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Altered chromosomal topology drives oncogenic programs in SDH-deficient GISTs

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Data Availability Statement

Sequencing data that support the findings of this study have been deposited in GEO with the accession code GSE107447.

The authors declare the following competing interests: B.E.B. is an advisor and equity holder for Fulcrum Therapeutics, 1CellBio, HiFiBio and Arsenal Biosciences, is an advisor for Cell Signaling Technologies, and has equity in Nohla Therapeutics. G.D.D. reports relationships with Novartis, Bayer, Pfizer, EMD-Serono, Sanofi, Ignyta, Roche, Loxo Oncology, AbbVie, Mirati Therapeutics, Epizyme, Daiichi-Sankyo, WIRB Copernicus Group, ZioPharm, Polaris Pharmaceuticals, M.J. Hennessey / OncLive, Adaptimmune, GlaxoSmithKline, Blueprint Medicines, Merrimack Pharmaceuticals, G1 Therapeutics, CARIS Life Sciences, Bessor Pharmaceuticals, ERASCA Pharmaceuticals, CHAMPIONS Oncology, Janssen, PharmaMar; in addition, G.D.D. has a Use patent on imatinib for GIST, licensed to Novartis, with royalties paid to the Dana-Farber Cancer Institute. Readers are welcome to comment on the online version of the paper.

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Abstract

Epigenetic aberrations are widespread in cancer, yet the underlying mechanisms and causality remain poorly understood¹⁻³. A subset of gastrointestinal stromal tumors (GISTs) lack canonical kinase mutations but instead have succinate dehydrogenase (SDH)-deficiency and global DNA hyper-methylation^{4,5}. Here we associate this hyper-methylation with changes in genome topology that activate oncogenic programs. To investigate epigenetic alterations systematically, we mapped DNA methylation, CTCF insulators, enhancers, and chromosome topology in KIT-mutant, PDGFRA-mutant, and SDH-deficient GISTs. Although these respective subtypes shared similar enhancer landscapes, we identified hundreds of putative insulators where DNA methylation replaced CTCF binding in SDH-deficient GISTs. We focused on a disrupted insulator that normally partitions a core GIST super-enhancer from the FGF4 oncogene. Recurrent loss of this insulator alters locus topology in SDH-deficient GISTs, allowing aberrant physical interaction between enhancer and oncogene. CRISPR-mediated excision of the corresponding CTCF motifs in an SDH-intact GIST model disrupted the boundary and strongly up-regulated *FGF4* expression. We also identified a second recurrent insulator loss event near the KIT oncogene, which is also highly expressed across SDH-deficient GISTs. Finally, we established a patient-derived xenograft (PDX) from an SDH-deficient GIST that faithfully maintains the epigenetics of the parental tumor, including hyper-methylation and insulator defects. This PDX model is highly sensitive to FGF receptor (FGFR) inhibitor, and more so to combined FGFR and KIT inhibition, validating the functional significance of the underlying epigenetic lesions. Our study reveals how epigenetic alterations can drive oncogenic programs in the absence of canonical kinase mutations, with implications for mechanistic targeting of aberrant pathways in cancers.

> The human genome is partitioned into physical domains, often termed topologicallyassociated domains (TADs), by chromosomal boundaries established by the DNA-binding insulator protein CTCF and cohesin⁶⁻⁹. Many proto-oncogenes and master regulators are isolated in such domains and thus protected from promiscuous enhancer interactions¹⁰.

Mutations of tricarboxylic acid cycle-related enzymes, including SDH and isocitrate dehydrogenase (IDH), are initiating events in many tumor types^{1,4,5}. These lesions cause accumulation of succinate and 2-hydroxyglutarate, respectively, which inhibit demethylases, and are associated with DNA hyper-methylation and other epigenetic alterations^{5,11,12}. The CTCF insulator is methylation-sensitive and may be displaced by DNA methylation¹³⁻¹⁵. We previously showed that the *PDGFRA* oncogene is aberrantly activated by insulator defects in *IDH*-mutant glioma¹⁶.

We hypothesized that SDH-deficiency alters chromosome topology to drive GIST tumorigenesis (Fig. 1a). GISTs are the most common gastrointestinal tract sarcoma. They are typically caused by gain-of-function mutations of the *KIT* or *PDGFRA* oncogenes that render these receptor tyrosine kinases (RTKs) active and ligand-independent¹⁷. However, ~15% of GISTs lack these defining mutations, and have instead lost SDH expression due to mutation or transcriptional silencing of *SDH* subunit genes¹⁸. We collected an initial cohort

of clinically-defined specimens, including 11 *KIT*-mutant, 2 *PDGFRA*-mutant and 8 SDHdeficient tumors (Supplementary Table 1). We used hybrid-selection bisulfite-sequencing to profile DNA methylation over 160,000 CTCF sites and representative promoters in 17 of these tumors and 2 normal stomach muscle samples (see Methods). Consistent with prior reports⁵, SDH-deficient GISTs exhibited CpG island hyper-methylation (Fig. 1b). In addition, a substantial fraction of CTCF sites were methylated in this GIST subtype (Fig. 1b).

We next identified candidate insulators and enhancers in these tumors by mapping CTCF and histone H3 lysine 27 acetylation (H3K27ac) by chromatin immunoprecipitation and sequencing (ChIP-seq). Overall patterns of enhancer acetylation were largely consistent across GISTs, relative to gastrointestinal carcinomas (Extended Data Fig. 1a). In contrast, comparison of genome-wide CTCF binding profiles revealed that ~5% of sites were specifically lost in SDH-deficient GISTs (Fig. 1c). CTCF loss was accompanied by striking increases in DNA methylation at these sites (Fig. 1c, Extended Data Fig. 1b,c). Given that DNA methylation has been established to prevent CTCF binding¹³⁻¹⁵, this suggest that hyper-methylation displaces CTCF from hundreds of candidate insulators in SDH-deficient tumors.

To investigate the impact of CTCF loss on genome topology, we used HiC to map TADs and TAD boundaries genome-wide in GIST-T1, a human cell line with an oncogenic *KIT* mutation and intact SDH expression¹⁹. We also used HiChIP⁹ to map CTCF loops and loop anchors, which correspond to TADs and boundaries, respectively^{.20} (Extended Data Fig. 1d). We used these maps to predict insulator losses likely to alter topology and gene expression. Of the 1,236 sites that lose CTCF and gain methylation in SDH-deficient GISTs, 688 corresponded to loop anchors. We reasoned that their disruption could alter topology and, in certain cases, permit aberrant enhancer-promoter interactions (Fig. 1a). We therefore further curated this list using enhancer maps and expression data. This highlighted 60 CTCF loop anchors that would normally have partitioned a large 'super-enhancer' from a gene, but that were lost in SDH-deficient GISTs (Fig. 1d,e; Supplementary Table 2). Top hits included lost CTCF insulators in the *FGF3/FGF4* locus (chromosome 11q13) and the *KIT* locus (chromosome 4q12) (Extended Data Fig 2a,b).

Although SDH-deficient GISTs lack *KIT* or *PDGFRA* mutations¹⁸, our insulator analysis raised the possibility that RTKs may instead be epigenetically deregulated. This prompted us to examine the expression of RTKs, ligands, and downstream signaling programs. First, we found that a signature for mitogen-activated protein kinase (MAPK) targets is highly expressed and suggestive of active RTK signaling in SDH-deficient GIST (Fig. 1f; see Methods). Second, a systematic analysis of tyrosine kinase gene expression revealed that *KIT* and FGF receptor 1 (*FGFR1*) are the most highly expressed RTKs in SDH-deficient GISTs (Fig. 1g). Third, we found that *FGF3* and *FGF4* were expressed at remarkably high levels, and were both specific to the SDH-deficient subtype (Fig. 1h). *FGF3* and *FGF4* are established oncogenes²¹, and FGF signaling could help explain the poor efficacy of KIT inhibitors in SDH-deficient GISTs^{22,23}. We therefore investigated the mechanisms underlying this striking and specific up-regulation of FGF ligands.

