ETV1 is a lineage-specific survival factor in GIST and cooperates with KIT in oncogenesis

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Abstract

Gastrointestinal stromal tumour (GIST) is the most common human sarcoma and is primarily defined by activating mutations in the KIT or PDGFRA receptor tyrosine kinases1,2. KIT is highly expressed in interstitial cells of Cajal (ICCs)—the presumed cell of origin for GIST—as well as in...
hematopoietic stem cells, melanocytes, mast cells and germ cells. Yet, families harbouring germline activating \textit{KIT} mutations and mice with knock-in \textit{Kit} mutations almost exclusively develop ICC hyperplasia and GIST\textsuperscript{4–7}, suggesting that the cellular context is important for \textit{KIT} to mediated oncogenesis. Here we show that the \textit{ETS} family member \textit{ETV1} is highly expressed in the subtypes of ICCs sensitive to oncogenic \textit{KIT} mediated transformation\textsuperscript{8}, and is required for their development. In addition, \textit{ETV1} is universally highly expressed in GISTs and is required for growth of imatinib-sensitive and resistant GIST cell lines. Transcriptome profiling and global analyses of \textit{ETV1}-binding sites suggest that \textit{ETV1} is a master regulator of an ICC-GIST-specific transcription network mainly through enhancer binding. The \textit{ETV1} transcriptional program is further regulated by activated \textit{KIT}, which prolongs \textit{ETV1} protein stability and cooperates with \textit{ETV1} to promote tumourigenesis. We propose that GIST arises from ICCs with high levels of endogenous \textit{ETV1} expression that, when coupled with an activating \textit{KIT} mutation, drives an oncogenic \textit{ETS} transcription program. This differs from other \textit{ETS}-dependent tumours such as prostate cancer, melanoma, and Ewing sarcoma where genomic translocation or amplification drives aberrant \textit{ETS} expression\textsuperscript{9–11} and represents a novel mechanism of oncogenic transcription factor activation.

Reasoning that transcription factors are likely to play critical roles in defining the cellular context, we utilized three expression datasets\textsuperscript{12,13} to search for GIST specific genes that might provide new molecular insights. We identified an eleven-gene signature exclusively associated with GIST in all three datasets that included the \textit{ETS} family transcription factor \textit{ETV1} (Fig. 1a, Supplementary Table 1). Examination of individual tumour samples revealed that \textit{ETV1} is highly expressed in all GISTs and at significantly higher levels than any other tumour type (Fig. 1b, Supplementary Fig. 1). \textit{ETV1} was of immediate interest since \textit{ETS} family transcription factors are well established oncogenes in Ewing sarcoma, melanoma, and prostate cancer\textsuperscript{9–11}.

Next, we assessed mRNA and protein levels of \textit{ETV1} in GIST and other sarcomas in clinical samples, GIST cell lines (imatinib-resistant GIST48 and imatinib-sensitive GIST882), the U2OS osteosarcoma cell line, and the LNCaP prostate cancer cell line known to overexpress \textit{ETV1} due to translocation\textsuperscript{14} (Fig. 1c, d). \textit{ETV1} mRNA and protein were highly and exclusively expressed in all GISTs and GIST cell lines, and in LNCaP cells. As expected, \textit{KIT} mRNA and protein were highly expressed in all GIST tumours and GIST cell lines, but not in other sarcomas or non-GIST cell lines, and phospho-KIT was only seen in GIST samples with activating \textit{KIT} mutations. Four additional GIST samples amenable to immunohistochemical analysis all showed strong nuclear \textit{ETV1} staining whereas a leiomyosarcoma control sample did not (Supplementary Fig. 2). These data show that \textit{ETV1} is universally highly expressed in all GISTs both at transcript and protein levels.

To explore the requirement of \textit{ETV1} in GIST pathogenesis, we performed RNAi experiments using two \textit{ETV1}-specific hairpins validated for both protein and mRNA suppression (Supplementary Fig. 3a). Infection with either hairpin resulted in growth inhibition of both GIST cell lines, but did not affect the growth of U2OS cells. Consistent with the level of \textit{ETV1} knockdown, \textit{ETV1}sh2 was more growth suppressive than \textit{ETV1}sh1 in both GIST cell lines (Fig. 1e). Cell cycle analysis showed that \textit{ETV1} knockdown resulted
in both decreased cell cycle progression and increased apoptosis (Supplementary Fig. 3b). 
ETO1 knockdown also impaired the tumourigenicity of GIST882 cells in SCID mouse xenografts, and those tumours that did grow had escaped ETO1 suppression (Fig. 1f).

