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Authors

Zhang, Wei

Bojorquez-Gomez, Ana

Velez, Daniel Ortiz

et al.

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A global transcriptional network connecting noncoding mutations to changes in tumor gene expression

Wei Zhang^{1,†,*}, Ana Bojorquez-Gomez^{1,†}, Daniel Ortiz Velez², Guorong Xu³, Kyle S. Sanchez¹, John Paul Shen¹, Kevin Chen², Katherine Licon¹, Collin Melton⁴, Katrina M. Olson^{1,5}, Michael Ku Yu¹, Justin K. Huang^{1,6}, Hannah Carter¹, Emma K. Farley^{1,5}, Michael Snyder⁴, Stephanie I. Fraley², Jason F. Kreisberg^{1,*}, and Trey Ideker^{1,2,6,*}

¹Department of Medicine, University of California, San Diego, La Jolla, California, USA

²Department of Bioengineering, University of California San Diego, La Jolla, CA, USA

³Center for Computational Biology and Bioinformatics, University of California, San Diego, La Jolla, California, USA

⁴Department of Genetics, Stanford University School of Medicine, Stanford, California, USA

⁵Division of Biological Sciences, University of California San Diego, La Jolla, CA, USA

⁶Bioinformatics and Systems Biology Program, UC San Diego, La Jolla, California, USA

Abstract

Although cancer genomes are replete with noncoding mutations, the effects of these mutations remain poorly characterized. Here we perform an integrative analysis of 930 tumor whole genomes and matched transcriptomes, identifying a network of 193 noncoding loci in which mutations disrupt target gene expression. These “somatic eQTLs” (expression Quantitative Trait Loci) are frequently mutated in specific cancer tissues, and the majority can be validated in an independent cohort of 3,382 tumors. Among these, we find that the effects of noncoding mutations on *DAAMI*, *MTG2* and *HYI* transcription are recapitulated in multiple cancer cell lines, and that increasing *DAAMI* expression leads to invasive cell migration. Collectively the noncoding loci

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*Correspondence should be addressed to T.I. (tideker@ucsd.edu), J.F.K. (jkreisberg@ucsd.edu), W.Z. (wez124@ucsd.edu).

†Wei Zhang and Ana Bojorquez-Gomez contributed equally to this work

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AUTHOR CONTRIBUTIONS

W.Z. and T.I. conceived the study. W.Z. designed and performed most of the analyses. G.X. performed mutation calling of 358 tumors. C.M. and M.S. provided mutation calling of 572 tumors. A.B., K.S.S., J.P.S., K.M.O. and E.K.F. performed the somatic eQTL reporter assays. A.B. and J.F.K. analyzed the flow cytometry and luciferase assay data. A.B., J.P.S. and K.L. performed protein electropherogram analysis. D.O.V., K.C. and S.I.F. performed 3D cell culture assays. M.K.Y. and H.C. helped W.Z. in designing the somatic eQTL analysis. J.K.H. helped W.Z. in network analysis. T.I., J.F.K. and W.Z. wrote the paper and formulated all figures.

COMPETING FINANCIAL INTERESTS

Trey Ideker is co-founder of Data4Cure, Inc. and has an equity interest. Trey Ideker has an equity interest in Ideaya BioSciences, Inc. The terms of this arrangement have been reviewed and approved by the University of California, San Diego, in accordance with its conflict of interest policies. No potential conflicts of interest were disclosed by the other authors.

converge on a set of core pathways, permitting a classification of tumors into pathway-based subtypes. The somatic eQTL network is disrupted in 88% of tumors, suggesting widespread impact of noncoding mutations in cancer.

Human cancers are fundamentally heterogeneous, with many distinct subtypes associated with differences in molecular, cellular and clinical characteristics. To gain insight into this complexity, projects such as The Cancer Genome Atlas (TCGA) and International Cancer Genome Consortium (ICGC) have used massively parallel DNA sequencing to construct large catalogs of somatic mutations in many types of tumors¹⁻³. Focusing initially on protein coding regions, several hundred genes were found to be recurrently mutated in cancer, a few of which are targetable therapeutically⁴.

Since coding regions account for less than 2% of the human genome, attention is now shifting to the greater number of somatic mutations in noncoding regions⁵. Thus far the clearest role for noncoding mutations in cancer has been in the promoter of the telomerase reverse transcriptase gene (*TERT*)⁶⁻⁸, with such mutations leading to increases in *TERT* expression levels in many types of tumors^{8,9}. Although whole-genome sequencing (WGS) of tumor-normal pairs has found recurrent somatic mutations at several other noncoding loci, assessing the function of these mutations, if any, has been challenging⁶⁻⁸. In this respect, the task of functional interpretation is greatly aided by recent efforts of consortia such as ENCODE^{10,11} and Roadmap^{12,13}, which have published extensive reference maps of non-coding regions and their likely transcriptional regulatory connections to genes. Here, we show that such networks provide critical information for identifying noncoding mutations with functional impacts among the many others that may be spurious⁶.

RESULTS

Genome-wide identification of somatic eQTLs in cancer

To identify noncoding mutations associated with functional effects, we performed a systematic analysis of 930 tumors integrating whole-genome sequences, matched mRNA expression profiles and reference transcriptional interaction maps. Using WGS of paired normal and tumor tissues in 930 patients across 22 types of cancer from TCGA¹ (Fig. 1a), we identified 3.5×10^7 sites with somatic single nucleotide variations (SNVs). We called these SNVs uniformly across all genomes using the MuTect suite¹⁴ according to GATK Best Practice recommendations^{15,16} and those of Melton *et al.*⁶ (Fig. 1b). Clusters of noncoding SNVs located within 50 base pairs of one another were grouped, defining recurrently mutated loci (Fig. 1c, Methods, Supplementary Fig. 1).

We then tested each locus for its association with changes in mRNA expression of target genes (Fig. 1d). This task made use of two additional datasets. First, enhancer-gene mappings in GeneHancer¹⁷ were used along with promoter-proximal regions, defined as sequences within 1 kb of each transcription start site (TSS), to link recurrently mutated loci to putative target genes considered to be under direct transcriptional control (Methods). Second, for the vast majority of patients with tumor genome sequences, tumor mRNA expression profiles were also available (Fig. 1a). From these data we developed a

Increasing *DAAMI* expression leads to cell invasion

We next sought to examine in more detail the somatic eQTL located –191 bp upstream of *DAAMI* (Fig. 2a, Methods), which is recurrently mutated in melanoma patients with metastatic disease in both cohorts (Figs. 2c, d). The *DAAMI* protein forms a complex with Dishevelled and RhoA to recruit the actin cytoskeleton, which is thought to increase the motility and invasiveness of cancer cells in response to Wnt signaling^{22–24}. Mutations at this somatic eQTL are associated with increased *DAAMI* mRNA expression levels due potentially to the loss of an E2F motif and the gain of an Ets motif (Fig. 3a, NC_000014.8:g.59655190G>A). To confirm a causal relationship between the somatic eQTL and gene expression level changes, wild-type and mutant *DAAMI* regulatory elements were inserted upstream of Green Fluorescent Protein (GFP) (Fig. 3b). Analysis by flow cytometry showed that the mutated regulatory element leads to a significantly higher percentage of cells expressing GFP in melanoma, sarcoma and breast cancer cell lines (Figs. 3c, d, Supplementary Fig. 4). Furthermore, the GFP-expressing cells had significantly higher levels of GFP expression with the mutant rather than the wild type *DAAMI* element in all four cell lines tested (Fig. 3d, Supplementary Figs. 4c, e, g).

