcoglycan alpha*, *MYOD1*, and *MYOG* [6]. All tumors consistently expressed keratins. Keratins are commonly expressed in rhabdomyosarcomas (staining was seen in 50% of alveolar rhabdomyosarcoma of our control cohort, Supplementary Table S4), although only focally, contrasting with the diffuse expression displayed by most *FET-TFCP2* rhabdomyosarcomas (Table 3). Among the other published cases, one case with *FUS-TFCP2* fusion is a “phenotypic outlier,” displaying a full myogenic profile (coexpression of caldesmon and smooth muscle actin (SMA)) and negative for keratins [10].

These tumors differ from the so-called “epithelioid rhabdomyosarcomas,” which display a pure solid architecture, a rare feature in *FET-TFCP2* tumors with only three purely epithelioid cases in this series [23]. Moreover, reported cases of epithelioid rhabdomyosarcomas are located in deep soft tissue and scarcely express epithelial markers contrasting with the features of *FET-TFCP2* rhabdomyosarcomas [23]. In the control cohort, all nine epithelioid rhabdomyosarcomas in soft tissue were negative for *TFCP2* rearrangement by FISH break-apart assay, further supporting this pathological subtype is distinct from *FET-TFCP2* rhabdomyosarcomas (Supplementary Table S2). We also report herein a new pattern in *FET-TFCP2* rhabdomyosarcoma with one case associated with a round cell pattern, in which tumor cells were arranged in trabeculae and embedded in an abundant fibromyxoid stroma (Fig. 2c, d).

All cases were associated with *TFCP2* rearrangements, fused in 5' to *FUS* ($n = 8/14$) or *EWSR1* ($n = 6/14$). *TFCP2*, also known as *Late Simian Virus 40 factor (LSF)*, located in 12q13.12 encodes an evolutionarily conserved transcription factor that targets the thymidylate synthase gene, involved in the enzymatic regulation of DNA synthesis [24, 25]. *TFCP2* contains an N-terminal DNA-binding domain and a C-terminal sterile alpha motif domain involved in protein dimerization [26] *TFCP2* has been formerly linked to cancer with evidence of oncogenic deregulation in hepatocellular carcinoma [27] mediated by C-MET signaling [28]. Breakpoints within *TFCP2* occur before the second exon, thereby preserving the DNA-binding domain of the protein. This fusion provides a further illustration of alternative involvement of *EWSR1* or *FUS* in a single tumor entity, as

these two genes are structurally and functionally related [29].

Our data show that *FET-TFCP2* rhabdomyosarcomas are associated with overall complex genetic profiles. This is in agreement with the features reported in one case characterized by genomic profiling (Affymetrix platform), which showed a tetraploid profile with numerous alterations, chromotripsis of 1p and 3p, and homozygous deletion of *CDKN2A* [20]. *CDKN2A* inactivation was indeed seen in all tested cases of our series (Supplementary Table S3). Concomitantly, three cases had genomic features consistent with unbalanced translocation.

Despite their complex genomic rearrangements, the *FET-TFCP2* rhabdomyosarcomas are transcriptionally homogeneous, as they clustered altogether. Moreover, adding more *FET-TFCP2* samples to the initial clustering analyses strengthen their clustering away from the other rhabdomyosarcoma subtypes, as shown by the clear consensus clustering (Supplementary Fig. S3).

In this series, all but three *FET-TFCP2* rhabdomyosarcomas overexpressed *ALK* both at transcriptional and protein levels, without underlying *ALK* fusion. According to RNA-sequencing and comparative genomic hybridization (CGH) data, this upregulation does not involve translocation nor amplification but correlated with *ALK* genomic deletion, presumably involving alternative transcription initiation [14] (Supplementary Fig. S4). No copy number alteration of *ALK* was detected in both *FET-TFCP2* tumors negative for *ALK* by immunohistochemistry studied by array-CGH. Interestingly, *ALK* deletion has been previously reported in one case of *FUS-TFCP2* sarcoma: the sequencing data showed that the upregulation did not involve exons 1–16, therefore in keeping with our finding [20]. Intriguingly, the expression of *ALK* has been shown with variable frequencies in other types of rhabdomyosarcomas, especially the alveolar subtype in up to 69% of cases [30, 31], but its underlying biological mechanism remains elusive. We confirmed this finding in our control cohort, as *ALK* staining was seen in 88.9% and 28.6% of alveolar and embryonal rhabdomyosarcomas, respectively (Supplementary Table S4). Nonetheless, *ALK* expression is rare in spindle cell rhabdomyosarcomas, present only in 23% of our control cases (Supplementary Table S4). Despite *ALK* overexpression, targeted therapy against *ALK* has not shown in vivo effects nor clinical benefits in alveolar rhabdomyosarcomas patients [32, 33], although the actual levels of expression of *ALK* were not reported in the studies. It is notable that at transcriptional level, the intensity of *ALK* upregulation is at least as high as in *ALK*-rearranged tumor types, such as *ALK*-fused inflammatory myofibroblastic tumors. Interestingly, one prospective patient of this series benefited from an *ALK* inhibitor after clinical progression under conventional chemotherapy, which

stabilized tumor size with available follow-up of 15 months under therapy (Table 2). Therefore, the potential benefits of ALK inhibition in *FET-TFCP2* rhabdomyosarcoma need to be further addressed, either as single therapy or in combination.

FET-TFCP2 rhabdomyosarcomas should not be confused with sarcomas with *EWSR1-PATZ1* fusions, as these tumors may develop in the bones and display a rhabdomyogenic phenotype [34]. The latter tumors have a hybrid spindle cell to round cell phenotype but lack the epithelioid and high-grade nuclear features of most *FET-TFCP2* rhabdomyosarcomas. *FET-TFCP2* tumors, especially those that have an exclusive spindle cell morphology, may be confused with inflammatory myofibroblastic tumors. Both tumor types commonly express keratins and ALK. Nonetheless, ALK overexpression in inflammatory myofibroblastic tumors is related to underlying *ALK* fusion with a variety of partner genes, in contrast with *FET-TFCP2* rhabdomyosarcomas. Intra-osseous rhabdomyosarcomas with *MEIS1-NCOA2* fusions represent another emerging entity with only two cases reported for now [8]. This variant displays pure spindle cell morphology and does not express keratins nor ALK [8]. The rhabdomyogenic phenotype and focal staining for S100 protein seen in a subset of cases may also raise suspicion for a malignant Triton tumor, although none of *TFCP2* patients had an underlying neurofibromatosis condition and the *SOX10* marker was negative in all tested cases. Furthermore, we did not find *TFCP2* fusions in control cases of malignant triton tumors (Supplementary Table S2).

In short, we report herein the largest and most comprehensive study of *FET-TFCP2* rhabdomyosarcomas. This new entity is associated with hybrid epithelioid and spindle cell features, a predilection for craniofacial bones, and expresses consistently keratins and ALK. Tumors harboring these features should therefore benefit from molecular investigations seeking for a *FET-TFCP2* fusion gene. Altogether, the clinical, morphological, and molecular data strongly suggest that these tumors represent a distinct rhabdomyosarcoma subtype.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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