Collectively, these observations indicate that ETO1 is required for GIST growth and survival.

Next, we addressed the mode of high ETO1 expression in GIST. FISH on 4 GIST samples and 2 GIST cell lines showed no evidence of amplification or “breakaway” between the 3′ and 5′ ends of ETO1 locus. qRT-PCR showed no evidence of differential exon expression, which is expected with ETO1 translocation (Supplementary Fig. 4). Furthermore, no focal ETO1 amplification was found in 40 GIST tumours and 6 GIST cell lines in a recent 250K SNP array study.15 The fact that high levels of ETO1 expression are consistently observed in the absence of obvious genomic alterations raises the possibility that the ICCs that give rise to GIST may endogenously express ETO1.

The musculature of the GI tract is organized into two principal layers—the inner circular muscle (CM) layer beneath the mucosa (M) and the outer longitudinal muscle (LM) layer.16 In the large intestine, myenteric ICCs (ICC-MY) form a network between the CM and LM layers surrounding the neuronal myenteric plexus, intramuscular ICCs (ICC-IM) are singly dispersed in the CM, and submucosal ICCs (ICC-SMP) form network surrounding the submucosal plexus (Fig. 2a). In the small intestine, ICC-IMs and ICC-SMPs are absent and ICC-DMPs form a network around the deep muscular plexus in the CM close to the mucosa (Supplementary Fig. 5a). All four ICC subtypes are identified by positive membrane expression of Kit16 (Fig. 2b and Supplementary Fig. 5b). In the large intestine, ICC-MYs and ICC-IMs but not ICC-SMPs stain with nuclear Etv1 (Fig. 2b). In the small intestine, ICC-MYs but not ICC-DMPs stain with nuclear Etv1 (Supplementary Fig. 5b). This finding is further supported by our analysis of a published ICC expression dataset from mouse small intestine showing that Etv1 is only highly expressed in ICC-MYs (Supplementary Fig. 5c). Notably, in the KitΔ558 mutant mice only ICC-MY and ICC-IM develop hyperplasia while ICC-SMP and ICC-DMP do not.8 These data suggest that ETO1 is a lineage-specific transcription factor for the ICCs that give rise to GIST.

We therefore asked if Etv1 is required for the normal development of ICCs by examining the GI tracts of Etv1−/− mice. Cross section and reconstructed whole-mount images from Etv1−/− mice showed significant loss of Kit-positive ICC-IMs and ICC-MYs in the large intestine (Fig. 2c–d, Supplementary Fig. 9, Supplementary Movies 1–2), small intestine, stomach, and cecum (Supplementary Figs. 6–9, Supplementary Movies 3–8). In contrast, ICC-DMPs and ICC-SMPs in the small and large intestine respectively were preserved, consistent with absent Etv1 expression in these ICC subtypes. These results were confirmed with a second ICC marker Ano119 (Supplementary Fig. 10). Immunostaining with the neuronal marker PGP9.5 confirmed the integrity of the myenteric plexus in Etv1−/− mice (Fig. 2c, Supplemental Figs. 6–8, 11). Collectively, these data indicate that Etv1 is selectively required for development of ICC-MY and ICC-IM and, by implication, a lineage-specific survival factor for the ICC-GIST lineage.
To identify ETV1 target genes in GIST, we analyzed the effect of shRNA-mediated ETV1 suppression on the transcriptomes of GIST48 and GIST882 cells. The overlap of genes perturbed by both ETV1-specific hairpins and across both cell lines was highly statistically significant, suggesting that ETV1 regulates a core set of genes in GIST (Supplementary Fig. 12). To minimize cell line-specific and off-target effects, we generated a ranked gene list based on the average change in gene expression induced by the two ETV1-specific hairpins in both GIST cell lines (Fig. 3a, b). We independently confirmed downregulation of 5 of these genes using real-time RT-PCR (Supplementary Fig. 13). Among the 48 genes suppressed >1.7-fold by ETV1 knockdown, 32 were expressed at higher levels in human GIST samples relative to other tumour types in the ExpO expression dataset (Fig. 3b). We performed gene set enrichment analysis (GSEA)20 of the “shETV1” ranked list using >3,000 gene sets in the Molecular Signature Database along with 5 custom gene sets defined by GIST-signature genes from the Segal, Nielsen, and ExpO datasets and by ICC-MY- and ICC-DMP-signature genes (Supplementary Table 1). All three GIST sets along with the ICC-MY set were among the most negatively enriched gene sets while the ICC-DMP set was not (Fig. 3c, Supplementary Fig. 14, and Supplementary Table 2). These data suggest that ETV1 is a master regulator of a transcriptional program conserved in ICC-IM/MYs and GISTs.

