Intracranial mesenchymal tumors with FET-CREB fusion are composed of at least two epigenetic subgroups distinct from meningioma and extracranial sarcomas

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INTRODUCTION

The newly recognized World Health Organization (WHO) tumor type, ‘intracranial mesenchymal tumor, FET-CREB fusion-positive’ [1], is a rare neoplasm of the central nervous system (CNS) that has been previously described as either intracranial angiomatoid fibrous histiocytoma (AFH) or intracranial myxoid mesenchymal tumor (IMMT) [2–5]. This tumor is molecularly defined by in-frame gene fusion of the FET family of RNA-binding proteins (EWSR1 or FUS) to the CREB (cyclic AMP response element binding protein) family of transcription factors, which includes ATF1, CREB1, and CREM. Notably, identical FET-CREB fusions are recurrently found in angiomatoid fibrous histiocytoma (AFH), clear cell sarcoma of soft tissue (CCS), and AFH of extracranial soft tissue. The first subgroup (Group A, 16 tumors) clustered nearest to but independent of solitary fibrous tumor and AFH of extracranial soft tissue, whereas the second epigenetic subgroup (Group B, 4 tumors) clustered nearest to but independent of CCS and also lacked expression of melanocytic markers (HMB45, Melan A, or MITF) characteristic of CCS. Group A tumors most often occurred in adolescence or early adulthood, arose throughout the neuroaxis, and contained mostly EWSR1-ATF1 and EWSR1-CREB1 fusions. Group B tumors arose most often in early childhood, were located along the cerebral convexities or spinal cord, and demonstrated an enrichment for tumors with CREM as the fusion partner (either EWSR1-CREM or FUS-CREM). Group A tumors more often demonstrated stellate/spindle cell morphology and hemangioma-like vasculature, whereas Group B tumors more often demonstrated round cell or epithelioid/rhabdoid morphology without hemangioma-like vasculature, although robust comparison of these clinical and histologic features requires future study. Patients with Group B tumors had inferior progression-free survival relative to Group A tumors (median 4.5 vs. 49 months, \( p = 0.001 \)). Together, these findings confirm that intracranial AFH-like neoplasms and IMMT represent histologic variants of a single tumor type (‘intracranial mesenchymal tumor, FET-CREB fusion-positive’) that is distinct from meningioma and extracranial sarcomas. Additionally, epigenomic evaluation may provide important prognostic subtyping for this unique tumor entity.

KEYWORDS
angiomatoid fibrous histiocytoma (AFH), ATF1, brain tumor, clear cell sarcoma, CREB1, CREM, EWSR1, intracranial mesenchymal tumor with FET-CREB fusion, intracranial myxoid mesenchymal tumor, molecular neuropathology, sarcoma
relationship of these intracranial mesenchymal tumors with FET-CREB fusions to AFH of extracranial soft tissue and other FET-CREB fusion-driven neoplasms has been uncertain.

We previously studied the clinical, radiologic, histologic, and genomic features of a cohort of 20 intracranial mesenchymal tumors with FET-CREB fusions [5]. We identified that these tumors occur with a female predominance (approximately 2:1 female/male ratio) in a wide age range, but most often occur in the second and third decades of life. They are predominantly extra-axial or intraventricular tumors which can arise throughout the neuroaxis, including the falx, tentorium, cerebral convexities, and lateral ventricles. They are typically contrast enhancing masses, well-circumscribed, with solid and cystic growth patterns, and often have pronounced peritumoral edema. Beyond the oncogenic FET-CREB fusions, they typically lack accompanying oncogenic mutations, amplifications, or deletions, and usually harbor near-diploid genomes. These tumors are associated with a propensity for local recurrence over time, but only a small subset have disseminated and resulted in patient mortality.

Here we have performed genome-wide DNA methylation profiling on our previously published cohort of 20 primary intracranial mesenchymal tumors with FET-CREB fusions to further study the ontology of these neoplasms and identify any clinically relevant epigenetic subgroups.

2 | MATERIALS AND METHODS

2.1 | Study population and tumor specimens

The study cohort consisted of 20 patients who underwent surgical resection of a primary intracranial neoplasm that was identified to harbor a gene fusion of EWSR1 or the related FUS together with a CREB family member (ATFI, CREBI, or CREM). The clinical features of these 20 patients and histopathologic features of the tumor cohort were previously reported [5].

2.2 | Genome-wide DNA methylation profiling

Tumor tissue was selectively scraped from unstained slides or punched from formalin-fixed, paraffin-embedded (FFPE) blocks using 2.0 mm disposable biopsy punches (Integra Miltex Instruments) to enrich for the highest tumor content possible. Genomic DNA was extracted from this macrodissected tumor tissue using the QIAamp DNA FFPE Tissue Kit (Qiagen). Genomic DNA was bisulfite converted using the EZ DNA Methylation kit following the manufacturer’s recommended protocol (Zymo Research). Bisulfite converted DNA was then amplified, fragmented, and hybridized to Infinium EPIC 850k Human DNA Methylation BeadChips following the manufacturer’s recommended protocol (Illumina).

