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Comprehensive genomic profiling of *EWSR1/FUS-CREB* translocation-associated tumors uncovers prognostically significant recurrent genetic alterations and methylation-transcriptional correlates

Josephine K. Dermawan, MD, PhD¹, Fabio Vanoli, PhD¹, Laurie Herviou, PhD¹, Yun-Shao Sung, MSc¹, Lei Zhang, MD¹, Samuel Singer, MD², William D. Tap, MD³, Ryma Benayed, PhD¹, Tejus A. Bale, MD¹, Jamal K. Benhamida, MD¹, Brendan C. Dickson, MD⁴, Cristina R. Antonescu, MD¹

¹Department of Pathology, Memorial Sloan Kettering Cancer Center, New York, NY, USA.

²Department of Surgery, Memorial Sloan Kettering Cancer Center, New York, NY, USA.

³Department of Medicine, Sarcoma Service, Memorial Sloan Kettering Cancer Center, New York, NY, USA.

⁴Department of Pathology and Laboratory Medicine, Sinai Health System, Toronto, Ontario, Canada

Abstract

To elucidate the mechanisms underlying the divergent clinicopathologic spectrum of *EWSR1/FUS-CREB* translocation-associated tumors, we performed a comprehensive genomic analysis of fusion transcript variants, recurrent genetic alterations (mutations, copy number alterations), gene expression, and methylation profiles across a large cohort of tumor types. The distribution of the *EWSR1/FUS* fusion partners – *ATF1*, *CREB1*, and *CREM* – and exon involvement was significantly different across different tumor types. Our targeted sequencing showed that secondary genetic events are associated with tumor type rather than fusion type. Of the 39 cases that underwent targeted NGS testing, 18 (46%) had secondary OncoKB mutations or copy number alterations (29 secondary genetic events in total), of which 15 (52%) were recurrent. Secondary recurrent, but mutually exclusive, *TERT* promoter and *CDKN2A* mutations were identified only in clear cell sarcoma (CCS) and associated with worse overall survival. *CDKN2A/B* homozygous

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Corresponding author: Cristina R. Antonescu, MD, Department of Pathology, Memorial Sloan Kettering Cancer Center, 1275 York Avenue, New York, NY 10065, antonesc@mskcc.org.

Author Contribution Statement

JKD performed study design, data acquisition, data analysis and interpretation, writing and revision of the paper. FV, LH, YS, LZ, SS, WT, RB, TB, JB, BD performed acquisition, analysis, and/or interpretation of data, and review of the paper. CRA performed study design and conception, analysis and interpretation of data, writing, review and revision of paper. All authors read and approved the final manuscript.

Ethics Approval / Consent to Participate

This study was approved by the Memorial Sloan Kettering Cancer Institute Institutional Review Board.

Conflict of Interest Statement

All authors report no conflict of interests related to this study.

deletions were recurrent in angiomatoid fibrous histiocytoma (AFH) and restricted to metastatic cases. mRNA upregulation of *MITF*, *CDH19*, *PARVB*, and *PFKP* was found in CCS, compared to AFH, and correlated with a hypomethylated profile. In contrast, *S100A4* and *XAF1* were differentially upregulated and hypomethylated in AFH but not CCS. A sarcoma methylation classifier was able to accurately match 100% of CCS cases to the correct methylation class; however, it was suboptimal when applied to other histologies. In conclusion, our comprehensive genomic profiling of *EWSR1/FUS-CREB* translocation-associated tumors uncovered mostly histotype, rather than fusion-type associated correlations in transcript variants, prognostically significant secondary genetic alterations, and gene expression and methylation patterns.

Keywords

CREB; *ATF1*; secondary recurrent genetic alterations; gene expression; methylation; fusion transcripts

Introduction

Recurrent gene fusions involving *EWSR1/FUS* with members of the cAMP response element binding protein (CREB) family (*ATF1*, *CREB1* and *CREM*) are shared amongst multiple tumor-types spanning a wide clinicopathologic spectrum. Despite sharing related gene fusions, members of the *EWSR1-CREB* family of translocation-associated tumors exhibit significantly different clinicopathologic characteristics. The prototypic example is angiomatoid fibrous histiocytoma (AFH) vs clear cell sarcoma (CCS)—two morphologically distinct tumors, the former mostly associated with a benign behavior, while the latter being an aggressive sarcoma with a high metastatic potential and poor outcome, as illustrated by the survival analysis of our cohort. Clear cell sarcoma-like tumor of gastrointestinal tract (GICCS, also known as gastrointestinal neuroectodermal tumor) and hyalinizing clear cell carcinoma of salivary gland (HCCC) had intermediate overall survival relative to AFH and CCS.

To elucidate the molecular mechanisms underlying their differences, we performed a comprehensive genomic analysis of fusion transcript variants, secondary recurrent genetic alterations (mutations, copy number alterations), gene expression and methylation profiles across a large cohort of tumour-types defined by *EWSR1/FUS-CREB* gene fusions. Specifically, the tumors included in this study encompassed AFH, CCS, GICCS, HCCC, clear cell odontogenic carcinoma (CCOC), malignant epithelioid neoplasm with predilection for mesothelial-lined cavities (ME), mesothelioma (Meso), myxoid mesenchymal tumor (MMT), and primary pulmonary myxoid sarcoma (PPMS).

Materials and Methods

Case selection and study cohort

After approval from the Institutional Review Board, cases were identified from the Memorial Sloan Kettering Cancer Center (MSKCC) surgical pathology archives, or from collaborating institutions, based on tumor types and/or presence of *EWSR1/FUS-ATF1*

CREB1/CREM fusions. The diagnosis of all 137 cases included molecular confirmation of both fusion partners: 37 cases by fluorescence in situ hybridization, 24 cases by reverse transcription PCR (RT-PCR), 29 cases by Memorial Sloan Kettering-Integrated Mutation Profiling of Actionable Cancer Targets (MSK-IMPACT) only, 10 cases by MSK-Fusion only, 10 cases by both MSK-IMPACT and MSK-Fusion, 21 cases by TruSight RNA Fusion Panel (Illumina, San Diego, CA), with the remaining cases based on NGS testing performed by referring institutions. The meta-analysis of the published literature was based on an exhaustive literature search on any fusions or gene rearrangements reported in all of the listed entities in Supplementary Table 1 that we could identify on PubMed.

DNA Seq and RNA Seq

Detailed descriptions of MSK-IMPACT workflow and data analysis, a hybridization capture-based targeted DNA NGS assay for solid tumor, and MSK-Fusion, an amplicon-based targeted RNA NGS assay using the Archer™ FusionPlex™ standard protocol, were described previously^{1,2}.

850k methylation array

Details of the methylation array protocol were described previously³. Briefly, for each sample, 250 ng of input DNA was used for bisulfite conversion (EZ DNA Methylation Kit; Zymo Research; catalog number D5002), followed by an FFPE restoration step using the Infinium HD FFPE DNA Restore Kit (Illumina; catalog number WG-321-1002). All samples were processed on the Infinium methylationEPIC 850k BeadChip array and scanned using the Illumina iScan. Each CpG site interrogated by the Infinium array was identified by a unique cg identifier in the format of cg#, where # is a number. The methylation level for each CpG site was quantified using β values (continuous values between zero and one), calculated as the ratio of methylated signal/total signal plus an offset. 850k methylation array profiling was performed in a total of 80 samples, including: 7 AFH, 4 CCS, 8 GICCS, and compared to 51 soft tissue tumors of various histotypes (4 angiosarcomas, 27 gastrointestinal stromal tumors, 1 HCCC, 1 ME, 3 Meso, 11 paragangliomas, 4 small blue round cell tumors) and 10 normal tissues (8 human peripheral white blood cell samples, 2 normal adrenal medulla). A minimum cutoff of \log_2FC (fold change) > 1.0 and $FDR < 0.01$ was used for statistical analysis of differential methylation analysis using t-test, focusing on comparison of the 7 AFH against all other samples and 4 CCS against all other samples. Unsupervised hierarchical clustering was performed using the pheatmap R package version 1.0.12 with Ward's linkage and Euclidean distance for clustering.