We also explored the functional relationship between increased *DAAMI* expression and cell motility, using an established 3D collagen hydrogel matrix model²⁵. Genome-wide mRNA sequencing was performed on cells grown within low or high density collagen, mimicking the stiffness of normal or tumor tissues and eliciting less or more invasive phenotypes, respectively^{26,27} (Methods). In these experiments *DAAMI* was one of the most up-regulated transcripts under invasive conditions²⁸ (Supplementary Fig. 5). To test whether invasion was functionally dependent on *DAAMI*, we quantified cell migration behavior after *DAAMI* expression was increased artificially by exogenous overexpression (OE, Fig. 3e, Supplementary Fig. 6e, Methods). When cells overexpressing *DAAMI* were embedded in the 3D collagen hydrogel, they migrated with significantly greater persistence than did wild type cells ($p = 0.008$, two-sided Mann–Whitney U test; Supplementary Fig. 6a). Cells overexpressing *DAAMI* also invaded for longer distances than wild type cells ($p = 0.01$, two-sided Mann–Whitney U test; Figs. 3f–h), while retaining the same velocities as wild type cells (Supplementary Fig. 6b, c). This invasive phenotype was observed in the absence or presence of additional Wnt5a signaling (Supplementary Fig. 6d). These results suggest that increased *DAAMI* expression levels allows cells to more efficiently invade the local microenvironment, thereby linking this noncoding mutation to *DAAMI* overexpression and cell invasion.

Noncoding mutations dysregulating *MTG2* and *HYI*

Beyond *DAAMI*, we examined two additional somatic eQTLs, one in the promoter of *MTG2* (+19 – +33) and another in the enhancer of *HYI* (+95097 – +95132) (Methods). The first eQTL was associated with decreased *MTG2* mRNA expression levels, likely due to the disruption of a HIF1b binding motif by the G-to-A mutation at +19 bp downstream of the TSS (Fig. 4a, NC_000020.10:g.60758100G>A). This somatic eQTL was present in several types of cancer including lung adenocarcinoma and sarcoma. Using another GFP-based reporter assay of promoter activity, we found that this G-to-A mutation greatly decreased reporter gene expression in both A549 lung epithelial carcinoma cells and U2OS bone

osteosarcoma cells (Fig. 4b). The second eQTL was present in 21% of melanomas (Fig. 2c) and was associated with increased *HYY* mRNA expression levels, likely due to G-to-A or GG-to-AA substitutions altering an Ets family binding motif (Fig. 4c, NC_000001.10:g.43824528G>A, NC_000001.10:g.43824529G>A, or NC_000001.10:g.43824528_43824529GG>AA). As this somatic eQTL was present in an enhancer region, we used a luciferase-based reporter assay where regulatory elements were cloned upstream of a mini-promoter and luciferase. We found that two out of the three *HYY* enhancer variants led to increased expression levels relative to wild type in both A375 melanoma cells and MDA-MB-231 breast cancer cells (Fig. 4d).

Noncoding and coding mutations converge on pathways

Next, we investigated the relationship between the 196 genes transcriptionally regulated by somatic eQTLs and the 138 genes previously documented to have recurrent coding mutations in cancer²¹. This combined set of genes was analyzed by Network-Based Stratification^{29,30} (NBS; Fig. 5a), which uses a reference molecular network to implicate network regions associated with the genetic alterations in a tumor and groups tumors into subtypes based on similarity of these implicated regions. As a reference molecular network we used ReactomeFI³¹, documenting 229,300 interactions among 12,177 human gene products pertaining to previously reported protein-protein, transcriptional and metabolic interactions.

This approach identified a collection of network regions (henceforth called “pathways” for simplicity) that stratified tumors into a hierarchy of increasingly specific subtypes (Fig. 5b). At a resolution of ten subtypes, each subtype was enriched in 2-5 tumor tissues and tumors of each tissue could be subdivided into 1-3 subtypes (Supplementary Fig. 7). Nonetheless, these subtypes differed significantly in their implications for disease-free survival, beyond the baseline survival for each tissue ($p = 3.3 \times 10^{-6}$, log likelihood ratio test controlling for the tissue types as covariates; Fig. 5c, Supplementary Fig. 8).

Subtypes aggregating noncoding and coding mutations

Among the ten subtypes, four were of particular interest as they contained a large proportion of patients with noncoding mutations (Fig. 5d). The “*CDKN2A-EGFR-TERT* subtype” (Figs. 5e, f) was defined by disruption of the *CDKN2A* coding sequence, sometimes in combination with noncoding mutations to the *TERT* promoter, *EGFR* activation, or *BRAF* activation. *CDKN2A* encodes p14^{ARF}, which can form a complex with Hif-1 α and inhibit HIF-1-mediated transcription of *TERT*^{32,33}. These loss-of-function mutations in *CDKN2A* may release a key brake on the activity of hTERT. Separately, gain-of-function mutations in *EGFR* may lead to increased levels of mTOR phosphorylation and activation³⁴, which can up-regulate telomerase activity by forming a complex with hTERT³⁵. The synergy between *BRAF* and *TERT* mutations has been previously noted and attributed to modulation of *TERT* transcription through *BRAF-RAS-ERK* signaling³⁶. This pathway was also linked to *DAAMI* promoter mutations (Fig. 5d), validated previously, as Daam1 forms a complex with Dishevelled (Dvl3)^{22,23}, which indirectly regulates transcription of *CDKN2A* and *EGFR* through inhibition of Notch1³⁷. This subtype was the most aggressive, with median disease free survival time at 13 months (Fig. 5c).

A second subtype of interest, the “*TERT-BRAF-IDH1* subtype” (Supplementary Fig. 9) was characterized by tumors with *TERT* noncoding mutations or amplifications, combined in some patients with coding alterations to functionally related genes such as *BRAF* and *SKP2*. Beyond the synergy between *BRAF* and *TERT* mutations as described above, Skp2 is essential for ubiquitination and degradation of p27^{Kip1} (encoded by *CDKN1B*)³⁸, which inhibits the activity of hTERT³⁹. Amplification of *SKP2* in this pathway may thus increase the activity of hTERT.

A third subtype, “*PIK3CA-PEX26-GATA3*” (Figs. 5g, h), integrated coding alterations activating *PIK3CA* and inactivating *GATA3* with noncoding alterations downregulating *PEX26*. In this pathway, members of peroxisomal biogenesis factor (Pex26 and Pex6) appear to indirectly interact with *PIK3CA* and *GATA3* through the binding of SMAD family members (Smad3 and Smad7)⁴⁰.

Finally, the fourth subtype, “*APOBEC2-ARID1A-CTNNB1*”, was characterized by the co-occurrence of noncoding mutations within an enhancer of *APOBEC2* and coding alterations in *ARID1A* and *CTNNB1*. *APOBEC2* encodes a nucleic-acid editing enzyme with well-known mutagenic effects in cancer⁴¹. Although *ARID1A* and *CTNNB1* are also known cancer drivers, the connections to *APOBEC* are unanticipated and create a compelling opportunity for further study.

DISCUSSION

Relative to coding changes, interpretation of noncoding mutations poses particular challenges due to the very large number of events and a limited understanding of their functional consequences. Dealing with these challenges requires strategies to boost signal-to-noise, which we have pursued here by integrating mutations with key structural and functional data on transcriptional networks. Structurally, maps of enhancer- and promoter-gene interactions amplify signal by selecting noncoding mutations within defined regulatory regions of specific target genes. These mutations are then characterized functionally as somatic eQTLs by requiring their presence to be significantly associated with expression changes in tumors. The result is a global network of transcriptional regulatory interactions in cancer supported by multiple lines of evidence. Given that most tumors we analyzed had noncoding mutations affecting some part of this network, such mutations appear to represent a widespread feature of cancer biology.

Of the approximately two hundred noncoding mutations that have been previously identified as recurrent in cancer^{6–8}, one third were also identified here as recurrently mutated loci (Fig. 1c), including well known mutations in the promoters of *PLEKHS1* and *DPH3*. Notably though, with the exception of *TERT*, these mutations did not associate significantly with mRNA expression level changes. This suggests that the effects of these mutations are through mechanisms outside of transcriptional regulation, or that the effects on mRNA expression are weaker than could be detected given our statistical power (Supplementary Fig. 2c). On the other hand, hundreds of somatic eQTLs were identified, all of which were unanticipated other than those in the promoter of *TERT*. Many of the affected genes are not

yet widely appreciated as cancer drivers, motivating further studies on the mechanistic basis of noncoding mutations in cancer.