To define the direct transcriptional targets of ETV1 in GIST, we performed genome-wide analyses of ETV1-binding sites using ChIP-Seq in GIST48 cells. We identified 14,741 ETV1-binding sites (ETV1 peaks) which are enriched in promoter regions (Fig. 3d). Motif analysis of the peaks identified the ETS core consensus motif, GGAA, in ~90% of peaks (Fig. 3f). Integrative analyses of the ETV1 ChIP-Seq data with the transcriptomes from shRNA-mediated ETV1 suppression in GIST cells showed that 38 of the top 48 shETV1 downregulated genes contain ETV1 peaks (Fig. 3b, e, Supplementary Fig. 15). Analysis of genes with 1.4-fold change by shETV1 knockdown revealed that both shETV1 upregulated and shETV1 downregulated genes are enriched for ETV1 peaks. Furthermore, enhancer binding and in particular enhancer and promoter binding is highly predictive of transcriptional activation by ETV1 (Fig. 3h). Since enhancers are in general cell-lineage specific21,22, our data suggest that these ICC-GIST-lineage specific genes are likely directly regulated by ETV1 binding to their enhancer regulatory elements.

The dual requirement of KIT and ETV1 in normal ICC development and GIST survival raise the possibility that KIT and ETV1 cooperate in GIST oncogenesis. Inhibition of KIT signalling by imatinib in imatinib-sensitive GIST882 cells resulted in rapid loss of ETV1 protein, without significant effect on ETV1 mRNA levels (Fig. 4a, b, Supplementary Fig. 16). Similar results were observed with the MEK inhibitor PD325901. This loss of ETV1 protein was faster than the natural degradation rate, as revealed by cyclohexamide experiments to inhibit protein synthesis, and was rescued from proteosomal degradation by MG132 (Fig. 4b). Therefore, KIT-MEK signalling stabilizes ETV1 protein. Consistent with this KIT-MEK-ETV1 signalling pathway model, the overlap between genes transcriptionally altered by imatinib treatment (KIT-regulated) and by ETV1 knockdown in GIST882 cells is highly significant (Fig. 4c). Furthermore, these ETV1 transcriptional targets preferentially
contain ETV1 enhancer peaks (Fig. 4d), indicating that KIT signalling influences the ETV1 transcriptional output of the tissue and lineage-specific genes in GIST.

Having established a signalling pathway from KIT to ETV1, we explored their potential cooperativity in tumourigenesis by expressing ETV1, wild-type KIT, KIT harbouring a common GIST mutation (KITΔ560) and control vectors in combination in NIH3T3 cells. KIT-dependent stabilization of ETV1 protein was recapitulated in this system (Fig. 4e). In anchorage independent colony formation assays, ETV1 significantly increased the number and size of colonies in KITΔ560 expressing cells but was insufficient to confer anchorage-independent growth on its own (Supplementary Fig. 17). Furthermore, KITΔ560 and ETV1 strongly cooperated in conferring tumourigenic growth in SCID mice (Fig. 4f, g).

Taken together, these findings establish an oncogenic role for ETV1 in GIST that differs from classical models of ETS-driven malignancies where structural alterations (e.g., TMPRSS2-ETV1 translocation in prostate cancer, ETV1 amplification in melanoma) lead to aberrant expression and promote tumourigenesis9,11. Rather, ETV1 expression in GIST is inherited from ICC-MY/IM cells, where ETV1 is also a survival factor. We further established that KIT activity, through MEK, stabilizes ETV1, providing a mechanism for KIT-ETV1 cooperativity (Fig. 4h). These observations provide an explanation for why patients and mice with germline activating KIT mutations develop neoplasia in only the ICC-MY/IM lineage. While the mechanism of ETV1-mediated oncogenesis in GIST differs from other ETS-driven cancers, we anticipate that the ETV1-dependent transcriptional program defined here may serve as a valuable resource for further understanding of other ETV1- and other ETS-driven transcriptional programs in various cellular contexts such as prostate cancer.