Sarcoma classification by DNA methylation profiling

Details of the DNA methylation-based machine learning sarcoma classification algorithm were described in Koelsche et al⁴. This random forest-based machine learning algorithm was developed at the German Cancer Research Center (DKFZ) in Heidelberg, Germany. Briefly, the method defines 62 methylation classes based on a reference cohort of 1077 samples encompassing a broad range of sarcomas. The classifier quantifies the confidence of the sample's assigned methylation class using a calibrated score between 0 and 1. The sum of all calibrated scores across all methylation classes is 1.0. A confident match is generally

considered > 0.9 and a poor match $< 0.5^4$. The 22 cases that underwent analysis with the DNA methylation classification algorithm corresponded to 6 AFH, 4 CCS, 8 GICCS, 1 ME, 3 Meso among the 80 samples that underwent methylation profiling.

Affymetrix microarray gene expression analysis

Details of the microarray protocol were described previously^{5,6}. RNA was isolated using RNawiz RNA isolation reagent (Ambion) and run through a column with RNase-free DNase (Qiagen). Ten micrograms of labeled and fragmented cRNA were then hybridized onto a Human Genome U133A expression array (Affymetrix, Santa Clara, CA). Post-hybridization staining, washing, and scanning were done according to instructions from the manufacturer (Affymetrix). The raw expression data were derived using the Affymetrix Microarray Analysis 5.0 (MAS 5.0) software. The data were normalized using a scaling target intensity of 500 to account for differences in the global chip intensity. The expression values were transformed using the logarithm base two. Affymetrix U133A gene expression array analysis was performed in a total of 58 samples, including 3 AFH, 4 CCS, 1 GICCS, and compared to 44 soft tissue tumors of various histotypes (3 adult fibrosarcomas, 5 angiosarcomas, 3 leiomyosarcomas, 10 gastrointestinal stromal tumors, 3 myxoid liposarcomas, 6 paragangliomas, 4 small blue round cell tumors, 4 solitary fibrous tumors, 3 synovial sarcoma, 3 undifferentiated pleomorphic sarcomas) and 6 normal tissues (adrenal gland, brain, kidney, small intestine, stomach, testis). For differential gene expression analysis, a minimum cutoff of \log_2FC (fold change) > 1.0 and FDR adjusted p-value < 0.01 were used for t-test. We compared one histotype against all other tumors for each respective analysis. For example: CCS (4 cases) vs all others (54 cases) in one analysis, and AFH (3 cases) vs all others (55 cases) in a different analysis. Unsupervised hierarchical clustering was performed using the pheatmap R package version 1.0.12 with Ward's linkage and Euclidean distance for clustering.

Integration of gene expression and methylation analysis

First, we performed differential gene expression and differential methylation analysis by setting a false discovery rate (FDR) adjusted p-value of 0.01 and a minimum \log_2FC (fold change) of 1.0 for t-test, comparing one histotype against all other tumors each time (e.g., CCS vs all others, AFH vs all others). Thereafter, for integration of transcriptomic and methylation data, we matched all the genes that were both differentially expressed based on $\log_2FC > 1.0$ and $FDR < 0.01$ and differentially methylated based on $\log_2FC > 2.0$ and $FDR < 0.01$ for the CCS vs all others and AFH vs all others comparisons. Out of the 3 AFH, 4 CCS and 1 GICCS on the Affymetrix U133 microarray, 1 AFH and 2 CCS did not overlap with the samples used for the methylation array.

Results

Clinicopathologic summary

A total of 137 cases were identified [76 females, 61 males, mean age 37 (range 2–86)], including: 40 CCS (29%), 36 AFH (26%), 20 GICCS (15%), 14 ME (10%), 10 HCCC (7%), 8 Meso (6%), 5 MMT (4%), 3 PPMS (2%), and 1 clear cell odontogenic carcinoma (CCOC) (1%) (Figure 1A). The mean ages in HCCC, PPMS and CCOC were higher than

those in AFH, CCS, GICCS, MMT, and ME (Table 1). As expected, the primary sites were predominantly soft tissue for AFH and CCS, gastrointestinal tract/pelvis for GICCS, brain for MMT, thoracic or abdominopelvic cavities for Meso and ME, lung for PPMS, and major and minor salivary glands for HCCC.

Fusion types and transcript variants by diagnosis

The distribution of the *EWSR1/FUS* fusion partners, *ATF1*, *CREB1*, and *CREM*, was significantly different across different tumor types (chi-square $P < 0.0001$) (Figure 1B). Specifically, *EWSR1-ATF1* fusions were the only fusion type in HCCC (100%) and the predominant fusion type in CCS (85%) and Meso (88%); *EWSR1-CREB1* fusions were the only fusion type in PPMS (100%) and the predominant fusion type in AFH (60%); *EWSR1/FUS-CREM* fusions were the predominant fusion type in ME (86%) and MMT (50%). *ATF1* and *CREB1* fusions were equally distributed in GICCS. The single case of CCOC had a *EWSR1-ATF1* fusion. Of the 137 cases, only 5 (4%) harbored *FUS* fusions: four were *FUS-CREM* fusions in ME, one was a *FUS-ATF1* fusion in a Meso.

The exon usage for the fusion transcript variants for each tumor type was derived from either MSK-Fusion and/or MSK-IMPACT testing and available in 48 cases (8 AFH, 18 CCS, 9 GICCS, 7 HCCC, 3 Meso, 1 PPMS, 2 ME) (Table 2). The predominant fusion transcript variants were *EWSR1*ex8-*ATF1*ex4 in CCS and GICCS; *EWSR1*ex7-*ATF1*ex5 and *EWSR1*ex7-*CREB1*ex7 in AFH; *EWSR1*ex11-*ATF1*ex3 in HCCC; *FUS*ex8-*CREM*ex5/7 for ME (Figure 2). Supplementary Table 1 summarizes the CREB family fusion variants of various tumor types derived from our meta-analysis of published studies in comparison to those detected in the current cohort.

Clinically significant recurrent genetic alterations

39 cases [6 AFH, 14 CCS, 9 GICCS, 5 HCCC, 3 Meso, 1 PMMS, 1 clear cell odontogenic carcinoma (CCOS)] were analyzed by MSK-IMPACT. Only clinically significant variants with OncoKB annotations (Chakravarty 2017) (or known recurrent hotspots) and secondary recurrent genetic alterations (events that occur > 1 in our cohort) were included. Variants of unknown significance were excluded. Of the 39 cases that underwent targeted NGS testing, 18 (46%) had OncoKB mutations or copy number alterations (29 secondary genetic events in total), of which 15 (52%) were recurrent. Specifically, *TERT* promoter hotspot mutations (n=5) and *CDKN2A* X51_splice and P81Lfs*30 mutations (n=2) were mutually exclusive and identified in CCS only. Other secondary recurrent genetic alterations identified were: *TP53* R248Q and T155Pfs*15 mutations (n=2, 1 CCS, 1 GICCS), 9p21.3 (*CDKN2A/CDKN2B*) copy number loss (homozygous deletion) (n=4, 2 AFH, 1 CCS, 1 HCCC), and *DIS3* D479G and D488N mutations (n=2, both GICCS) (Figure 3A). No secondary recurrent genetic alterations were identified in any of the 3 Meso, 1 PMMS, or 1 CCOC. The type of secondary recurrent genetic alterations did not correlate with the *EWSR1/FUS* fusion partner type (Supplementary Figure 1).

