

Epigenetic control of oncogene signaling

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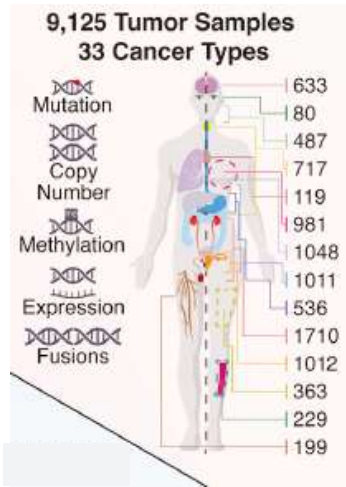
Memorial Sloan Kettering Cancer Center

April 16, 2026

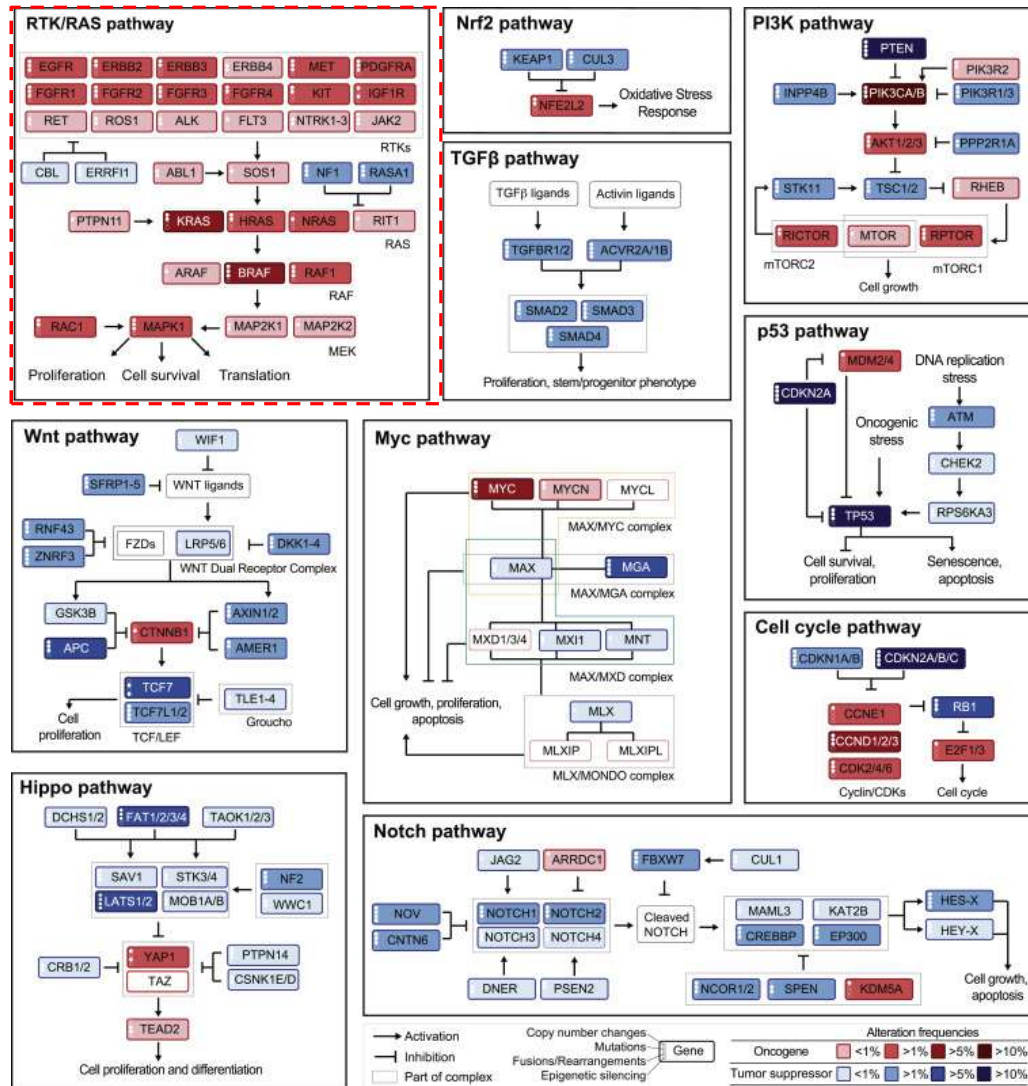


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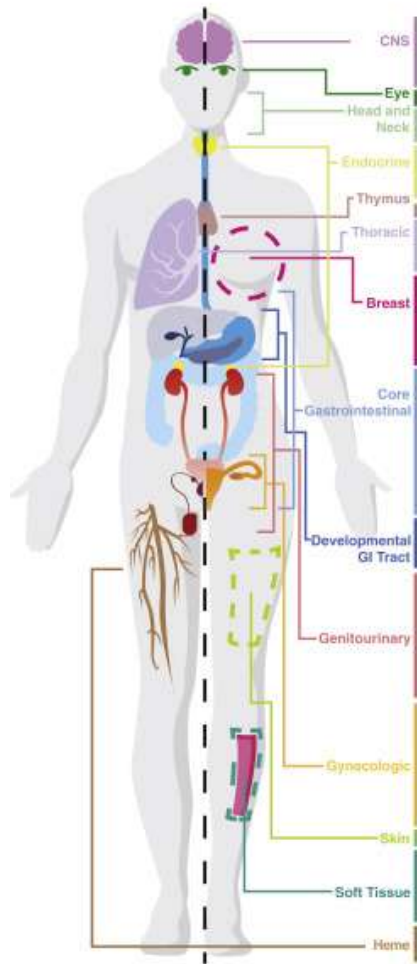
Oncogenic signaling pathways in cancer



89% ≥1 driver alteration in the pathways
 57% ≥1 druggable targets
 30% multiple druggable alterations



RTK signaling in cancer

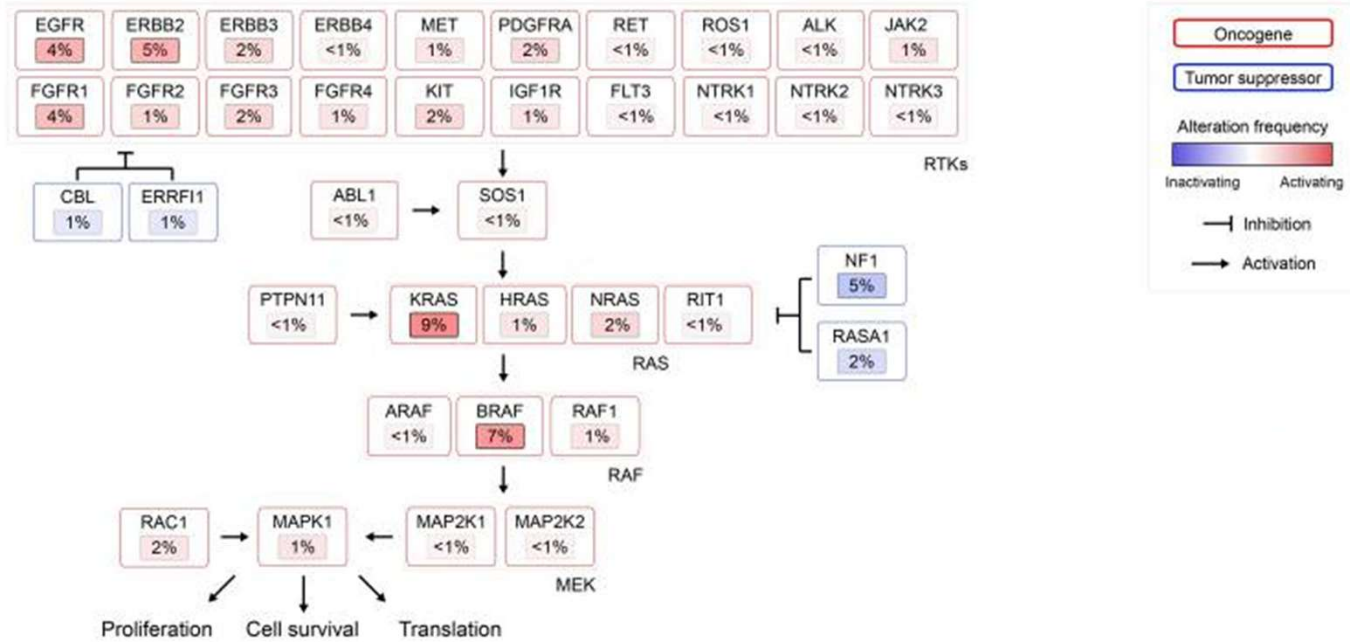


Alteration frequencies

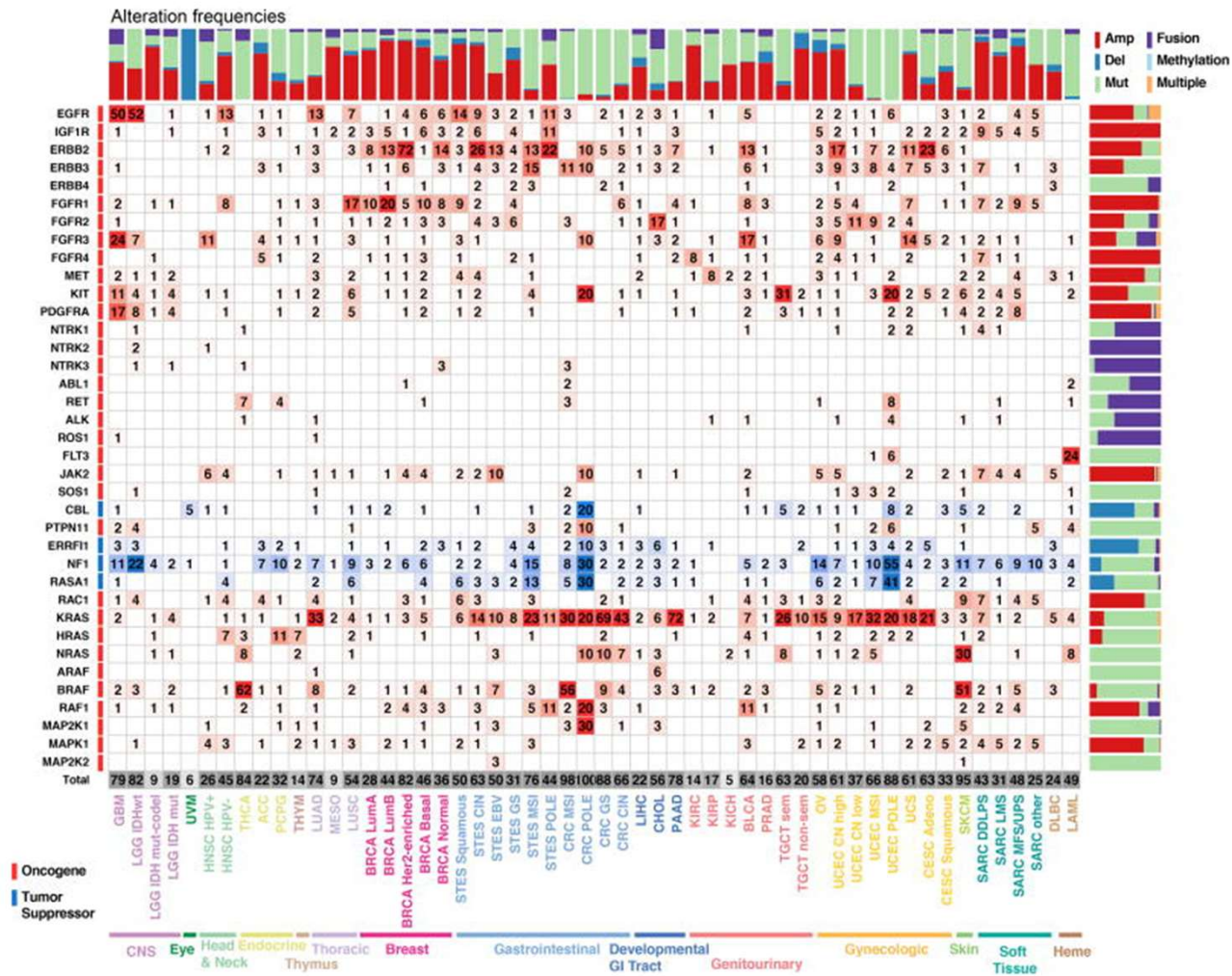
										CIN	Foa	TMB	
GBM	77	86	57	48	18	8	6	10	2	119	0.37	4.3	
LGG IDHwt	82	64	47	29	27	5	1	5		68	0.25	4.3	
LGG IDHmut-codel	9	45	22	5	26	66	3	99	50	34	0.21	0.8	
LGG IDHmut	19	28	15	92	24	8	10	92	21	1	60	0.17	0.8
UVM	6	6	4	2	10	1	2	10		1	65	0.28	0.4
HNSC HPV+	26	32	60	11	25	8	4	8	11	1	83	0.32	3.3
HNSC HPV-	45	86	39	82	36	16	20	42	13	13	108	0.43	4.6
THCA	84	14	4	1	4	13	2	1	2		16	0.03	0.4
ACC	22	30	16	28	11	41	7	5	1		146	0.78	1.8
PCPG	32	15	6	6	11	10	1	4	1		81	0.33	0.3
THYM	14	9	4	7	5	7	1	7	3	2	26	0.09	0.6
LUAD	74	56	38	61	21	19	23	23	10	15	118	0.48	8.2
MESO	9	54	13	21	9	6	7	40	2		98	0.41	0.8
LUSC	54	79	68	86	31	18	12	28	11	25	158	0.61	7.7
BRCA LumA	28	31	62	25	14	15	12	5	4	1	65	0.34	2.0
BRCA LumB	44	48	48	49	25	31	26	15	10	2	211	0.60	2.0
BRCA Her2-enriched	82	40	60	78	18	17	29	10	8	1	230	0.53	4.3
BRCA Basal	46	51	53	91	38	11	39	14	8	4	246	0.67	2.7
BRCA Normal	36	36	33	31	3	6	19	3			53	0.16	1.5
STES Squamous	50	89	53	96	38	13	22	21	13	23	189	0.59	3.1
STES CIN	63	74	33	76	21	26	21	16	23	2	222	0.58	3.4
STES EBV	50	100	80	13	83	67	7	10	17		52	0.22	4.1
STES GS	31	39	18	24	31	20	12	4	20	2	66	0.10	2.1
STES MSI-POLE	71	64	64	49	79	70	19	54	57	2	85	0.19	37.1
CRC MSI-POLE	99	74	68	49	74	95	52	64	55	1	45	0.09	56.9
CRC GS	88	45	53	19	29	90	21	10	38	5	55	0.23	2.9
CRC CIN	66	36	32	84	23	91	17	8	22	1	115	0.54	2.9
LIHC	22	69	25	37	26	43	19	12	7	7	121	0.45	2.9
CHOL	56	53	17	19	8	17	19	17	3	6	100	0.58	1.8
PAAD	78	70	19	69	14	12	14	7	41		62	0.26	3.5
KIRC	14	14	17	6	8	7	5	5	3	3	49	0.25	1.6
KIRP	17	12	8	4	12	9	6	11	1	6	49	0.35	2.2
KICH	5	23	15	32	3	3	2	3	5		77	0.80	0.9
BLCA	64	81	46	62	42	20	18	26	9	9	150	0.50	6.8
PRAD	15	28	32	21	13	35	11	5	6	1	92	0.16	1.3
TGCT sem	63	8	11	6	6		2				70	0.54	0.4
TGCT non-sem	20	7	5	5	16	2		10	2		99	0.67	0.4
OV	58	48	49	96	28	10	40	21	5	5	316	0.79	2.4
UCEC CN high	61	43	86	90	32	18	31	13	5	5	296	0.67	1.9
UCEC CN low	37	9	95	10	14	54	10	7	1	5	42	0.15	2.1
UCEC MSI-POLE	71	31	98	42	64	70	30	55	31	19	30	0.08	71.2
UCS	61	70	79	91	54	18	27	16	4	4	247	0.71	3.5
CESC Adeno	63	21	56	19	30	14	16	14	21	5	95	0.36	3.6
CESC Squamous	32	19	59	12	35	12	5	33	11	10	101	0.44	5.2
SKCM	94	77	33	28	27	23	10	25	7	1	131	0.53	22.1
SARC DDLPS	43	83	20	85	17	15	7	9	7		450	0.36	1.1
SARC LMS	31	55	33	71	14	11	4	4	1	4	177	0.69	1.8
SARC MFS/UPS	48	74	32	68	34	20	8	21	6	4	328	0.66	3.0
SARC other	25	30	15	5	5	5		10			104	0.33	1.2
DLBC	24	76	8	19	70	70	14	35	14		90	0.29	3.5
LAML	49	17	3	9	18	11	2	3	1	1	28	0.05	1.1
	46	45	33	29	23	15	11	10	7	1			
RTK/RAS													
Cell cycle													
P13k													
p53													
Notch													
Wnt													
Myc													
Hippo													
TGFβ													
Nrf2													