The fact that ETV1 is universally highly expressed in all GISTs makes it immediately useful as a candidate diagnostic biomarker, since the current standard of KIT immunoreactivity is negative in about 5% of all GISTs23. While transcription factors has classically been considered “undruggable”, reports of successful inhibition of the NOTCH transcription factor complex and AR activity by blocking coactivator binding have challenged this paradigm24,25. Due to established requirements of ETV1 in subsets of prostate cancer and melanoma, efforts to find ETV1 inhibitors are underway and may yield novel therapeutic agents for imatinib-resistant GIST.

**Methods Summary**

**Expression data mining, microarray analysis and ChIP-Seq**

All mined datasets were downloaded Gene Expression Omnibus (GSE2109, GSE7809, GSE2719, GSE3443, GSE8167, GSE17743) and were analyzed by Oncomine™ or using Genespring 10. GIST-signature genes from three datasets containing both GIST and non-GIST malignancies met the following two criteria: 1) q<0.05, and 2) a Z-score expression difference >1.5 between GIST and non-GIST tumours. Expression profiling of GIST cell lines with different shRNA conditions was performed in duplicate on Illumina Human HT-12 array. GSEA was performed using MSigDB C2, MSigDB C4, and the GIST and ICC signature gene sets. For ChIP-Seq, sheared chromatin enriched by ETV1 IP was sequenced.
on Solexa Genome Analyzer, aligned using ELAND alignment software. Peaks were identified by MACS using input DNA as control using a FDR <1%.

Materials

GIST48 and GIST882 cells were established in the Fletcher laboratory (DFCI). All other cells were obtained from ATCC. 

Etv1−/− mice, with targeted deletion of the ETS domain, was obtained from the Jessell laboratory (Columbia) and CB17-SCID mice was from Taconic. Antibody sources are: ETV1, ANO1, PGP9.5 (Abcam), KIT for WB, P-Tyr703-KIT (Cell Signaling), P-Tyr204-ERK, GAPDH (Santa Cruz), and anti-mouse Kit for IF (clone ACK2, E-Biosciences).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This work is supported in part by the NCI (K08CA140946, YC), (5F32CA130372, PC), (CA47179, CRA, RGM), (CA148260, RGM), US NIMH (R21MH087840, DZ), NCI-ASCO Cancer Foundation Clinical Investigator Team Leadership Supplemental Award (RGM), ASCO YIA (PC), Doris Duke (CLS), Charles H Revson (YC), the Charles A. Dana (YC) Foundations, ACS MRSG CCE-106841 (CRA), P01CA47179 (CRA, RGM), Life Raft Group (CRA), GIST Cancer Research Fund (CRA), Shuman Family Fund for GIST Research (CRA, RGM), Cycle for Survival (RGM) and Startup Funds from Albert Einstein College of Medicine (DZ). We thank International Genomics Consortium (IGC) for generating ExpO data. We thank G. Wang, P. Iaquinta, and H. Hieronymus for discussions, and especially T. M. Jessell and J. N. Betley for providing and breeding 

Etv1−/− mice.

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Nature. Author manuscript; available in PMC 2011 April 14.
Figure 1. **ETV1 is universally highly expressed and required for tumour growth and survival in GIST**

- **a.** Venn diagram of GIST-signature genes from three datasets. **b.** Expression of *ETV1* in multiple tumour types from the ExpO dataset. Box, 25–75 percentile; error bar, 10–90 percentile; dots, outliers. **c.** *ETV1* and *KIT* mRNA levels by qRT-PCR of GIST and non-GIST samples, whose details are described in Full Methods. Mean±SD, n=3. **d.** Immunoblotting of selected tumour tissues and cell lines from c. **e.** Growth curves of GIST and U2OS cells after shRNA-mediated *ETV1* suppression compared to control. Mean±SEM, n=3. **f.** Tumour volume over time in SCID mice implanted with GIST882 cells after shRNA-mediated *ETV1* suppression compared to scrambled shRNA controls. Mean±SEM, *p*<0.05; n=7, 10, 8 for scrambled, ETV1sh1, and ETV1sh2 respectively. **g.** *ETV1* mRNA levels of preimplanted GIST882 cells and explanted xenografts at week 10. Mean±SD.
Figure 2. *Etv1* is expressed in the subtypes of ICCs susceptible to oncogenesis and is required for their development

**a**, Schematic showing localization of ICC-MY (yellow arrowheads), ICC-IM (yellow arrows) and ICC-SMP (white arrowheads) in the large intestine. M: mucosa, CM: circular muscle, LM: longitudinal muscle. All three ICC subtypes express Kit (red).