Interestingly, AFH cases with *CDKN2A/CDKN2B* homozygous deletion (n=2, 33%) were exclusively found in metastatic cases, whereas the remaining *CDKN2A/CDKN2B* non-altered AFH cases were non-metastatic (Figure 3B). On the other hand, CCS cases with

TERT promoter mutations and *CDKN2A* loss-of-function mutations (frameshift and splice site mutations) (n=7, 50%) were significantly correlated with decreased overall survival (Mantel Haenszel chi-square $P=0.0196$) (Figure 3C), with a median survival of 5.13 vs 22.85 months in non-altered CCS cases (n=7, 50%). The presence of *DIS3* mutations were not correlated with metastatic nor survival status in GICCS.

Methylation and gene expression correlation

Gene expression profiling were performed on the Affymetrix U133A expression array comparing 3 AFH, 4 CCS cases, and 1 GICCS to a group of 44 soft tissue tumors of various histotypes and 6 normal tissue samples (Supplementary Figure 2 and Supplementary Table 2). Methylation profile testing was performed comparing 7 AFH, 4 CCS, 4 GICCS to a group of 51 soft tissue tumors of various histotypes and 10 normal tissue samples on the 850k methylation array (Supplementary Figure 3 and Supplementary Table 3). The goal is to identify correlates between differential gene expression ($1.5 \log_2$ FC, FDR 0.01) and differential methylation ($4 \log_2$ FC and FDR 0.01) for *EWSR1-ATF1*-rearranged CCS and *EWSR1-CREB1*-rearranged AFH, respectively, in relation to other tumor types. Gene expression profiling revealed upregulation of *PMP22*, *MITF*, *SLC7A5*, *CDH19*, *WIP1*, *FYN*, *PARVB*, and *PFKP* in CCS but not AFH, and upregulation of *SGK1*, *S100A4*, *XAF1* and *LY96* expression in AFH but not CCS. However, despite differential gene expression, CREB family translocation tumors mostly cluster together in terms of methylation profile in relation to other tumor types (Supplementary Figure 3 and Supplementary Table 3).

Thereafter, differentially expressed genes were matched to CpG sites based on chromosomal locations. We matched all the genes that were both differentially expressed based on \log_2 FC > 1.0 and FDR < 0.01 and differentially methylated based on \log_2 FC > 2.0 and FDR < 0.01. We focused on upregulated genes with corresponding hypomethylation. Our analyses revealed genes (*MITF*, *CDH19*, *PARVB*, and *PFKP*) with increased expression and hypomethylation in CCS but not AFH (Figure 4A), and genes (*S100A4*, *XAF1*) with increased expression and hypomethylation in AFH but not CCS (Figure 4B). *MITF* is involved in melanogenesis and overexpressed in CCS as part of its core gene signature^{5,7}. *CDH19* and *PARVB* are involved in cell adhesion and were highly expressed in primary melanoma⁸. *S100A4* has been implicated in cell migration and cancer metastases⁹. *XAF1* is a proapoptotic tumor suppressor gene¹⁰.

Tumor type prediction by the Sarcoma Methylation Classifier

The DNA-methylation based sarcoma classification algorithm described in Koelsche et al⁴ was applied to 22 cases (6 AFH, 4 CCS, 8 GICCS, 1 ME, and 3 Meso) (Table 3). This algorithm was able to accurately match 100% of four CCS cases to the correct methylation class (calibrated score = 0.99 in all cases), but only 33% (2 of 6) of AFH cases (calibrated score = 0.75 and 0.33, respectively). GICCS was not a methylation class in the original classifier. Interestingly, the algorithm matched 1 GICCS to AFH (calibrated score 0.56) and 2 GICCS to CCS (calibrated score = 0.65 and 0.96, respectively).

Survival analysis

The overall survival across AFH, CCS, GICCS, HCCC was significantly different (log rank $P=0.023$), with CCS associated with the worse survival (median survival 15 months), followed by HCCC (median survival 36 months) and then GICCS (median survival 43 months). All AFH patients remained alive across the follow-up period of 42 months (Figure 5).

Discussion

The *EWSRI*-CREB family of translocation-associated tumors encompasses a wide and heterogeneous clinicopathologic spectrum. To understand the pathogenesis that sets them apart, we performed comprehensive genomic analysis of fusion variants, secondary recurrent genetic alterations (mutations, copy number alterations), gene expression and methylation profiles across a large cohort of *EWSRI*-CREB family of translocation-associated tumors, with emphasis on AFH, CCS, GICCS, and HCCC.

Although the analysis of fusion transcript variants in our cohort largely paralleled the published literature, some new interesting findings emerged. For AFH, the most common reported fusion transcript variant is *EWSRI-CREB1* (ex7-ex7) (58%)^{7,11–15}. However, we identified a significant proportion (39%) of AFH cases with *EWSRI-ATF1* (largely ex7-ex5). A minority (3%) of AFH harbored *EWSRI-CREM* fusions. Interestingly, MMT^{16–21}, which remains disputed by some authors to be related to a myxoid, intracranial variant of AFH^{22–27}, harbor roughly equal proportions of *EWSRI-ATF1* and *EWSRI-CREB1* fusions, with a significant minority harboring *EWSRI-CREM*. For CCS, the predominant fusion transcript is *EWSRI*ex8-*ATF1*ex4^{15,28–34}. This pattern is mirrored by a subtype of Meso, initially described by our group and occurs in younger patients without asbestos exposure history, which are driven predominantly by *EWSRI-ATF1*ex5^{35–37}. Of interest, in contrast to prior data, GICCS showed similar proportions of *EWSRI-ATF1* (mostly ex8-ex4) and *EWSRI-CREB1* fusions^{5,34,38}. On the other hand, the recently described distinct tumor type, so-called ‘malignant epithelioid neoplasm with predilection for mesothelial-lined cavities’³⁹ and subsequently validated by Shibayama et al⁴⁰, most commonly harbor either fusions between *EWSRI* or *FUS* and exon 7 of *CREM*. In contrast, PPMS is almost exclusively driven by *EWSRI-CREB1* (mostly ex7-ex7)^{41–48} except for a rare case with *EWSRI-ATF1*⁴⁹. Some authors proposed that PPMS and AFH exist on a morphologic and molecular spectrum^{43,49}. Finally, both HCCC^{50–55} and CCOC^{56–60} harbor only *EWSRI-ATF1*, supporting the notion that HCCC and CCOC are likely related tumors. Nevertheless, it is evident from our meta-analysis of the published literature and from the current study that there is significant intertumoral overlap as well as intratumoral heterogeneity of fusion transcript variants and exon usage across the different CREB family translocation tumors. Here, intratumoral heterogeneity refers to variation of fusion transcript variants, e.g., *EWSRI-ATF1* and *EWSRI-CREB1* in GICCS, and exon usage within the same histotype, e.g., *FUS-CREM* exon 5 and *FUS-CREM* exon 7 in ME.

This is the first study to report secondary recurrent genomic alterations in CCS, AFH and GICCS. In CCS, we identified the presence of recurrent *TERT* promoter and *CDKN2A* hotspot mutations, which were mutually exclusive but in combination strongly associated

with worse overall survival. *TERT* promoter somatic mutations and amplifications are frequently found across multiple tumor types^{61,62}. In soft tissue tumors, *TERT* promoter mutations have been identified in myxoid liposarcomas⁶³, atypical fibroxanthoma/pleomorphic dermal sarcomas⁶⁴, chondrosarcomas⁶⁵, and solitary fibrous tumors⁶⁶; this is reported to be associated with a worse prognosis in the latter. Our findings suggest that *TERT* promoter and *CDKN2A* mutations may serve as prognostic biomarkers for worse survival in CCS.