FGF3 and *FGF4* reside in a ~250kb TAD flanked by boundaries that contain CTCF binding sites (Fig. 2a). The adjacent TAD on the 11q side contains a large cluster of enhancers or 'super-enhancer' This super-enhancer overlaps the gene *ANO1*, which encodes the GIST clinical biomarker also known as DOG-1 ('Discovered on GIST-1')²⁴. It is highly acetylated and *ANO1* is highly expressed in all GIST subtypes (Extended Data Fig. 2a). Notably, the TAD boundary that partitions this super-enhancer from the FGF genes, which we refer to as the 'FGF insulator', contains several CTCF binding sites (Fig. 2a).

We hypothesized that disruption of CTCF binding could compromise the FGF insulator and allow the *ANO1* super-enhancer to contact and activate the FGF genes. The FGF insulator contains two strong and several weak CTCF binding sites. Although these sites are consistently bound in *KIT*- and *PDGFRA*-mutant tumors and normal stomach controls, all five are markedly reduced in SDH-deficient GISTs (Extended Data Fig. 2c). The strongest CTCF binding site, which is closest to the *ANO1* super-enhancer, is almost completely lost in the SDH-deficient samples (Extended Data Fig. 2d). Consistently, it is methylated specifically in SDH-deficient tumors. This suggests that the FGF insulator has switched to a methylated state that occludes CTCF binding.

To assess the impact of CTCF loss on boundary integrity, we performed circularized chromatin conformation capture sequencing (4C-seq) on 4 SDH-intact and 3 SDH-deficient GISTs. We designed a 'viewpoint' primer that enabled us to quantify contacts between a central position in the *ANO1* super-enhancer and other genomic positions at high-resolution (Fig. 2b). In SDH-intact tumors and stomach muscle control, we detected robust interactions throughout the *ANO1* TAD, but not beyond its boundaries, consistent with robust FGF insulator function. In SDH-deficient tumors, however, the same super-enhancer viewpoint physically interacted with sequences well beyond the boundary, including with the *FGF3* and *FGF4* genes, which are ~200 kb from the viewpoint (Fig. 2b, Extended Data Fig. 3a-c). These data suggest that *FGF3/4* locus topology is profoundly altered in SDH-deficient GISTs, with CTCF insulator loss allowing aberrant contacts between *ANO1* super-enhancer and FGF ligand genes.

To directly assess whether FGF insulator loss affects FGF gene expression, we used genome editing to disrupt the insulator in GIST-T1 cells, which harbor a GIST-like enhancer landscape and retain CTCF binding and boundary function. We used CRISPR/Cas9 and short guide RNAs (sgRNAs) to edit the motifs underlying the two strongest CTCF sites in the FGF insulator (Extended Data Fig. 3d). This resulted in a 6-fold induction of *FGF4*, and a 35-fold induction of *FGF3* (Fig. 2c,d). These data directly link insulator loss to the striking up-regulation of FGF ligands in SDH-deficient GISTs.

Notably, a switch between CTCF-bound and DNA-methylated insulator states underlies genomic imprinting^{13,14}. FGF insulator loss might therefore also represent a stable epigenetic event or 'epimutation' that effects a single allele. In five of our SDH-deficient samples, we identified heterozygous SNPs within an *FGF3* or *FGF4* exon. In four of these cases, analysis of the informative SNP in RNA-seq data revealed that the FGF ligand gene was mono-allelically expressed (Extended Data Fig. 4a-c). In contrast, *ANO1* was bi-allelically expressed, suggesting that the biased FGF expression reflected allele-specific

insulator loss. Consistently, in one SDH-deficient tumor with a heterozygous SNP near the CTCF site, we confirmed that only one allele of the FGF insulator was methylated (Extended Data Fig 4d). In a second tumor with an informative SNP near the *ANO1* superenhancer 4C-seq viewpoint, we confirmed that the aberrant interaction between superenhancer and *FGF4* was also strongly biased to one allele (Extended Data Fig. 4e,f). These data suggest that insulator loss, topological reorganization and FGF induction reflect a stable epigenetic alteration propagated in the malignant clone.

In addition to the FGF insulator, our screen identified a top-ranked CTCF insulator loss in the *KIT* locus. This hit was of interest given that KIT is an established GIST oncogene, and given prior reports of cross-talk between FGF and KIT signaling^{22,23}. HiC and HiChIP data reveal that the *KIT* gene is contained within a ~600 kb TAD (Fig. 3a). This large TAD contains within it a smaller insulated domain (~100 kb) that is flanked by CTCF sites. This smaller domain harbors a large super-enhancer that is highly acetylated in all GIST specimens examined (Extended Data Fig. 2b). It is partitioned from KIT by a topological boundary that we refer to as the KIT insulator.

The KIT insulator contains two strong CTCF sites separated by ~7 kb. Both sites gain methylation and lose CTCF binding in SDH-deficient GISTs (Extended Data Fig. 2e,f). To determine whether CTCF loss is associated with altered *KIT* locus topology, we performed 4C-seq using a viewpoint primer in the insulated super-enhancer (Fig. 3b). In SDH-intact tumors, the super-enhancer engages in robust interactions throughout the small insulated domain, but not beyond its boundaries (Extended Data Fig. 5a,b). In SDH-deficient tumors, however, the super-enhancer interacts with sequences well beyond the KIT insulator (Extended Data Fig. 5c,d), consistent with loss of CTCF binding and boundary function. Remarkably, quantification of interaction signals in SDH-deficient tumors indicated that ~15-20% of interactions made by this super-enhancer viewpoint are with the *KIT* promoter and gene, compared to ~1-5% in SDH-intact tumors (Extended Data Fig. 5d).

To test directly whether CTCF loss alters *KIT* locus topology, we edited the two CTCF motifs in the KIT insulator in GIST-T1 cells (Extended Data Fig. 5e) and evaluated locus topology by 4C-seq. Although the KIT insulator boundary was clearly evident in control GIST-T1 cells (Fig. 3a), it was compromised in the edited cells, as demonstrated by frequent contacts between super-enhancer and KIT (Fig 3c). We also considered the impact of insulator loss on *KIT* expression. Although GIST-T1 cells already highly express a constitutively active form of this oncogene, we found that insulator disruption further increased *KIT* expression by ~50%, under culture conditions that partially mimic SDH-deficiency (Fig. 3d; Extended Data Fig. 6). Although this proportional increase is modest, it corresponds to a substantial increase in transcriptional output given high baseline *KIT* expression in GIST-T1 cells.

Our hypothesis that FGF and KIT insulator losses drive SDH-deficient GIST predicts that these insulators should be recurrently disabled, and the corresponding oncogenes consistently expressed across tumors. We therefore examined insulator methylation across 32 GIST specimens from our original cohort and an additional validation cohort. Both insulators were highly methylated in all SDH-deficient cases, but not in any SDH-intact

tumors or normal controls (Fig. 3e,f). Consistently, CTCF binding to these insulators was compromised in all six SDH-deficient GISTs evaluated, but retained in all SDH-intact tumors and normal stomach controls (Extended Data Fig. 2c-f). Furthermore, these CTCF sites were consistently unmethylated across multiple non-malignant cell and tissue types, including a population enriched for interstitial cells of Cajal (ICCs), the presumed GIST cell-of-origin (Extended Data Fig. 7a). Finally, *FGF4* is consistently expressed across SDH-deficient tumors yet it is only expressed at very low or undetectable levels in KIT-mutant GISTs, PDGFRA-mutant GISTs, and ICCs (Fig. 1h, Extended Data Fig. 7b). The recurrence and specificity of these insulator losses support their functional significance in SDH-deficient GISTs.