Given an association between gene expression changes and a somatic mutation, it is important to consider whether this association reflects a causal relationship. Although it is tempting to assume that the occurrence of a mutation drives gene expression changes, the opposite could be true, where the change in gene expression levels drives the appearance of the mutation (*e.g.*, by increased opening and exposure of chromatin). It is also possible that both effects could be due to a third causal factor. However, the three examples we tested experimentally do support a causal link from mutation to expression changes. These results include transcriptional alterations of *DAAMI*, impacting cell migration (Fig. 3, Supplementary Fig. 4); *MTG2*, which encodes a GTPase that regulates mitochondrial ribosomes⁴² (Figs. 4a, b); and *HYI*, which encodes a putative hydroxypyruvate isomerase and may be involved in carbohydrate transport and metabolism⁴³ (Figs. 4c, d).

Finally, the somatic eQTL analysis introduced here contrasts with germline eQTL studies in several key aspects. First, in GWAS and germline eQTL studies, testing of multiple SNPs is complicated by the strong codependencies among neighboring SNPs at a genomic locus – so-called linkage disequilibrium^{44,45}. In contrast, somatic mutations near to one another in the genome are not in linkage disequilibrium since these alterations, by definition, arise independently in each tumor. Second, population stratification caused by racial diversity has been a major confounder in analysis of germline variants^{44,45}. It is less of a concern for somatic variants, since these are derived from comparisons between tumor and normal genomes from the same individual, eliminating many, if not all, effects due to ancestry. Nonetheless, we controlled for racial diversity and found the impact on somatic eQTL discovery was minimal. Given these aspects, somatic eQTL analysis may have future interest alongside classical eQTLs as a general mode of mapping transcriptional regulatory architecture.

METHODS

Calling and clustering of somatic noncoding mutations

Somatic noncoding mutations of 930 tumors were called as described in the main text. Clusters of noncoding mutations within $d = 50$ bp from each other were merged using BEDTools⁴⁶ until no such locus was located within d bp from any other. Loci with mutations in $k < 5$ tumors were removed from further analyses. The above parameters d and k were chosen to aggregate mutations within short distance with a modest requirement of recurrence. We achieved very similar results when d was within the range of 20 to 60 (inclusive). Whenever a subset of 930 tumors was used in subsequent analyses (Fig. 1a), this set was again filtered to remove those altered in fewer than k tumors within the subset. We also calculated a “concentration score” to penalize loci where mutations were spread over a large region rather than concentrated at a single base pair, as might be expected for sites affecting gene transcription. Within each locus, we selected the mutated position present in the largest number of patients. The proportion of patients affected at that position (out of all patients affected by mutations at that locus) was defined as the concentration score. Loci scoring $< 35\%$ were removed from further study. It is worth noting that the threshold of

concentration score is somewhat arbitrary and could lead to missing certain loci with multiple closely located somatic mutations. It should also be noted that by clustering noncoding mutations into loci, we assume that all SNVs in a locus act in a similar way. This assumption is consistent with the previously identified SNVs in *TERT* promoter. Our analysis does not attempt to detect loci in which different SNVs alter gene expression in opposite directions.

RNA-seq, CNA and DNA methylation data processing

RNA-seq, CNA (SNP 6.0) and DNA methylation (Illumina HM450) data for TCGA tumors were downloaded from Firehose (see URLs). The data were processed as follows. First, for RNA-seq the RSEM count for a gene (RNA-Seq by Expectation Maximization)⁴⁷ was normalized by dividing by the 75th percentile of RSEM values within the tumor sample and multiplying by 1000, as per TCGA practice (see URLs). Genes were retained if the normalized RSEM > 1 in > 50% of tumors, resulting in 16,413 expressed genes. Normalized RSEM values were \log_2 transformed and z-score standardized for subsequent analyses. Second, for copy number alterations (CNAs) we used the output of GISTIC2, which indicates gene-level CNAs for all samples. The CNAs are in units of (copy number – 2), so that normal copy number (no amplification or deletion) has a value of 0, whereas genes with amplifications have positive values and genes with deletions have negative values. A gene is assigned the highest amplification or the lowest deletion value among the markers it covers. Among the 783 patients with both mRNA expression and genome sequence, 761 also had copy number data available. The remaining patients were assigned 0 for all CNAs. Third, methylation probes were mapped to the promoter regions of genes (\pm 1 kb from TSS), and each gene was assigned the mean methylation (beta) values of these probes. Among the 783 TCGA patients with both mRNA expression and genome sequences, 605 had methylation data available. Methylation data for the remaining patients were imputed using mean values for the DNA methylation of each gene.

Linking recurrently mutated loci to transcriptional target genes

Our recurrently mutated loci were extended by 100 bp on each side when mapping to the promoters or enhancers. Transcriptional regulatory interactions from recurrently mutated loci to target genes were defined whenever a locus had 50% of its sequence overlap with either the promoter region of a gene (\pm 1 kb from its TSS) or a gene enhancer region defined by GeneHancer¹⁷. In case an enhancer is shorter than a locus, the mapping was performed when 50% of the enhancer sequence overlaps with the locus.

Somatic eQTL analysis using multivariate linear regression

For each gene target linked to recurrently mutated loci, we fit a regression model of the normalized gene expression level e as a function of l , the alteration status of its recurrently mutated loci (1 = mutated; 0 = wt), controlling for the impact of CNA status c (0 = wt; positive values for amplifications; negative values for deletions), DNA methylation m (mean beta-values), 21 tumor tissues t (binary variables), 3 races r (binary variables: Asian; Black or African American; White), gender g (1 = Female; 0 = male) and 20 hidden factors h (real values) as covariates:

$$e = \beta_0 + \beta_1 l + \beta_2 c + \beta_3 m + \beta_4 t + \beta_5 r + \beta_6 g + \beta_7 h \quad (\text{Eq. 1})$$

The hidden factors h were identified using probabilistic estimation of expression residuals (PEER)^{48,49}, while accounting for the effect of known covariates t , r and g . The number of hidden factors was determined by the posterior variance of the factor weights, as previously recommended⁴⁹. The parameters β were estimated from data from 783 tumors with matched RNA-seq and WGS data. Somatic eQTLs were identified as follows. First, for each gene, we selected features by adding an L1-norm to the objective function based on least squared error between true and predicted gene expression level.

$$(e - \hat{e})^2 + \lambda \|\beta\|_1 \quad (\text{Eq. 2})$$

The sparsity parameter λ was optimized by cross validation. For genes in which the L1-norm resulted in $\beta_1 = 0$ for all loci, we decreased λ to include at least one locus. Second, to assess whether mutation status of any locus contributed significantly to gene expression, the accuracy of the complete model was compared to that of a simple model under null hypothesis of no genetic associations (*i.e.*, $\beta_1 = 0$ for all loci). The F-test p -value between the two nested models was used as the test statistic. Third, having derived an F-test p -value for each gene, q -values were calculated using the Storey approach⁵⁰ with a threshold of FDR 20%. And finally, for each gene that passed the selection, this threshold was mapped back to the equivalent F-test p -value of each locus. Loci with F-test p -value below or equal to this threshold were included in the final list and defined as somatic eQTLs.

We elected to perform one test per gene for three reasons. First, in GWAS and typical (germline) eQTL studies, linkage disequilibrium complicates the simultaneous testing of multiple SNPs in a single model, because these SNPs are usually codependent. Unlike inherited SNPs, somatic mutations observed in a tumor population are not in linkage disequilibrium no matter how closely they are located. Therefore, a simple F-test, which assumes independent influences of multiple factors, is sufficient to simultaneously test whether any loci are associated with gene expression. Second, for each gene, all eQTLs share the same set of covariates along with the associated phenotype of mRNA expression level. If multiple eQTLs are associated with gene expression levels, they can be covariates of one another. It is then convenient to fit them all in a single model and enjoy the benefit of gene-based approaches such as feature selection by L1 regularization. Third, there is precedent in the literature to fit gene-level models in eQTL studies^{51,52}.