Sanchez-Vega, F., et al., Cell 2018

RTK signaling in cancer



Genetic mechanisms of RTK activation in cancer



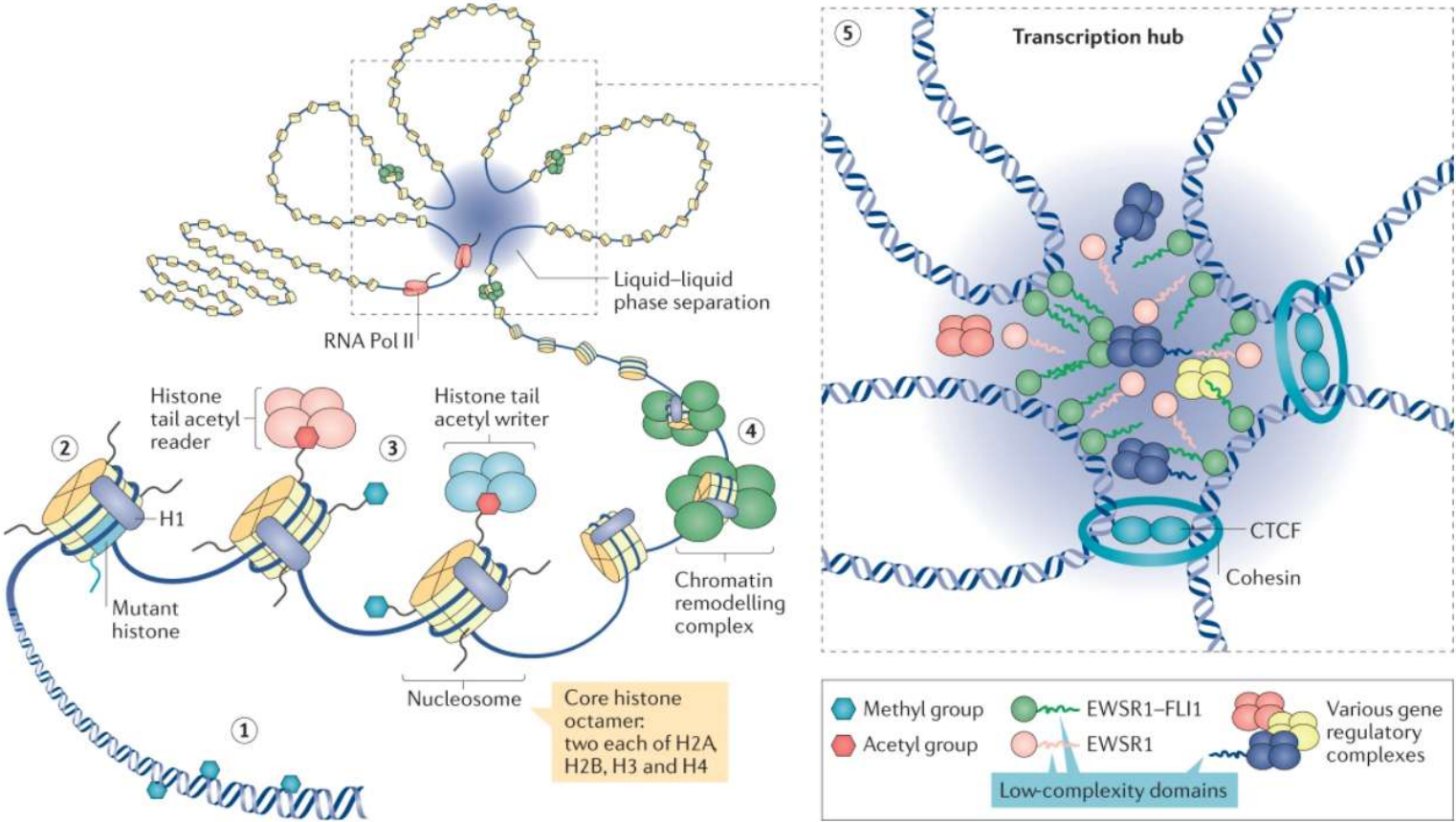
Genetic alterations

- Amplification
- Fusion
- Mutations
- Multiple

Sanchez-Vega F., et al., Cell 2018

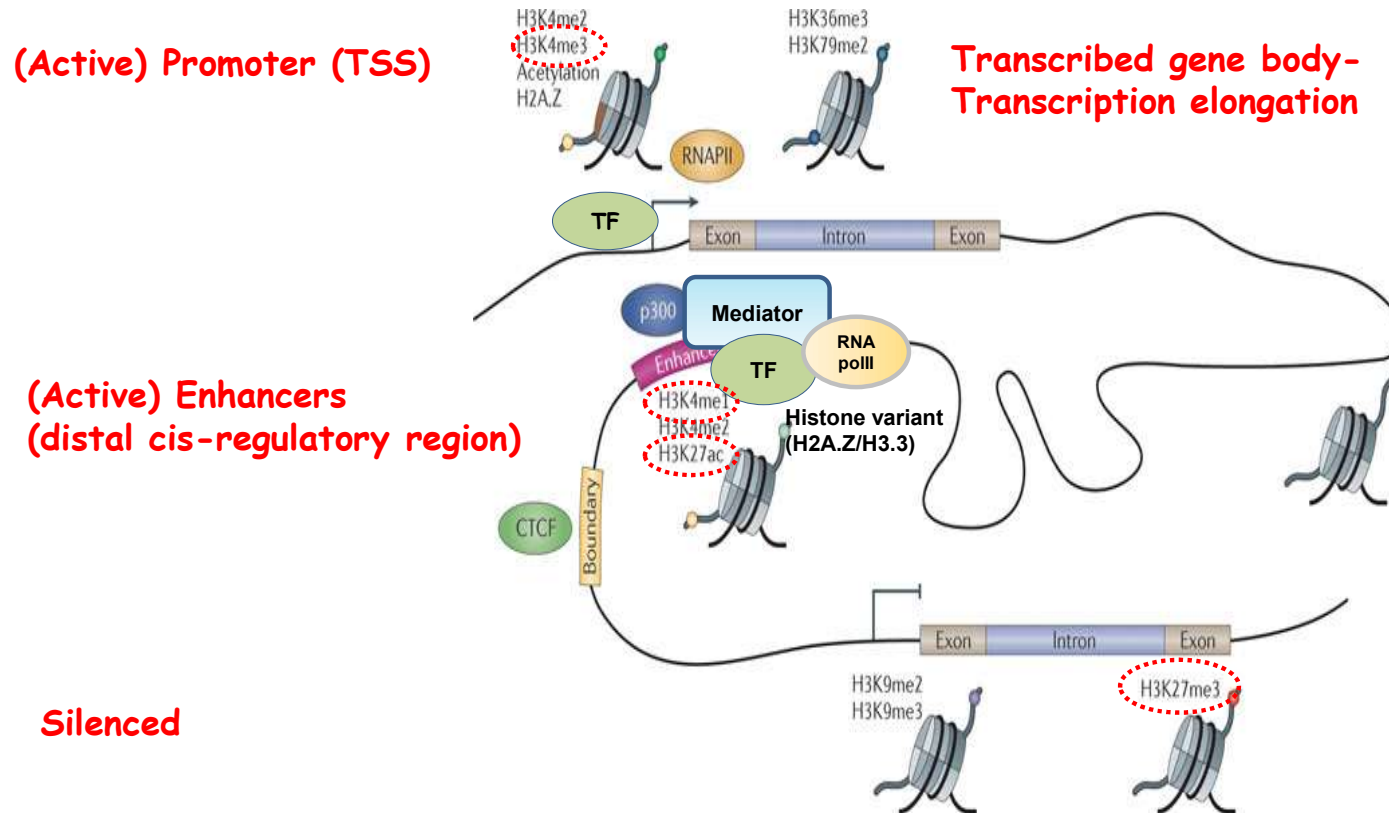


Epigenetic control of transcription



Nacev B et al., The epigenomics of sarcoma, NRC 2020

Features of transcription modules



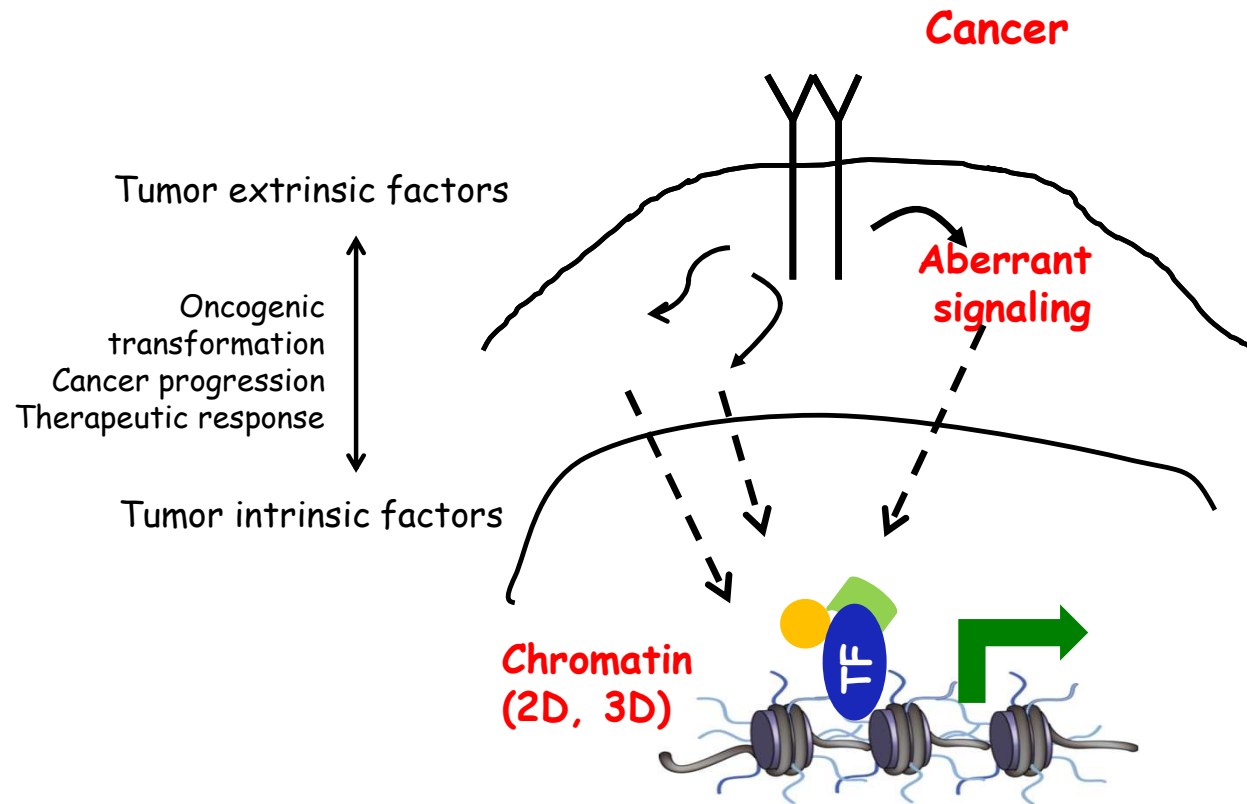
Nature Reviews | Genetics

Adapted Zhou, Goren and Bernstein, NRG 2011; Calo and Wysocka, Mol Cell 2013



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Genetic-epigenetic interaction in cellular context-dependent tumorigenesis



Three main themes:

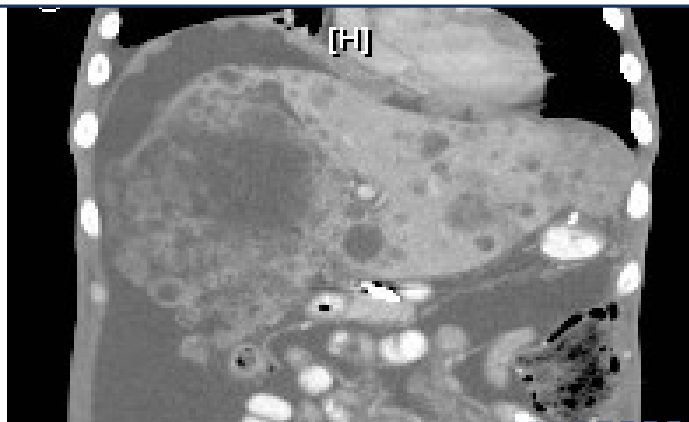
- Oncogenic TF networks.
- Chromatin modifiers complex perturbations.
- 3D chromatin organization and alternative promoter usage in cancer.



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Sarcomas (GIST, MPNST, Angiosarcomas, etc.),
Melanomas (uveal, cutaneous), others

Gastrointestinal stromal tumor (GIST)



Management:

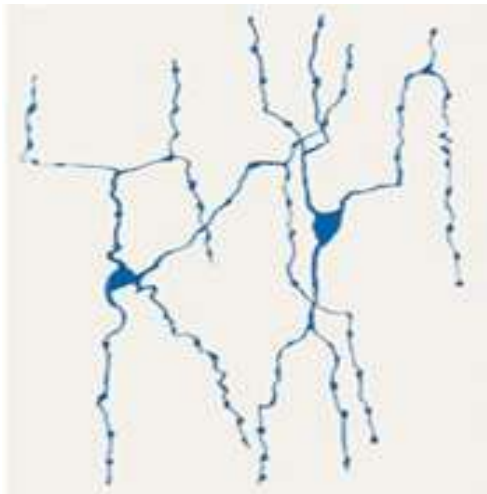
- Surgery mainstay treatment
- Recurrence or metastatic disease - fatal
- Refractory to chemotherapy and radiation
- Imatinib is the first line therapy for past 25 years

- ~5,000 cases diagnosed per year in the US.
- One of the most common subtypes of soft tissue sarcomas, the most common mesenchymal neoplasm in the GI tract.
- Can arise anywhere from the entire GI tract; stomach is the most common primary site (2/3), then small bowel (1/4), esophagus/colon/rectum (the rest).
- Originates from Interstitial Cells of Cajal (ICCs).
- The majority of GISTs are characterized by KIT or PDGFRA activating mutations, other mutations include BRAFV600E, NF1 loss, SDH-deficiency, etc.
- Peak incidence 50-65-year-old for sporadic GIST. Familial syndromes with early onset.

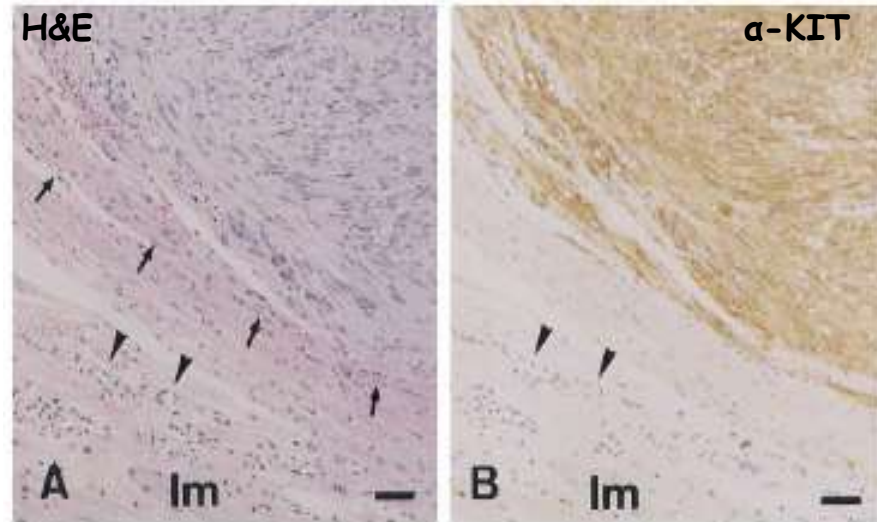
GIST originates from ICC naturally express KIT

- Originates from the Interstitial Cells of Cajal (ICCs) of the GI tract
- Characterized by KIT positive IHC and activating mutations in KIT or PDGFRA

**Interstitial Cell of Cajal (ICC)-
Pacemaker cells of the GI tract**



GIST of stomach



Huizinga, J.D. et al., *Nature*, 1995; Hirota, S., et al., *Science*, 1998



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EFFICACY AND SAFETY OF IMATINIB MESYLATE IN ADVANCED GASTROINTESTINAL STROMAL TUMORS

GEORGE D. DEMETRI, M.D., MARGARET VON MEHREN, M.D., CHARLES D. BLANKE, M.D.,
ANNICK D. VAN DEN ABEELE, M.D., BURTON EISENBERG, M.D., PETER J. ROBERTS, M.D., MICHAEL C. HEINRICH, M.D.,
DAVID A. TUVESON, M.D., PH.D., SAMUEL SINGER, M.D., MILOS JANICEK, M.D., PH.D., JONATHAN A. FLETCHER, M.D.,
STUART G. SILVERMAN, M.D., SANDRA L. SILBERMAN, M.D., PH.D., RENAUD CAPDEVILLE, M.D., BEATE KIESE, M.Sc.,
BIN PENG, M.D., PH.D., SASA DIMITRIJEVIC, PH.D., BRIAN J. DRUKER, M.D., CHRISTOPHER CORLESS, M.D.,
CHRISTOPHER D.M. FLETCHER, M.D., AND HEIKKI JOENSUU, M.D.