Finally, we evaluated directly whether signaling through FGFR and/or KIT is required for tumor growth. Although we are unaware of any *in vitro* SDH-deficient GIST models, we successfully established an in vivo patient-derived xenograft (PDX) model from one of our SDH-deficient GIST specimens (*KIT/PDGFRA*-wild-type) (Fig. 4a). Model and parental tumor have remarkably similar RNA expression, H3K27ac enhancer landscapes, methylation and CTCF binding profiles (Fig. 4b-e). The PDX also maintains characteristic enhancers and CTCF insulator losses in the FGF and KIT loci, and strongly expresses *FGF3*, *FGF4* and *KIT* (Extended Data Fig. 8a,b). These data support the fidelity of this SDH-deficient GIST model.

We therefore tested the efficacy of FGFR and KIT inhibitors in this model. We used BGJ-398, a potent and selective inhibitor of FGFR1-4 in clinical development²⁵, and sunitinib, a drug approved for GIST with potent activity against unmutated KIT²⁶. We dosed PDX mice for 28 days with BGJ-398 (20 mg/kg), sunitinib (40 mg/kg) or the combination (Fig. 4f). Single agent sunitinib minimally suppressed tumor growth, consistent with the drug-resistant phenotype of this GIST subtype and prior reports that cross-talk between FGF and KIT signaling confers resistance to KIT inhibition^{22,23}. In contrast, single agent BGJ-398 completely suppressed tumor growth throughout the dosing period, strongly supporting a critical role for FGF signaling in tumorigenesis (Fig. 4g, Supplementary Table 3). Sensitivity to FGFR inhibition is specific to this GIST subtype as BGJ-398 lacks efficacy against SDH-intact PDX models²³. Notably, the combination of FGFR and KIT inhibitors resulted in the most durable response, with growth suppression well beyond the dosing period (Fig. 4g, Extended Data Fig. 8c,d). These pre-clinical data indicate that both RTK signaling pathways drive SDH-deficient GIST, and strongly support the significance of the underlying epigenetic lesions.

In conclusion, we identify multiple epigenetic lesions that converge to activate RTK signaling and proliferation in SDH-deficient GIST. We show that the characteristic hypermethylation in these tumors is associated with pervasive insulator losses, topological reorganization of the *FGF* and *KIT* loci, and particularly potent induction of the *FGF4* and *FGF3* oncogenes. Although our data do not address the precise cellular contexts in which these lesions arise, it is notable that KIT signaling regulates proliferation of the presumed GIST cell-of-origin, ICC²⁷. Similarly, the *ANO1* gene whose super-enhancer aberrantly drives *FGF3/4* expression encodes an ion channel that is highly expressed and essential for ICC^{28,29}. Hence, topological changes that deregulate *FGF* and *KIT* expression could lead to

unchecked signaling in these precursors. Although the corresponding loci are genetically wild-type in SDH-deficient GIST, the functional significance of their deregulation is supported by the prevalence of gain-of-function KIT mutations in SDH-intact GISTs and by a recent report that FGF4 is genetically amplified in a rare subset of KIT/ PDGFRA/SDH/RAS quadruple wild-type GIST^{17,30}. Our pre-clinical PDX data substantiate their significance and establish proof-of-concept for therapeutic intervention. Given that few stable epigenetic events have been established as drivers of tumorigenesis², our nomination of two novel functional lesions represents an important advance.

Methods

Primary GIST specimens and cell culture models.

Epigenetically-characterized clinical samples were obtained as frozen specimens from Brigham and Women's Hospital or from the Massachusetts General Hospital Pathology Tissue Bank. The validation cohort was obtained as FFPE samples from the BWH tissue bank. All samples were acquired with Institutional Review Board approval, and were deidentified before receipt. *PDGFRA* and *KIT* mutational status were confirmed through Sanger sequencing for frozen specimens, while SDH status was determined by immunohistochemistry (details below).

Samples were also examined via RNAseq and ChIP-seq input controls (details below) in order to look for mutations or copy number changes in all FGF ligand and receptor genes – no copy number alterations were found and no sequence variants were detected other than known annotated SNPs.

GIST-T1 cell line was obtained from Cosmo Biosciences¹⁹. Cells were passaged in DMEM with 10% serum, 1X antibiotics and 1X Glutamax (Life Technologies). For pseudohypoxia experiments, cells were treated with 200 μ M deferoxamine mesylate (Sigma) or vehicle control (water) for 72 hours. For succinate conditions, cells were cultured in 20 μ M dimethylsuccinate (Sigma), which was slowly added to a cell culture dish containing standard media and allowed to dissolve prior to addition of cells.

Chromatin Immunoprecipitation.

ChIP-seq was performed as described previously. In brief, cultured cells or minced frozen tissue were crosslinked in 1% formaldehyde and snap frozen in liquid nitrogen before storage at -80°C for at least overnight. Sonication of samples were calibrated such that DNA was sheared to between 300 and 700 base pair fragment length. CTCF was precipitated with a monoclonal rabbit CTCF antibody, clone D31H2 (Cell Signaling #3418). Histone H3K27 acetylation was immunoprecipitated with antibody from Active Motif (#39133). ChIP DNA was used to generate sequencing libraries by end repair (End-It DNA repair kit, Epicentre), 3' A base overhang addition via Klenow fragment (NEB) and ligation of barcoded sequencing adapters. Barcoded fragments were amplified via PCR. Libraries were sequenced as 38-base paired end reads on an Illumina NextSeq500 instrument. Processed genomic data has been deposited into dbGaP. See also Supplementary Table 4.

Reads were aligned to the reference genome (hg38) using BWA aln version $0.7.4^{31}$, removing reads with mapping quality score < 10. For H3K27 acetylation ChIPseq and input controls, PCR duplicates were removed by Picard toolkit 2.9.2. Peaks were called with HOMER 4.9^{32} against input controls. To call all H3K27ac peaks, we used 'histone' settings. To call super-enhancers³³, we used 'super' settings and no local filtering. CTCF peaks were called with 'factor' settings. To measure H3K27ac correlations, signal at the union of the peaks (5kb window around the center) was calculated by featureCounts $1.6.2^{34}$. We downloaded and reprocessed publicly available data of other GI tumors for comparison (GSM1969645³⁵, GSM1969657³⁵, GSM2058055³⁶, GSM2058056³⁶; GSM2131266³⁷, GSM2131280³⁷). The dendrogram is based on unweighted average distance linkage of the Pearson correlations between the 10,000 most variable peaks, although analysis results were similar when comparing correlations over all peaks.

Hybrid selection bisulfite sequencing.