Power analysis

Statistical power depends on various parameters, including the number of samples, the eQTL effect size, the noise, and the significance threshold. Instead of a simulation based on a model of noise, we evaluated statistical power using the actual data. All locus-gene pairs were plotted in Supplementary Fig. 2c, evaluated by the number of patients with mutations (x-axis) versus the change in gene expression given the mutation (y-axis; defined by

$W = \frac{\text{Coefficient}}{\text{Residual standard deviation}}$; one unit of W represents one standard deviation of change in residual gene expression). Power was defined as $1 - P(\text{Type II error})$ at a significance level of $P(\text{Type I error}) = 0.0085$, which is approximately at 20% FDR. We calculated power using the “pwr.f2.test” function in R, where the f^2 effect size was calculated based on the proportion of explained variance by two nested models ($f^2 = \frac{R^2_{\text{alternative}} - R^2_{\text{null}}}{1 - R^2_{\text{alternative}}}$). Our somatic eQTL analysis has 50% power to detect a somatic eQTL with 5 mutations if $W > 1.2$ or with 10 mutations if $W > 0.9$.

Independent validation of recurrence

To validate the recurrence of mutations in the identified somatic eQTLs, we downloaded simple somatic mutations (substitutions) called from the WGS of $n = 3,382$ publicly available non-US donors from the ICGC². For each eQTL, the number of mutated patients k was used as the test statistic. To determine whether k was greater than expected due to the background mutation rate (BMR), we developed an approach for estimating BMR that was conceptually similar to MutSigCV⁵³. First, a large pool of 20,000 candidate background sequences was created by randomly reassigning (without replacement) the location of the eQTL to the same type of noncoding genomic regions (promoters or putative enhancers¹⁷), while retaining the eQTL’s length. Each of these 20,000 sequences was placed in a three-dimensional feature space taking into account nucleotide content, DNA replication timing and gene expression. Nucleotide content was represented as the percentages of all possible mono- (A/T vs. C/G), di- (e.g., AA, AC and AG) and tri-nucleotides (e.g., AAA, AAC and AAG), encoded as a 44-dimensional vector. This information was then compressed into a single feature representing nucleotide content, using the Pearson’s correlation between the vector of the candidate sequence and the vector of the original eQTL. DNA replication timing was obtained from ENCODE via the UCSC Genome Browser⁵⁴. To create a single replication timing feature, we used the average wavelet-smoothed signal from the following 14 cell lines: BJ, GM06990, GM12801, GM12812, GM12813, GM12878, HeLa-S3, HepG2, HUVEC, IMR-90, K-562, MCF-7, NHEK and SK-N-SH, according to the method of Melton and colleagues⁶. For gene expression, the median expression value of the nearest gene (\log_2 transformed RNA-seq data, 783 TCGA patients) was used as a feature. The above three features were z-score standardized. Within this feature space, the top 5% (1,000 out of 20,000) background sequences with the smallest Euclidean distance to the eQTL of interest were selected. For each patient, a patient-specific BMR was estimated as the number of sequences with at least one mutation in that patient out of the 1,000 selected sequences. Finally, we estimated the probability of having observed k or more mutations in n patients in the eQTL of interest using a Poisson binomial model:

$$P(K \geq k) = \sum_{l=k}^n \sum_{A \in F_l} \prod_{i \in A} p_i \prod_{j \in A^c} (1 - p_j) \quad (\text{Eq. 3})$$

where F_l is the set of all subsets of k integers that can be selected from $\{1, 2, \dots, n\}$, p_i or p_j is the probability that patient i or patient j is mutated, A is a set of k integers that can be

selected from $\{1, 2, \dots, n\}$ and A^c is the complement of A . In practice, we used an approximation for the Poisson binomial in the `poibin` python package (see URLs).

Transcription factor binding motif analysis

Each reference and somatically altered nucleotide site, along with ± 7 bp flanking sequence, was analyzed using HOMER⁵⁵ (see URLs). HOMER searches for matches within a library of 319 vertebrate motifs (position weight matrices). Specifically, we ran the “findMotifs.pl” program with default parameters to find motifs from FASTA files. The reference and altered sequences were used as the background for each other to control the nucleotide context. The command line is: `findMotifs.pl seqList_mappable_alt.fa fasta log/ -fastaBg seqList_mappable_ref.fa -p 16 -find ~/soft/homer/data/knownTFs/vertebrates/known.motifs.`

The list of somatic eQTLs disrupt or create transcription factor binding motifs in four or more patients were reported in Supplementary Table 2.

Prioritizing somatic eQTLs for subsequent functional validation

The three somatic eQTLs selected for functional studies (*DAAMI*, *HYI1* and *MTG2*) were chosen based on the specific biological interest of the authors and several rules-of-thumb:

1. The somatic eQTL alters a known transcription factor binding motif in many patients;
2. The somatic eQTL falls in open chromatin in previously mapped cell lines and conditions (e.g., in regions with markers such as H3K27ac and H3K4me1)¹¹;
3. The affected target gene has high endogenous mRNA expression levels in cell lines⁵⁶ that match where the somatic eQTL was detected; and
4. The somatic eQTL is not present in a region with repetitive DNA.

Note that none of this information was used to filter loci prior to somatic eQTL analysis, as it is not complete, conclusive or cancer-specific.

Generation of reporter plasmids

To examine the effect of the *DAAMI* somatic eQTL on gene expression levels, the wild type and mutant regulatory regions, from -233 bp to $+148$ bp relative to the TSS, including the somatic eQTL at $-202 - -191$ bp, were synthesized and cloned upstream of GFP (Fig. 3b). For *MTG2*, the cloned region spanned -200 bp to $+200$ bp relative to the TSS, including the somatic eQTL at $+19 - +33$ bp.

For the somatic eQTL located in the *HYI* enhancer, the region corresponding to $+94931$ bp to $+95332$ bp relative to the TSS, including the somatic eQTL at $+95097 - +95132$ bp, was cloned into the firefly luciferase reporter plasmid pGL4.23 (Promega). The mutations were generated using the Q5 Site-Directed Mutagenesis Kit (New England BioLabs). All inserts for the GFP and luciferase reporter plasmids were confirmed to match the human reference genome hg19 by Sanger sequencing.

Promoter and enhancer activity assays

Cell lines used to evaluate of promoter activity were plated in 6-well dishes at 300,000 cells per well, three replicates per group. The next day, plasmid DNA (1 μ g) was transfected using Lipofectamine 3000 (ThermoFisher). Forty-eight hours after transfection, cells were harvested and suspended in ice cold PBS with 1% fetal bovine serum. GFP expression was measured by flow cytometry on a FACSCalibur or FACSCanto (BD Biosciences). Flow cytometry data were analyzed with FlowJo v10 (BD Biosciences). Cells with typical forward (size) and side (granularity) scatter properties were further analyzed for GFP expression. As a negative control, cells were transfected with an empty lentiGuide-Puro plasmid (Addgene) for the *DAAMI* experiments (Figs. 3c, d; Supplementary Fig. 4) or a promoter-less GFP plasmid (pRMT-tGFP, Origene) for the *MTG2* experiments (Fig. 4b). As a positive control for all GFP experiments, we used a plasmid with the cytomegalovirus promoter upstream of GFP. All flow cytometry experiments were performed at least three times. Early pilot experiments were often performed on single or duplicate samples with then the final triplicate version often performed at least twice.