2.3 | Processing and quality assessment of DNA methylation data

Methylation data were preprocessed using the minfi package (v.1.30.0) in R Bioconductor (version 3.5.3) [17]. The detection p-value for each sample was computed, and CpG sites with detection p values above 0.05 were discarded from the analysis. Additional quality control was performed by calculating the median log (base2) intensities for methylated and unmethylated signals for each array. All samples had unmethylated and methylated median intensity values above 10 that were used for analysis. Functional normalization with NOOB background correction and dye-bias normalization was performed [18, 19]. Probe filtering was performed after normalization. Specifically, probes located on sex chromosomes, containing nucleotide polymorphisms (dbSNP132 Common) within five base pairs of and including the targeted CpG site, or mapping to multiple sites on hg19 (allowing for one mismatch), as well as cross reactive probes were removed from analysis.

2.4 | Unsupervised hierarchical clustering of DNA methylation data

We performed unsupervised hierarchical clustering with the hclust function in Rstats (v3.6.0) to assess variation in DNA methylation patterns and determine any relevant epigenetic subgrouping among the 20 tumors. The lmFit function from the Limma package (v.3.40.6) was applied on a log-transformed β-value matrix to identify the 20,000 most differentially methylated CpG probes across the tumor cohort. Then K-means clustering utilizing the Pearson distance matrix with complete linkage was used to determine the optimal number of clusters, through 500 re-sampling interactions of the dataset for K-means of 2, 3, 4, or 5. Visualization was performed using the R package ComplexHeatmap (v.2.0.0) [20].

We also compared the DNA methylation patterns of the 20 intracranial mesenchymal tumors with FET-CREB fusion against the DNA methylation patterns of two extracranial sarcoma entities with identical FET-CREB fusions (clear cell sarcoma of soft tissue and angiomatoid fibrous histiocytoma) that we recently generated as part of our development of the DKFZ Sarcoma Methylation Classifier (www.molecularsarcomapathology.org) [21]. This included 7 reference cases of clear cell sarcoma of soft parts (tumor ID’s: 956, 957, 958, 959, 960, 961, and 962), which were all located in the soft tissue of the upper or lower extremities with median patient age of 56 years (range 18–78 years). Also included
were 8 reference cases of angiomatoid fibrous histiocytoma (tumor ID’s: 340, 341, 360, 361, 362, 363, 364, 1066), which were all located in extracranial soft tissue with median patient age of 11 years (range 6–13 years). The Limma function from the Limma package (v.3.40.6) was applied on a log-transformed β-value matrix to identify the 20,000 most differentially methylated CpG probes across the cohort of 20 intracranial mesenchymal tumors with FET-CREB fusion together with the 7 reference cases of clear cell sarcoma of soft parts and 8 reference cases of angiomatoid fibrous histiocytoma of extracranial soft tissue. K-means clustering utilizing the Pearson distance matrix with complete linkage was then used to determine the optimal number of clusters, through 500 re-sampling interactions of the dataset for K-means of 2, 3, 4, or 5. Unsupervised hierarchical clustering of DNA methylation data was performed using the heatmap function in Rstats (v3.6.0). Visualization was performed using the R package ComplexHeatmap (v.2.0.0).

2.5 | tSNE dimensionality reduction of DNA methylation data

The DNA methylation profiles of the 20 intracranial mesenchymal tumors with FET-CREB fusions were assessed together with 210 reference tumors spanning 17 sarcoma and CNS tumor entities previously generated at DKFZ [21, 22], which were selected based on tumor types with a similar meningeal/extra-axial location, tumor types that might enter into the differential diagnosis based on overlapping morphologic appearance, and extracranial sarcomas driven by EWSR1 or FUS gene fusions. These included 10 angiomatoid fibrous histiocytoma of extracranial soft tissue, 30 atypical teratoid/rhabdoid tumor (belonging to the three different epigenetic subgroups: MYC, SHH, and TYR), 10 clear cell sarcoma of soft tissue, 10 chordoma, 10 extraskeletal myxoid chondrosarcoma, 10 H3 K27M-mutant diffuse midline glioma, 10 desmoplastic small round cell tumor, 10 Ewing sarcoma, 10 IDH-wildtype glioblastoma of the mesenchymal epigenetic subclass, 10 low-grade fibromyxoid sarcoma, 30 meningioma (belonging to the various epigenetic subgroups), 10 alveolar rhabdomyosarcoma, 10 embryonal rhabdomyosarcoma, 10 CIC-altered sarcoma, 10 DICER1-mutant sarcoma, 10 solitary fibrous tumor, and 10 synovial sarcoma. Since the reference cohort contained methylation data generated using the Infinium Human Methylation 450k BeadChips, the approximately 450,000 overlapping CpG sites between the EPIC 850k and 450k BeadChips were used in the analysis. A beta value matrix with approximately 390,000 CpG probes was used for all downstream analysis. Row-wise standard deviation was calculated for each probe across all samples, and the 20,000 most differentially methylated probes with standard deviation >0.216 were selected. Dimensionality reduction using t-distributed stochastic neighbor embedding (tSNE) was performed by Rtsne (v.0.15) using the following analysis parameters: dims = 2, max_iter = 3000, theta = 0, perplexity = 20, eta = 200. The tSNE plot was visualized with ggplot2 (v.3.2.0) [https://ggplot2.tidyverse.org/].