Hybrid selection probes were designed to capture ~160,000 CTCF binding sites, and ~5,000 promoters. CTCF bind sites lists were collated from ENCODE (as downloaded from UCSC genome browser, table wgEncodeRegTfbsClusteredV3, Release 4) as well as additional CTCF maps of primary cholangiocarcinoma and glioma¹⁶. Total genomic DNA was isolated using the DNAeasy Blood & Tissue Kit (Qiagen) and sheared using the Covaris LE220. Ampure XP beads (Agencourt) were used to size select gDNA fragments within 150-320bp and sheared distribution was verified via BioAnalyzer (Agilient). 1 microgram of gDNA was end repaired, 3' A base tailed (KAPA Hyper Prep Kit #KK8502) and ligated to sequencing adapter (Roche SeqCap Epi Enrichment System). Ligated products were purified using Ampure XP beads. Following bead clean-up, products were bisulfite converted using the EZ DNA Methylation-Lightning Kit (Zymo Research) and then PCR amplified using KAPA HiFi U+ HotStart ReadyMix (KAPA #KK2800). Equal concentrations of each library were then combined in sets of three or four along with SeqCap Epi universal & indexing oligos and bisulfite capture enhancer (SeqCap Epi Accessory Kit). Each pool was lyophilized using TOMY Micro-Vac (MV100), resuspended in hybridization buffer (SeqCap Epi Hybridization and Wash Kit), and then hybridized to SeqCap Epi Probe Pool (Roche) for 72 hours at 47C in a thermocycler. Following the 72-hour incubation, captured bisulfiteconverted libraries were recovered (SeqCap Pure Capture Bead Kit) at 47C in a thermocycler for 45 minutes, with intermediate vortexing. Capture beads were washed (SeqCap Hybridization and Wash Kit) in a 47C water bath and room temperature, respectively. Captured bisulfite-converted libraries were then amplified via PCR (SeqCap Epi Accessory Kit). Libraries were sequenced with 10% PhiX spike-in as 100-base paired end reads on the HiSeq2500 in rapid run mode.

HSBS data were processed by methylCtools 0.9.4³⁸, utilizing BWA mem version 0.7.12, and aligned to human reference hg38. Due to the sizes of the captured fragments, probe capture resulted in an effective coverage of about 600bp around CTCF sites. PCR duplicates were removed by Picard toolkit 2.9.2. DNA methylation levels were called by methylCtools 0.9.4 in loci covered by at least 5 reads. Methylation at 36,281 CTCF binding sites that are bound in GIST tumors and covered by the assay were used for downstream analysis.

HiC and HiChIP.

In-situ HiC was performed as described⁹. CTCF HiChIP was performed as described²⁰. In brief, 3 tubes of ~5 million GIST-T1 cells were crosslinked in 1% formaldehyde (2 replicates for HiChIP, 1 for HiC). For HiChIP, cells were lysed using HiC lysis buffer as described. Chromatin was digested with 375 U MboI restriction enzyme (NEB, R0147). After heat inactivation of restriction enzyme and marking of ends with biotin-14-dATP (Life Technologies, 19524-016), DNA was ligated using T4 buffer (NEB, B0202). Chromatin was sheared by Covaris LE220. Chromatin immunoprecipitation was performed with 30 ul of monoclonal rabbit CTCF antibody, clone D31H2, (Cell Signaling 3418). Protein was bound by Protein G beads and after washing was incubated in DNA elution buffer. Eluant was treated with proteinase, crosslinks were reversed and the sample was Zvmo purified (Zvmo DNA Clean and Concentrator D4003). Biotinylated DNA was pulled down with M280 Streptavidin beads (Invitrogen 11205D) and the DNA was fragmented with Tn5 and libraries were constructed with Nextera kit (Illumina). For HiC, cells were lysed in HiC lysis buffer and chromatin was digested with 200 U MboI (NEB, R0147) overnight. Ends were marked and DNA was ligated as in HiChIP. DNA was precipitated and sheared by Covaris E220. Biotinylated DNA was pulled down by T1 Streptavidin beads (Life Technologies, 65602). End-repair, A-tailing and adapter ligation were performed as described⁹. Libraries were prepared using Phusion High-Fidelity DNA Polymerase (NEB, M0530). HiChIP and HiC libraries were sequenced as 75-base paired end reads on an Illumina NextSeq500 instrument. Data were processed using HiC-Pro³⁹ and visualized by the WashU EpiGenome Browser⁴⁰ and the R package Sushi⁴¹.

The two HiChIP replicates showed high similarity, and therefore were merged for the rest of the analysis. CTCF-CTCF loops were called from HiChIP data with hichipper $0.7.3^{42}$. Only loops with FDR < 5% and supported by at least 5 reads were considered for downstream analysis.

4C analysis.

4C analysis was performed using methods adapted from published protocols⁴³. In brief, ~10 million cells from culture or frozen minced tumor specimens were crosslinked in 2% formaldehyde. Fixed samples were lysed in lysis buffer containing protease inhibitor cocktail and mechanically disrupted using a Biomasher tissue grinder (Kimble Chase). Lysis was confirmed using methyl green-pyronin staining. Following lysis, nuclei were digested with NlaIII (NEB) overnight at 37°C in a thermomixer set to 950 RPM. After heat inactivation of restriction enzyme, diluted nuclei were ligated using T4 DNA ligase (NEB) overnight at 16°C, followed by RNase and proteinase K treatment. Isolated DNA was then digested overnight in Csp6I (Thermo) at 37°C, diluted and ligated overnight at 16°C in order to circularize fragments. Efficacy of each ligation and digestion step was verified via agarose gel electrophoresis. Purified circularized DNA was used as input in PCR reactions to create sequencing libraries. 16 reactions per sample were performed, each using 200ng of circularized 4C DNA (3.2 µg total) in 50ul using Q5 high-fidelity PCR mastermix (NEB). Primers contained sequencing adapters and barcodes, and annealing sections were as follows: KIT Enhancer viewpoint primer: 5' - TTTCTATTTGCTCGTTCATG - 3'; KIT non-viewpoint primer: 5' - GGAAACTTCCAAAGTAGGCT - 3'; ANO1 Enhancer

viewpoint primer: 5' – ATGTCGCCCTCCTGCATG – 3'; ANO1 non-viewpoint primer: 5' – AGACAAATGAGGCCTGGACG – 3'; ANO1 viewpoint primer for SNP detection: 5' – CTCAAACAGACACTCACATG – 3'; ANO1 non-viewpoint primer for SNP detection: 5' – TCTTTTTGGTTGGATTGTAGGAGT – 3'. Standard 4C sequencing libraries were sequenced as 38-base paired end reads on an Illumina NextSeq500, although only the first read (the viewpoint primer read) was utilized for further processing. 4C sequencing libraries for detecting SNPs were read as 75-base paired end reads on the same machine, and the second read was used for SNP detection. Data was analyzed via 4Cseqpipe⁴⁴, and median normalized data with a main trend resolution of 22.5kb were visualized in R.

RNA-seq.

Total RNA was isolated from clinical GIST samples using the RNeasy Plus Kit (Qiagen) and quality was determined via Bioanalyzer (Agilent). Libraries were prepared using the TruSeq Stranded mRNA Library Prep Kit (Illumina), and equimolar multiplexed libraries were sequenced with single-end 75-bp reads on an Illumina NextSeq 500.

Reads were aligned using STAR 2.5.3⁴⁵ to the human reference (hg19). RNA-seq data for SDH intact GISTs were previously published⁴⁶. Gene expression was estimated by featureCounts 1.6.2³⁴. TPM values were calculated for these datasets⁴⁷. RNA-seq TPM values for normal stomach was downloaded from GTEx v7⁴⁸.

Statistical analysis and reproducibility.