N Engl J Med, Vol. 347, No. 7 · August 15, 2002

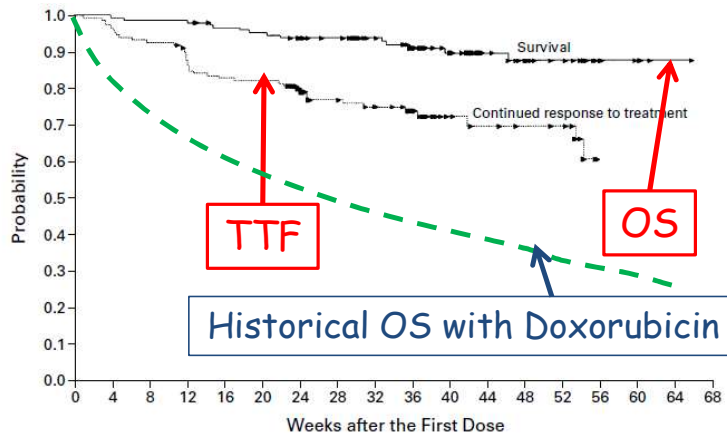
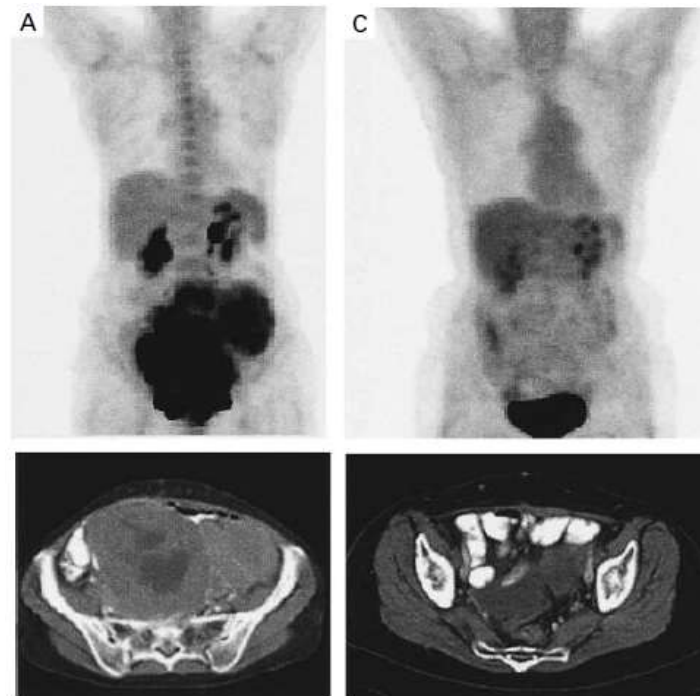


Figure 1. Kaplan-Meier Estimates of Overall Survival and Time to Treatment Failure for All Patients. Each arrowhead represents the point at which a patient's data were censored.



EORTC-62005
Phase III Trial (n = 377)⁶⁹

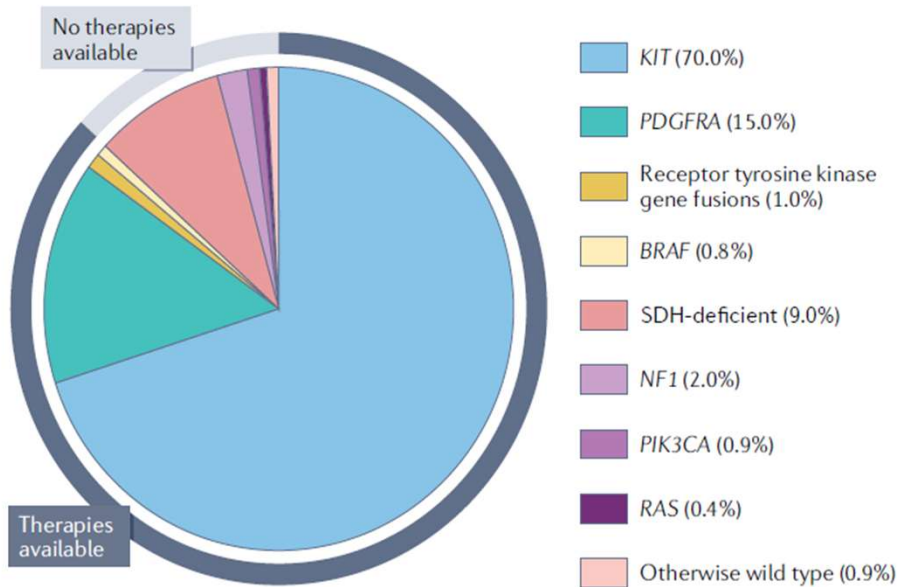
SWOGS0033/CALGB150105
Phase III Trial (n = 428)⁷⁰

**Imatinib (Gleevec)-FDA approved as 1st line therapy for GIST 2002!
1st targeted therapy in solid tumor.**

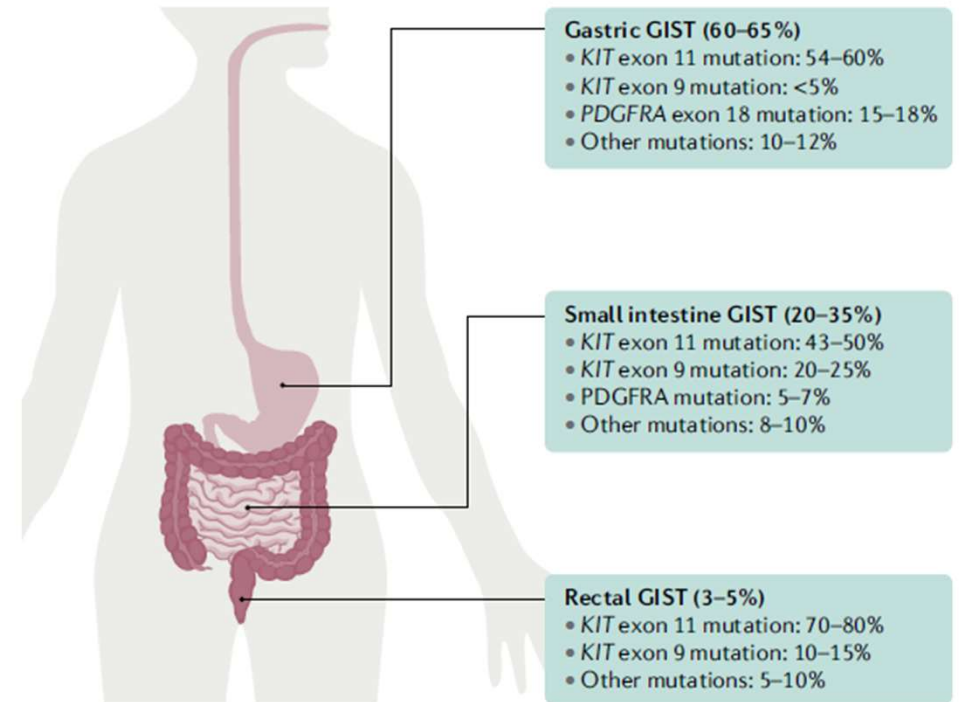


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Molecular characterization of GIST



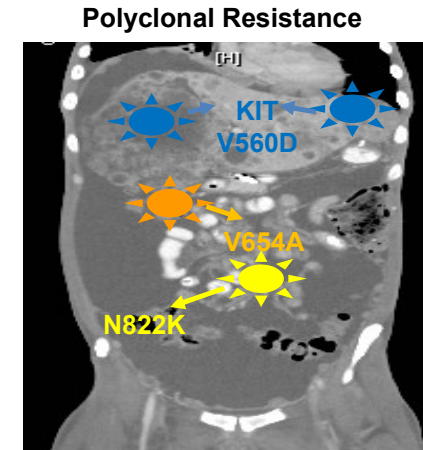
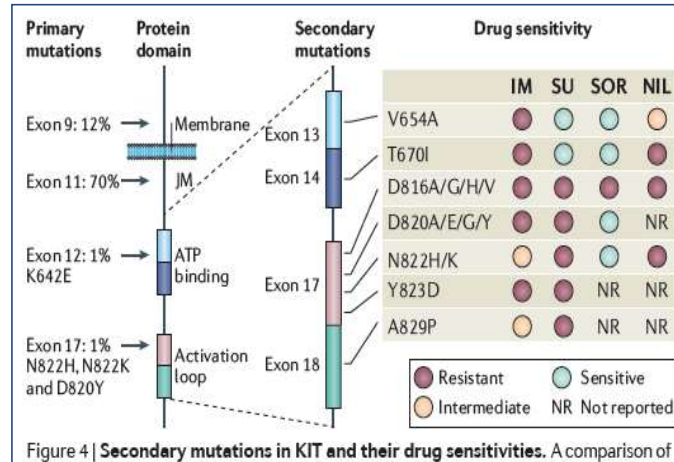
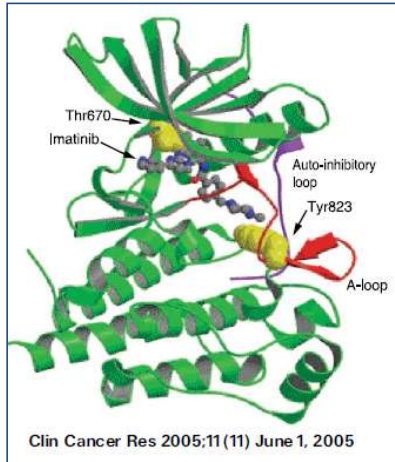
Klug LR et al., Nat Rev Clin Oncol, 2022



Blay JY et al., Nat Rev Dis Primers, 2021

Clinical challenges- heterogeneous imatinib resistance mechanisms

14% - Primary resistance; 50% - Develop imatinib resistance within 2 years



Resistance Mechanisms:

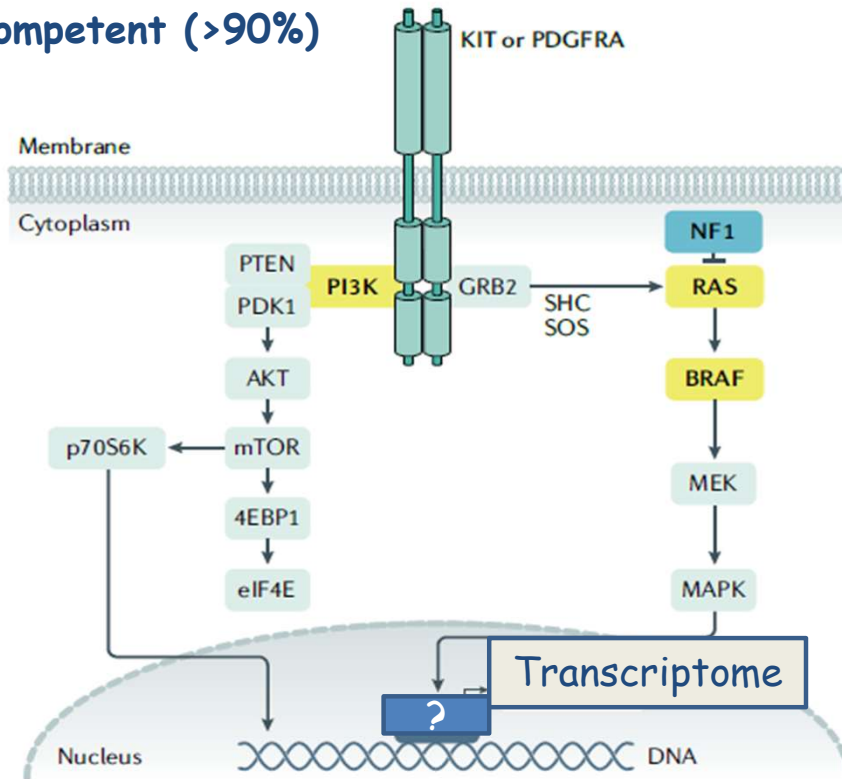
1. Various secondary mutations (50-65%)
2. Genomic Amplification of RTKs
3. Activation alternative signaling pathways
4. Kit-low, imatinib-resistant GIST stem/progenitors
5. Tumor adaptation and persistence of disease
6. Tumor heterogeneity and polyclonal resistance
7. Others...



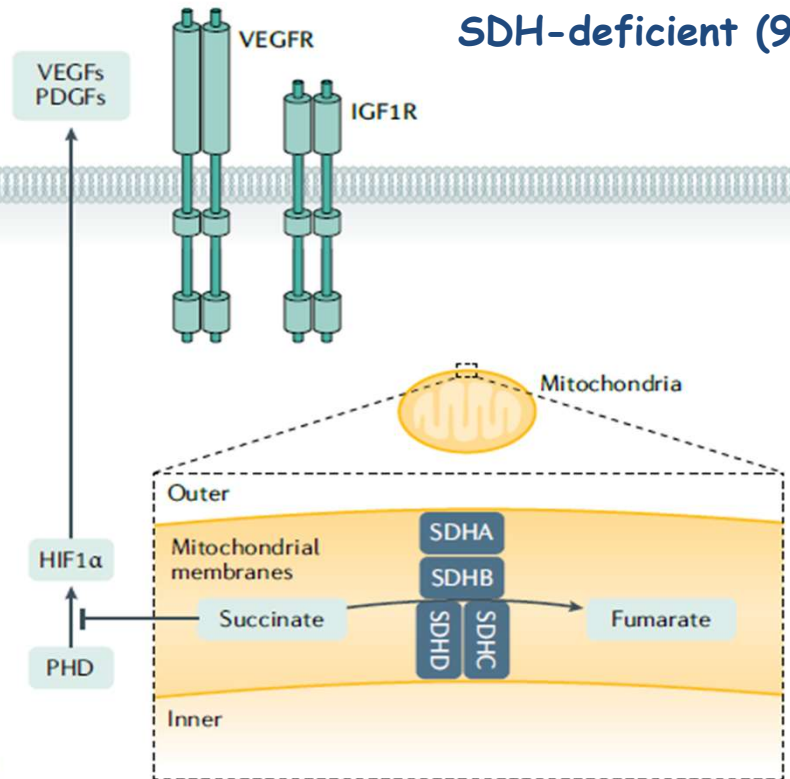
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Molecular pathogenesis of GIST

SDH-competent (>90%)