In AFH, we identified *CDKN2A/CDKN2B* homozygous deletions exclusively amongst cases with metastasis. Genomic profiling of multiple sarcoma types has revealed secondary recurrent *CDKN2A* alterations^{67,68}, with a role suggested as a biomarker for poor prognosis⁶⁷. Compared to other CREB family translocation tumors, AFH is a soft tissue tumor of borderline malignant potential and a relatively good prognosis; metastasis is usually < 2%. In fact, all the patients whose AFH were sequenced in our cohort remain alive at the time of reporting. Our finding of *CDKN2A/CDKN2B* deletions in the two AFH cases with biopsy-proven metastasis, but not in the non-metastatic AFH cases, raises the possibility of *CDKN2A/CDKN2B* deletion testing as a biomarker for metastatic potential. Although not a recurrent abnormality in this cohort, one of the metastatic AFH case showed a co-existing *BRAF*V600E mutation which was confirmed by immunohistochemistry to have diffuse and strong VE1 expression. This was the only case in our cohort to have a *BRAF* mutation detected.

Gene expression profiling revealed differential gene expression in AFH vs CCS, which clustered in distinct genomic groups by unsupervised analysis. A number of genes involved in melanocyte regulation and cellular membrane/migration were upregulated in CCS compared to AFH, including *PMP22*, *MITF*, *SLC7A5*, *CDH19*, *WIPI1*, *FYN*, *PARVB*, and *PFKP*, while upregulation of *SGK1*, *S100A4*, *XAF1* and *LY96* mRNA expression was detected in AFH but not CCS. An expression profiling analysis of CCS cell lines revealed upregulation of *S100A11* (encoding for S100 protein), *MITF* (microphthalmia-associated transcription factor), and *Pmel17* (*SILV*) (silver mouse homologue-like melanosomal protein detected by the IHC marker HMB45)⁶⁹. Moreover, in an *in vitro* CCS induced pluripotent stem cell model, Komura et al reported expression of several Schwann cell marker genes, such as *P75^{NTR}*, *S100b*, *Mbp*, *Plp1*, and *Pmp22*⁷⁰. They proposed that S100-expressing peripheral nerve cells could be a cell of origin for *EWS/ATF1*-induced CCS. In a recent study using human embryonic stem (hES) cell models, hES cells driven by *EWSR1-CREB1* and *EWSR1-ATF1* fusions recapitulate the core gene signatures, respectively, of AFH (*SGK1* and *MXRA5* upregulation) and GICCS (*SGK1*, *MXRA5*, *SOX10*, and *DUSP4* upregulation)⁷¹. Our gene expression profiling of patient samples validates a subset of the findings of these preclinical studies.

Our methylation analysis did not reveal significant differential methylation profiles among different tumor types within the CREB family translocation-associated tumors. Rather, these tumors clustered together and displayed methylation profiles distinct from other soft tissue tumor types and normal tissue. However, when matching the differentially expressed genes to the corresponding methylation probes/CpG sites, we found significant correlations between upregulated genes that were hypomethylated in CCS but not AFH.

These genes included *MITF*, *CDH19*, *PARVB*, and *PFKP* in CCS. *MITF* is involved in melanogenesis and found to be overexpressed in CCS as part of its core gene signature, but not in AFH or GICCS^{5,7}. More recently, a Cre-loxP-induced *Ewsr1-Atf1*-driven CCS model demonstrated that *Mitf* and *Myc* can contribute to sarcomagenesis⁷². Both *CDH19* and *PARVB* are involved in cell adhesion and were highly expressed in primary melanoma, associated with worse survival⁸. It is interesting how they were found to show increased expression and hypomethylation in CCS in our study. On the other hand, we identified upregulation and hypomethylation of *S100A4* and *XAF1* in AFH but not CCS. *S100A4* protein is a member of the S100 calcium binding protein family, also known as metastasin, and has been implicated in cell migration and cancer metastases⁹. *XAF1* is a proapoptotic, interferon-stimulated tumor suppressor gene that suppresses tumorigenesis¹⁰. While *XAF1* is usually hypermethylated and downregulated in most cancers, it was found to be paradoxically hypomethylated in glioblastoma with adaptive temozolomide resistance⁷³. These findings serve as a proof-of-concept example of how integrative gene expression and methylation profiling may provide interesting biological insights into the different pathogenesis underlying tumors sharing the same driver gene fusions. Integrated DNA methylation and gene expression studies have been performed in Ewing sarcoma⁷⁴, pediatric rhabdomyosarcomas⁷⁵, myxoid, dedifferentiated, and pleomorphic liposarcomas⁷⁶, which identified sets of genes with inverse methylation and gene expression relationship. In a comprehensive molecular and genomic study of undifferentiated sarcomas (USARC), DNA methylation profiling failed to identify distinct USARC subgroups and did not correlate with gene expression, but showed *MSH2* and *TERT* promoter hypermethylation⁷⁷. On the other hand, DNA methylation profiling also revealed epigenetic heterogeneity within the same tumor type, e.g., Ewing sarcoma⁷⁸. Unfortunately, the sample size of individual tumor types used for methylation profiling in the current study is insufficient to perform differential methylation analysis within the same tumor type, which could be explored in future studies.

Genome-wide DNA methylation profiling has largely been performed for tumor classification purposes in a wide range of mesenchymal tumors, with varying degree of success. These include: benign and malignant nerve sheath tumors⁷⁹, osteosarcomas⁸⁰, undifferentiated small round blue cell tumors⁸¹, *CIC*-rearranged undifferentiated sarcomas⁸². Most recently, a Random Forest machine learning sarcoma classifier from the German Cancer Research Center (DKFZ) in Heidelberg were developed to classify a wide spectrum of 66 soft tissue and bone tumors using a large reference and validation cohort⁴. The limitations of using methylation profiling alone to differentiate soft tissue tumors were illustrated by the inability of the Heidelberg methylation classifier to accurately classify tumor entities in our cohort, with the exception of CCS. There are several major shortcomings to the applicability of this methylation classifier for soft tissue tumors with *EWSR1/FUS* fusions with CREB family transcription factors. First, several tumor types, including GICCS and Meso, were not included in the reference cohort that was used to develop the classifier. Second, although the reference cohort included 8 cases of AFH, only 1 case was used in the validation cohort, which was misclassified as desmoplastic small round cell tumor⁴. In our study, the classifier was able to correctly classify one-third of the AFH cases. On the other hand, the methylation classifier performed very well, both in the Koelsche et al study and in our experience, in the classification of CCS: classifying 100%

of the cases correctly. It is also interesting that when we applied the classifier to GICCS, 2 cases were classified as CCS and 1 case as AFH, illustrating their overlapping methylation profiling as described above. All 3 of these GICCS cases were located in the gastrointestinal tract (1 stomach, 2 small bowel), and were diffusely and strongly positive for S100 and negative for HMB45. The combined clinical and immunohistochemical profile essentially excludes CCS and AFH. These findings highlight the existing limitation of methylation profiling in soft tissue tumor classification, which may require further algorithm refinement as well as larger reference and validation cohorts⁸³.

In addition to these molecular mechanisms, the nature of the initial stem cell host in which the fusion, and its degree of commitment / plasticity, arose may also play a significant role in ultimate tumor type (i.e., depending on location / extent of totipotency). These are interesting questions that are beyond the scope of the current study. Recent studies using *Cre* inducible mouse and human embryonic stem cell models have begun to address these questions^{71,72}.

