Novel GATA6-FOXO1 Fusions in a Subset of Epithelioid Hemangioma

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Abstract

The genetic hallmark of epithelioid hemangioma (EH) is the presence of recurrent gene fusions involving FOS and FOSB transcription factors, which occur in one third of the cases. Certain clinical, pathologic and genotypic correlations have been described, with FOS-related fusions being more often detected in skeletal and cellular variants of EH, while FOSB gene rearrangements are more commonly associated with atypical histologic features and penile location. These fusions are infrequently detected in the cutaneous or head and neck EH. Overall, two-thirds of EH lack these canonical fusions and remain difficult to classify, especially when associated with atypical features and/or clinical presentations. Triggered by an index case of an intravascular soft tissue EH with a novel GATA6-FOXO1 gene fusion by targeted RNA sequencing (Archer® FusionPlex® Sarcoma Panel), we have investigated 27 additional EH cases negative for FOS and FOSB gene rearrangements for this novel abnormality to determine its recurrent potential and its association with clinical and pathologic features. Four additional EH cases were found to display GATA6-FOXO1 fusions (18%). There were 3 females and 2 males, with a mean age of 32 years old. Three lesions occurred in the head and neck (dura, nasopharyngeal, cheek), one in the back and one in the leg. Two of these lesions were cutaneous and one was intravascular in the subcutis of the leg. Microscopically, the tumors showed a variegated morphology, with alternating vasoformative and solid components, extravasated red blood cells and mild to moderate cytologic atypia. None showed brisk mitotic activity or necrosis. Tumors were negative for FOS and FOSB by immunohistochemistry. In conclusion, we report a new GATA6-FOXO1 fusion in a subset of EH, with a predilection for skin and head and neck location. The relationship of this novel molecular subset with the more common FOS/FOSB-fusion positive EH remains to be determined.
Keywords
epithelioid hemangioma; fusion; FOXO1; GATA6

INTRODUCTION
Epithelioid hemangioma (EH) is an uncommon but distinctive vascular neoplasm displaying well-formed vascular channels lined by prominent epithelioid endothelial cells[1]. The diagnosis of EH remains challenging due to its wide histologic spectrum, including a cellular/solid growth, a heavy inflammatory infiltrate, and occasional intravascular growth. The diverse histologic appearances are also reflected by the variety of its previous designations, such as intravenous atypical vascular proliferation[2], angiolymphoid hyperplasia with eosinophilia (ALHE)[3-5], inflammatory angiomatous nodule[6], and histiocytoid hemangioma[7]. Moreover, a subset of cases, particularly the ones arising in the bone, display an aggressive clinical and radiographic presentation, being associated with a high local recurrence rate and regional multifocality, which may suggest malignancy, however, they lack distant metastatic potential or tumor-related deaths[8]. Although the pathogenesis of EH has been long controversial, with some early reports suggesting that in particular skeletal EH represents a variant of epithelioid hemangioendothelioma (EHE), thus having metastatic potential[9], it is now widely recognized that EH is a benign, locally aggressive neoplasm[1, 10].

The discovery of recurrent FOS and FOSB gene fusions has strengthened its stand-alone pathogenesis, distinct from other look-alike malignant epithelioid vascular lesions, such as epithelioid hemangioendothelioma or angiosarcoma. However, only one-third of EH display FOS/FOSB gene alterations[11, 12], although the incidence varies significantly with anatomic locations, with skeletal EH having the highest proportion of fusion positive (>70%)[13]. The pathogenesis of FOS/FOSB-fusion negative EH cases remains undetermined, especially in cases that occur at non-skeletal locations. In this study we investigate a large cohort of EH cases lacking these canonical alterations, for the presence of a novel GATA6-FOXO1 fusion.