2.6 | Differentially methylated region and gene ontology analysis

Differentially methylated regions (DMRs) between epigenetic tumor classes were identified using DMRcate (v.1.20.0) [23], which ranks the most differentially methylated genomic regions using gaussian smoothing across adjacent CpG sites. For DMRcate, a model with coefficients using the following parameters and thresholds was created: lambda (genomic window length) = 1000 nucleotides, C (scaling factor) = 2, and probe false discovery rate (FDR) cutoff of less than 0.05. Results were ranked by Fisher’s multiple comparison statistic and filtered for those DMRs with both FDR and Stouffer scores less than 0.001. DMRs were then annotated to the nearest gene transcriptional start sites, based on ENSEMBL genome annotations. Gene Ontology (GO) analysis of differentially methylated gene regions was performed using the gene function in the missMethyl package [24].

2.7 | Immunohistochemistry

Immunohistochemistry was performed on whole formalin-fixed, paraffin-embedded tissue sections using the following antibodies: desmin (Cell Marque, clone D33, undiluted, ER1 antigen retrieval), epithelial membrane antigen (EMA, Leica, clone GPL1.4, undiluted, ER1 antigen retrieval), CD99 (Signet, clone CD99, 1:400 dilution, ER1 antigen retrieval), S100 (DAKO, polyclonal, 1:2000 dilution, no antigen retrieval), MUC4 (Millipore, clone 8G7, 1:500 dilution, ER1 antigen retrieval), somatostatin receptor 2A (SSTR2A, Abcam, clone UMBI, 1:2000 dilution, ER2 antigen retrieval), OLG2 (Innunu Bio Labs, polyclonal, 1:200 dilution, ER1 antigen retrieval), glial fibrillary acidic protein (GFAP, DAKO, polyclonal, 1:3000 dilution, no antigen retrieval), synaptophysin (Cell Marque, polyclonal, 1:100 dilution, ER2 antigen retrieval), CD68 (Leica, clone 514H12, undiluted, ER2 antigen retrieval), cytokeratin AE1/AE3 (Dako, clone AE1/AE3, 1:100 dilution, ER1 antigen retrieval), cytokeratin CAM5.2 (Becton Dickinson, clone CAM5.2, 1:100 dilution, ER1 antigen retrieval), HMB45 (Dako, clone HMB45, undiluted, CC1 antigen retrieval), Melan A (Dako, clone A103, 1:10 dilution, ER1 antigen retrieval), MITF (Dako, clone D5, 1:200 dilution, ER1 antigen retrieval), myogenin (Cell Marque, clone A103, 1:10 dilution, ER1 antigen retrieval), and Ki-67 (Dako, clone Mib1, 1:50 dilution, ER2 antigen retrieval). Immunostaining for desmin, EMA, CD99, S100, MUC4,
SSTR2A, OLIG2, GFAP, synaptophysin, CD68, cytokeratin AE1/AE3, cytokeratin CAM5.2, Melan A, MITF, and Ki-67 was performed in a Leica BOND-III automated stainer. Immunostaining for HMB45 and myogenin was performed in a Ventana BenchMark Ultra automated stainer. Diaminobenzidine was used as the chromogen, followed by hematoxylin counterstain.

2.8 Kaplan-Meier survival plots and statistical analyses

Clinical outcomes were studied by Kaplan-Meier analysis using GraphPad Prism. The Kaplan-Meier survival analysis was stratified by epigenetic subgroup, and p value was calculated by Log-rank (Mantel-Cox) test. Statistical comparison of histologic and immunohistochemical features was performed by Mann-Whitney unpaired two-tailed t test using GraphPad Prism.

3 RESULTS

3.1 DNA methylation profiling reveals two epigenetic subgroups

We performed genome-wide DNA methylation profiling on our previously published cohort of 20 primary intracranial mesenchymal tumors with FET-CREB fusions. The clinical and radiologic features of these 20 patients and the histopathologic and genomic features of the tumor cohort were previously reported in open access format – https://onlinelibrary.wiley.com/doi/10.1111/bpa.12918 [5]. Unsupervised hierarchical clustering of the DNA methylation profiles segregated these tumors into two epigenetically distinct subgroups: Group A consisting of 16 tumors and Group B consisting of 4 tumors (Figure 1A and Table 1).