To evaluate the activity of the enhancer region of *HYY*, A375 and MDA-MB-231 cells were plated in white, opaque, 96-well plates at 10,000 cells per well, four replicates per group. Cells were transfected 24 hours later using lipofectamine 3000 with 33 ng of total DNA: 27.5 ng of the firefly pGL4.23 constructs and 5.5 ng of control Renilla pGL4.75 (Promega) plasmid. Firefly and Renilla activity were measured 48 hours after transfection using the Dual-Glo Luciferase Assay System (Promega) as per the manufacturer's instructions. Luciferase values were collected on a BioTek Synergy HT, and data collected via software Gen5 2.01.14. To calculate relative luciferase values, background signal was first subtracted from each channel. Then firefly luminescence was divided by Renilla luminescence. The average value for the wildtype enhancer was set to 1, and the mutated samples were evaluated in comparison to this control. Experiments in both cell lines were performed three times, with each experiment consisting of samples in quadruplicate.

DAAMI overexpression

Wild type MDA-MB-231 breast cancer cells were transfected with a plasmid encoding the full *DAAMI* gene cDNA (Origene catalog number: RC217675). Cells were then selected using G418 (500 μ g/mL) for 7 days to ensure stable expression of the *DAAMI* construct. *DAAMI*-overexpression was verified by extracting total protein and quantitating using the Wes electropherogram (Proteinsimple) with anti-*DAAMI* antibody (clone WW-3, cat# sc-100942, lot# B1815, Santa Cruz, 1:250 dilution) and anti-tubulin antibody (clone YL1/2, cat# MAB1864, lot# 2886723, Millipore, 1:250 dilution). *DAAMI* expression was 5.5 fold greater in cells with *DAAMI*-overexpression construct relative to wild type (Supplementary Fig. 6e).

3D collagen cell migration assay

Collagen matrices were prepared by mixing cells suspended in culture medium and 10 \times reconstitution buffer, one-to-one with soluble rat tail type I collagen in acetic acid (Corning)²⁵. Sodium hydroxide was used to normalize pH (pH 7.0, 10 – 20 μ L 1M NaOH), and the mixture was placed in 48-well culture plates for polymerization at 37°C. Final gel

volumes were approximately 200 μL with final collagen concentration set to 2.5 mg/mL. The polymerized cell-laden hydrogels were incubated for 24 hours under a standard cell culture environment before imaging. Gels were then transferred to a microscope stage-top incubator, and cells were imaged at low magnification (10 \times) every 2 minutes for 48 hours. Coordinates of the cell location at each time frame were determined using image recognition software (Metamorph/Metavue, Molecular Devices). Tracking data were processed to calculate cell speed using an extension of previously published scripts⁵⁷. Cell migration assays (Fig. 3f-h) were performed two times and both attempts showed the same trend.

RNA sequencing from cells in 3D culture

In Supplementary Figure 5, cell migration assays were performed using wild type MDA-MB-231 breast cancer cells and HT-1080 fibrosarcoma cells. 3D collagen I gels were seeded in 3 independent experiments and harvested after 24 hours of culture for RNA extraction and directly homogenized in Trizol reagent (ThermoFisher). Total RNA was purified using High Pure RNA Isolation Kit (ROCHE) and the integrity of the sample verified using RNA Analysis ScreenTape (Agilent Technologies). Total RNA samples were sequenced using the TruSeq Stranded mRNA Sample Prep Kit (Illumina) and the Illumina MiSeq platform at a depth of > 25 million reads per sample. Paired-end reads were aligned to the hg19 UCSC human genome reference using Bowtie2⁵⁸ and streamed to eXpress⁵⁹ for transcript abundance quantification.

Tumor genetic profiles integrating noncoding and coding alterations

Integrated genetic alteration profiles were constructed for the 810 tumors with WGS, WES and CNA data (Fig. 1a) as follows. Known oncogenes or tumor suppressors²¹ were combined with the set of target genes of eQTLs identified by the somatic eQTL analysis (see above); each of these genes was then classified as wild type (0) or altered (1) in each tumor, constituting its tumor genetic profile. In this profile, an alteration was defined as follows: Most oncogenes (*e.g.*, *EGFR*) were considered altered (activated) if impacted by a missense mutation, in-frame indel, or copy-number amplification. For oncogenes typically altered only by amplification²¹ (*CCND1*, *MDM2*, *MDM4*, *MYC*, *MYCL*, *MYCN*, *NCOA3* and *SKP2*), only copy number amplifications were considered as alterations and not SNVs or indels. Tumor suppressors (*e.g.*, *CDKN2A*) were considered altered (inactivated) if there was any type of non-silent mutation or a copy number deletion. For each target gene, we defined a dominant direction of regulation $d \in \{+1, -1\}$ as the sign of the coefficient (β_1 in Eq. 1) of its most significantly associated eQTL. Noncoding mutations in eQTLs that lead to a transcriptional change in the dominant direction were considered alterations of such genes. For *TERT*, copy number amplifications in the coding region was also considered as alterations, since both promoter mutations and gene amplifications have been associated with growth advantage of tumor cells and poor prognosis of patients^{60,61}.

Network-Based Stratification (NBS) to identify tumor subtypes

Network propagation²⁹ was used to compute the pairwise similarities among tumor genetic alteration profiles (see above) within the Reactome functional interaction network (ReactomeFI)³¹. Each tumor genetic profile was propagated across this network based on a random walk model (equivalent to heat diffusion) with a restart probability of 0.5. After

convergence, the score of each gene (temperature) represents its network proximity to genetic alterations. The top 70 principal components of these scores, representing the tumor's network-transformed profile (Fig. 5a), were analyzed using the `sklearn.cluster.SpectralClustering` package⁶² (affinity = *k*-Nearest-Neighbors, assign-labels = discretize, n_clusters = [2..10]). This method first constructs a similarity graph on all pairs of tumors, where each tumor is connected to the *k* others with shortest Euclidean distance. We chose *k* = 170, which ensures the similarity graph to be connected, as previously recommended⁶². Next this graph is analyzed to partition tumors into subtypes at different resolutions (number of subtypes *n* = [2..10]). Following spectral clustering, each set of *n* (parent) subtypes was compared to the *n* + 1 (child) subtypes to track the similarity of tumor assignments (Fig. 5b). An arrow is drawn from a parent to child subtype if they share 18 tumors.

Characterizing tumor subtypes with signature genes and subnetworks

For each subtype, we defined a set of “signature genes” as those that had higher network-transformed scores in that subtype than others (t-test, Benjamini Hochberg FDR < 0.1) and, among these, were more frequently altered in that subtype (Fisher Exact Test, FDR < 0.05, Figs. 5b-e). To identify subnetworks, this set was expanded to include “intermediate genes” with relatively high network-transformed score (t-test, FDR < 0.05) that lie on the shortest paths between each pair of signature genes. The union of the signature and intermediate genes was used to induce a subnetwork within ReactomeFI³¹, referenced in the main text as the corresponding “pathway” impacted in that subtype (Figs. 5d). An additional filter was applied in Fig. 5e and Supplementary Fig. 9a, where we only visualized the signature genes with 10 or more mutations and the shortest paths among them with at most one intermediate gene. All networks were visualized in Cytoscape⁶³.

Survival analysis

We used the “coxph” package in the R statistics software to fit Cox proportional hazard models⁶⁴. *P*-values were calculated by log likelihood ratio test. To evaluate whether the subtype classifications provide additional prognostic power beyond the baseline survival expectancy due to cancer tissue, we compared the likelihood for the complete model, including NBS-derived molecular subtypes *s* and cancer tissues *c* as covariates, against that of a null model that includes cancer tissues *c* only:

$$\text{Complete model: } \lambda(t|s, c) = \lambda_0(t)\exp(\beta_0 + \beta_1s + \beta_2c) \quad (\text{Eq. 4})$$

$$\text{Null model: } \lambda(t|s, c) = \lambda_0(t)\exp(\beta_0 + \beta_2c) \quad (\text{Eq. 5})$$

where $\lambda_0(t)$ is the baseline hazard function. Then a log-likelihood ratio statistic was defined as:

$$D = -2\ln\left(\frac{\text{likelihood for null model}}{\text{likelihood for complete model}}\right) \quad (\text{Eq. 6})$$

Finally, a chi-squared test p -value was calculated based on D with the number of degrees of freedom equal to the number of NBS-derived molecular subtypes.