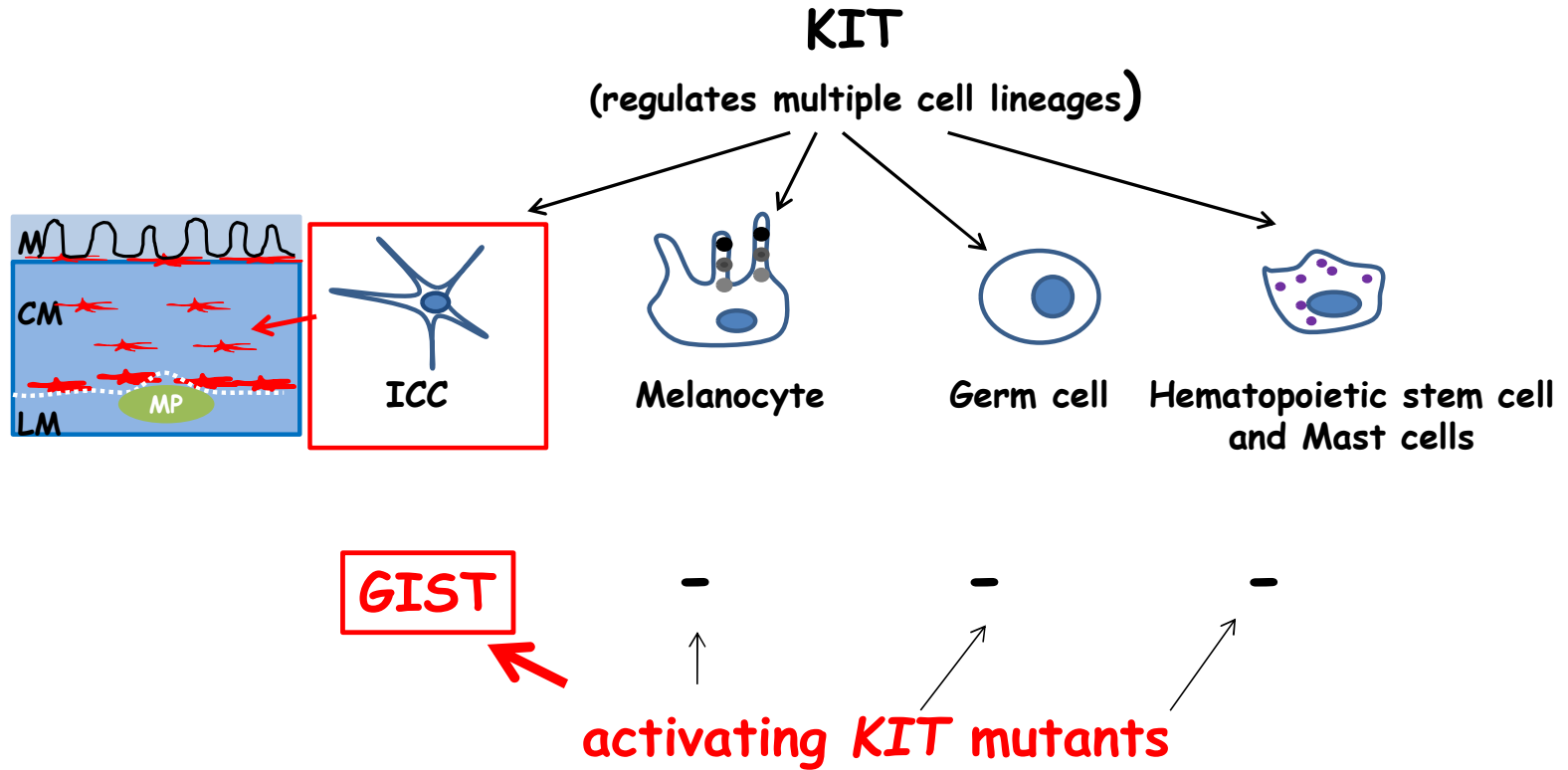


SDH-deficient (9%)

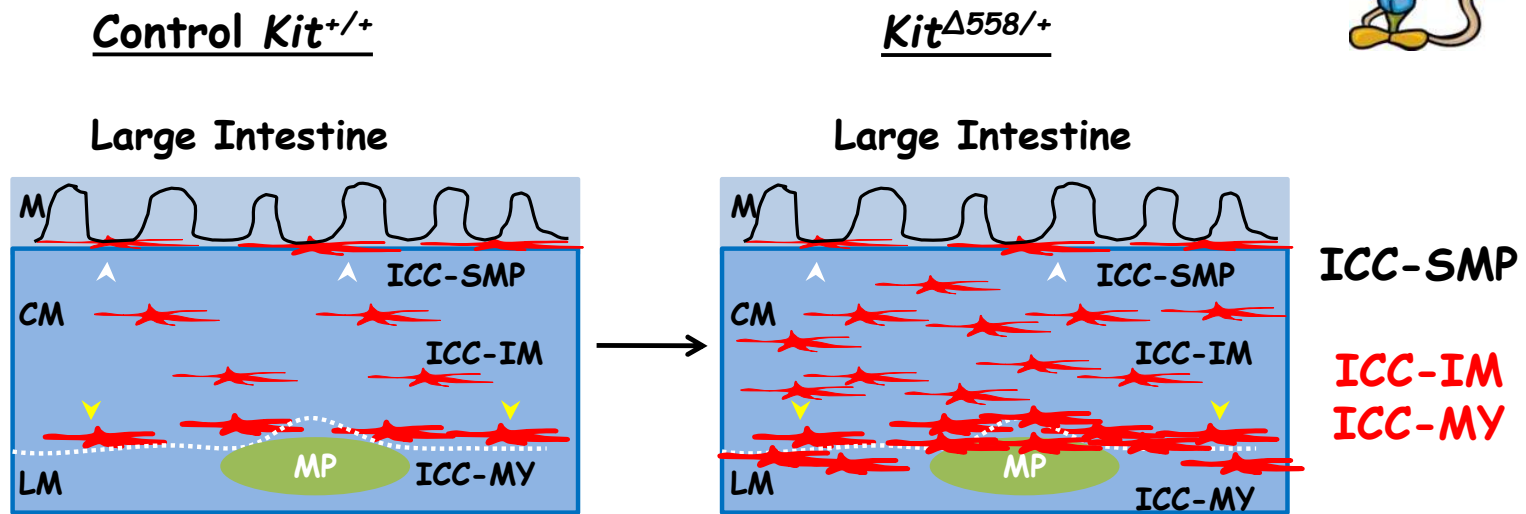


Corless, et al, Nat Rev Cancer, 2011; Blay JY et al., Nat Re Dis Primers, 2021

Familial gastrointestinal stromal tumor (GIST)

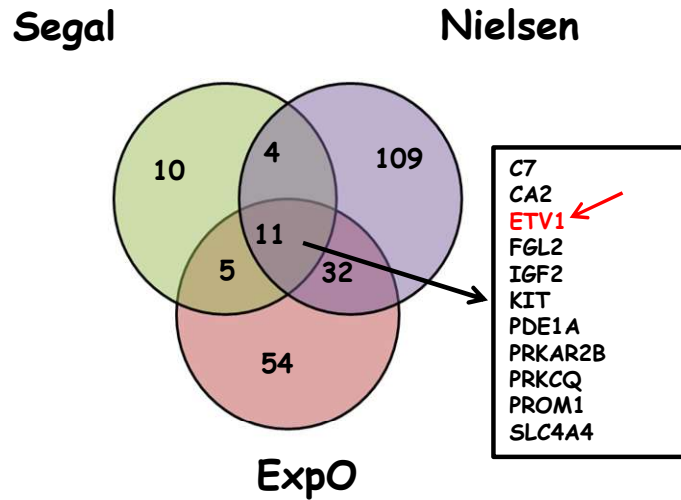


Familial GIST mouse models (Kit V558del, K641E, D818Y)

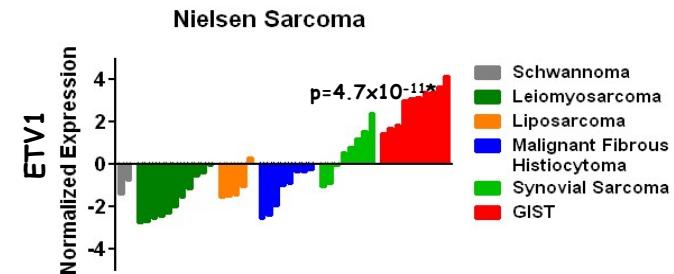
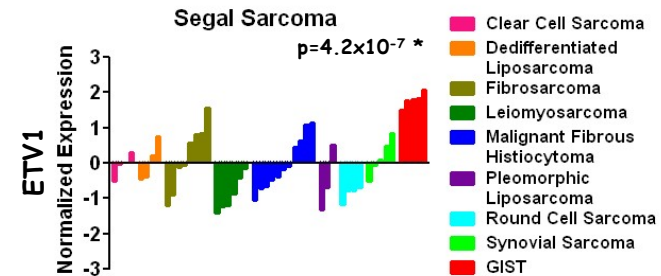
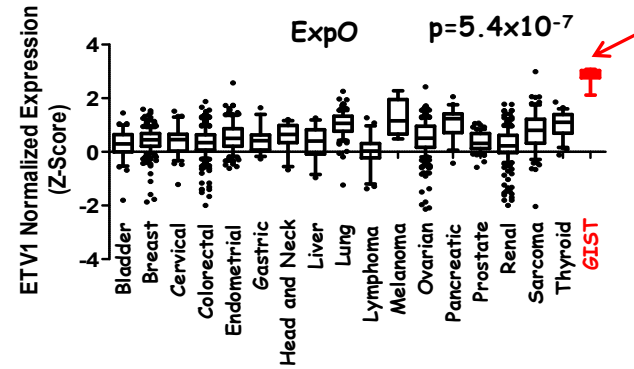


Sommer G et al, PNAS 2003; Rubin BP et al, Cancer Res 2005; Nakai N et al, J Pathol 2008; Kwon JG et al, Gastroenterology 2009

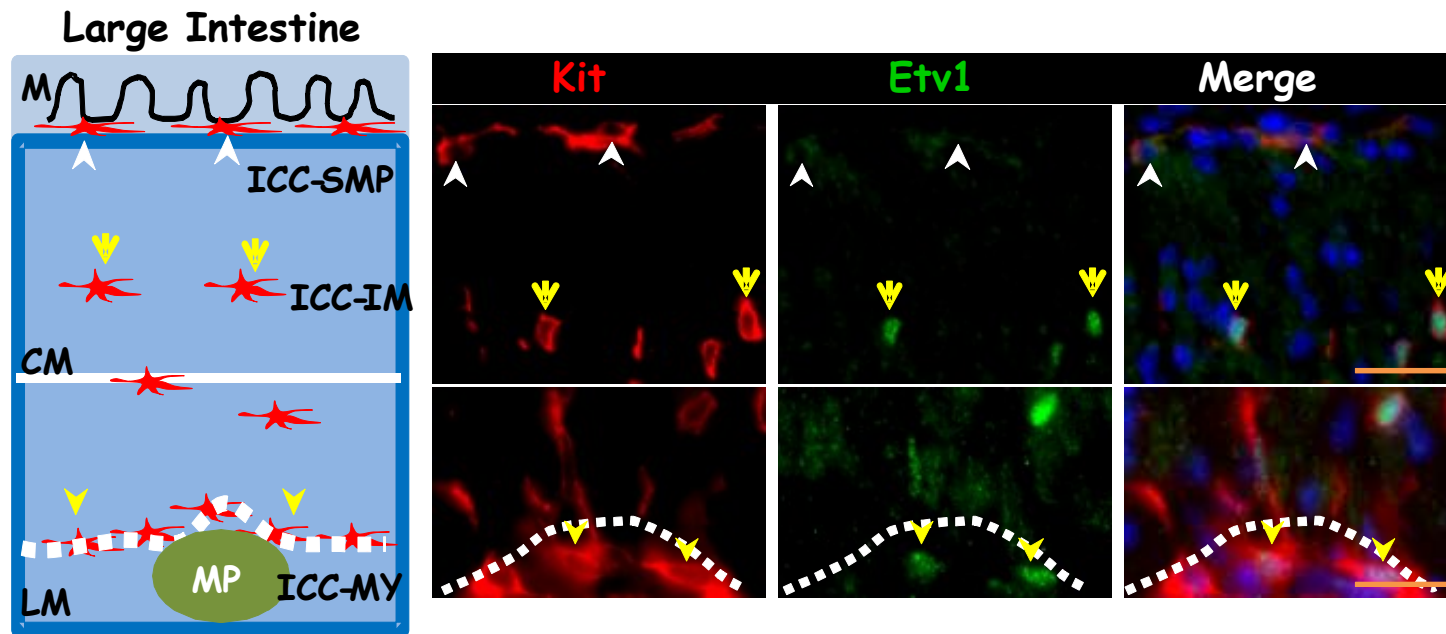
ETV1 is differentially highly expressed in GIST



$q < 0.05$
 $Z \text{ difference} > 1.5$



ETV1 is expressed in specific ICCs susceptible to oncogenesis

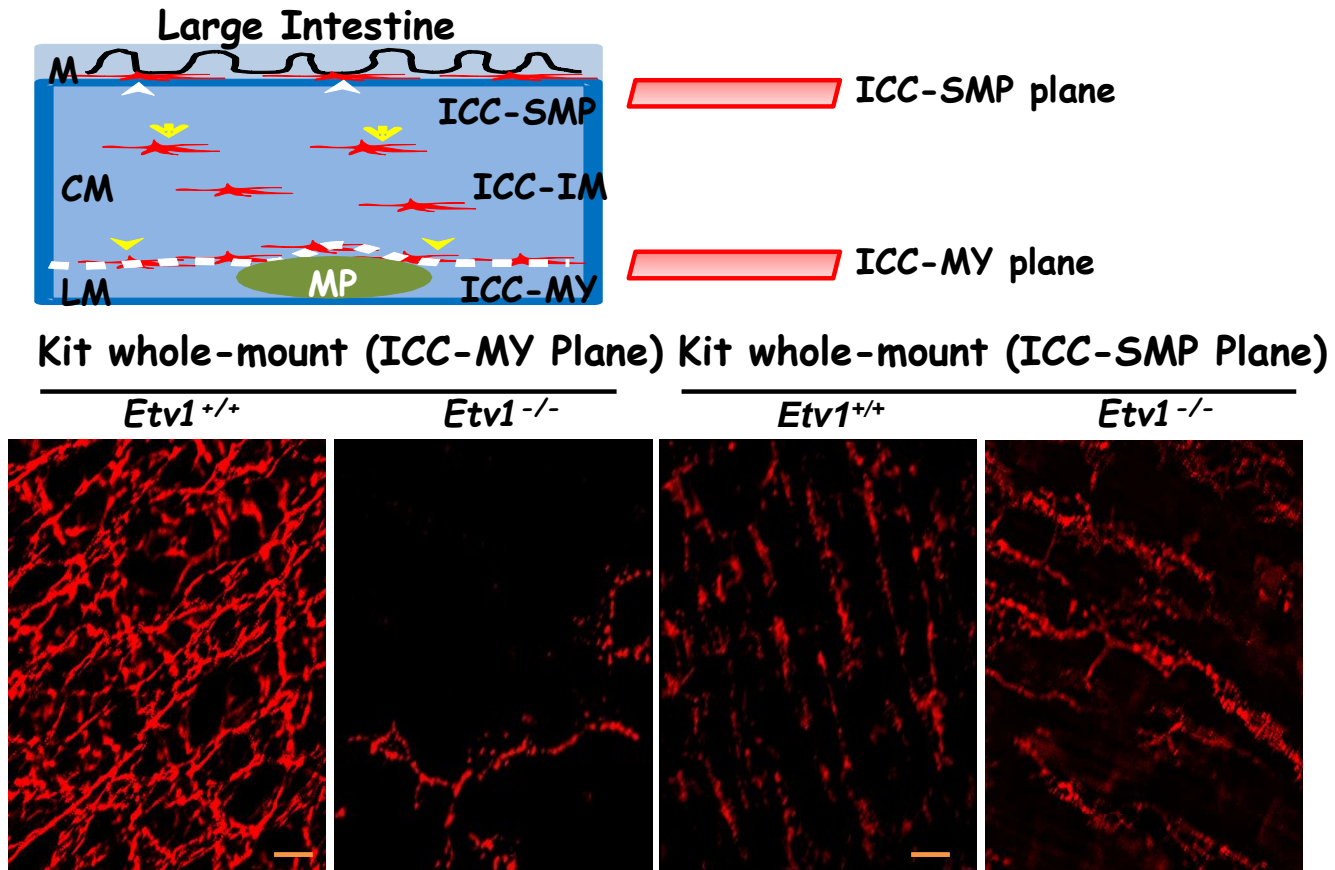


Chi P, Chen Y, et al, Nature, 2010



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ETV1 is required for the development of ICC-IM /ICC-MY networks



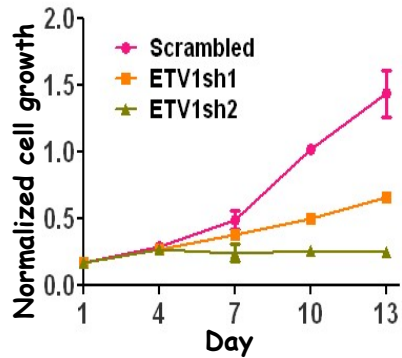
Chi P, Chen Y, et al, Nature, 2010



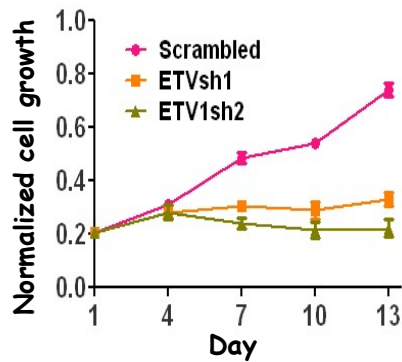
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ETV1 is required for GIST growth and survival

GIST882 cell (imatinib sensitive)