3.2 Clinical and molecular characteristics of the two epigenetic subgroups

Group A tumors were from 11 females and 5 males with a median age at diagnosis of 15 years (range 12–70 years) (Tables 1 and 2, Figure 1B). The tumors were located throughout the neuroaxis, including the cerebral convexities (n = 4), tentorium (n = 2), falx (n = 2), lateral ventricles (n = 4), and cerebellopontine angle (n = 4) (Figure 1C). Fusion partners were EWSRI-ATFI (n = 7), EWSRI-CREB1 (n = 7), and EWSRI-CREM (n = 2) (Figure 1D). The 4 Group B tumors were exclusively from females with a median age at diagnosis of 7 years (range 4–15 years). The tumors were located along the cerebral convexities (n = 3) or spinal cord (n = 1). Fusion partners were EWSRI-ATFI (n = 1), EWSRI-CREM (n = 2), and FUS-CREM (n = 1). As previously reported, these tumors all had near-diploid genomes [5], and we did not observe any recurrent chromosomal copy number changes among either epigenetic subgroup.

3.3 Differential gene methylation analysis between two epigenetic subgroups

We next determined all differentially methylated genomic regions (DMR) with a mean beta value difference of at least 0.3 between the two epigenetic subgroups, which yielded nearly 1100 such DMR (Tables S1 and S2). Gene Ontology analysis of the most differentially methylated genes between the two epigenetic subgroups revealed networks involved in Rho GTPase signaling (Figure 1E, Table S3), a pathway known to control cell growth, motility, and actin cytoskeletal remodeling.

3.4 Histologic and immunophenotypic features of the two epigenetic subgroups

We have previously described the wide morphologic spectrum of intracranial mesenchymal tumors with FET-CREB fusion, ranging from stellate/spindle cell to epithelioid/rhabdoid cytology along with variable stromal mucin content and hemangioma-like vasculature [5]. When comparing histologic features between the two epigenetic subgroups, no statistically significant differences were apparent (Figure 2, Table 3). The presence of a myxoid stroma did not strictly correlate with the epigenetic subgroups – 11/16 (69%) of the Group A tumors and 1/4 (25%) of the Group B tumors demonstrated a mucin-rich stroma. Hemangioma-like vasculature was only encountered in the Group A tumors (10/16 [63%]). Stellate/spindle cell cytromorphology was also only encountered in Group A tumors (10/16 [63%]).

The Ki-67 labeling index in this tumor cohort was generally low (less than 5%, 8 of the 15 evaluated tumors), but occasionally was elevated up to 15%–25% (7 of the 15 evaluated tumors) [5]. There was no significant difference in Ki-67 labeling index between the two epigenetic subgroups, though three of the four Group B tumors were those with elevated Ki-67 labeling index.

We also examined for any differences in immunophenotype between the two epigenetic subgroups (Table 4). Desmin, EMA, CD99, and CD68 expression was nearly ubiquitous among all tumors belonging to both subgroups, and MUC4 and synaptophysin expression was also frequent in both subgroups. Among the seven Group A tumors evaluated for MUC4 expression, three demonstrated diffuse strong staining, one demonstrated focal positivity, and two were negative. Among the four Group B tumors evaluated for MUC4 expression, three demonstrated focal positivity and one was negative. Among the seven Group A tumors evaluated for synaptophysin expression, three demonstrated patchy variable
intensity staining while the other four were negative. Both of the two Group B tumors evaluated for synaptophysin expression demonstrated patchy variable intensity staining. None of the examined tumors belonging to either subgroup was positive for somatostatin receptor 2A (SSTR2A) expression or markers of melanocytic differentiation (MITF, Melan A, and HMB45). None of the investigated proteins had significantly different expression levels between Group A and Group B tumors, and further studies are necessary to identify potential immunohistochemical surrogates for segregating the two epigenetic subgroups.

3.5 | Clinical outcomes of the two epigenetic subgroups

The complete clinical data including extent of resection, treatment regimen, and outcome data from the twenty patients were previously reported [5]. Kaplan-Meier analysis of progression-free survival (PFS) stratified by epigenetic subgroup revealed inferior outcomes of Group B tumors relative to Group A tumors (median PFS of 4.5 vs. 49 months, \( p = 0.001 \)) (Figure 3). Only three of the 20 patients succumbed to disease during the period of clinical follow-up, all of whom harbored EWSR1-ATF1 fusions, of which two (ATF1 #6 and ATF1 #7) belonged to Group A and one (ATF1 #2) belonged to Group B. Kaplan-Meier analysis of overall survival stratified by epigenetic subgroup did not reveal a significant difference (data not shown).