Data availability

Somatic mutations of 930 tumors are publicly available (see URLs). RNA-seq data is accessible through the GEO Series accession number GSE101209.

Code availability

Custom code for annotating mutations, somatic eQTL analysis, validation of recurrence, motif analysis and Network-Based Stratification are available through GitHub (see URLs).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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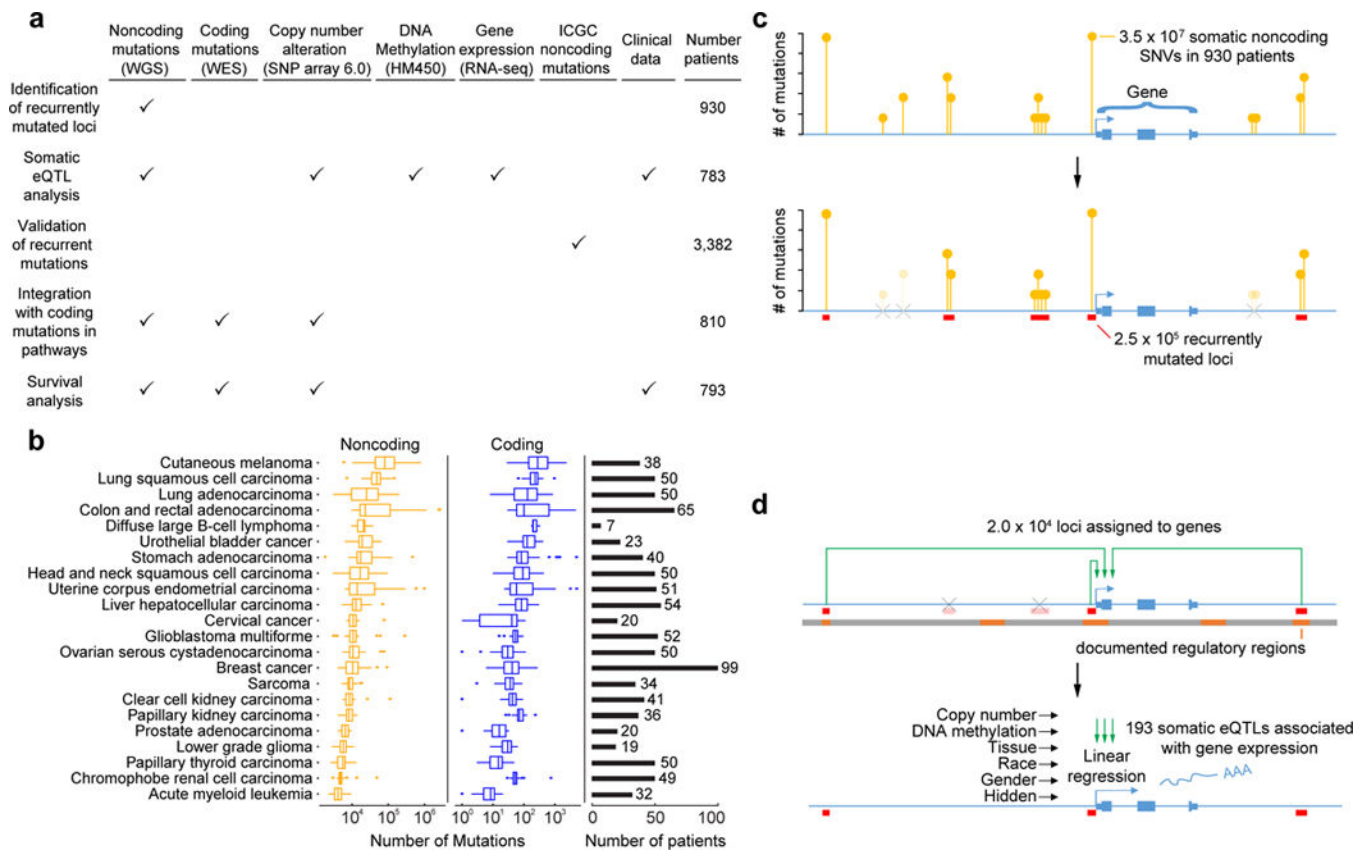


Figure 1. Mutation calling and somatic eQTL analysis

(a) Types of data and numbers of tumors used in this study. (b) Number of mutations called per tumor. Boxplots show the distribution of this number within tumors of each tissue type (center line, median; upper and lower hinges, first and third quartiles; whiskers, highest and lowest values within 1.5 times the interquartile range outside hinges; dots, outliers beyond 1.5 times interquartile range). The number of tumors of each type (sample size) is shown on the right panel. (c) Clustering of somatic noncoding mutations resulting in identification of recurrently mutated loci. (d) Workflow of somatic eQTL analysis.