CTCF peaks of all GISTs were merged by bedtools merge 2.26⁴⁹, including only peaks with a score > 50 and in the top 50,000 as reported by HOMER. Peaks were then centered around CTCF motif where found by FIMO (MEME 4.7)⁵⁰ at 100bp window around the peak center, based on JASPAR 2014 CTCF motif MA0139.1 bases 4-19⁵¹. If multiple motifs were detected, we kept the one with the highest score. Reads were counted by HTSeq $0.6.1^{52}$. CTCF profiles were normalized by copy number estimates and across samples by standard median ratio. Copy number values were estimated from input by CNVnator 0.3⁵³, with 5kb bins. CTCF sites bound in all samples were used for median ratio normalization as implemented by DESeq2⁵⁴, where a site is considered bound in any given sample if its signal is at least 0.25 of the median of the top 10,000 sites for that sample. Normalization factors were used to scale CTCF signal for visualization (Figures 2 and 4) and differential analysis. Differential CTCF binding was called by DESeq254, identifying 2106 CTCF binding sited that significantly lose CTCF binding in our cohort (FDR < 5%, fold-change > 2). To estimate methylation at lost insulators sites we measured average methylation over 250bp window around the peak center. We identified 4502 sites that significantly gain methylation (FDR < 5% as determined by t-test, methylation increase > 25%). Sites that both significantly lose CTCF binding and gain DNA methylation were considered 'perturbed' for downstream analysis. We focused on CTCF-CTCF loops that overlap with a perturbed CTCF site and either 1) the loop contains a promoter that is insulated from a super-enhancer (<500kb away) by the perturbed CTCF site; or 2) the loop contains a superenhancer (at least half of the super-enhancer resides in the loop) that is insulated from a promoter (<500kb away) by the perturbed CTCF site. Here we only considered superenhancers that scored in at least two SDH-deficient GISTs. This resulted in the identification

of 60 putatively lost insulators with loss of CTCF and > 50% methylation gain, and 167 putatively deregulated genes with >1 TPM median expression across SDH deficient GIST, 25 of which with > 100 TPM (Fig. 1e, Supplementary Table 2).

To test correlations between methylation and CTCF binding we focused only on peaks detected in at least one GIST sample and with an annotated CTCF motif (see above). To empirically estimate the null distribution of the correlation coefficient, CTCF binding sites were permuted 100 times (Extended Data Fig. 1c).

To derive GIST MAPK activity score, we utilized previously identified MAPK biomarkers⁵⁵, and published data of Imatinib treatment of a GIST line⁵⁶. We picked biomarkers that were downregulated by the Imatinib treatment (p < 0.01 by t-test, fold change > 2), and expressed in all primary GISTs (>5 TPM). This yielded four biomarkers: DUSP6, ETV5, SPRY2, and SPRY4. Final MAPK score was computed by summing z-scores of the 4 genes and dividing by 2, as suggested⁵⁵.

For Fig. 2a and Fig. 3a, representative ChIP-seq traces were selected from ChIP-seq profiles for the 11 *KIT* mutant and 6 SDH-deficient GISTs characterized, all of which displayed similar results (see Supplementary Table 1 and Extended Data Fig. 2).

For all CRISPR insulator deletions, viral introduction of CRISPR/Cas9+sgRNA vector was repeated three times with separate viral preparations and infections to generate biologically independent replicates.

For epigenomic and transcriptomic characterization of clinical tissue (e.g. ChIP-seq, 4C, RNA-seq), multiple clinical specimens were analyzed, but technical replicates could not be performed on individual samples due to limited availability of material.

For analysis of clinical tissue, no statistical methods were used to predetermine sample size; rather, all available SDH-deficient tumor specimens with validated SDH loss and enough material available for analysis were tested. For mouse studies, no specific statistical calculations were performed; rather, sample size was determined based on prior experience with similar PDX trials.

Immunohistochemistry.

Immunohistochemistry was performed on 4-µm-thick paraffin embedded tissue sections using a mouse anti-SDHB monoclonal antibody (clone 21A11AE7; 1:200 dilution; Abcam, Cambridge, MA), a rabbit anti-PDGFRA monoclonal antibody (clone D13C6; 1:300 dilution; Cell Signaling Technology, Danvers, MA), and a rabbit anti-KIT polyclonal antibody (1:150 dilution; Dako, Carpinteria, CA). Pressure cooker antigen retrieval in citrate buffer (pH 6.1; Dako Target Retrieval Solution) was used for PDGFRA and SDHB. Dako Envision+ secondary antibody was used. The sections were developed using 3,3'diaminobenzidine as substrate and counterstained with Mayer's hematoxylin.

CRISPR/Cas9 insulator disruption.

The following CRISPR sgRNAs were cloned into the LentiCRISPR vector⁵⁷: sgGFP 5' -GAGCTGGACGGCGACGTAAA - 3'; sgKIT_CTCFpeak1 5' -GTCTCTCTTCTGCCAGCAGG - 3'; sgKIT CTCFpeak2 5' -GACTTCCCTGACACTAGATG - 3', sgFGF_CTCFpeak1: 5' -GTCCCACTGCCACCACAAGA - 3'; sgFGF_CTCFpeak2: 5' -GGGCCAGGCCCGCCAGG - 3'; sgSDHB 5' - GTGTCTCTTTCAGGCATCTG - 3'. sgRNAs were designed to either the GG PAM in the consensus CTCF motifs for CTCF disruption, or to a PAM near the 5' splice junction of exon 4 of the SDHB gene. GIST-T1 cells were infected with the relevant lentivirus(es) for 48h. Cells were then selected in 2 µg/ml puromycin for 4 days, with puromycin-containing media refreshed every 2 days. Cells were allowed to recover from puromycin for one week prior to analysis. Genomic DNA was then isolated and the region of interest was amplified using primers with sequencing adaptors and the following annealing regions: KIT CTCF Peak1 Forward 5' -TTTGGGATTCGAGTGACCAC - 3'; KIT CTCF Peak1 Reverse 5' -TTCAGGGCTCAACAGCTTCA - 3'; KIT CTCF Peak2 Forward 5' -GGAAATAACCTCAACCGGTG - 3'; KIT CTCF Peak2 Reverse 5' -GACTCGGTCTTGCTCCTCTAA - 3'. Libraries were sequenced on an Illumina NextSeq500 as 38 base paired-end reads, and analyzed for editing efficiency. Crosslinked cells were also harvested for ChIP analysis to verify loss of CTCF binding.

Quantitative real-time polymerase chain reaction (qPCR).

Total RNA was isolated from GIST-T1 cells using the RNeasy minikit (Qiagen) and used to synthesize cDNA with the SuperScriptIII system (Invitrogen). cDNA was analyzed using SYBR masternix (Applied Biosystems) on a 7500 Fast Real Time system (Applied Biosystems). Gene expression primers were as follows: KIT forward 5' -GCACAATGGCACGGTTGAAT - 3'; KIT reverse 5' - GGTGTGGGGATGGATTTGCT -3'; SCF/KITLG forward 5' - AGCGCTGCCTTTCCTTATGA - 3'; SCF/KITLG reverse 5' - CCGGGGGACATATTTGAGGGT - 3'; HIF2a/EPAS1 forward 5' -CCACCAGCTTCACTCTCTCC - 3'; HIF2a/EPAS1 reverse 5' -TCAGAAAAAGGCCACTGCTT - 3'; FGF4 set 1 forward 5' -CCAACAACTACAACGCCTACGA; FGF4 set 1 reverse 5' -CCCTTCTTGGTCTTCCCATTCT - 3', FGF4 set 2 forward 5' -GCAGCAAGGGCAAGCTCTAT - 3'; FGF4 set 2 reverse 5' -CGGTTCCCCTTCTTGGTCTT - 3'; FGF3 forward 5' - ATGCTTCGGAGCACTACAGC - 3'; FGF3 reverse 5' - CACGTACCACAGTCTCTCGG - 3'. All gene expression results were normalized to primers for ribosomal protein, large, P0 (RPLP0) as follows: forward 5' - TCCCACTTGCTGAAAAGGTCA - 3'; reverse 5' - CCGACTCTTCCTTGGCTTCA -3'.

Tyrosine Kinome Tree Visualization.