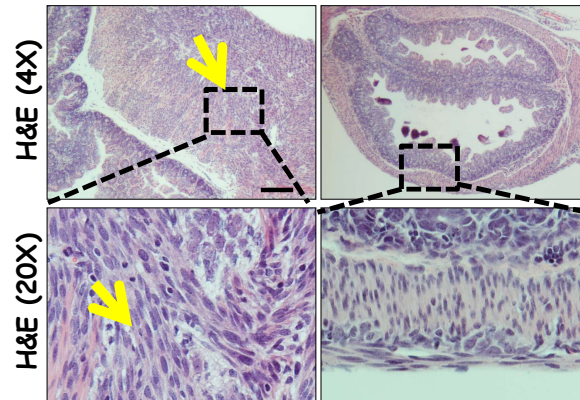


GIST48 cell (imatinib resistant)



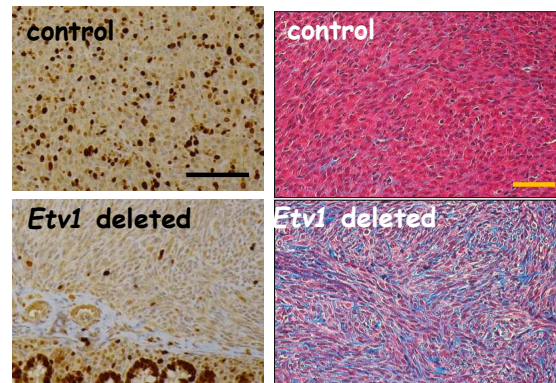
GEMM: *Kit*^{Δ558/+}

Cecum (*Etv1*^{+/+}; *Kit*^{V558Δ/+}) Cecum (*Etv1*^{-/-}; *Kit*^{V558Δ/+})

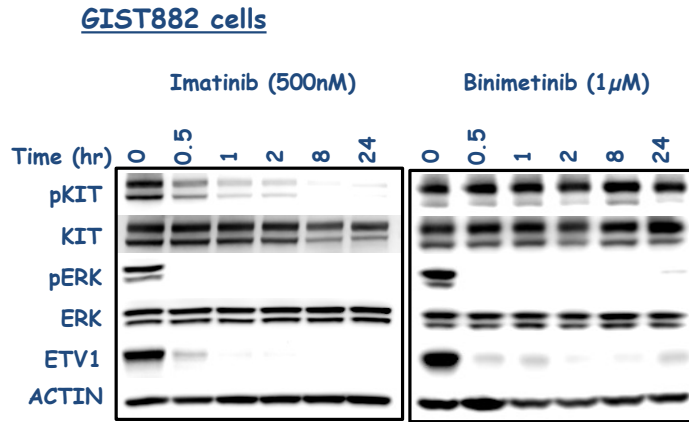


GEMM: *Kit*^{Δ558/+}; *Etv1*^{flox/flox}; *Rosa26*^{CreERT2/CreERT2}

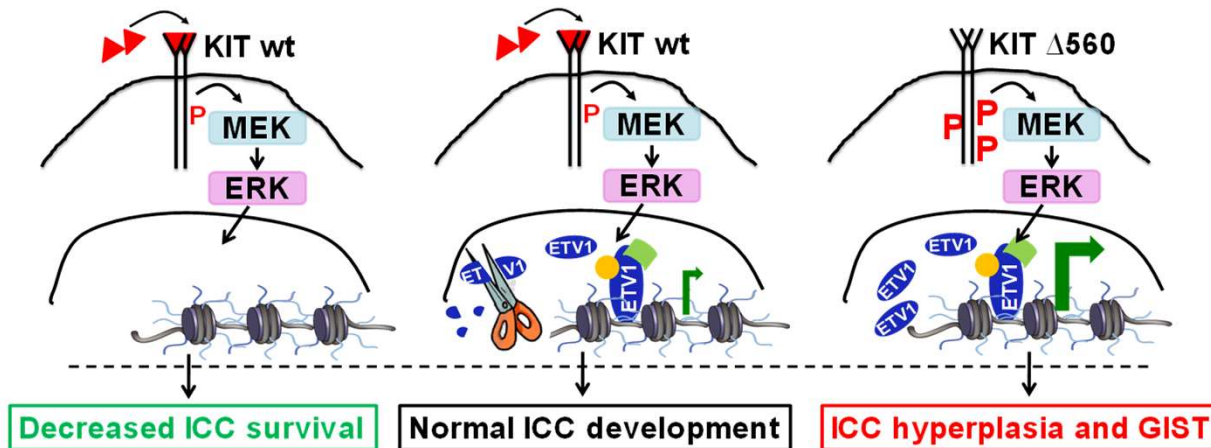
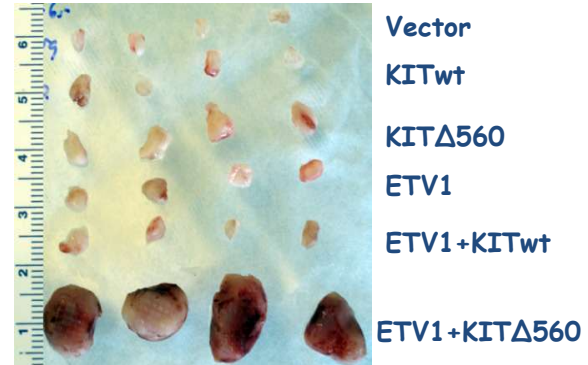
Cecal tumor (Ki67 IHC) Cecal tumor (trichrome)



Transcriptional upregulation of KIT by stabilized ETV1



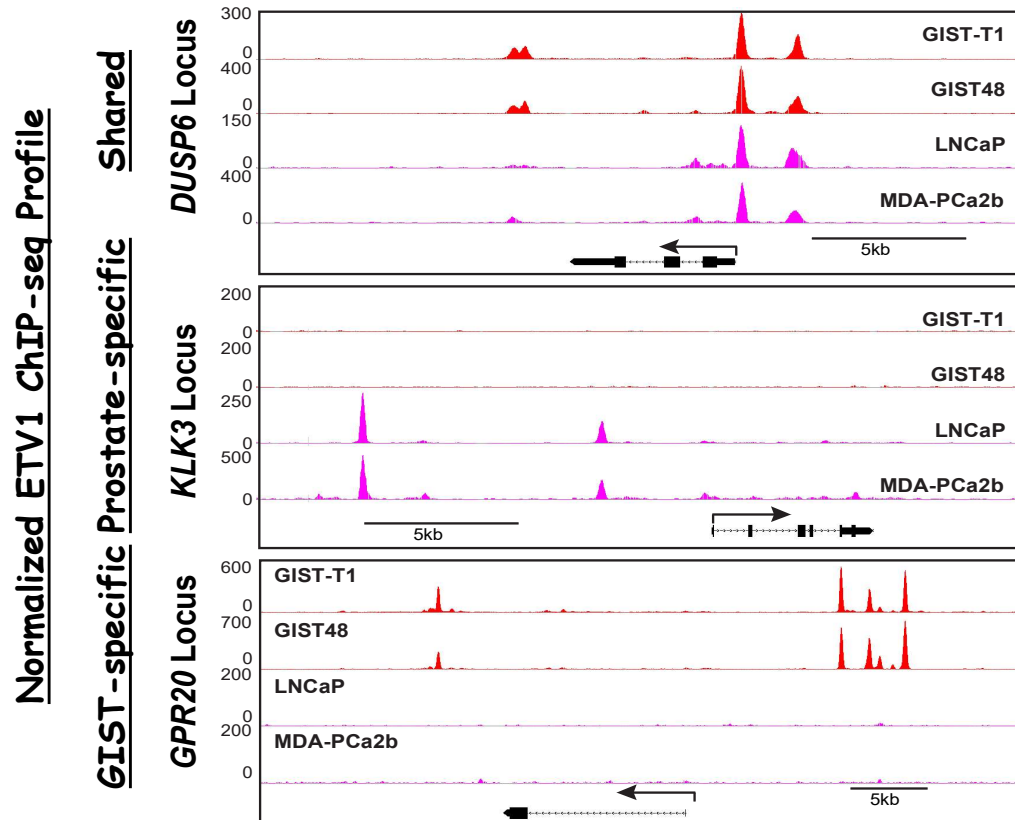
Excised 3T3 allograft tumors



Chi, P, Chen, Y et al, Nature 2010; Ran L et al., Cancer Discovery, 2015

How does ETV1 regulate lineage-specific oncogenic transcriptome in cancer?

- GIST (ICC/GIST lineage-specific)
- Prostate Cancer (Prostate lineage-specific)

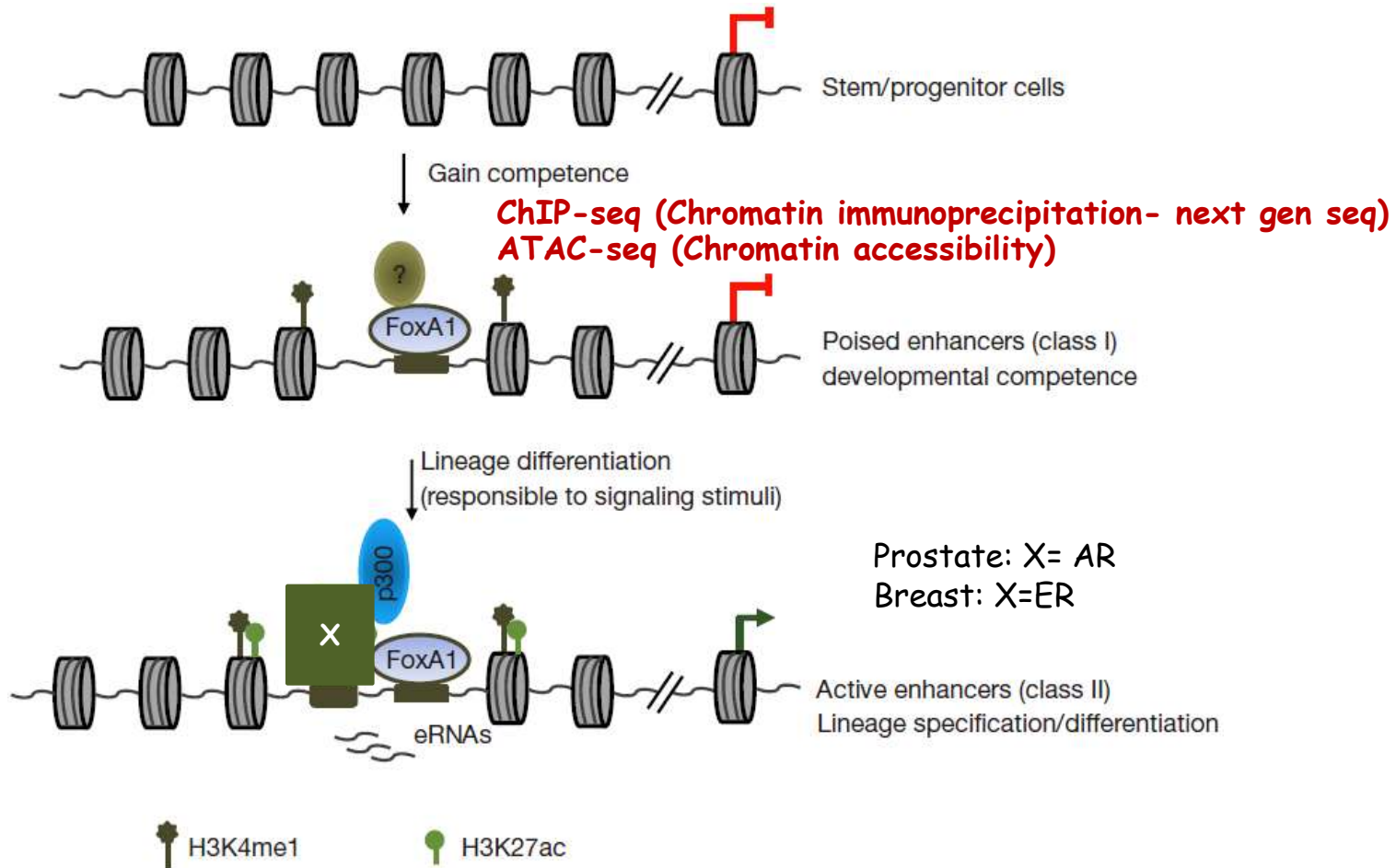


Chi P, Chen Y, et al, Nature, 2010; Chen Y, Chi P, et al., Nat Medicine 2013; Ran L, Chen Y, et al, Cancer Discovery 2015, 2018



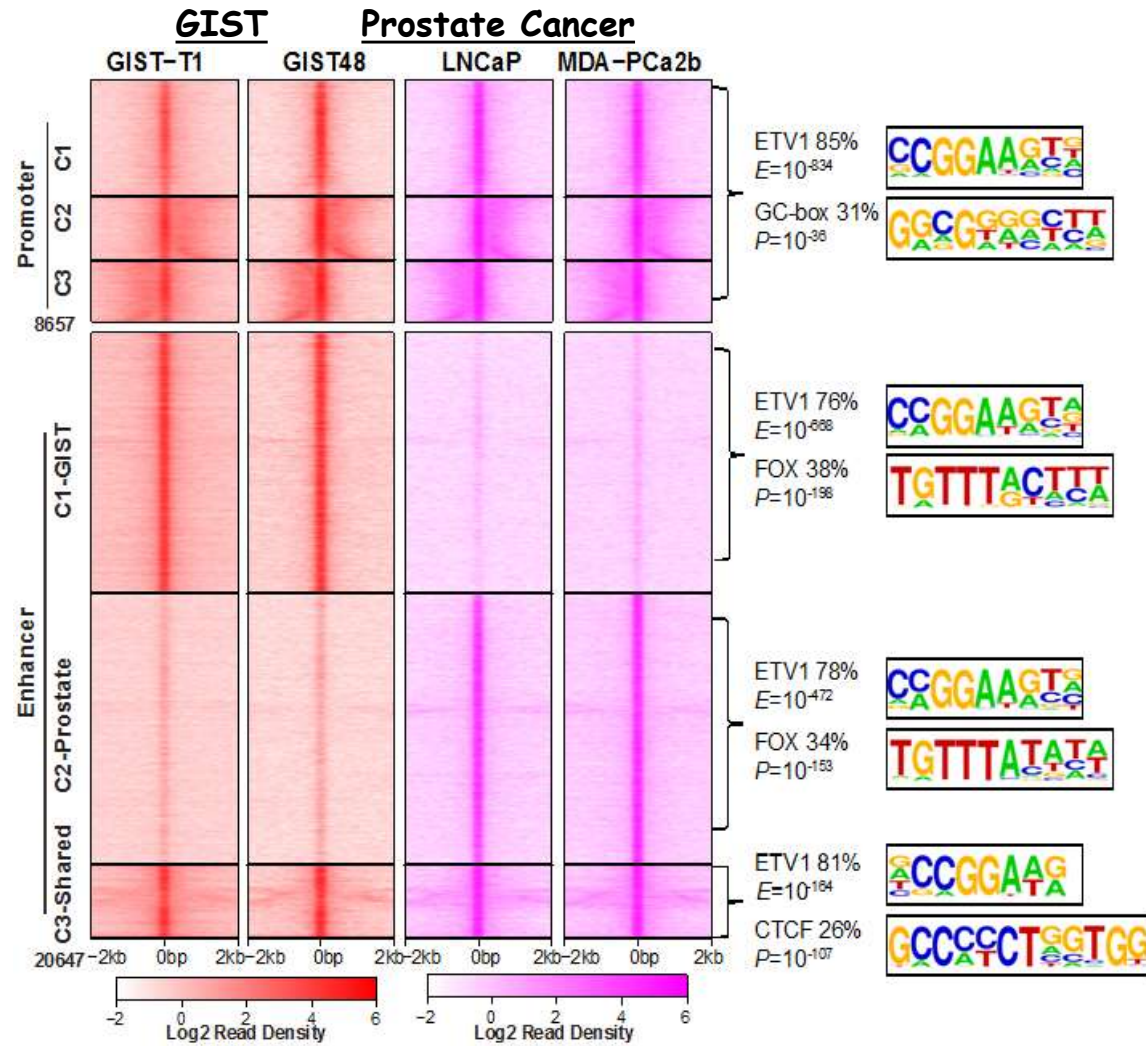
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Enhancers define cell lineage