The lack of consistency in the sample sizes of the cases with each technique is a major drawback of our paper. Further studies focusing on specific molecular profiling techniques with deeper genomic characterization utilizing a larger sample size of some of the rarer histotypes would be beneficial to validate or expand on our findings.

In conclusion, our comprehensive genomic profiling of *EWSR1/FUS-CREB* translocation-associated tumors uncover fusion transcript variant heterogeneity, prognostically significant secondary recurrent genetic alterations, and differentially hypomethylated and upregulated genes. These findings underscore the utility of integrative genomic approaches in the study of translocation-associated tumors with diverse clinicopathologic features, and whether some of the entities in this family could be unified under the same morphologic/molecular spectrum (e.g., CCS and GICCS, AFH and PPMS).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data Availability Statement

The datasets generated or analyzed during this study are included in this published article [and its supplementary information files]. The raw Affymetrix and Illumina 850k

methylation array data generated are not publicly available due to lack of access to indefinite hosting capabilities, but are available from the corresponding author on reasonable request.

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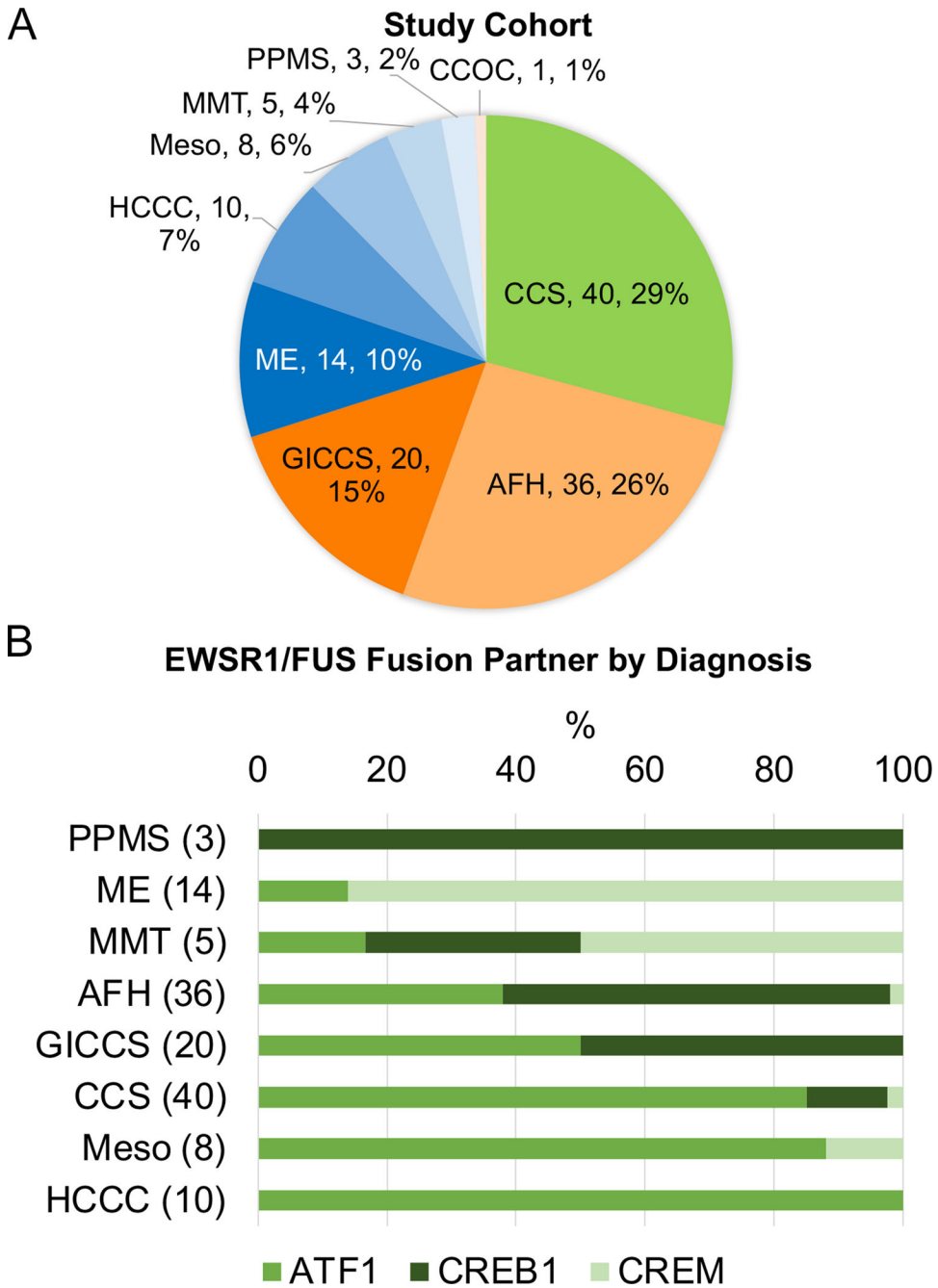


Figure 1. **A**, Study cohort showing number of cases and percentages by tumor type. **B**, Distribution of *EWSR1/FUS* fusion partners (*ATF1*, *CREB1*, *CREM*) by diagnosis (number of cases for each histotype indicated between parenthesis). Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; CCOS: clear cell odontogenic carcinoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract; HCCC: hyalinizing clear cell carcinoma of salivary gland; ME: malignant epithelioid neoplasm with predilection for mesothelial-lined cavities; Meso: mesothelioma; PPMS: primary pulmonary myxoid sarcoma.