Tyrosine kinase phylogeny data was downloaded from kinase.com⁵⁸. Phylogenetic tree was visualized using the R package ggtree⁵⁹. Expression data of each tyrosine kinase was

average across the SDH-deficient GIST, and then plotted on the tree, with the area of the red circles corresponding to the average TPM value in SDH-deficient GIST.

Interrogation of public normal tissue, GIST, and ICC expression data.

Data for normal tissue expression was obtained from ENCODE⁶⁰. Mouse interstitial cell of Cajal expression data was previously processed and published⁶². Raw Affymetrix microarray data (CEL files) of human ICC and GIST samples were downloaded from GEO, under accessions GSE56670⁶¹, GSE77839⁶³, GSE17743⁶⁴, and GSE20708⁶⁵. CEL files were imported, normalized, and RMA values exported using the R/Bioconductor package affy⁶⁶.

Flow cytometry enrichment and analysis of ICCs.

Fresh benign stomach muscle tissue was obtained from the MGH Pathology Tissue Bank and dissected from the gastric epithelium. Tissue was initially manually mechanically dissociated with sterile scalpel, and then subjected to fine mechanical dissociation through three cycles of one minute each in a Miltenyi gentleMACS dissociator, resulting in a singlecell suspension. A small portion was removed from the cell suspension to serve as the unlabeled and unpermeabilized control to set size gates and test viability. The remainder of the cell suspension was then incubated in a permeabilizing flow cytometry buffer and stained with ANO1-Alexa488 (Santa Cruz Biotechnology clone C-5), KIT-PE (Biolegend Clone 104D2) or CD45-APC (BD Biosciences clone 2D1) for 30 minutes at 4°C. Nonpermeabilized control cells were treated with propidium iodide immediately prior to analysis. Stained cells were analyzed and collected on a Sony SH800S cell sorter. Compensation parameters were determined using single-labeled UltraComp eBeads (ThermoFisher). Approximately 1.5 million cells were sorted, of which 2,000 were collected as CD45-/KIT+/ANO1+ (ICC-enriched). Cells were lysed in a small volume of TAE/DTT and treated with Proteinase K. Genomic DNA in the lysed cell mixture was then bisulfite converted using the EZ DNA Methylation Gold Kit (Zymo), subjected to locus PCR, and then sequenced on an Illumina NextSeq500.

Patient-Derived Xenograft (PDX) Model Generation and efficacy studies.

The PDX model was generated from surgical resection tissue from an SDH-deficient GIST patient who consented to research use of material under an IRB-approved protocol. The surgical sample was implanted subcutaneously in female NSG mice and allowed to grow. Tumor growth was monitored by caliper measurements. Once tumors grew to a size of 1000 mm³, tumors were isolated and cut into pieces of approximately $3 \times 3 \times 3$ mm, dipped in Matrigel (Corning Life Science, MA) and transplanted subcutaneously in additional NSG mice. Tumors were passaged for no more than 10 times. Tumor samples from all passages were banked by viably freezing in Bambanker freezing media (Fisher Scientific) and used for further studies. For efficacy studies, tumor fragments were implanted into 8-week old NSG mice. Tumors were allowed to establish to 192 ± 35.7 mm³ in size before randomization into various treatment groups with n=8/group as: vehicle control (0.1M citrate buffer, pH 4.5), 40 mg/kg sunitinib (LC Laboratories, 0.1M citrate buffer, pH 4.5), 20 mg/kg BGJ-398 (LC Laboratories, acetate buffer, pH 4.6 and PEG300 in 1:1 ratio) or the combination of BGJ-398 and sunitinib. Mice were treated orally once daily for 28 days with

these agents. In the BGJ-398 treatment group, 4 of 8 mice, and in the combination treatment group, 7 of 8 mice, lost >15% body weight requiring drug holidays (1 to 3 days of drug holidays in the single agent BGJ-398 group and 1 to 15 days of drug holidays in the combination group). Mice were re-started on treatment once body weight recovered to at least >85% of initial body weight. Treatment groups were censored when the tumor volume reached the maximum permissible size of 2000 mm³in any single mouse in that group. Statistics were determined by two-way ANOVA.

All relevant ethical regulations regarding research in animal models were followed.

All animal experiments and study protocols were approved by the Dana Farber Cancer Institute Institutional Animal Care and Usage Committee (IACUC). The endpoint criteria for mice were if the total tumor burden/size reaches 2 cm in any direction or tumor volume exceeds 2000 mm³, and/or if the tumor mass interferes with basic/vital bodily functions or becomes persistently ulcerated, and these criteria were followed for all mice in the study.

Extended Data



Extended Data figure 1. Epigenomic characterization of GIST.

(a) ChIP-seq profiles for H3K27ac were compared for GIST specimens and other gastrointestinal tract tumor specimens (GAC: gastric adenocarcinoma, CRC: colorectal cancer, PDAC: pancreatic ductal adenocarcinoma). Heatmap depicts pairwise Pearson correlations between the top 10,000 most variable peaks (yellow indicates high correlation; blue indicates low correlation). Dendrogram (left) was derived by unweighted average distance linkage. Enhancer patterns are relatively consistent across GIST subtypes, compared to other tumor types. (b) DNA methylation levels in the vicinity of CTCF sites were profiled genomewide by hybrid-selection bisulfite sequencing. CTCF sites are binned according to the amount their methylation increased in SDH-deficient GISTs, relative to SDH-intact GISTs (methylation change computed over 250 bp window centered on the motif). For each bin, bar graphs depict the percentage of sites that lose CTCF binding in SDH-deficient GISTs, per ChIP-seq. Separate plots are shown for CTCF sites whose motifs do or do not contain a CpG. Increased methylation over CTCF sites is associated with more frequent loss of CTCF binding, even when the CTCF motif lacks a CpG. (c) Plot depicts correlation between CTCF occupancy and DNA methylation in SDH-deficient GISTs. Red points show Spearman correlations between CTCF ChIP-seq signal and methylation of CpGs at indicated positions relative to the center of the CTCF motif. Red line reflects correlation to average methylation over 10bp windows. Randomly permuted data (black) are shown for comparison. Anti-correlation between CTCF occupancy and methylation is evident over a ~250 bp binding footprint. (d) Genomic views of a representative 10 MB region on chromosome 21 depict chromosome topology (HiC, red), CTCF binding (ChIPseq, orange) and CTCF-CTCF loop interactions (HiChIP, black) for the SDH-intact GIST model, GIST-T1. TADs are visible as triangles of enhanced interaction in HiC data, flanked by boundaries that correspond to loop interactions in HiChIP data. Genes (blue) are also indicated.



Extended Data figure 2. Super-enhancers and insulators in GIST.

(a) Traces depict H3K27ac ChIP-seq signal for normal stomach muscle (NSM) and GISTs of the indicated subtype over the FGF/ANO1 locus. (b) Traces depict H3K27ac ChIP-seq signal for NSM and GISTs of the indicated subtype over the PDGFRA/KIT locus. Genes are indicated in blue, and superenhancer locations are indicated by green bars. For (a,b), traces are representative of 11 KIT-mutant and 6 SDH-deficient tumors with similar results. (c) Traces depict CTCF binding over the FGF insulator in normal stomach muscle (NSM) and GIST clinical specimens. (d) Plot depicts CTCF ChIP-seq signal over the strongest CTCF peak in the FGF insulator in normal stomach muscle (NSM, n=4), and KIT mutant (n=11), PDGFRA mutant (n=2) and SDH-deficient GISTs (n=6). (e) Traces depict CTCF binding over the strongest CTCF peak in the KIT insulator in normal stomach muscle (NSM) and GIST clinical specimens. (f) Plot depicts CTCF ChIP-seq signal over the KIT insulator in normal stomach muscle (NSM, n=4), and KIT mutant (n=2) and SDH-deficient GISTs (n=6). (n=11), PDGFRA mutant (n=2) and SDH-deficient GISTs (n=6). For (d) and (f), P-values indicate significance of CTCF loss in SDH-deficient GIST, as determined by Walt test (via DEseq2⁵⁴). All n values represent number of biologically independent clinical specimens.