Adapted from Li and Huang, 2016

ETV1 cistrome analysis

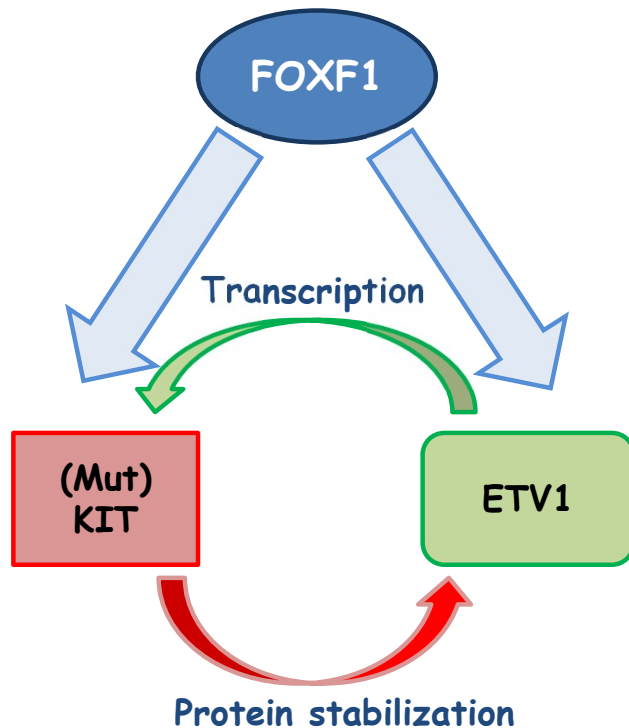


Ran L, Chen Y, et al., Cancer Discovery 2018



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FOXF1 functions as a “pioneer factor” that enforces lineage-dependency in GIST pathogenesis



Multilevel transcriptional regulation

- Direct transcriptional activation of KIT
- Transcriptional regulation of masterregulator-ETV1 (Cistrome regulation, Chromatin accessibility, integrate with transcription)

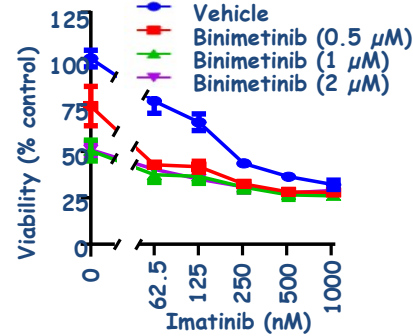
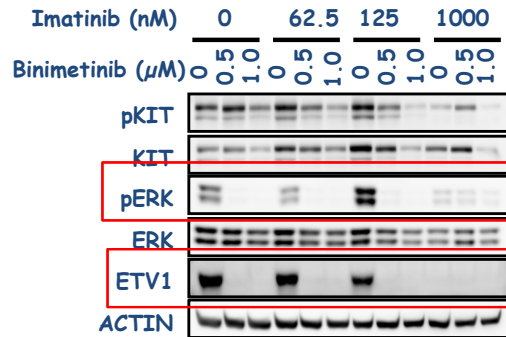
GIST-High lineage addiction

- dependent on KIT/PDGFR α signaling
- <1% dedifferentiation in treatment-refractory GIST

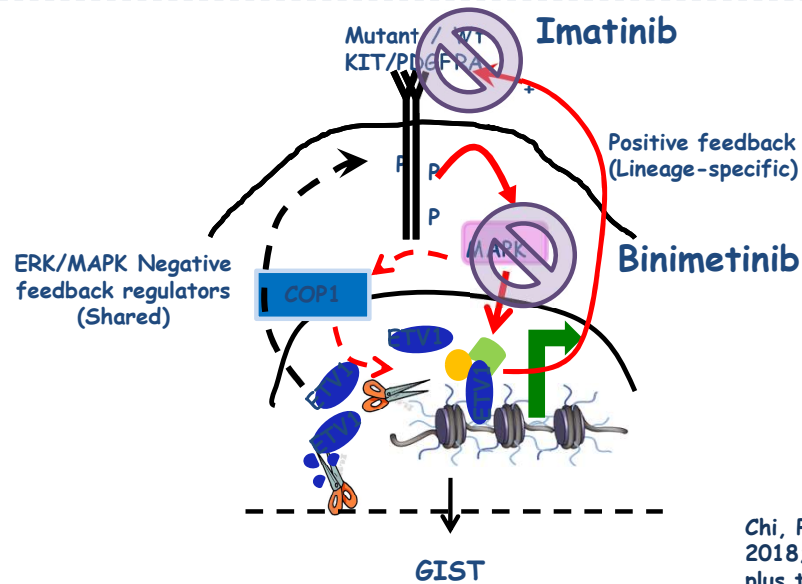
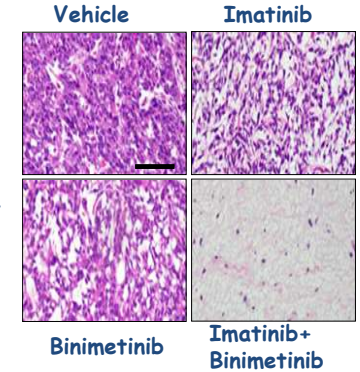
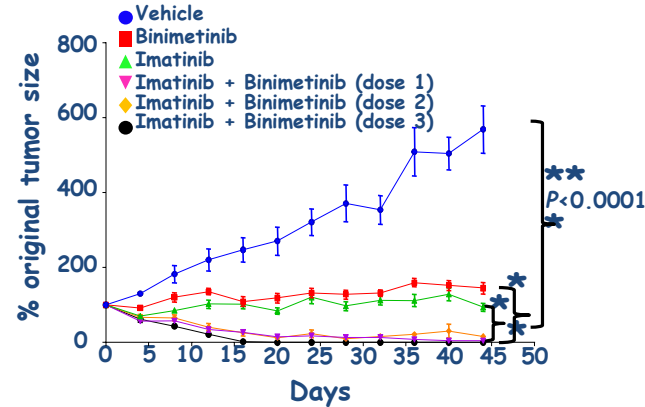


Synergy of combined MAPK and KIT pathway inhibition

GIST882



GIST882 xenografts



Advantage of targeting lineage dependence:

- Bypasses multiple upstream resistance mechanisms
- Break the positive feedback circuit of ETV1/KIT-target KIT expression regardless of mutations
- Block early adaptation and forestall resistance development and induces enhanced depth of cytotoxicity

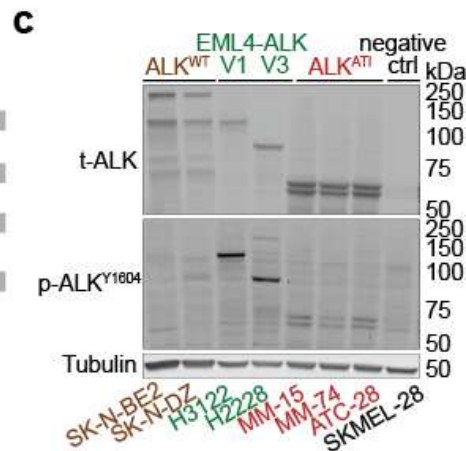
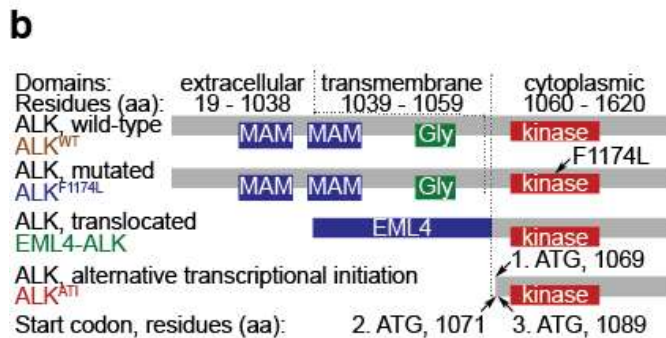
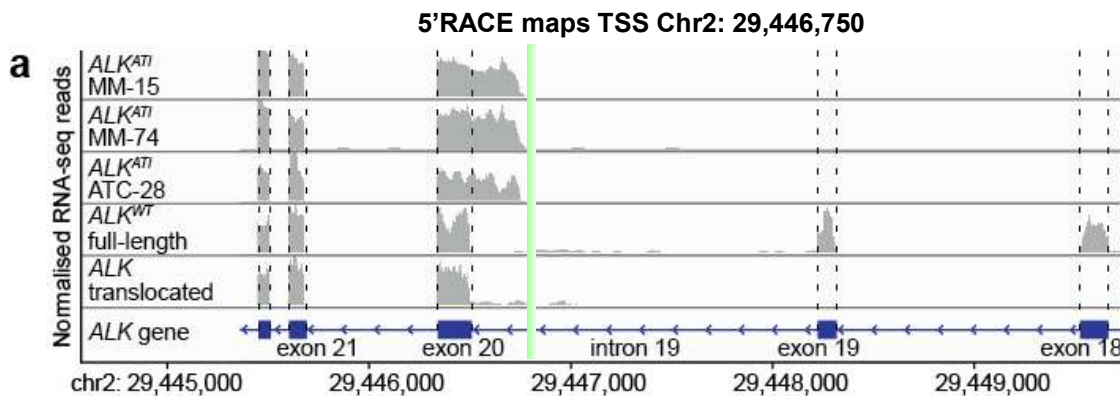
A phase Ib/II trial of imatinib plus binimetinib in advanced GIST - Positive trial ! Combination is also effective in SDH-deficient GIST

Chi, P, Chen, Y et al, Nature 2010; Ran L et al., Cancer Discovery, 2015, 2018; Xie et al, JCI 2018; Gupta A et al., Mol Cancer Ther, 2021 (riporetinib plus trametinib); Chi P et al., CCR 2022; Chi P et al., JCO 2022.



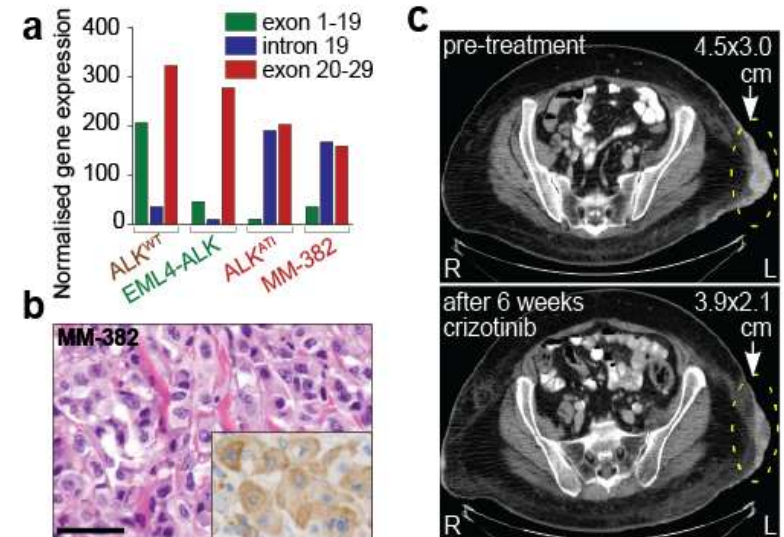
Memorial Sloan Kettering Cancer Center.

A novel oncogenic ALK^{ATI} arises through *de novo* alternative transcriptional initiation



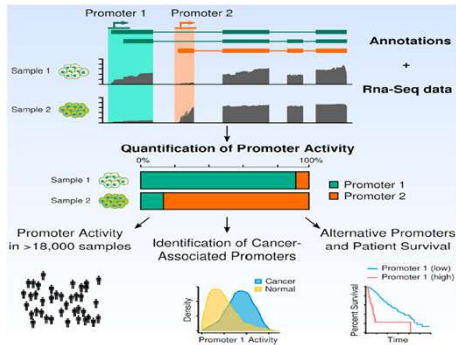
Alternative transcription initiation leads to expression of a novel oncogenic ALK isoform in cancer

- ALK^{ATI} present in 3.4% of all TCGA tumors; 12% of melanomas and sporadically other cancers.
- ALK^{ATI} functions as an independent driver, can co-occur with *BRAF/NRAS/KIT* - mutant melanoma.



Drs. Klaus Busam, Michael Postow & Charlotte Ariyan

Alternative promoter usage in cancer



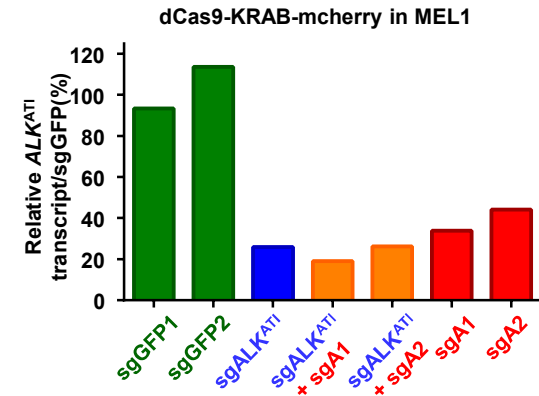
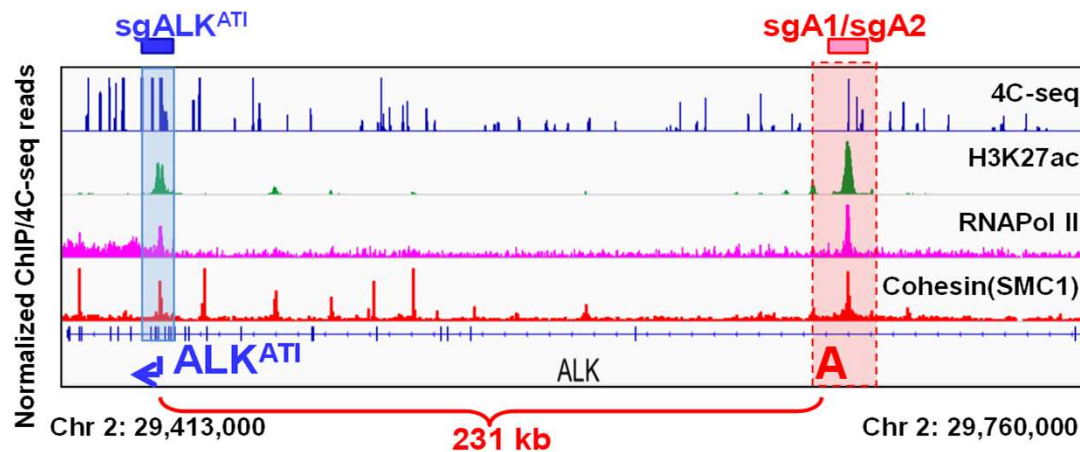
Demircioglu D., et al, Cell 2019

Pan-cancer transcriptome analysis reveals pervasive regulation through **alternative promoters**

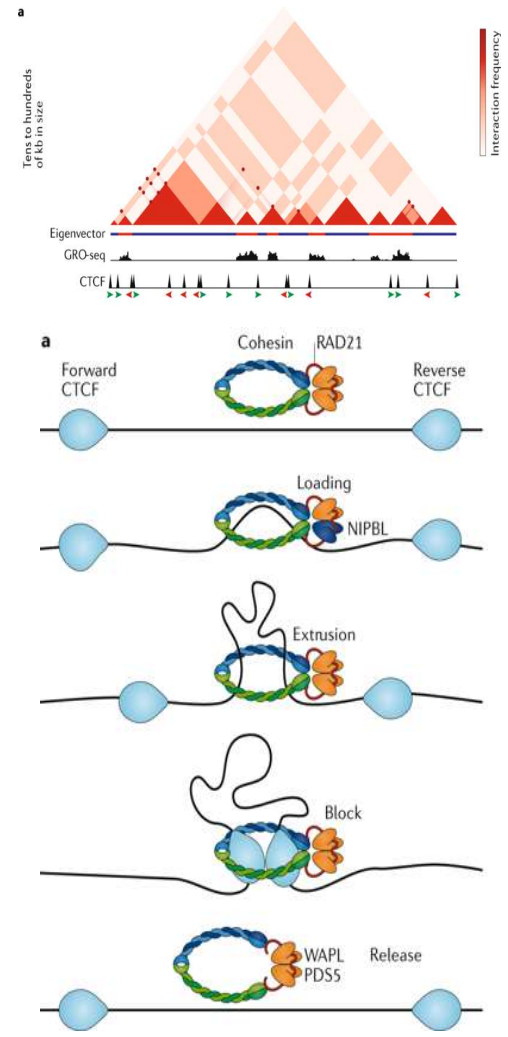
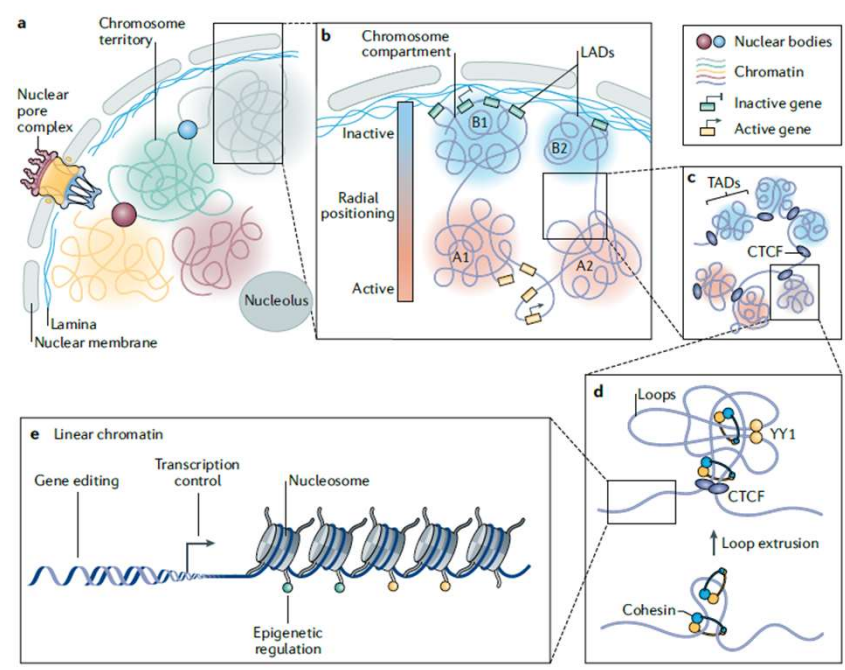
- Estimation of promoter activity in 18,468 RNA-seq, 42 cancer types
- Alternative promoters display tissue-specific regulation and impact isoform diversity
- Cancer associated promoters alter the transcriptome independent from gene expression
- Patient-to-patient variation in alternative promoter is associated with survival

What regulates alternative promoter usage in cancer?