MATERIAL AND METHODS
Patient Selection and Pathologic Review
The study was initiated by an index case of an EH lacking the canonical FOS and FOSB gene rearrangements, but instead harboring a novel gene fusion. The case occurred in a 42 year-old female presenting with a painful lesion in her right leg. The tumor was located superficially in the subcutis and measured 1 cm in greatest dimension (case 1, Table 1). Prompted by this result, we have then searched the MSKCC Pathology Department and the personal consultation files of the corresponding author (CRA) for the diagnosis of EH during a 15-year period (2005-2020), which had molecular results or material available for genetic work-up. A total of 28 EH were selected that were negative for gene alterations in FOS and FOSB genes by FISH or targeted RNA sequencing methods. The study was approved by the Institutional Review Board.
Hematoxylin and eosin-stained slides and immunohistochemical stains were re-reviewed. The tumors were assessed for growth pattern, degree of vasoformation, presence of blister cells (or intra-cytoplasmic vacuoles), cytomorphology, cellularity, nuclear features including nuclear contour, chromatin pattern, pleomorphism and presence of nucleoli, mitotic activity, and necrosis. In addition, the lesions were examined for the presence and type of inflammatory infiltrate and extracellular stroma. The endothelial differentiation was confirmed by CD31, CD34 and ERG immunohistochemistry.

**Fluorescence In Situ Hybridization**

Subsequent to the results obtained from targeted RNA sequencing in the index case, all tumors were tested by fluorescence in situ hybridization (FISH) for FOXO1 and GATA6 gene abnormalities. Custom probes made by bacterial artificial chromosomes (BAC) clones flanking the FOXO1 and GATA6 genes of interest according to UCSC genome browser (http://genome.ucsc.edu) and obtained from BACPAC sources of Children’s Hospital of Oakland Research Institute (Oakland, CA; https://bacpacresources.org/) (Supplementary Table 1). DNA from each BAC was isolated according to the manufacturer’s instructions. The BAC clones were labeled with fluorochromes (fluorescent-labeled dUTPs, Enzo Life Sciences, New York, NY) by nick translation and validated on normal metaphase chromosomes. The 4 μm-thick FFPE slides were deparaffinized, pretreated, and hybridized with denatured probes. After overnight incubation, the slides were washed, stained with 4',6-diamidino-2-phenylindole, mounted with an antifade solution, and then examined on a Zeiss fluorescence microscope (Zeiss Axioplan, Oberkochen, Germany) controlled by Isis 5 software (Metasystems). Two hundred successive nuclei were examined using a Zeiss fluorescence microscope (Zeiss Axioplan, Oberkochen, Germany), controlled by Isis 5 software (Metasystems, Newton, MA). Cases were first tested for FOXO1 break-apart signals. If positive, a follow-up 3-color FISH fusion assay was performed, using 2-color for GATA6 (red, centromeric; green, telomeric) and one-color for FOXO1 (orange centromeric). To avoid false negative results due to an abundant inflammatory component intermixed in some cases, a total of 300 cells were counted in screening areas that were carefully circled on H&E as enriched in lesional cells. A positive score was interpreted when at least 20% of the nuclei showed either a break-apart or a come-together signal. Nuclei with incomplete set of signals were omitted from the score.

**Targeted RNA Sequencing**

Only one case had adequate material to be studied by RNA sequencing. The index case was tested by Anchored Multiplex Polymerase Chain Reaction Based NGS using standard protocols according to the manufacturer. RNA extraction from archival material, library preparation, and semiconductor sequencing were performed as described previously[14, 15]. In short, tumor tissue was macrodissected to achieve a histological tumor cell content of at least 15%. The extraction was carried out with the automated Maxwell® 16 Research extraction system (Promega, Madison, WI, USA) and the Maxwell® 16 FFPE Plus LEV RNA Purification Kit following the manufacturer’s instructions. The sample was then digested with the TURBO DNA-free™ Kit (Thermo Fisher Scientific, Waltham, MA, USA) to obtain DNA-free RNA. The concentration of RNA was measured fluorimetrically (QuBit 2.0 RNA high sensitivity kit (Thermo Fisher Scientific).
For enriched cDNA library preparation, the multiplex PCR-based Ion Torrent AmpliSeqTM technology (Life Technologies, Thermo Fisher Scientific) with the Archer® FusionPlex® Sarcoma kit (Archer® DX, Boulder, CO, USA) was used according to the manufacturer’s instructions and as described previously[16].