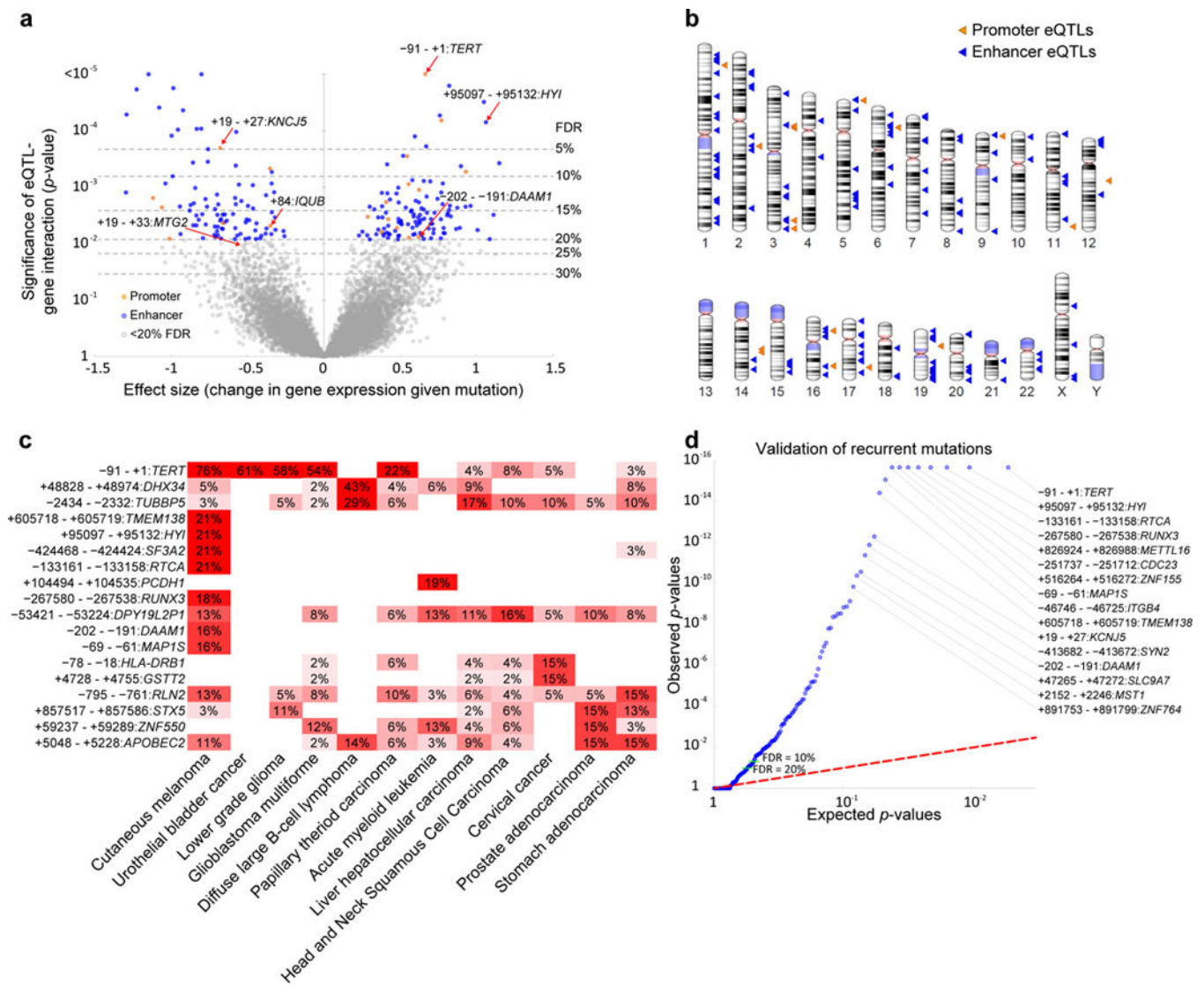


Figure 2. Effect size and recurrence of somatic eQTLs

(a) Volcano plot of associations between somatic eQTLs and the expression levels changes of their target genes, evaluated by significance (y-axis, F-test p -value, $n = 783$ tumors) versus effect size (x-axis). One unit on the x-axis represents one standard deviation of change in gene expression. FDR is calculated using the Storey approach⁵⁰. Selected somatic eQTLs are labeled by coordinates in base pairs relative to the TSS of the target gene. (b) Ideogram of the 193 significant somatic eQTLs at FDR < 20%. (c) Heatmap showing the percentage of patients in various cancer tissues with alterations in each somatic eQTL. Somatic eQTLs and cancer tissues with 15% mutation rates are shown. (d) Validation of somatic eQTL recurrence in a pan-cancer cohort from ICGC. The quantile-quantile plot shows the observed empirical p -values of mutation recurrence ($n = 3,382$ tumors) compared to the random expectation for the 193 somatic eQTLs. FDR is calculated using the Benjamini-Hochberg approach.

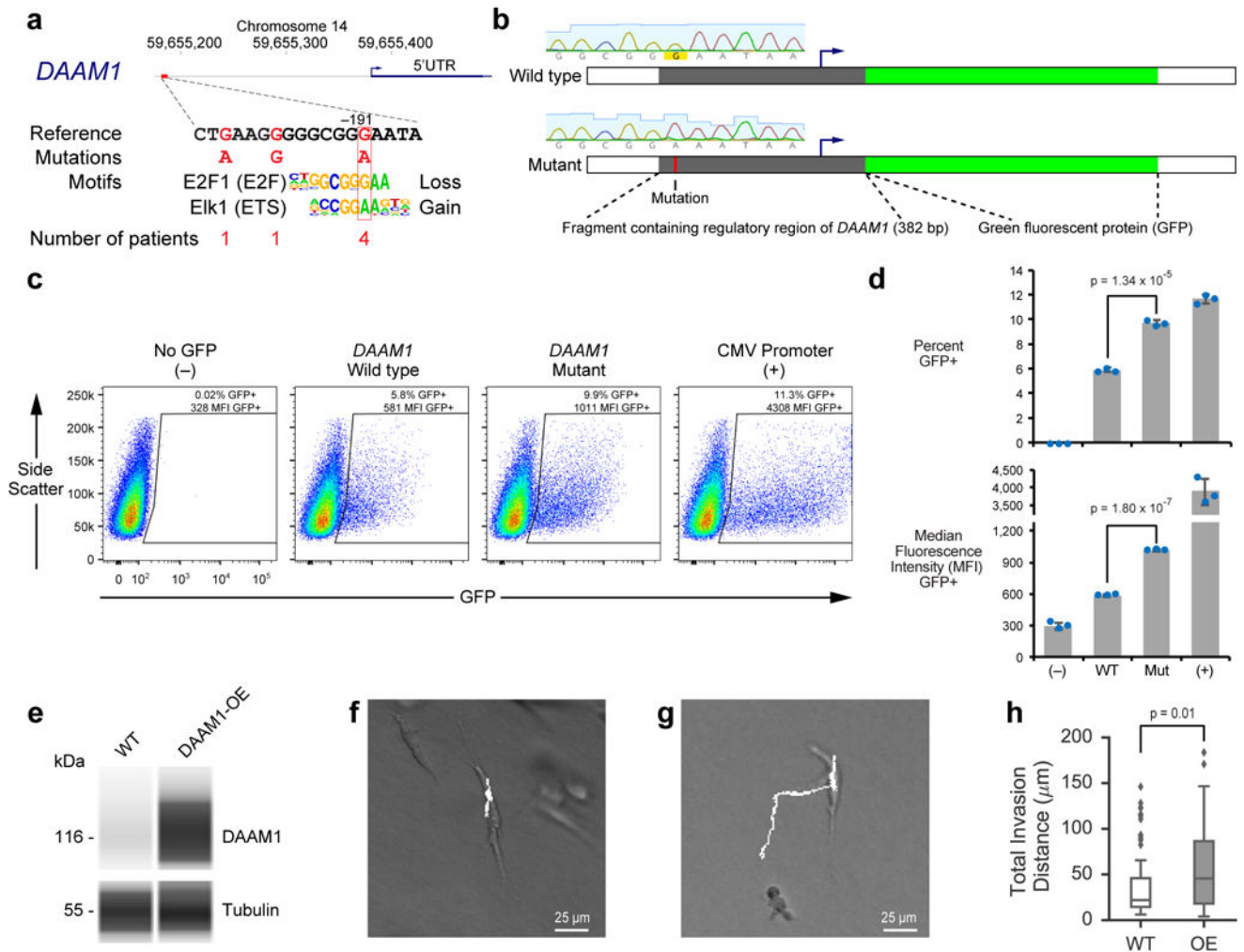


Figure 3. Functional validation of the mutated *DAAM1* regulatory element

(a) A somatic eQTL in the *DAAM1* promoter region is associated with increased mRNA expression levels. (b) Schematic of wild type and mutant GFP reporter constructs along with the Sanger sequencing traces confirming the sequence of the key nucleotide. (c) Flow cytometry analysis of A375 human melanoma cells 48 hours after transient transfection. The polygon delineated by black lines shows the gated region used to define GFP+ cells. (d) Bar graphs (average \pm standard deviation across 3 cell culture replicates; p -values from two-tailed t -tests) showing the percentage of GFP+ cells and the median fluorescence intensity of the GFP+ cells. Individual data points are in Supplementary Table 5. (e) Protein electropherogram analysis of wild type and *DAAM1* overexpressing MDA-MB-231 cells using the antibodies against DAAM1 and tubulin. The complete electropherogram is in Supplementary Fig. 6e. The image is representative of two independent cell culture experiments. (f, g) Sample trajectories of (f) wild type and (g) *DAAM1*-overexpressing cells embedded in 2.5 mg/mL 3D collagen hydrogels. (h) Total invasion distance travelled by individual cells (p -value from two-tailed Mann–Whitney U test; 95% confidence intervals of mean are (32.3 μ m, 48.2 μ m) and (47.6 μ m, 67.0 μ m) for wild type and *DAAM1*-overexpressing cells, respectively). Imaging and quantitation was performed on 74 and 83

cells in the wild type and *DAAMI*-overexpression groups, respectively. Box-plot elements are defined as Fig. 1b.