ALK^{ATI} transcriptional activation is regulated by **epigenetic mechanisms** (e.g., biallelic expression, no genomic alteration at the ALK locus).

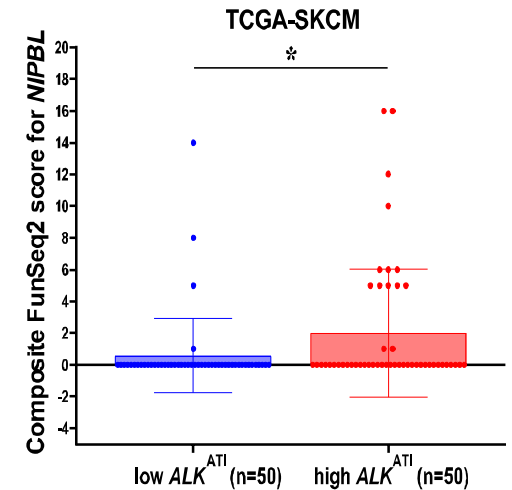


Cohesin and CTCF establish and maintain the 3D genome in the nucleus



Pan-cancer driver genes and mutations analysis identifies **NIPBL** as a TSG (3.4%)

Bailey MH et al., Cell 2018

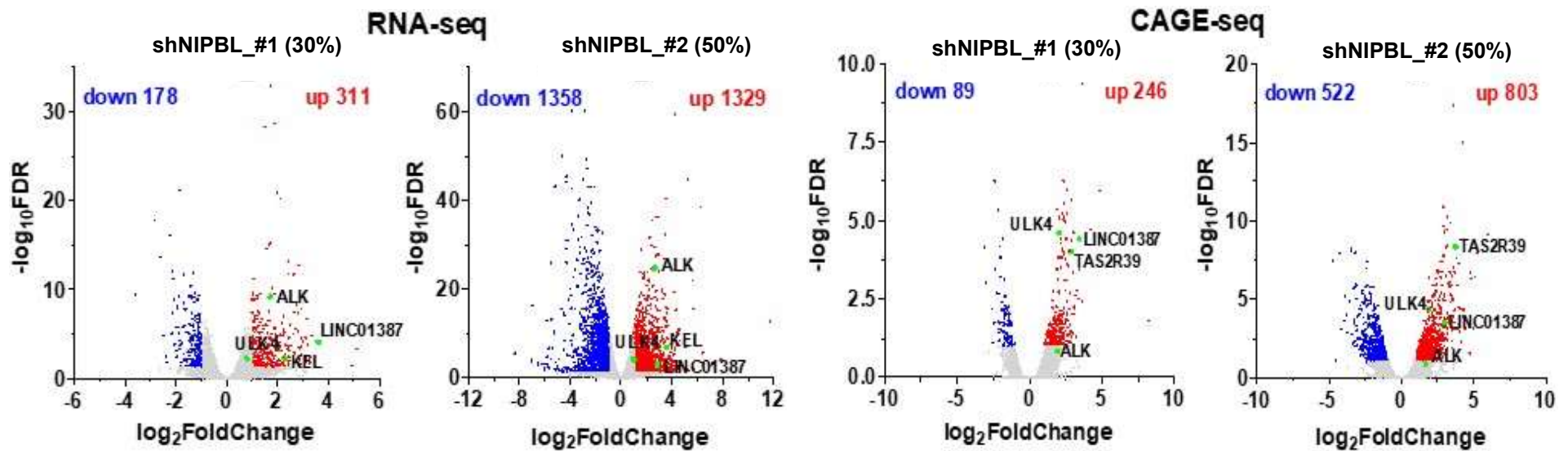


Fu Y et al., Genome Biol 2014

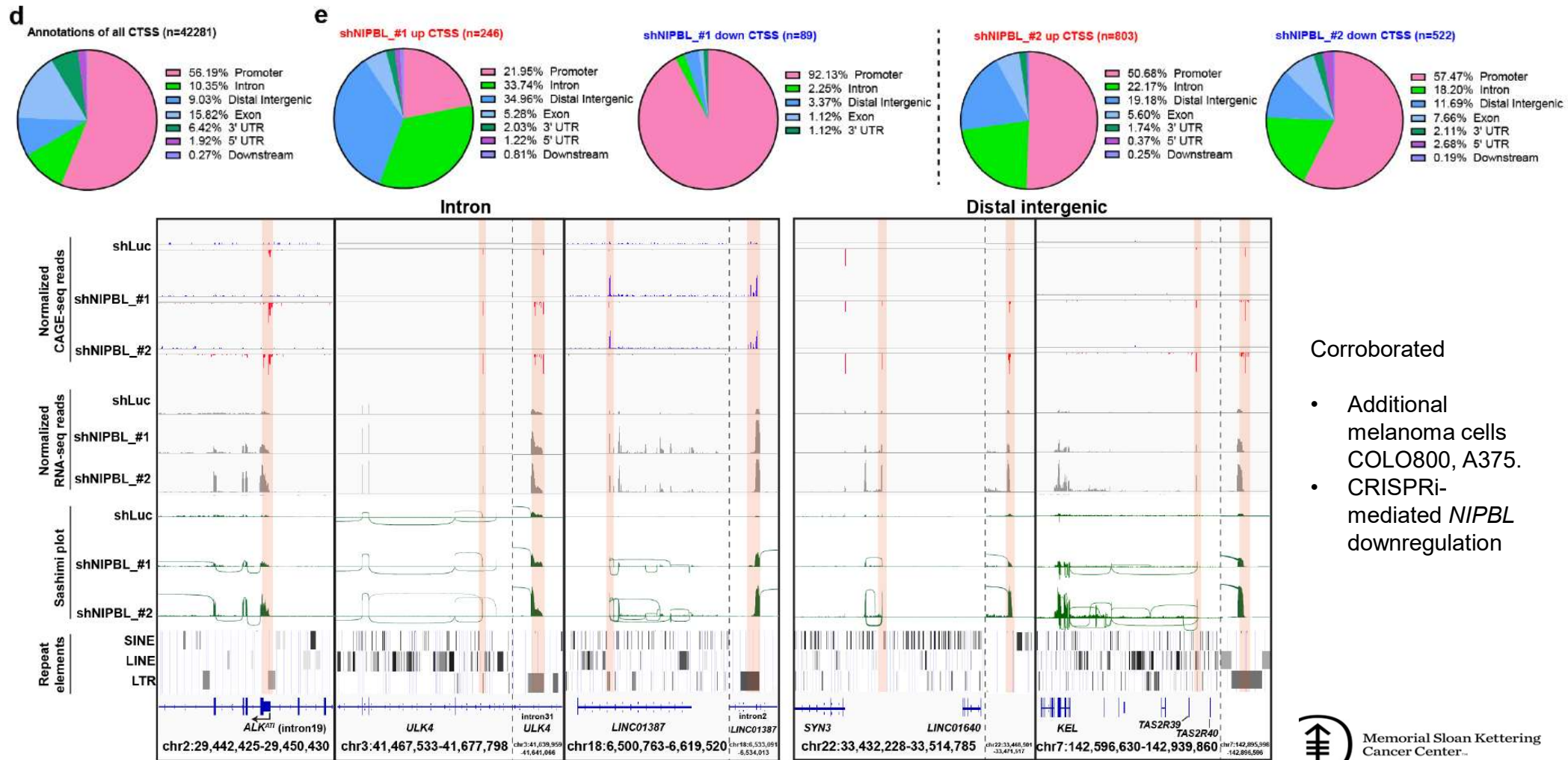
Compartments: 10-100 Mb
TADs: ~200kb-2Mb
Loops: ~10kb-1Mb

Rowley and Corces, NRG 2018
Wang H, Han M and Qi LS, NRG 2021

Haploinsufficiency of *NIPBL* leads to global alternative promoter usage



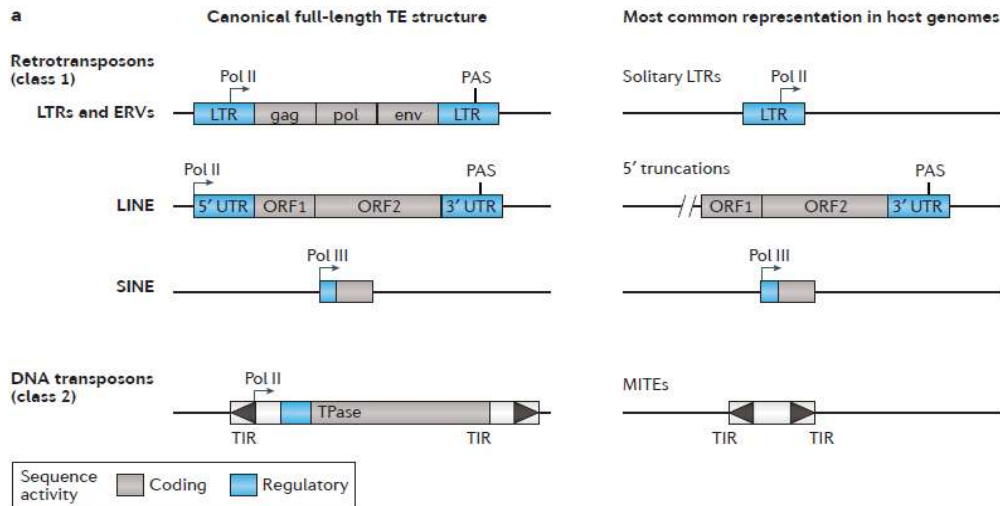
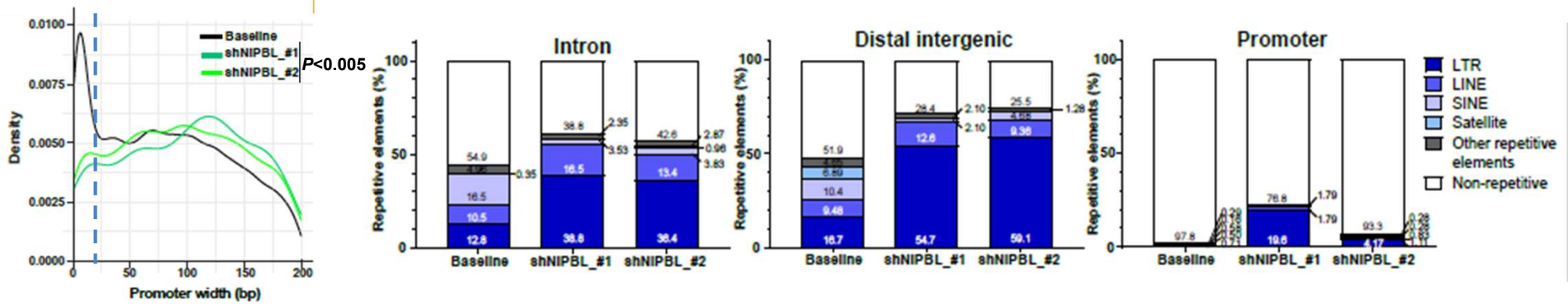
NIPBL perturbation leads to alternative promoter activation in introns and intergenic regions



Corroborated

- Additional melanoma cells COLO800, A375.
- CRISPRi-mediated *NIPBL* downregulation

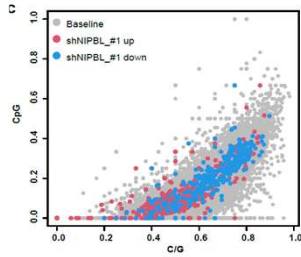
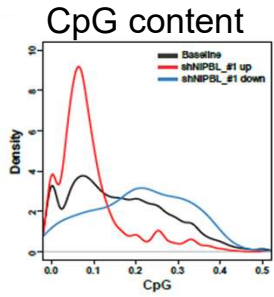
NIPBL perturbation leads to alternative promoter usage arising from LTR



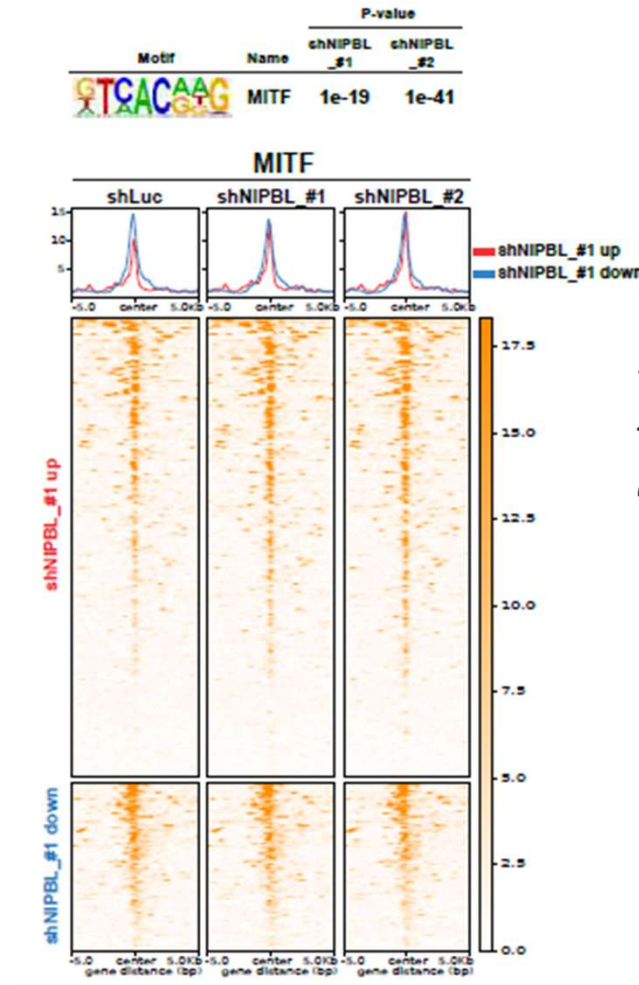
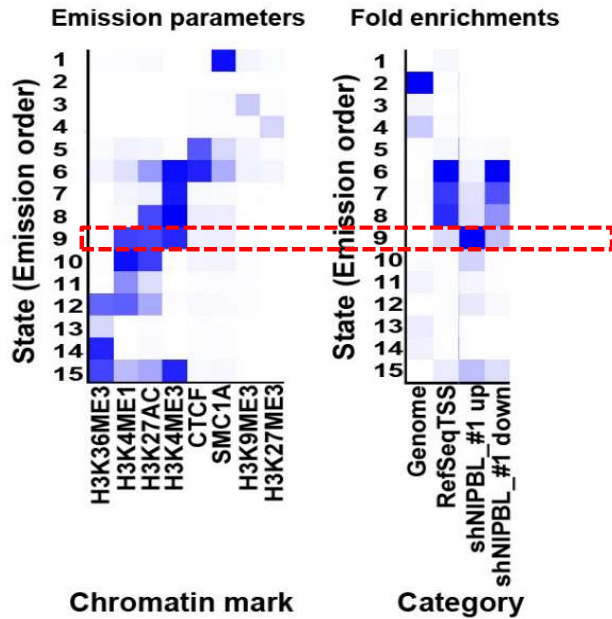
- Transposable elements: DNA sequences that once can move around in the genome
- Retrotransposons: replicate in genome via transcription and reverse transcription i.e., copy-and-paste
- LTRs account for ~8% of human genome
- LTRs harbor regulatory regions e.g., RNA Pol II and TFBS, that required for transcription
- subset of LTRs function as primary or alternative promoters in embryo and pluripotent stem cells, and sometimes in cancer
- Usually silenced by DNA methylation and repressive chromatin modification (e.g., H3K9me3)



NIPBL perturbation-mediated LTR alternative promoters are characterized by open chromatin and MITF binding