The data was analyzed with the Archer® analysis software (Version 5.1; Archer® DX) for the presence of gene fusions. The sequence quality for each sample was assessed by the following criteria: >10% or at least 150,000 unique fragments, >50 average unique RNA start sites per gene specific primer 2 (GSP2) control and on-target deduplication ratio < 40[16].

RESULTS

We have identified 28 cases of EH from our files that were negative for FOS and FOSB gene rearrangements and had tissue for additional molecular work-up. Of these, 5 (18%) cases showed the presence of a GATA6-FOXO1 fusion. Additionally, 6 ALHE cases, mostly from the head and neck location, were also tested.

Clinical and Histologic Features of GATA6-FOXO1-fusion positive EHs

There were 3 females and 2 males, with an age range at diagnosis of 15-42 years (mean, 32; median, 33). Three lesions occurred in the head and neck, including: dura, nasopharyngeal, and cheek. One case each occurred in the back and in the leg. Two of these lesions were cutaneous and one was intravascular (index case #1, involving subcutis of the leg). The tumor size ranged from 0.5-3.0 cm (mean 1.2 cm)(Table 1). All except one of the lesions were surgically resected, two of them with negative margins and two were incompletely or intra-lesionally excised. Follow-up was available in 2 of these patients, being with no evidence of disease at 6 and 36 months follow-up (Table 1). One tumor (case #4, a nasopharyngeal lesion) was only biopsied and the patient has been followed with periodical imaging, with only minimal increase in size from 2.4 to 3.0 cm over the 4 year follow-up.

Microscopically, the tumors showed a variegated morphology, with alternating vasoformative areas and solid components. In fact, all except one case showed biphasic architectural patterns, with large solid sheets of epithelioid cells adjacent to areas that were predominantly vasoformative (Figs 1-2, Table 1). The vasoformation was represented by well-formed vascular channels with irregular lumina or inter-anastomosing sinusoidal vessels (Figs. 1,2). One tumor showed very scant evidence of vascular channel formation. Most cases showed ample evidence of extravasated red blood cells (Fig 1,2). Other typical microscopic features of EH, such as the so-called ‘tombstone pattern’ of hobnailed endothelial cells protruding into the vascular lumina were not observed. One case showed a focal spindle cell component, composed of bland ovoid to short fusiform cells with vesicular nuclei and pale eosinophilic cytoplasm (case #1, Fig 1). Moreover, most cases lacked blister cells, or intracytoplasmic vacuoles as well as eosinophils. Only one case showed abundant inflammation, which was comprised by sheets of plasma cells and lymphocytes (index case #1)(Fig 1). Only one case showed very rare eosinophils, while none were detected in the remaining 4 cases. None of the cases showed prominent or distinctive stromal component,
such as hyalinization or chondromyxoid extracellular matrix, as seen with other epithelioid vascular lesions.

All except one case showed some degree of cytologic atypia, including nuclear enlargement, mild to moderate cytologic atypia, and prominent nucleoli (Figs 1, 2, Table 1). However, despite these worrisome features, none showed a brisk mitotic activity or necrosis. Overall the lesions had a low mitotic count with 1-2 MF/10 HPFs.

Targeted RNA sequencing using the Archer® FusionPlex® Sarcoma Panel was performed in the index case. The supporting reads covering the fusion junction showed the presence of a fusion transcript composed of GATA6 (NM_005257) exon 6 fused to FOXO1 (NM_002015) exon 2 (Fig. 3). This result was also confirmed by FISH. Moreover, FISH break-apart assay performed in the remaining cases showed the presence of FOXO1 gene rearrangement, as well as come-together signal with GATA6 using the FISH fusion assay (Supplem Fig 1).

**Immunohistochemistry for FOS and FOSB is negative in the GATA6-FOXO1 subset of EH**

In order to evaluate the potential relationship between our cohort and the more common EH groups with FOS or FOSB gene rearrangements, we have performed immunohistochemistry for these 2 markers, using 4-μm-thick formalin-fixed paraffin-embedded whole-tissue sections following pressure cooker heat-induced epitope retrieval. Immunohistochemistry for FOSB was performed using Target Retrieval Solution (pH 6.1; Dako, Carpinteria, CA) and a rabbit monoclonal antibody (1:2000 dilution, clone 5G4; Cell Signaling Technology, Danvers, MA) and the Envision Plus detection system (Dako), as previously reported[17]. For the FOS immunohistochemistry we used a polyclonal c-FOS antibody targeting the N-terminus (1:500 dilution; clone ABE457, Merck Millipore, MA)[18].