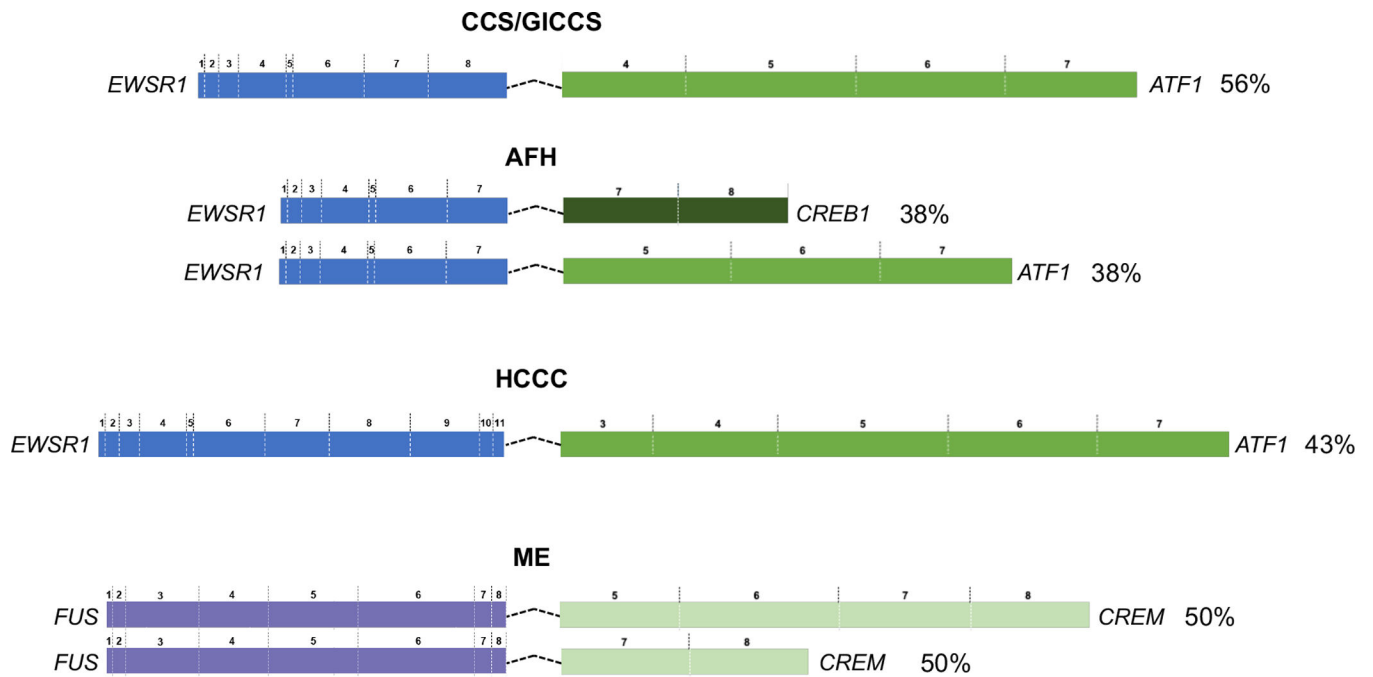


Figure 2. Schematics of predominant fusion transcript variants, for AFH, CCS, GICCS, HCCC, and ME. Exon numbers were based on canonical transcripts for each gene. Percentage indicates frequency of the fusion transcript variant within the corresponding histotype subgroup. RefSeq accession number: *ATF1* (NM_005171); *CREB1* (NM_134442); *CREM* (NM_181571); *EWSR1* (NM_005432); *FUS* (NM_004960). Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract; HCCC: hyalinizing clear cell carcinoma of salivary gland; ME: malignant epithelioid neoplasm with predilection for mesothelial-lined cavities; Meso: mesothelioma.

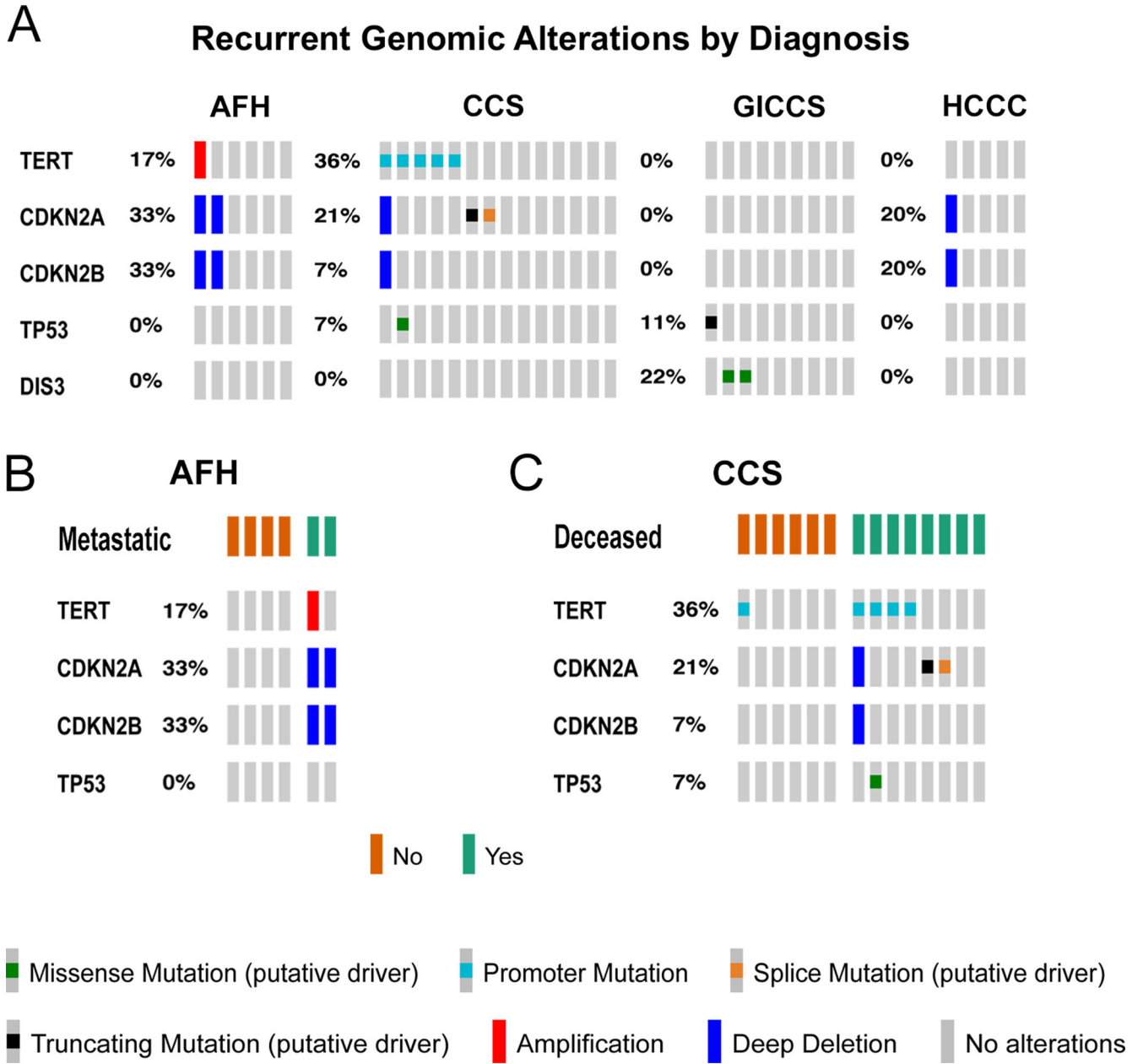


Figure 3.
A, Recurrent genomic alterations identified by MSK-IMPACT, including OncoKB mutations and copy number alterations⁸⁴, in 6 AFH, 14 CCS, 9 GICCS, and 5 HCCC. Only genomic alteration events occurring > 1 were included. **B**, Presence of *TERT*, *CDKN2A*, and *CDKN2B* alterations in AFH with or without metastatic disease. **C**, Presence of *TERT*, *CDKN2A*, *CDKN2B* and *TP53* alterations in living vs deceased CCS patients. Data generated from cBioPortal and visualized using OncoPrint⁸⁵. Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract; HCCC: hyalinizing clear cell carcinoma of salivary gland.