Bisulfite-free enzymatic methyl sequencing (EM-seq)
No global DNA methylation changes by NIPBL perturbation

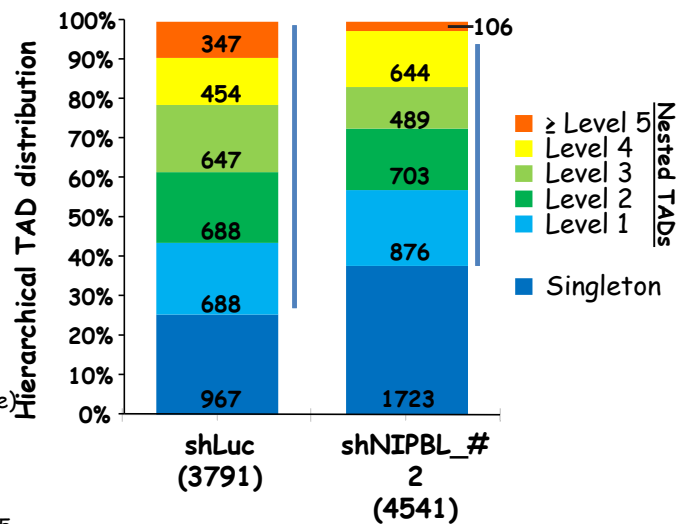
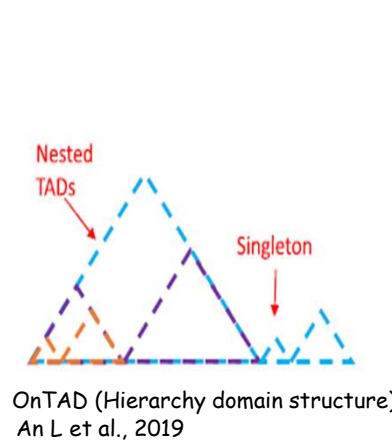
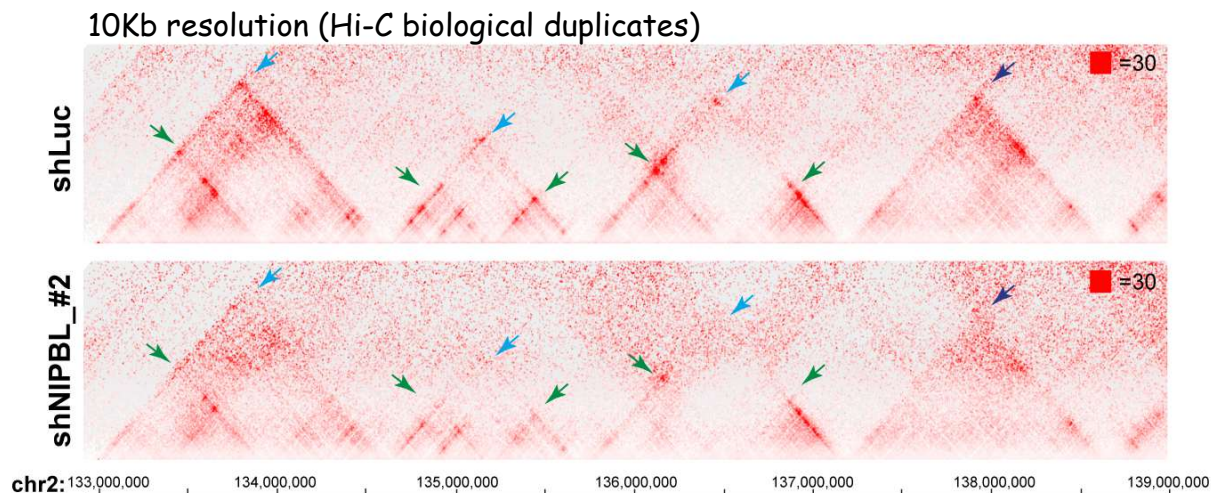


What is restricting the LTR transcriptional activation by MITF at baseline?

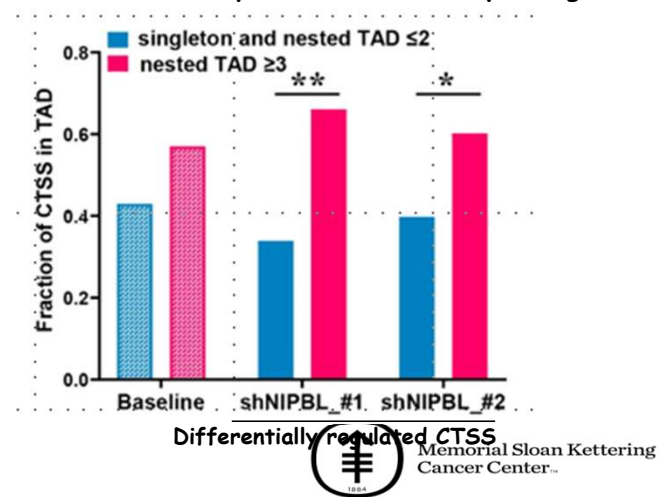


Memorial Sloan Kettering Cancer Center

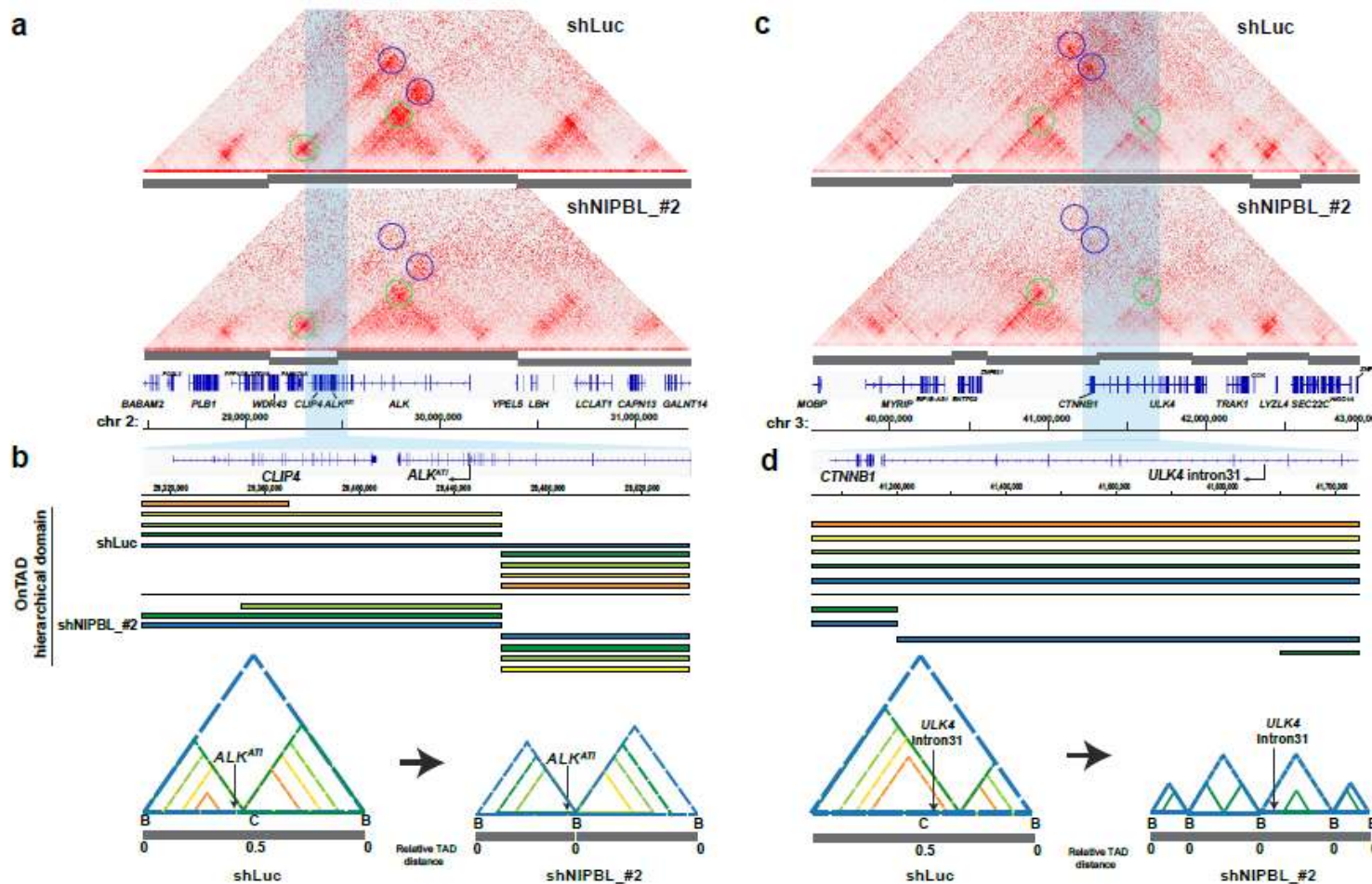
NIPBL perturbation leads to TAD hierarchy loss and alternative promoter activation



Integration of CAGE-seq and TAD hierarchy changes

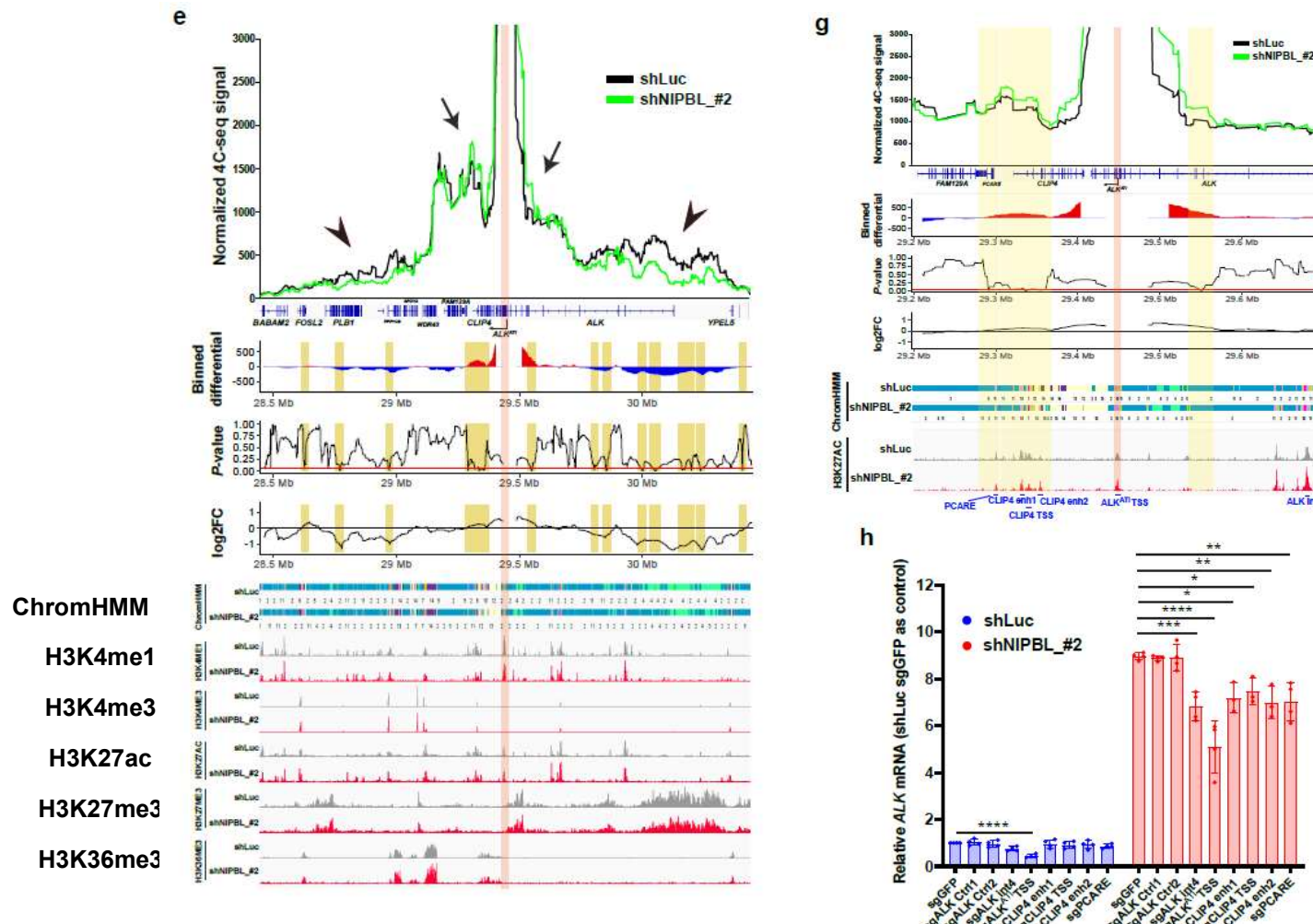


TAD hierarchy loss results in alternative promoter activation from intron/intergenic LTRs

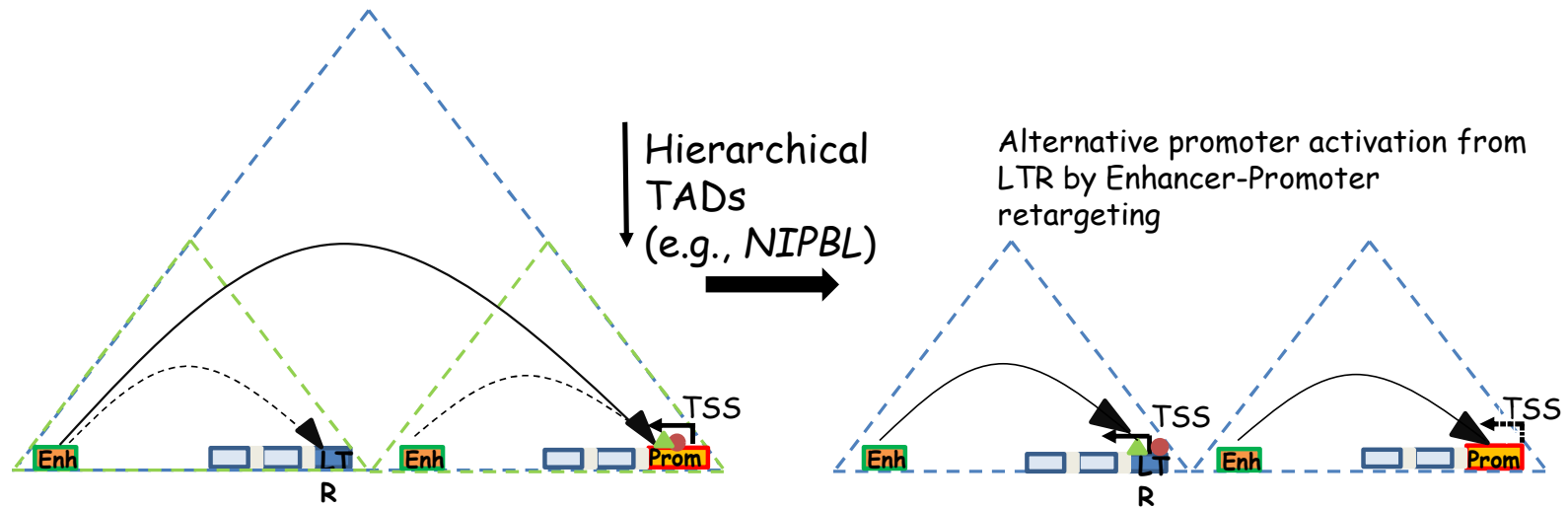


Corroborated by CRISPR/CRISPRi-mediated TAD boundary perturbation, and alternative promoter activation.

TAD hierarchy loss results in alternative promoter activation from E-altP retargeting (switch from long-range E to short-intermediate range E)



Alternative promoter usage in cancer via perturbation in 3D chromatin hierarchy



Hierarchical 3D chromatin organization and effects on transposable elements (TE) regulation

Physiologically - Hierarchical TAD organization suppresses LTR-mediated spurious transcription and regulate LTR co-option. A new layer of regulatory mechanism of TE expression beyond DNA and chromatin modification in human genome.

Pathologically - Perturbation in hierarchical TAD organization can lead to context-dependent oncogene activation, transcriptional diversity/plasticity and neopeptide expression through LTR co-option in cancer.




Paper discussion...

Discussion Paper: <https://pubmed.ncbi.nlm.nih.gov/31666694/>

Flavahan WA et al., Altered chromosomal topology drives oncogenic programs in SDH-deficient GISTs. Nature 2019

Published: 23 December 2015

Insulator dysfunction and oncogene activation in *IDH* mutant gliomas

William A. Flavahan, Yotam Drier, Brian B. Liau, Shawn M. Gillespie, Andrew S. Venteicher, Anat O. Stemmer-Rachamimov, Mario L. Suvà & Bradley E. Bernstein 



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Cancer Center..