3.6 | Epigenetic comparison with other CNS tumor entities and extracranial sarcomas

We next performed tSNE dimensionality reduction of the DNA methylation profiles for the 20 intracranial mesenchymal tumors with FET-CREB fusion together with 210 reference tumors spanning 17 sarcoma and CNS tumor entities previously generated at DKFZ [21, 22] (Table S4). The intracranial mesenchymal tumors with

![Figure 1](image-url)
FET-CREB fusion resolved into two distinct epigenetic subgroups that were both divergent from all other analyzed intracranial neoplasms and soft tissue sarcomas, including meningioma, Ewing sarcoma, extraskeletal myxoid chondrosarcoma, clear cell sarcoma of soft tissue (CCS), and AFH of extracranial soft tissue (Figure 4, top panel). The two epigenetic subgroups identified by unsupervised hierarchical clustering (Figure 1A) were recapitulated by the tSNE dimensionality reduction analysis, with the same 16 tumors aligning with Group A and same 4 tumors aligning with Group B by both analyses (Figure 4, bottom panel). The Group A tumors clustered nearest to but independent of solitary fibrous tumor and AFH of extracranial soft tissue, whereas the Group B tumors clustered nearest to but independent of CCS and the mesenchymal subclass of IDH-wildtype glioblastoma. By random forest classification using both the online DKFZ Sarcoma Classifier tool version 12.2 and the online DKFZ Brain Tumor Classifier tool version 11b4, only 3 of the 16 Group A tumors aligned with the methylation class “Angiomatoid fibrous histiocytoma” with a calibrated score of greater than 0.9, whereas the remainder of the 13 tumors did not reliably classify as “Angiomatoid fibrous histiocytoma” or any other reference methylation class of sarcoma or CNS tumor (Table S5). None of the 4 Group B tumors reliably classified as “Angiomatoid fibrous histiocytoma”, “Clear cell sarcoma of soft tissue”, or any other reference methylation class of sarcoma or CNS tumor (Table S5).

We next further compared the DNA methylation patterns of the 20 intracranial mesenchymal tumors with FET-CREB fusion against two extracranial sarcoma entities with identical FET-CREB fusions (clear cell sarcoma of soft tissue and angiomatoid fibrous histiocytoma) that were recently generated as part of the DKFZ Sarcoma Methylation Classifier [21]. Unsupervised hierarchical clustering was performed on the 16 Group A tumors together with 8 reference cases of angiomatoid fibrous histiocytoma, which were all located in extracranial soft tissue with a median patient age of 11 years (range 6–13 years). This unsupervised clustering analysis segregated the 24 total tumors into two groups
– one composed of the 16 Group A tumors and the other composed of the 8 AFH of extracranial soft tissue (Figure 5A). We next determined all differentially methylated genomic regions (DMR) with a mean beta value difference of at least 0.3 between the Group A intracranial mesenchymal tumors and AFH of extracranial soft tissue, which yielded nearly 600 such DMR (Tables S6 and S7). Gene Ontology analysis of the most differentially methylated gene regions between the Group A tumors and AFH of extracranial soft tissue revealed networks involved in muscle structure development and axial mesoderm formation (Figure 5B, Table S8).

Unsupervised hierarchical clustering was also performed on the 4 Group B tumors together with 7 reference cases of clear cell sarcoma, which were all located in the soft tissue of the upper or lower extremities with median patient age of 56 years (range 18–78 years). This unsupervised clustering analysis segregated the 11 total tumors into two groups – one composed of the 4 Group B tumors and the other composed of the 7 clear cell sarcomas (Figure 6A). The one intracranial mesenchymal tumor with EWSR1-CREM fusion (CREM #1) that clustered somewhat nearer to CCS than the other three Group B tumors on tSNE dimensionality reduction segregated together with the Group B tumors and not CCS by this unsupervised hierarchical clustering analysis. We next determined all differentially methylated genomic regions (DMR) with a mean beta value difference of at least 0.3 between the Group B intracranial mesenchymal tumors and CCS of soft tissue, which yielded nearly 700 such DMR (Tables S9 and S10). One of the most differentially methylated genomic regions was the MITF gene, which was substantially hypermethylated in the Group B intracranial mesenchymal tumors versus hypomethylated/unmethylated in the CCS tumors (Figure 6B). MITF encodes the microphthalmia-associated transcription factor (MITF).

| Clinical features   | Group A | Group B | All tumors |
|--------------------|---------|---------|------------|
| Age at diagnosis (years) |         |         |            |
| Median             | 15      | 7       | 14         |
| Range              | 12–70   | 4–15    | 4–70       |
| Sex                |         |         |            |
| Male               | 5       | 0       | 5          |
| Female             | 11      | 4       | 15         |
| Tumor location     |         |         |            |
| Cerebral convexity | 4       | 3       | 7          |
| Tentorium          | 2       | 0       | 2          |
| Falx               | 2       | 0       | 2          |
| Lateral ventricle  | 4       | 0       | 4          |
| CP angle           | 4       | 0       | 4          |
| Spinal cord        | 0       | 1       | 1          |
| Fusion type        |         |         |            |
| EWSR1-ATF1         | 7       | 1       | 8          |
| EWSR1-CREB1        | 7       | 0       | 7          |
| EWSR1-CREM         | 2       | 2       | 4          |
| FUS-CREM           | 0       | 1       | 1          |