**Clinical and Histologic Features of fusion negative EHs**

Twenty-three EH cases were negative for GATA6 and FOXO1 gene rearrangements by FISH. This molecular negative cohort included the following locations: 8 soft tissue extremities, 2 skin, 6 head and neck, 5 bone, and 2 visceral (testis). No distinct differences in morphology were noted with the fusion positive group, although most cases showed more overt vasoformation and readily identifiable intra-cytoplasmic vacuoles. Additionally, all 6 ALHE cases tested were negative for GATA6 and FOXO1 gene abnormalities.

**DISCUSSION**

In this study we report a small series of 5 EH lesions associated with a novel GATA6-FOXO1 fusions. The tumors occurred in young adults with no gender predilection, with three cases occurring in the head and neck location. Tumors showed mostly a superficial location, involving either skin or subcutis, and had an intravascular growth in one case arising in the leg. Their clinical presentation is somewhat distinct from the more common molecular subsets of EH associated with either FOS or FOSB gene fusions, which occur preferentially in the bone, deep-seated soft tissues and outside the head and neck location[11-13].
The morphologic spectrum of EH exhibits a wide range of appearances, including cellular/solid proliferation, atypical cytology, prominent inflammatory infiltrate, and intravascular growth[19]. Due to this diverse histomorphology, the diagnosis of EH remains challenging, being confused at one end of the spectrum with inflammatory conditions (e.g., Kimura disease), while at the other end with malignant epithelioid vascular tumors, such as epithelioid hemangioendothelioma (EHE) and epithelioid angiosarcoma (AS)[20]. Moreover, the distinction from malignant lesions is further complicated by the aggressive clinical presentation, mostly encountered with intra-osseous EH, showing destructive growth and/or multifocality[8]. The current study group showed significant histologic overlap with the FOS and FOSB rearranged EH, including alternating solid and vasoformative component, predominant epithelioid cytology, hemorrhagic changes within the stroma. However, the GATA6-FOXI1-molecular subset of EH rarely exhibited blister cells or evidence of intracytoplasmic vacuoles, tombstone pattern or a stromal eosinophil infiltrate. Atypical histologic features have been previously described in the setting of EH, especially those associated with ZFP36-FOSB fusions, which often showed solid components with increased cellularity, the presence of mild to moderate degree of nuclear pleomorphism and areas of necrosis[12]. Similarly, some of the cases in our study group showed areas of solid growth and cytologic atypia, but lacked increased mitotic activity or necrosis.

It is intriguing to speculate that some gene fusions are more prone to occur in certain anatomic locations, with FOS-related fusions being common in EH of bone[11, 13], FOSB in the penis[12] and GATA6-FOXI1 fusions showing a preferential distribution in the skin and head and neck. Of note, the initial descriptions of EH, albeit described under different terminologies, including ALHE, showed a predilection for the skin and subcutis of the head and neck[3-5, 7]. Recent molecular studies have shown that lesions in the morphologic spectrum of ALHE, typically defined by variable epithelioid endothelial cells associated with a vascular ‘blow-out’ pattern and prominent inflammation, including eosinophils, do not harbor FOS and FOSB fusions[11]. Our current study, however, highlights the heterogeneity of the vascular lesions with epithelioid endothelial changes occurring in the skin and head and neck, by identifying a small subset with novel fusions which highly resembles classic EH rather than ALHE. Moreover, the current subset did not show an enriched inflammatory component in eosinophils, as commonly seen in ALHE.

GATA-6 is the only member of the GATA family of zinc-finger transcription factors expressed in vascular smooth muscle cell (VSMC), being involved in inhibiting proliferation and promoting a contractile phenotype[21]. Some of the GATA6 target genes include genes encoding the angiotensin type1a receptor, endothelin-1, and vascular cell adhesion molecule-1 (VCAM-1). The role of GATA6 in cancer remains under investigation, being different in different organs and tissues, with both tumor suppressor and oncogenic roles being proposed[22].