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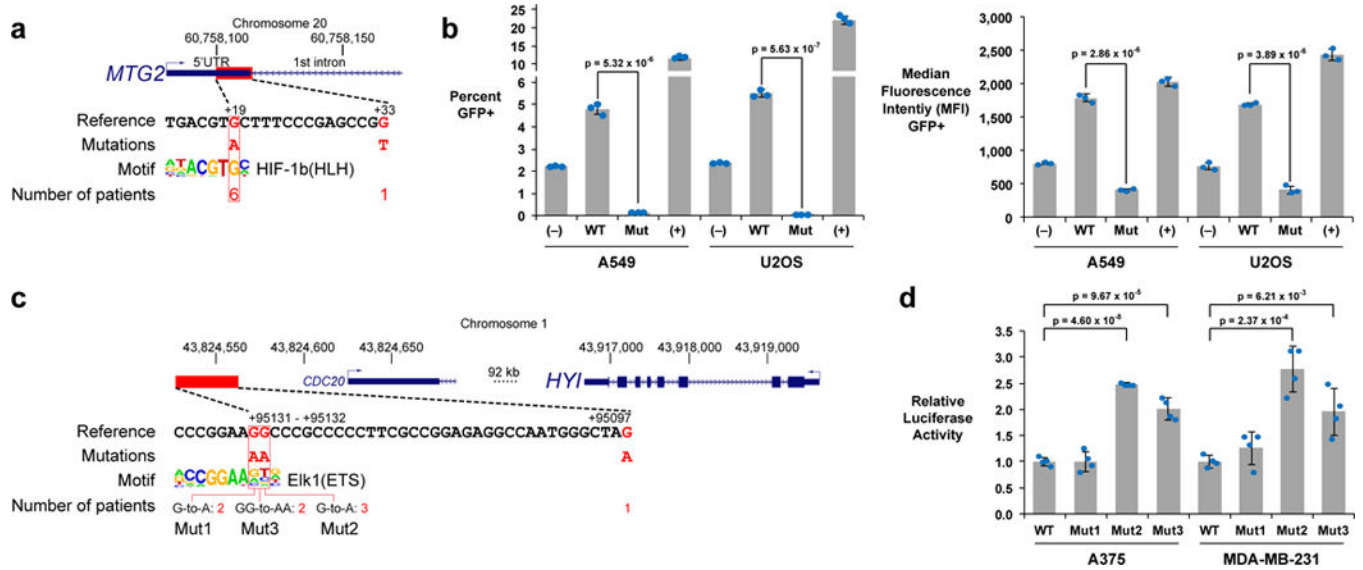


Figure 4. Additional case studies

(a) The somatic eQTL associated with downregulation of *MTG2* is located in its 5' UTR and frequently alters a potential HIF-1b binding motif. (b) Flow cytometry analysis of A549 lung epithelial carcinoma cells and U2OS bone osteosarcoma cells 48 hours after transient transfection with *MTG2* GFP reporter constructs. Bar graphs (mean \pm standard deviation across three cell culture replicates; *p*-values from two-tailed *t*-tests) showing the percentage of GFP+ cells and the median fluorescence intensity of GFP+ events. (c) The somatic eQTL associated with upregulation of *HYI* is located 95 kb downstream of the TSS and frequently alters a potential Ets family binding motif. (d) Luciferase assay results (mean \pm standard deviation across four cell culture replicates; *p*-values from two-tailed *t*-tests) for the *HYI* somatic eQTLs 48 hours after transient transfection in A375 melanoma cells and MDA-MB-231 breast cancer cells. Individual data points are available in Supplementary Tables 5 and 6.

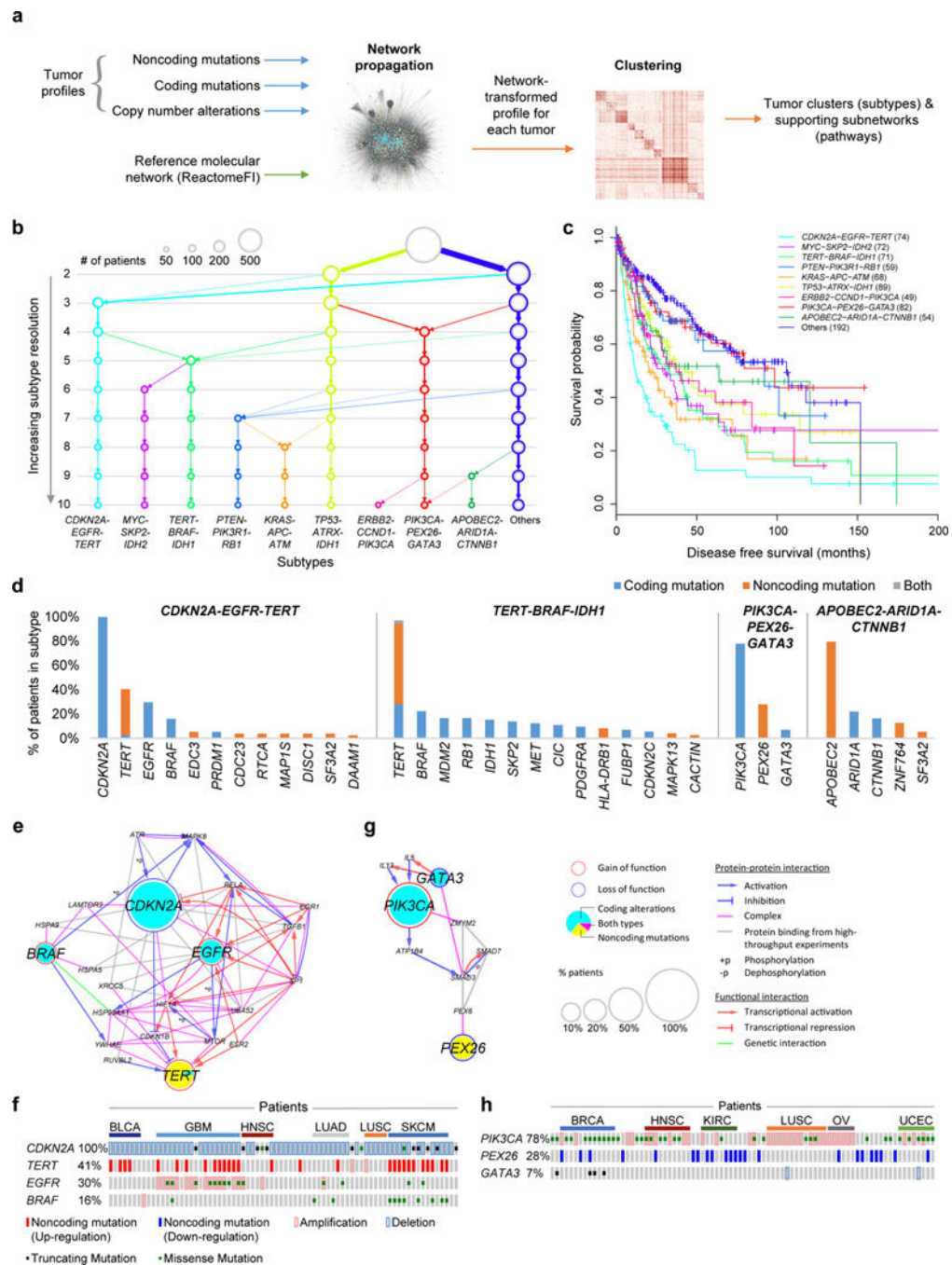


Figure 5. Identification of molecular networks and associated tumor subtypes incorporating noncoding mutations

(a) Workflow of Network-Based Stratification. (b) Resulting hierarchy of subtypes, at increasing resolution from 2-10 subtypes. (c) Disease-free survival probabilities (y-axis) are plotted against time after diagnosis in months (x-axis) for each of the identified cancer subtypes (colors). Patients with censored survival data are indicated by a “+” at the censoring time (last follow-up). (d) Signature genes are shown for each subtype with a large proportion of patients with noncoding mutations (x-axis), ordered by the percent of patients

with alterations (y-axis). **(e, g)** Pathways characterizing **(e)** *CDKN2A-EGFR-TERT* or **(g)** *PIK3CA-PEX26-GATA3* subtypes, defined as subnetwork regions extracted from ReactomeFI by Network-Based Stratification. **(f, h)** Mutation matrix of the **(f)** *CDKN2A-EGFR-TERT* or **(h)** *PIK3CA-PEX26-GATA3* pathway subtypes showing individual tumors (columns, ordered by cancer tissues) with indicated types of mutations on signature genes for that subtype (rows).