**Figure 2**  
Histologic features of intracranial mesenchymal tumors with FET-CREB fusion belonging to the two epigenetic subgroups. Hematoxylin and eosin-stained sections from three representative tumors of the two epigenetic subgroups.
factor on chromosome 3p13 that functions as a critical transcription factor for specifying melanocytic differentiation, and is robustly expressed in clear cell sarcoma of soft tissue (formerly referred to as melanoma of soft parts) that is pathologically defined by its expression of melanocytic markers including MITF, HMB45, and Melan A [25, 26]. In contrast to CCS, we found an absence of MITF expression by immunohistochemical staining in all four of the Group B intracranial mesenchymal tumors with FET-CREB fusion, as well as other melanocytic markers (Figure 6C). In addition to MITF, Gene Ontology analysis of the most differentially methylated gene regions between Group B intracranial mesenchymal tumors and CCS of soft tissue revealed networks involved in roof of mouth development and embryonic eye development (Figure 6D, Table S11).

### DISCUSSION

Here we have interrogated the epigenomic landscape of intracranial mesenchymal tumors harboring FET-CREB fusion and correlated the results together with clinical and histopathologic features. Our findings reveal that these tumors segregate into two discrete epigenetic...
subgroups which we have termed Group A and Group B. Group A tumors most often occurred in adolescence or early adulthood, arose throughout the neuroaxis, and consisted of a mix of EWSR1-ATF1, EWSR1-CREB1, and EWSR1-CREM fusions. Group B tumors arose most often in early childhood, were located along the cerebral convexities or spinal cord, and demonstrated an enrichment for tumors with CREM as the fusion partner (either EWSR1-CREM or FUS-CREM). Group A tumors more often demonstrated stellate/spindle cell morphology and hemangioma-like vasculature, whereas Group B tumors more often demonstrated round cell or epithelioid/rhabdoid morphology without hemangioma-like vasculature, although these differences did not reach statistical significance. The presence of a myxoid stroma did not correlate with epigenetic subgrouping, as both epigenetic subgroups contained some tumors with and some without a mucin-rich background. No immunohistochemical differences between the two epigenetic subgroups were identified, and further studies are necessary to test potential immunohistochemical surrogates for segregating the two epigenetic subgroups, perhaps utilizing the list of most differentially methylated genes as a starting point (Tables S1 and S2). Analysis of patient outcomes demonstrated worse progression-free survival of Group B tumors relative to Group A tumors (median PFS of 4.5 vs. 49 months, respectively), although the cohort size of this study is small and this finding requires further confirmation in larger patient cohorts.

Notably, one of the four tumors that we assigned as belonging to Group B (CREM #1 located in the spinal cord of a 15-year-old female harboring EWSR1-CREM fusion) demonstrated a somewhat divergent epigenetic profile relative to the other three tumors within Group B by both unsupervised hierarchical clustering (Figure 1A) and tSNE dimensionality reduction (Figure 4). Despite being in closer proximity to the reference cluster of CCS on the tSNE plot, this tumor CREM #1 more closely grouped with the other Group B tumors than CCS by unsupervised hierarchical clustering (Figure 6A), demonstrated hypermethylation of the MITF locus similar to other Group B tumors (Figure 6B), and lacked expression of melanocytic protein markers similar to other Group B tumors (Table 4). Whether this solitary tumor
intracranial mesenchymal tumors with FET-CREB fusion to angiomatoid fibrous histiocytoma (AFH) of extracranial soft tissue. (A) Unsupervised hierarchical clustering of DNA methylation data showing segregation of the 16 Group A tumors from 8 reference cases of angiomatoid fibrous histiocytoma arising in extracranial soft tissue. Differentially methylated genomic regions are annotated in Tables S6 and S7. (B) Differential methylation-based gene ontology analysis for Group A intracranial mesenchymal tumors versus AFH of extracranial soft tissue, represented in a bar plot of $-\log_{10} p$ values for the most differentially methylated gene networks.