As FOS and FOSB related fusions are the common genetic alterations seen in EH, we further investigated our cohort using immunohistochemistry for these antibodies for a potential shared pathogenic relationship. However, none of the cases showed positivity. Moreover, FOSB fusions have been also described in a different vascular neoplasm,
pseudomyogenic hemangioendothelioma (PHE), which can display a mixture of spindle and epithelioid phenotype and may be considered in the differential diagnosis[23]. However, PHE typically lack any evidence of vasoformation and are associated with a fibrotic rather than hemorrhagic and inflammatory background.

In contrast, FOXO1 transcription factor has been well documented to be involved in a number of fusions driving mesenchymal neoplasms. FOXO1 fusions with either PAX3 and PAX7 represent the molecular hallmark of alveolar rhabdomyosarcomas[1]. Moreover, FOXO1 fusions associated with SRF have been reported in a distinctive subset of well-differentiated rhabdomyosarcoma[24]. And more recently, a single case report of a OGT-FOXO1 fusion was described in a so-called ‘myoepithelioma-like hyalinizing epithelioid tumor’, arising in the foot of a 40 year-old female[25]. However, our report appears to be the first example of FOXO1-related fusions driving the pathogenesis of an endothelial neoplasm. FOXO1 belongs to the forkhead box protein O (FoxO) transcription factor family and typically retains its transactivation domain in the fusion oncprotein. FOXO1 has important roles in regulating proliferation and differentiation of various cell types[26].

The index case showed the fusion of two transcription factors, GATA6 and FOXO1, similar to the PAX3-FOXO1 fusion in alveolar rhabdomyosarcoma[27]. In the latter fusion event, the DNA binding sites of PAX3 are retained and thus enable FOXO1-mediated stimulation of a multitude of known target genes across the genome[28][29]. Interestingly, in the GATA6-FOXO1 fusion event, all functional domains of the GATA6 protein are retained, including its DNA-binding sites. FOXO1, however, is truncated: only the distal half of the forkhead DNA-binding domain (DBD) as well as the other C-terminal domains, including nuclear localization (NLS), nuclear export signal (NES) and the transactivation domain (TAD) are retained[30, 31]. The loss of the N-terminal FOXO1 sequence prevents PKB/Akt phosphorylation at the conserved region 1 (CR1), thereby abrogating negative regulation of transcriptional activity, and may also negatively affect DNA binding. We noted a similar genetic phenomenon in a couple of other GATA6 fusion events reported in the scientific literature[32][33, 34]. Notably, these occurred in different tumor types and with different fusion partners. Collectively, these observations suggest a chimeric fusion protein that hijacks DNA-binding domains of GATA6 and may use catalytic properties of both proteins. Functional studies are required to investigate this further.

In summary, this study identified a novel molecular EH subset, which occurs at various anatomic locations and spans a wide morphologic spectrum, including atypical features that can mimic malignant vascular tumors. Of interest none of these lesions occurred in the bone, a common site for EH, which are often characterized by FOS and FOSB fusions. Although overall findings are in keeping with an epithelioid hemangioma family of tumors, this subset displayed somewhat distinct features, including a predominant solid growth, paucity of blister cells and the characteristic tombstone nuclear projection of endothelial nuclei into the lumen. Despite lack of brisk mitotic activity or necrosis, most lesions showed enlarged nuclei with small but prominent nucleoli and mild to moderate nuclear pleomorphism, worrisome for a malignant process. This study illustrates the practical utility of RNA sequencing in classifying challenging neoplasms by identifying novel gene fusions. Further
studies are clearly necessary to establish the relationship of this molecular subset with the
more common EH with FOS and FOSB gene fusions.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Morphologic spectrum of EH cases with GATA6-FOXO1 fusions.