Extended Data figure 3. FGF locus 4C-seq data and insulator deletion.

(a) Traces depict 4C-seq data at FGF locus, as in Fig. 2b., except graphed on the same axis to allow direct comparison. (b,c) Barplots quantify 4C-seq interactions between the superenhancer viewpoint and FGF4 (b) or FGF3 (c). Expression of these genes in the corresponding SDH-deficient GIST specimens is also shown. (d) Traces depict CTCF ChIPseq signal in GIST-T1 cells infected with CRISPR/Cas9 and either a control sgRNA directed at GFP (black, top) or sgRNAs directed against the two indicated CTCF motifs in the FGF insulator (second row, red).



Extended Data figure 4. Allelic imbalance in FGF3/FGF4 activation.

(a) Two heterozygous SNPs in FGF4 (both 3' UTR) enabled us to evaluate allelic expression in three SDH-deficient GISTs (tumors S6, S1, and S4). Both alleles for each SNP were detected in DNA sequencing data for these tumors, but only one allele was detected in RNAseq data of tumors S1 and S6, indicative of mono-allelic FGF4 expression. Both alleles are detected in tumor S4, indicating biallelic expression of FGF4. (b) Heterozygous SNPs in FGF3 exons (both synonymous base substitutions) enabled us to evaluate allelic expression in the SDH-deficient GISTs (tumors S2 and S5). In both cases, DNA sequencing confirmed heterozygosity at the genome level (C/A and T/C, respectively), but RNA-seq data demonstrated mono-allelic FGF3 expression. (c) Both alleles of heterozygous SNPs in ANO1 exons were found in the RNA-seq data derived from SDH-deficient GIST samples, confirming bi-allelic expression of ANO1. Similarly, both alleles of heterozygous SNPs were found in the histone H3K27ac ChIP-seq data, confirming the biallelic nature of the superenhancer (not shown). (d) One SDH-deficient GIST sample was heterozygous for a SNP (rs386829467) located about 50bp from the CTCF motif of Peak 2 in the FGF insulator. Allele-agnostic methylation data confirmed 43% methylation of the CTCF peak in

this tumor, while essentially no methylation was detected in the SDH-intact tumors (left). Separation of the two alleles using the heterozygous SNP revealed strong allelic bias in the SDH-deficient tumor: one allele was largely unmethylated (\sim 3% methylation), while the other was highly methylated (~75% methylation), consistent with monoallelic methylation of the CTCF site (right). (e) Schematic depicts 4C-seq experimental protocol and primer design for detecting SNPs. DNA elements in close physical proximity are crosslinked and restricted with an enzyme that leaves nucleotide overhangs. These overhangs are then proximity ligated to crosslinked fragments. A second restriction enzyme (with different restriction sites) is then used to circularize the ligated fragments, allowing for inverse PCR. Here we selected restriction enzymes and designed a custom read 2 primer to capture a heterozygous SNP within the super-enhancer. This second read is normally non-informative as contact frequencies are determined through the viewpoint primer (read 1), but in this case enabled us to detect the SNP and assign each ligated fragement to a specific allele. (f) Left trace (gray) depicts standard 4C-seq data (allele agnostic), which demonstrates strong interaction between super-enhancer viewpoint and FGF4. However, the SNP covered in the non-viewpoint read enabled us to distinguish interactions involving the minor (top right) or major (bottom right) allele. This revealed that the major allele (purple) is responsible for ~97% of superenhancer-FGF4 interactions.



Extended Data figure 5. KIT locus 4C-seq data and insulator deletion.

(a) Traces depict KIT locus 4C-seq data, as in Fig. 3b., except graphed on the same axis to allow direct comparison. (b-d) Barplots quantify 4C-seq interactions (top, reproduced from Fig. 3b) between the super-enhancer viewpoint and positions within the super-enhancer TAD (b), sequences just beyond the KIT insulator (c), or the KIT gene itself (d). P-values indicate significance of difference between SDH-intact and SDH-deficient, by two-sided T test. (e) Traces depict CTCF ChIP-seq signal in GIST-T1 cells infected with Cas9 and either a control sgRNA directed at GFP (black, top) or sgRNAs directed against the two bound CTCF motifs in the KIT insulator (second row, red).



Extended Data figure 6. Hypoxia marker induction in GIST-T1 cells.

(a) Bar plot depicts levels of the TET product, 5-hydroxymethyl-Cytosine (5-hmC), measured by ELISA in GIST-T1 cells infected with CRISPR/Cas9 and either a short guide RNA targeting GFP (sgGFP) or SDHB. Cells were cultured in either control media or media supplemented with 20uM of dimethylsuccinate (DMS), a membrane-permeable ester of succinate, for 3 days, as indicated. (**b**,**c**) Plots show relative expression of pseudo-hypoxia-associated genes EPAS1/HIF2 α^{67} and IGF1 R^{68} (b), and KITLG/SCF⁶⁹ (c) in control GIST-T1 cells (black), SDH-deficient GIST-T1 cells generated by CRISPR/Cas9 knockout of SDHB and cultured with exogenous succinate (red), or GIST-T1 cells treated with the iron chelator DFX to simulate hypoxia (blue). Up-regulation of KIT ligand due to pseudo-hypoxia or tumor hypoxia may supplement FGF ligands in promoting RTK signaling in SDH-deficient GIST.



Extended Data figure 7. FGF and KIT insulator methylation and expression in GIST subtypes and non-malignant cells.

(a) Barplot depicts methylation of FGF insulator CTCF peak 2 (top) and KIT insulator CTCF peak 2 (bottom) in 34 tissues and primary cells available through ENCODE⁶⁰. Values are average methylation of CpGs nearest the CTCF motifs, determined by WGBS (KIT insulator: 3 CpGs; FGF insulator: 6 CpGs). Methylation of these sites is also shown for SDH-intact and SDH-deficient GISTs (see Fig. 3e,f), and for flow sorted CD45-/ ANO1+/KIT+ (ICC-enriched) cells from normal stomach muscle tissue (n values represent biologically-independent specimens). (b) Left: Table depicts FPKM values of relevant genes in mouse ICCs isolated from jejunum or colon⁶². Right: Dot plot depicts expression of FGF4 in either flow sorted ICCs (green) or GISTs of the indicated subtype: KIT mutant in black, PDGFRA mutant in blue, SDH-deficient in red, and KIT-/PDGFRA-wildtype in purple^{61, 63-66}. SDH status of the latter group is unknown, but SDH-deficient GIST represent a significant portion of KIT-/PDGFRA-wildtype tumors. Data are drawn from the indicated GEO series and publications.



Extended Data figure 8. PDX trial of FGFR/KIT combination therapy in SDH-deficient GIST. (**a,b**) Genomic views depict RNA expression (black), H3K27ac (green) and CTCF occupancy (orange) over the FGF (a) and KIT (b) loci for the S1 primary tumor and PDX. Genes (blue), super-enhancers (green bar and shade) and lost CTCF insulators (orange shade) are indicated. (**c**) Plot depicts tumor volume during treatment and observation periods of experiment, as in Figure 4g, except with time axis extended until final group reached censor point (one tumor in the group > 2000 mm³). Points represent mean tumor size, error bars represent S.E.M., and shading represents range of tumor sizes for n=8 biologically independent xenograft-bearing mice per group. For statistics, see Fig. 4g. (**d**) Kaplan-Meier plot depicts survival until clinical endpoint (tumor size > 2000 mm³) for the same PDX trial. Median and range survival are indicated for each group. P-values reflect difference in survival between groups as calculated by logrank test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data generated for this study are available through the Gene Expression Omnibus (GEO) under accession number GSE107447.