FIGURE 5 Epigenetic comparison of Group A intracranial mesenchymal tumors with FET-CREB fusion to angiomatoid fibrous histiocytoma (AFH) of extracranial soft tissue. (A) Unsupervised hierarchical clustering of DNA methylation data showing segregation of the 16 Group A tumors from 8 reference cases of angiomatoid fibrous histiocytoma arising in extracranial soft tissue. Differentially methylated genomic regions are annotated in Tables S6 and S7. (B) Differential methylation-based gene ontology analysis for Group A intracranial mesenchymal tumors versus AFH of extracranial soft tissue, represented in a bar plot of $-\log_{10} p$ values for the most differentially methylated gene networks.

in our cohort is representative of epigenetic heterogeneity amongst Group B tumors or alternatively might represent a third distinct epigenetic subgroup of intracranial mesenchymal tumors with FET-CREB fusion remains uncertain. Future studies with larger patient cohorts are necessary to reveal the full biologic spectrum and clinically relevant subgrouping of these tumors. Overall however, our finding of at least two distinct epigenetic subgroups is similar to a recent report of epigenomic characterization performed on a cohort of 11 primary intracranial mesenchymal tumors with FET-CREB fusion, which identified that 6 of their 11 cases formed a unique epigenetic cluster, whereas the other 5 cases were epigenetically heterogeneous and unclassifiable [27].

For the time being, we believe that these are best considered as two epigenetic subtypes under the single overarching tumor type ‘intracranial mesenchymal tumor, FET-CREB fusion-positive’. This conclusion is based on our cohort of 20 tumors and the fact that there were not statistically significant differences in histomorphology (including stromal mucin content), immunophenotype, fusion partner, patient age, sex, tumor anatomic location, or other features between the two epigenetic subgroups that would enable definitive segregation into two or more distinct tumor types/entities at this point beyond epigenomic signature. However, future studies encompassing larger patient cohorts may potentially indicate and be used to provide support that these actually represent two or more distinct tumor types.

There has been ongoing uncertainty as to the relationship of tumors diagnosed as “intracranial angiomatoid fibrous histiocytoma” and those diagnosed as “intracranial myxoid mesenchymal tumor”, and we previously proposed the unifying terminology of “intracranial mesenchymal tumor, FET-CREB fusion-positive” for this group of neoplasms [5], which has been adopted in the 5th edition of the WHO Classification of Tumors of the Central Nervous System [1]. There has also been ongoing uncertainty as to the relationship of these intracranial mesenchymal tumors with FET-CREB fusion to meningiomas, as well as to the myriad of extracranial neoplasms harboring identical FET-CREB fusions, which include angiomatoid fibrous histiocytoma, clear cell sarcoma of soft tissue, clear cell sarcoma of the gastrointestinal tract, primary pulmonary myxoid sarcoma, hyalinizing clear cell carcinoma of the salivary gland, and a subset of malignant mesotheliomas lacking BAP1 and NF2 alterations [6–16]. Our epigenomic profiling has shed substantial light on these issues that we discuss herein.

First, our epigenomic data, together with the differential immunophenotype (e.g. lack of somatostatin receptor 2A [SSTR2A] expression, presence of desmin and MUC4 expression), further differentiate intracranial
mesenchymal tumors with FET-CREB fusion from meningiomas. However, we cannot exclude a shared cell of origin with meningiomas and/or meningeal solitary fibrous tumors, potentially with epigenetic reprogramming driven by the FET-CREB fusion causing the unique epigenetic signature we found for this tumor type. Further studies are required to define the specific cell of origin of these tumors, which we speculate to be a mesenchymal cell within the meningeal covering of the brain. In support of this hypothesis, electron microscopy performed on intracranial mesenchymal tumors with FET-CREB fusion has reportedly demonstrated ultrastructural features overlapping with those typically seen in meningioma including interdigitating cell processes lined by well-formed desmosomes and abundant extracellular collagen [27].

Second, as both epigenetic subgroups of intracranial mesenchymal tumors with FET-CREB fusion contained some tumors with a mucin-rich stroma resembling “intracranial myxoid mesenchymal tumor” (or the so-called “myxoid variant of AFH”) and also some with a...
mucin-poor stroma resembling “intracranial angioma-toid fibrous histiocytoma”, we conclude that these rep-resent histologic variants of a single overarching tumor type. We believe these results further substantiate the unifying nosology of ‘intracranial mesenchymal tumor, FET-CREB fusion-positive’. However, as previously discussed, future studies encompassing larger patient cohorts may potentially indicate and be used to provide support that these epigenetic subgroups among intra-cranial mesenchymal tumors with FET-CREB fusions actually represent two or more distinct tumor types.

Lastly, we have compared the epigenomic signature of these intracranial mesenchymal tumors with two of the extracranial neoplasms harboring identical FET-CREB fusions for which DNA methylation profiles have been generated to date. Our results reveal that intracranial mesenchymal tumors with FET-CREB fusion are epi-genetically distinct from both AFH of extracranial soft tissue and CCS of soft tissue. Given both their distinct epigenetic signature and divergent anatomic site of origin, our results provide further evidence that these tu-mors should be considered a distinct tumor entity, and not merely regarded as intracranial occurrence of these two sarcomatous neoplasms which characteristically occur in extracranial soft tissue, most often in the extremities. This is particularly true for CCS of soft tissue, given the differential methylation of the MITF gene we identified and the absence of melanocytic marker expression in intracranial mesenchymal tumors with FET-CREB fusion.