**b**, Co-immunofluorescence (divided into two microscopy fields) of Kit (red), *Etv1* (green) and DAPI (blue) of the large intestine of wild-type mice.

**c**, Co-immunofluorescence of Kit (red), Pgp9.5 (green), and DAPI (blue) of the large intestine of *Etv1*+/+ and *Etv1*−/− mice.

**d**, Representative deconvoluted whole-mount Kit-immunofluorescence images of the large intestine of *Etv1*+/+ and *Etv1*−/− mice. A single microscopy field focused to the ICC-MY and ICC-SMP planes are shown. The entire Z-stacks are shown in Supplemental Movies 1, 2. Scale bar, 20 μm.
Figure 3. ETV1 regulates GIST-signature genes predominantly through enhancer binding

a. Ranked list of ETV1 regulated genes was generated based on the average fold-change by the two ETV1 hairpins in two cell lines. b. Heatmap of expression of the 48 genes with average downregulation >1.7-fold. For each gene, table shows p-value of GIST vs. other tumour types from the ExpO dataset, calculated by Oncomine™ (NS: p>0.05), and the presence of ETV1 binding sites from ChIP-Seq analysis. c. GSEA plots of the shETV1 ranked list using three gene sets: GIST signature genes from ExpO dataset, ICC-MY and ICC-DMP signature genes in mouse small intestine. ES, enrichment score; FDR, false discovery rate. d. Pie charts of genomic structure and distribution of ETV1 ChIP-Seq peaks. TSS, transcription start site; TES, transcription end site. e. Representative ChIP-Seq reads in top ETV1 target genes. f. The consensus sequence motif identified in the ETV1 binding sites by the MEME program. g. Pie chart of genes with ETV1 binding sites divided into promoter

Nature. Author manuscript; available in PMC 2011 April 14.
only, enhancer only and both. h, Plot of percent of all genes, genes averagely downregulated 1.4-fold by shETV1 (n=410), and genes averagely upregulated 1.4-fold by shETV1 (n=380) with promoter only, enhancer only and both promoter and enhancer ETV1 binding. Fold enrichment over all genes is shown above the plots.
Figure 4. KIT signalling synergizes with ETV1 in GIST tumourigenesis by stabilization of ETV1 protein

a. Immunoblots of GIST882 cells treated with the imatinib (1 μM) and PD325901 (100 nM) for the indicated time points. b. Immunoblots of GIST882 cells treated for 2 hours with imatinib or PD325901 in combination with cyclohexamide (10 μg/ml) or MG132 (10 μM).

c. Venn diagram of genes downregulated by 1.4-fold by shETV1 and by imatinib in GIST882 cells. P-value: Fisher’s exact test based on number of expressed genes. d. Percent of all genes, imatinib-downregulated genes, shETV1-downregulated genes, and overlapping genes with ETV1 enhancer peaks.

e. Immunoblot of NIH3T3 cells expressing ETV1 and either KITwt or KITΔ560 two hours after treatment with PD325901, imatinib, or MG132. f. Growth of xenografts of engineered NIH3T3 cells stabilizing the indicated genes (n=12, Mean ±SEM). g. Photograph of 4 representative explanted xenografts at 4 weeks after implanting. Scale bar 1 cm.

h. Model of the role of ETV1 in ICC maintenance and GIST oncogenesis. Normal level of KIT activation by KIT ligand (red triangle) stabilizes ETV1 transcription factor through the MAPK pathway, and results in physiological ETV1 transcriptional output critical for ICC development (middle). In the absence of ETV1, there is decreased ICC development, which phenocopies genetic loss of KIT signalling (left). Activating mutation of KIT (e.g. KITΔ560) leads to constitutive activation of the KIT-
MAPK signalling pathway, increased stabilization and augmented ETV1 transcriptional output that promotes tumourigenesis (right).