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Figure 1. Insulator dysfunction in SDH-deficient GIST.

(a) Proposed mechanism of epigenetic oncogene activation. Left: Oncogene shielded from super-enhancer by CTCF insulator, which creates a topological boundary. Right: CTCF insulator displaced by DNA methylation, allowing super-enhancer to contact and induce oncogene. (b) Violin plots depict DNA methylation levels over the 10,000 most variable CpG island promoters (top) and CTCF sites (bottom) in normal stomach muscle (NSM; n=2), and KIT mutant (n=9), PDGFRA mutant (P; n=2) and SDH-deficient GISTs (n=6). Yellow bars indicate mean. (c) Volcano plot depicts differential CTCF occupancy between SDH-deficient (n=6) and SDH-intact (n=8) GISTs. Sites that gain DNA methylation in SDH-deficient GISTs are indicated in red (>25% increase, two-sided t-test FDR < 5%). (d) Plot depicts H3K27ac peaks near lost CTCF insulators (y-axis) rank ordered by signal strength. (e) Scatter plot depicts genes (points) separated from a super-enhancer by a CTCF loop anchor that is lost in SDH-deficient GIST. Genes are positioned according to their relative (y-axis) and absolute median expression (x-axis) in SDH-deficient GISTs. Potentially deregulated gene targets (outliers) include oncogenes FGF3, FGF4 and KIT (red); see also Supplementary information. (f) Box plot depicts average expression of MAPK signature genes in RNA-seq data for normal stomach (n=262), and KIT mutant (n=10), PDGFRA mutant (n=3), and SDH-deficient GIST (n=8). Boxes depict 25th, 50th and 75th percentiles, and whiskers depict extreme values. (g) Radial phylogenetic tree depicts tyrosine kinase gene expression in SDH-deficient GISTs. Each branch is one tyrosine kinase, arranged by similarity, and with major families depicted by color. The area of each red circle is proportional to the average expression of the kinase. (h) Scatter plot depicts

average expression of FGF ligands in SDH-intact (x-axis) and SDH-deficient (y-axis) GISTs. For all panels, n values indicate number of biologically-independent specimens.



Figure 2. FGF3/4 locus topology reorganized in SDH-deficient GIST.

(a) Genomic views of the *FGF3/4* and *ANO1* loci depict baseline chromosome topology (HiC, red), genes (blue), CTCF-CTCF loop interactions (HiChIP, arcs, with darkness indicating significance), CTCF binding (ChIP-seq, orange) and candidate enhancers (H3K27ac ChIP-seq, green). HiC/HiChIP data are for the SDH-intact model GIST-T1, while CTCF and H3K27ac data are for representative clinical specimens (see also Extended Data Fig. 2). *ANO1* super-enhancer (green bar) and FGF insulator (orange shading) are indicated.
(b) Traces depict 4C-seq interaction frequency (y-axis) between the ANO1 super-enhancer viewpoint (dashed white line) and genomic positions in the FGF3/4-ANO1 locus (x-axis). Data shown for SDH-intact GISTs (n=4; top), normal stomach muscle (n=1; gray line, top) and SDH-deficient GISTs (n=3; bottom). CTCF binding profiles for representative SDH-intact (top) and SDH-deficient (bottom) tumors also shown (orange). Genes (blue) and CTCF sites in the FGF insulator (orange) are highlighted. (c,d) Plots depict relative *FGF4* (c) and *FGF3* (d) expression in GIST-T1 cells expressing CRISPR/Cas9 and control sgRNA (black) or sgRNAs targeting the two CTCF sites in the FGF insulator (red). Bar indicates mean of 3 biologically-independent replicates (dots). P-values by two-sided T-test.



Figure 3. KIT/PDGFRA locus topology reorganized in SDH-deficient GIST.

(a) Genomic views of *PDGFRA* and *KIT* loci depict baseline chromosome topology (HiC, red), genes (blue), CTCF-CTCF loop interactions (HiChIP, arcs), CTCF binding (ChIP-seq, orange) and candidate enhancers (H3K27ac ChIP-seq, green). HiC/HiChIP data are for the SDH-intact GIST model GIST-T1, while CTCF and H3K27ac data are for representative clinical specimens (see also Extended Data Fig. 2). KIT super-enhancer (green bar) and KIT insulator (orange shading) are indicated. (b) Traces depict 4C-seq interaction frequency (yaxis) between the KIT super-enhancer viewpoint (dashed white line) and genomic positions in the KIT/PDGFRA locus (x-axis). Data shown for SDH-intact GISTs (n=6, top), and SDH-deficient GISTs (n=3, bottom). CTCF profiles for representative SDH-intact (top) and SDH-deficient (bottom) tumors also shown. Genes (blue) and CTCF binding sites in the KIT insulator (orange) are highlighted. (c) Traces depict 4C-seq interaction signal between the KIT super-enhancer viewpoint and the KIT gene in GIST-T1 cells expressing Cas9 and control (black) or KIT insulator targeting sgRNAs (red). (d) Plot depicts relative KIT expression in GIST-T1 cells expressing Cas9 and control (black) or KIT insulator targeting sgRNAs (red). Bar indicates mean of three biologically-independent replicates (dots). Pvalues by two-sided T-test. (e,f) FGF and KIT insulator methylation evaluated in an expanded cohort of GIST tumors by locus-specific bisulfite sequencing. (e) Bar plot depicts average methylation levels across six CpGs within FGF insulator CTCF peak 2 in normal stomach muscle (n=2), SDH-intact GISTs (n=17) and SDH-deficient GISTs (n=11). (f) Bar plot depicts average methylation across four CpGs within KIT insulator CTCF peak 2 in normal stomach muscle (n=2), SDH-intact GISTs (n=20) and SDH-deficient GISTs (n=12) (n values indicate number of biologically-independent tumors).



Figure 4. SDH-deficient GIST PDX trial confirms dependence on FGF signaling.

(a) Specimen collection and generation of PDX model. (b) Scatter plot compares expression of genes (points) between primary tumor S1 (x-axis) and PDX (y-axis) per RNA-seq. Pearson correlation indicated. (c) Venn diagram depicts overlap between strong H3K27ac ChIP-seq peaks in primary tumor (black) and PDX (green). (d) Venn Diagram depicts overlap between hypermethylated CpG islands in primary tumor (black) and PDX (red) per bisulfite-sequencing. (e) Scatter plot depicts principle component analysis (PCA) on top 1,000 differential CTCF sites for primary tumors (colored by subtype) and PDX (star). PDX and originating tumor (S1) both cluster with SDH-deficient GISTs (red). (f) Experimental design of pre-clinical trial. Following xenograft implantation and growth, mice were randomized to four treatment groups and treated with the indicated regimen daily for 28 days. Observation was continued until clinical endpoint (tumor volume of $2,000 \text{ mm}^3$). (g) Plot depicts tumor volume during treatment and observation periods. Points represent mean tumor volume, error bars represent S.E.M., and shading represents range of tumor volumes for n=8 biologically-independent xenografts per group. Relative tumor volume immediately following treatment cessation (day 29) is indicated at right. P-values reflect difference in tumor growth between group per two-way ANOVA (* P < 0.05, ** P < 0.001, *** P <0.0001).