**A-D.** Intravascular EH presenting as a 1 cm painful nodule in the subcutis of the right leg of a 42 year-old female (case 1, index case). A. Whole mount showing an intravascular lesion with variegated cellularity and central hemorrhage. B. High power of the increased cellularity component showing well-formed branching vascular channels, surrounded by brisk lympho-plasmacytic infiltrate. C. Central hemorrhagic area is composed of epithelioid endothelial cells arranged in cords, sheets or ill-defined lumina, obscured by abundant extravasated erythrocytes. D. Tumor cells are strongly and diffusely positive for ERG.

**E-F (case 2, cheek).** Cutaneous EH showing a biphasic architectural pattern, with well-defined vascular channels in subepidermal location, while the solid component present in the deeper dermis.

**G-I (case 3, dura).** EH with focal vasoformative growth composed of sinusoidal, inter-anastomosing pattern (G), and large areas of solid growth (H,I). Tumor cells show abundant eosinophilic cytoplasm and enlarged nuclei with irregular nuclear contours, open chromatin, small but distinct nucleoli, and mild to moderate nuclear pleomorphism.
Figure 2. Additional histologic features of EH with GATA6-FOXO1 fusions.
A-C. (case 4) EH involving nasopharyngeal mucosa shows a distinctive multinodular growth (A). Higher power shows solid sheets of epithelioid cells with light eosinophilic cytoplasm and enlarged nuclei with mild atypia, vesicular chromatin and prominent nucleoli. Scattered mitotic figures can be seen, but no necrosis present (B). Extravasated red blood cells are seen intermixed within the solid component. A focal spindle cell component is noted (C).
D-F. Cutaneous EH (case 5, back) showing a solid nodular growth, associated with overlying ulceration and delineated by a vague collarette. Tumor displays a biphasic growth, with alternating vasoformative areas composed of sinusoidal and irregular-contour vascular channels (E), and solid components (F). The latter finding reveals sheets of plump ovoid to epithelioid cells with abundant eosinophilic cytoplasm and irregular nuclei with open chromatin and moderate cytologic atypia. No necrosis is identified.
Figure 3. 
A. Diagrammatic representation of the chromosomal locations of \textit{GATA6} on 18q11.2 and \textit{FOXO1} on 13q14.11. Vertical red bars indicate the exact genomic break, while orange and green arrows indicate the direction of transcription for each gene. 
B. Breakpoints on the two genes (red vertical line) showing the exonic composition of the fusion transcript and the retained coding regions of the encoded fusion protein. Protein domains are also represented.
Table 1.
Clinicopathologic features of EH characterized by *GATA6-FOXO1* fusions

| EH# | Age/Sex | Site/Size/Procedure/FU | Microscopic Features | IHC positive |
|-----|---------|------------------------|----------------------|--------------|
| 1   | 42/F    | R leg, subcutis 1 cm Resection, negative margins | Biphasic, vasoformation and solid growth Intravascular Abundant inflammation (lympho-plasma cells), Extravasated red blood cells 1 MF/10 HPF, mild atypia | CD31, CD34, ERG |
| 2   | 32/M    | Cheek, skin 1.2 cm; Resection, negative margins; NED 6 months | Biphasic, vasoformation and solid growth Scant to no inflammation Limited extravasation 2 MF/10 HPFs, mild atypia | CD31, ERG |
| 3   | 33/F    | Dura 3 x 2.5 x 1 cm aggregate; Craniotomy, intralesional resection; NED 36 months | Biphasic, vasoformation and solid growth Epithelioid, ovoid to spindling (short fascicles) Abundant extravasated blood Scant acute inflammation 2 MF/10 HPFs, moderate atypia | CD31, CD34 |
| 4   | 42/F    | H&N, nasopharyngeal 2.4 cm Biopsy only AWD 48 months | Scant vasoformation Epithelioid, solid, nodular Scant to no inflammation Moderate extravasated blood, 1 MF/10 HPFs, mild atypia | CD31, ERG |
| 5   | 15/M    | Left upper back, skin 0.5 x 0.3 cm Incomplete resection | Biphasic, vasoformation and solid growth Moderate extravasated blood Scattered eosinophils 2 MF/10 HPF, moderate nuclear atypia | CD31, ERG |

FU, follow-up; NED, no evidence of disease; AWD, alive with disease.