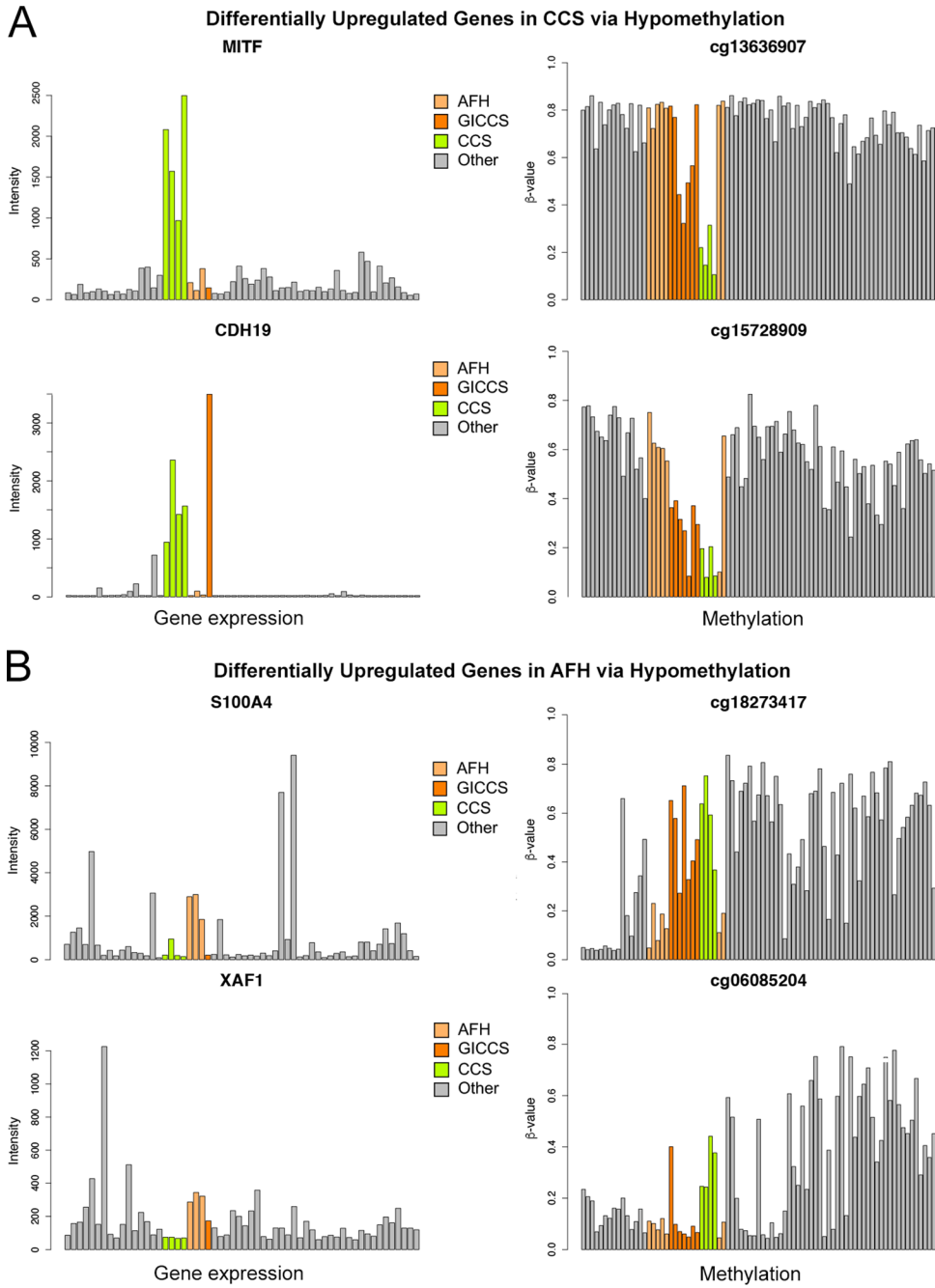
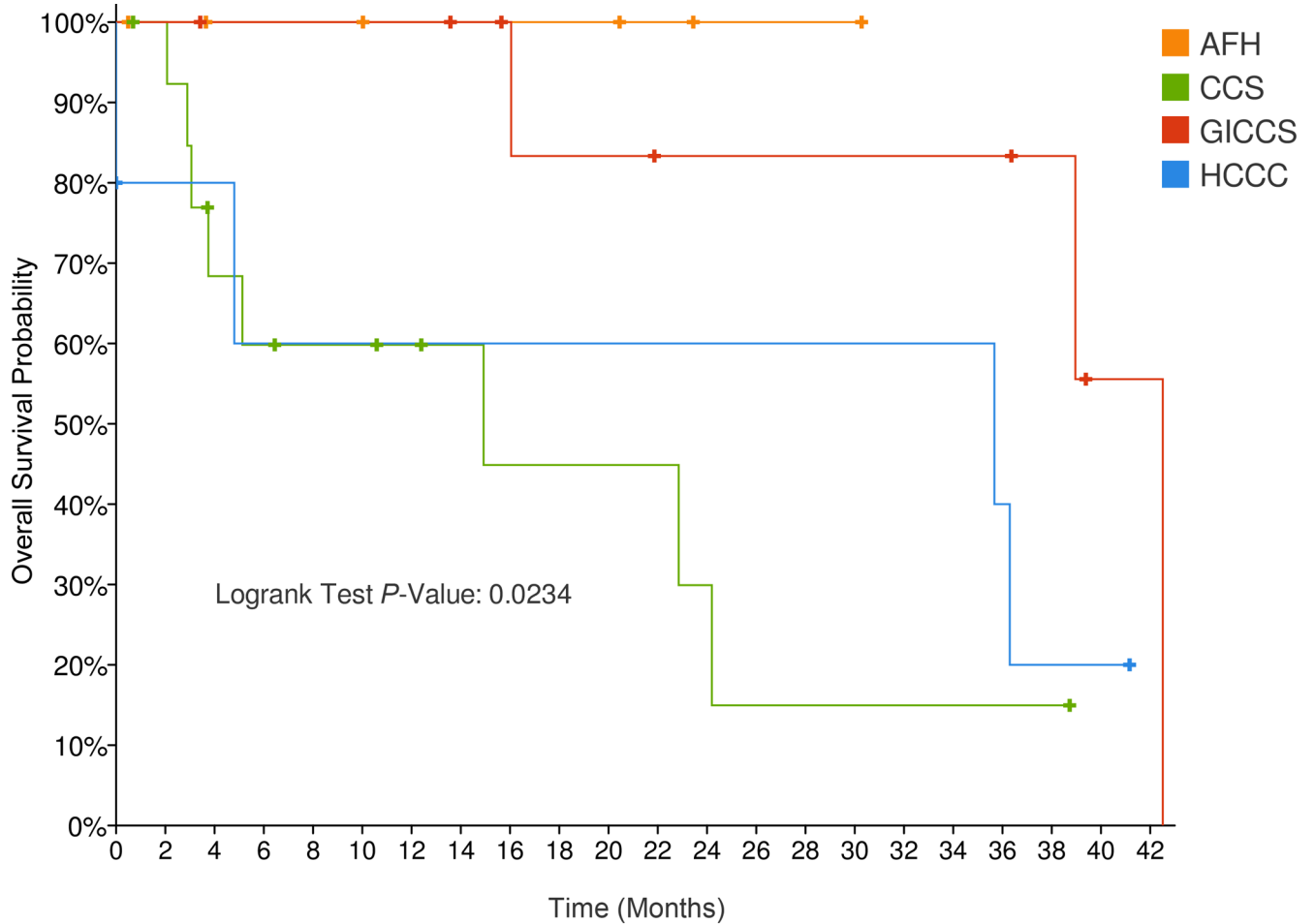


Figure 4. Differential gene upregulation corresponding to hypomethylation on matched CpG sites in CCS (A) and AFH (B). Affymetrix U133A was performed comparing 3 AFH, 4 CCS cases, and 1 GICCS to a group of 44 tumors of various histotypes and 6 normal tissues, using log₂-fold change threshold of 1 and *P* < 0.01. Infinium 850k methylation array was performed comparing 7 AFH, 4 CCS, 4 GICCS to a group of 29 tumors of various histotypes and 8 normal tissues, using a log₂-fold change threshold of 2 and *P* < 0.01. Differentially expressed genes were matched to CpG site identified by a unique cg identifier in the format of cg#. The numbers of CpG sites assigned to each of these 4 genes on the 850k array

were as follows: 8 for *S100A4*, 14 for *XAF1*, 27 for *MITF*, and 3 for *CDH19*. Out of these, the numbers of CpG sites that showed negative correlation with gene expression are as follows: 3 for *S100A4*, 6 for *XAF1*, 3 for *MITF*, and 3 for *CDH19*. The most representative CpG site from each gene is displayed in this figure. Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract.