In summary, we have epigenetically characterized intracranial mesenchymal tumors with FET-CREB fu-sion, revealing at least two distinct epigenetic subgroups with potential prognostic significance. Our epigenomic results also provide substantial clarification for the on-tology of these unique intracranial neoplasms.

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CONFLICT OF INTEREST
A patent for a DNA methylation-based method for clas-sifying brain tumors has been applied for by DKFZ-Heidelberg University with A.v.D. as an inventor. B.K.D., F.J.R., D.W.E., A.P., and D.A.S. are on the editorial board of Brain Pathology, but were not involved with the assessment or decision-making process for this manuscript. The other authors declare that they have no competing interests.

AUTHOR CONTRIBUTIONS
Emily A. Sloan performed the DNA extractions. Rohit Gupta, Christian Koelsche, Andreas von Deimling, and David A. Solomon performed the epigenomic analysis. Emily A. Sloan and David A. Solomon per-formed pathologic assessment. Javier E. Villaneuva-Meyer performed radiologic assessment. Emily A. Sloan, Jason Chiang, Sandra Alexandrescu, Jennifer M. Eschbacher, Wesley Wang, Manuela Mafra, Nasir Ud Din, Emily Carr-Boyd, Michael Watson, Michael Punsoni, Angelica Oviedo, Ahmed Gilani, Bette K. Kleinschmidt-DeMasters, Dylan J. Coss, M. Beatriz Lopes, Alyssa Reddy, Soo-Jin Cho, Andrew E. Horvai, Julieann C. Lee, Melike Pekmezci, Tarik Tihan, Andrew W. Bollen, Fausto J. Rodriguez, David W. Ellison, Arie Perry, Susan M. Chang, Mitchel S. Berger, and David A. Solomon provided clinical care and contributed to the patient cohort. Emily A. Sloan and David A. Solomon conceptualized the study, reviewed all data, prepared the figures, and wrote the manuscript. Susan M. Chang, Mitchel S. Berger, and David A. Solomon secured funding to support this research study. All authors critically reviewed the manuscript and approved its submission.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE
This study was approved by the Committee on Human Research of the University of California, San Francisco, with a waiver of patient consent.

DATA AVAILABILITY STATEMENT
DNA methylation array data files for the 20 intracranial mesenchymal tumors with FET-CREB fusion generated as part of this study are available from the Gene Expression Omnibus (GEO) repository under accession number GSE164994 (https://www.ncbi.nlm.nih.gov/geo/). Scanned image files of H&E stained sections from 19 of the tumors in this cohort are available for downloading and viewing at the following link: https://figshare.com/projects/Intracranial_mesenchymal_tumors_with_EWS-CREB_fusion/88661.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher’s website.

Table S1-S11

TABLE S1 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypermethylated in Group A tumors relative to Group B tumors

TABLE S2 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypermethylated in Group B tumors relative to Group A tumors

TABLE S3 Gene Ontology analysis ranked by differentially methylated genes in epigenetic subgroup A versus B of intracranial mesenchymal tumors with FET-CREB fusion
TABLE S4 Sarcoma and CNS tumor reference samples used for tSNE dimensionality reduction in Figure 4
TABLE S5 Clinical features of the 20 patients with ‘intracranial mesenchymal tumor, FET-CREB fusion-positive’, DNA methylation array data identifiers, epigenetic subgroup, and calibrated scores from random forest classification using the online DKFZ Sarcoma Classifier v12.2
TABLE S6 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypermethylated in AFH of extracranial soft tissue versus Group A intracranial mesenchymal tumors with FET-CREB fusion
TABLE S7 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypomethylated in AFH of extracranial soft tissue versus Group A intracranial mesenchymal tumors with FET-CREB fusion
TABLE S8 Gene Ontology analysis ranked by differentially methylated genes in angiomatoid fibrous histiocytoma (AFH) of extracranial soft tissue versus epigenetic Group A of intracranial mesenchymal tumors with FET-CREB fusion
TABLE S9 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypermethylated in clear cell sarcoma of soft tissue (CCS) versus Group B intracranial mesenchymal tumors with FET-CREB fusion
TABLE S10 Differentially methylated regions ranked by mean beta-value fold change for those regions which are hypomethylated in clear cell sarcoma of soft tissue (CCS) versus Group B intracranial mesenchymal tumors with FET-CREB fusion
TABLE S11 Gene Ontology analysis ranked by differentially methylated genes in clear cell sarcoma of soft tissue (CCS) versus epigenetic Group B of intracranial mesenchymal tumors with FET-CREB fusion

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