	Number of Cases	Number of Events	Median Survival (Months)
AFH	6	0	N/A
CCS	14	8	14.93
GICCS	9	3	42.51
HCCC	5	4	35.67

Figure 5. Comparison of overall survival in 6 AFH, 14 CCS, 9 GICCS, and 5 HCCC. Median survival time (months) for each tumor type listed beneath Kaplan-Meier curves. Hazard ratios compared using log-rank analysis. Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract; HCCC: hyalinizing clear cell carcinoma of salivary gland.

Clinicopathologic summary of the study cohort

Table 1.

Diagnosis	Female, n (%)	Male, n (%)	Mean age in years (range)	Median tumor size in cm (range)	Top primary site	Metastasis, n (%)	Top metastatic site	Total
Angiomatoid fibrous histiocytoma	19 (53)	17 (47)	28.3 (2–79)	3.4 (1.0–10.5)	Lower extremities, upper extremities, head and neck, trunk	4 (11)	Lung, bone, adrenal	36
Clear cell odontogenic carcinoma	1 (100)	0	57.0	2.7	Head and neck	0	N/A	1
Clear cell sarcoma	16 (40)	24 (60)	32.9 (7–71)	4.7 (2.5–12.0)	Lower extremities, upper extremities	18 (45)	Lymph node, lung	40
Clear cell sarcoma-like tumor of the gastrointestinal tract	14 (70)	6 (30)	39.0 (18–76)	4.5 (2.5–6.0)	Gastrointestinal tract	8 (40)	Liver, abdominopelvic	20
Hyalinizing clear cell carcinoma of salivary gland	8 (80)	2 (20)	67.6 (55–86)	2.1 (1.3–3.3)	Salivary gland	3 (30)	Lymph node, trunk, abdominopelvic	10
Malignant epithelioid neoplasm with predilection for mesothelial-lined cavities	7 (50)	7 (50)	37.2 (9–63)	9.0 (4.5–15.0)	Abdominopelvic	3 (21)	Lymph node, liver abdominopelvic	14
Mesothelioma	5 (62)	3 (38)	39.0 (15–78)	3.8 (2.6–8.5)	Abdominopelvic, pleura	2 (25)	Lymph node, abdominopelvic	8
Myxoid mesenchymal tumor	3 (60)	2 (40)	27.0 (12–48)	2.9 (2.8–3.0)	Brain	0	N/A	5
Primary pulmonary myxoid sarcoma	3 (100)	0	59.0 (43–82)	1.4	Lung	1 (33)	Lymph node	3
Total	76	61	36.4 (5–86)	4.0		39 (28)		137

Table 2.

Distribution of the most prevalent fusion transcript variants by exon usage in the current study

Histotype	Fusion transcript variant	Number	Percentage
Angiomatoid fibrous histiocytoma (AFH)	<i>EWSR1-ATF1</i> ex7-ex5	3	37.5
	<i>EWSR1-ATF1</i> ex7-ex7	1	12.5
	<i>EWSR1-CREB1</i> ex7-ex7	3	37.5
Clear cell sarcoma (CCS)	<i>EWSR1-ATF1</i> ex7-ex4	1	5.6
	<i>EWSR1-ATF1</i> ex7-ex5	3	16.7
	<i>EWSR1-ATF1</i> ex8-ex4	10	55.6
	<i>EWSR1-ATF1</i> ex9-ex4	2	11.1
	<i>EWSR1-CREB1</i> ex7-ex7	1	5.6
	<i>EWSR1-CREM</i> ex8-ex7	1	5.6
Clear cell sarcoma-like tumor of the gastrointestinal tract (GICCS)	<i>EWSR1-ATF1</i> ex7-ex5	1	11.1
	<i>EWSR1-ATF1</i> ex8-ex4	5	55.6
	<i>EWSR1-CREB1</i> ex6-ex6	1	11.1
	<i>EWSR1-CREB1</i> ex7-ex6	1	11.1
	<i>EWSR1-CREB1</i> ex7-ex7	1	11.1
Hyalinizing clear cell carcinoma (HCCC)	<i>EWSR1-ATF1</i> ex7-ex4	1	14.3
	<i>EWSR1-ATF1</i> ex8-ex4	1	14.3
	<i>EWSR1-ATF1</i> ex9-ex2	1	14.3
	<i>EWSR1-ATF1</i> ex10-ex3	1	14.3
	<i>EWSR1-ATF1</i> ex11-ex3	3	42.9
Malignant epithelioid neoplasm with predilection for mesothelial-lined cavities (ME)	<i>FUS-CREM</i> ex8-ex5	1	50.0
	<i>FUS-CREM</i> ex8-ex7	1	50.0
Mesothelioma (Meso)	<i>EWSR1-ATF1</i> ex7-ex5	1	33.3
	<i>EWSR1-ATF1</i> ex13-ex5	1	33.3
	<i>EWSR1-CREM</i> ex10-ex5	1	33.3

Transcripts: *ATF1* (NM_005171); *CREB1* (NM_134442); *CREM* (NM_181571); *EWSR1* (NM_005243); *FUS* (NM_004960).

Table 3.Tumor type prediction by sarcoma methylation classifier^{*}

Diagnosis	Age/Sex	Site	Fusion	Calibrated score	Matching methylation class
AFH	21/M	groin	EWSR1-CREB1	0.33	angiomatoid fibrous histiocytoma
AFH	16/M	scalp	EWSR1-CREB1	0.75	angiomatoid fibrous histiocytoma
AFH	79/F	knee	EWSR1-ATF1	< 0.3	no matching methylation class
AFH	22/F	inguinal	EWSR1-CREB1	< 0.3	no matching methylation class
AFH	34/M	foot	EWSR1-ATF1	< 0.3	no matching methylation class
AFH	17/M	axilla	EWSR1-CREB1	0.74	squamous cell carcinoma (cutaneous)
CCS	20/F	heel	EWSR1-ATF1	0.99	clear cell sarcoma of soft parts
CCS	30/M	knee	EWSR1-ATF1	0.99	clear cell sarcoma of soft parts
CCS	24/M	arm	EWSR1-ATF1	0.99	clear cell sarcoma of soft parts
CCS	46/F	hip	EWSR1-ATF1	0.99	clear cell sarcoma of soft parts
GI CCS	42/F	small bowel	EWSR1-CREB1	0.56	angiomatoid fibrous histiocytoma
GI CCS	47/F	small bowel	EWSR1-CREB1	0.65	clear cell sarcoma of soft parts
GI CCS	57/M	stomach	EWSR1-CREB1	0.96	clear cell sarcoma of soft parts
GI CCS	42/F	small bowel	EWSR1-CREB1	0.51	neurofibroma (plexiform)
GI CCS	19/F	mesentery	EWSR1-ATF1	< 0.3	no matching methylation class
GI CCS	76/F	colon	EWSR1-CREB1	< 0.3	no matching methylation class
GI CCS	18/F	small bowel	EWSR1-ATF1	< 0.3	no matching methylation class
GI CCS	25/M	stomach	EWSR1-CREM	0.68	sclerosing epithelioid fibrosarcoma
ME	20/F	peri-rectal	EWSR1-CREM	< 0.3	no matching methylation class
Meso	26/F	peritoneal	FUS-ATF1	< 0.3	no matching methylation class
Meso	34/F	pleura	EWSR1-ATF1	< 0.3	no matching methylation class
Meso	79/F	mediastinum	EWSR1-ATF1	< 0.3	no matching methylation class

* Percentage matched: AFH 33.3%, CCS 100%, GICCS/ME/meso (methylation class doesn't exist in the classifier)

Abbreviations—AFH: angiomatoid fibrous histiocytoma; CCS: clear cell sarcoma; GICCS: clear cell sarcoma-like tumor of gastrointestinal tract; ME: malignant epithelioid neoplasm with predilection for mesothelial-lined cavities; Meso: mesothelioma.