# Genomic basis for RNA alterations in cancer

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Transcript alterations often result from somatic changes in cancer genomes<sup>1</sup>. Various forms of RNA alterations have been described in cancer, including overexpression<sup>2</sup>, altered splicing<sup>3</sup> and gene fusions<sup>4</sup>; however, it is difficult to attribute these to underlying genomic changes owing to heterogeneity among patients and tumour types, and the relatively small cohorts of patients for whom samples have been analysed by both transcriptome and whole-genome sequencing. Here we present, to our knowledge, the most comprehensive catalogue of cancer-associated gene alterations to date, obtained by characterizing tumour transcriptomes from 1,188 donors of the Pan-Cancer Analysis of Whole Genomes (PCAWG) Consortium of the International Cancer Genome Consortium (ICGC) and The Cancer Genome Atlas (TCGA)<sup>5</sup>. Using matched whole-genome sequencing data, we associated several categories of RNA alterations with germline and somatic DNA alterations, and identified probable genetic mechanisms. Somatic copy-number alterations were the major drivers of variations in total gene and allele-specific expression. We identified 649 associations of somatic single-nucleotide variants with gene expression in cis, of which 68.4% involved associations with flanking non-coding regions of the gene. We found 1,900 splicing alterations associated with somatic mutations, including the formation of exons within introns in proximity to Alu elements. In addition, 82% of gene fusions were associated with structural variants, including 75 of a new class, termed 'bridged' fusions, in which a third genomic location bridges two genes. We observed transcriptomic alteration signatures that differ between cancer types and have associations with variations in DNA mutational signatures. This compendium of RNA alterations in the genomic context provides a rich resource for identifying genes and mechanisms that are functionally implicated in cancer.

For a more extensive study of cancer genome alterations, particularly in non-coding regions, the PCAWG project was formed to analyse the large number of whole-genome samples that were contributed to the ICGC and TCGA projects<sup>5</sup>. Individual projects did not use the same methods for key analyses; therefore, a major focus for each of the 16 PCAWG Working Groups was the unified analysis of the PCAWG data. For example, the PCAWG Technical Working Group led raw data collection, realignment of whole-genome sequencing data and implemented core somatic mutation calling pipelines<sup>5</sup>. Other PCAWG working groups focused on unified analyses of copynumber variation<sup>6</sup>, structural variants<sup>7,8</sup>, germline variants<sup>5</sup>, mutational signatures<sup>9</sup> and identification of driver genes<sup>8</sup>, among others<sup>5</sup>. Here, we report the joint analysis of available matched transcriptome and genome profiling for 1,188 samples from 27 tumour types by the PCAWG Transcriptome Working Group<sup>5</sup>, providing the largest, to our knowledge, resource of RNA phenotypes and their underlying genetic changes in cancer so far (Extended Data Fig. 1, Methods, Supplementary Results, Supplementary Table 23). We demonstrate the importance of transcriptomics data in understanding how different dimensions of specific DNA alterations contribute to carcinogenesis and map out the landscape of cancer-related RNA alterations.

## Cancer-specific germline cis-eQTLs

To investigate the underlying mechanisms of different types of RNA alteration, we first focused on changes in the gene expression level (Extended Data Fig. 2). We initially considered common germline variants (minor allele frequency  $\geq$  1%) proximal to individual genes ( $\pm$ 100 kb), and mapped expression quantitative trait loci (eQTL) across the cohort (Extended Data Fig. 3, Supplementary Table 1). This pan-cancer analysis identified 3,532 genes with an eQTL (false discovery rate (FDR)  $\leq$  5%, hereafter denoted eGenes) (Supplementary Table 2), enriched in proximal regions of transcription start sites (TSSs) (Extended Data Fig. 3).

To identify cancer-specific regulatory variants, we compared our eQTLs to eQTLs from the Genotype-Tissue Expression (GTEx) project  $^{10}$ , adopting previous strategies to assess eQTL replication  $^{11}$ , and probed lead eQTL variants for marginal significance in GTEx tissues ( $P \! \leq \! 0.01$ , Bonferroni-adjusted). Although most lead variants could be detected in GTEx samples (3,110 out of 3,532 eQTL variants), we identified 422 eQTLs that did not correspond to GTEx tissues, which suggests cancer-specific regulation (Extended Data Fig. 4, Supplementary Table 3). The corresponding eQTL lead variants were enriched for heterochromatic regions (Fig. 1a). Overall, this analysis revealed that the germline framework of gene expression regulation is largely conserved in cancer tissues.

### Somatic cis-eQTLs in non-coding regions

Previous studies have described the landscape of non-coding mutations in cancer<sup>1</sup>, particularly in promoter regions, and also their regulatory effects on gene expression<sup>12,13</sup>. Here, we looked at possible somatic DNA changes, across the whole genome, that underlie alterations in gene expression. We estimated local mutation burdens by aggregating single-nucleotide variants (SNVs) in 2-kb intervals adjacent to genes (flanking), as well as in exons and introns (Extended Data Figs. 2, 5, 6). Next, we decomposed the expression variation of individual genes, considering common mutation burdens in *cis*, as well as *cis* germline variants and somatic copy-number alterations (SCNAs). This identified SCNAs as the major driver of expression variation (17%), followed by somatic SNVs in gene flanking regions (1.8%) and germline variants (1.3%) (Fig. 1b).

We also tested for associations between all common mutation burdens and gene expression across the whole genome. We identified 649 genes with a somatic eQTL (FDR  $\leq$  5%) (Supplementary Table 5). Of these, 11 associations were located in introns or exons of the respective eGene, including genes with known roles in the pathogenesis of specific cancers such as CDK12 in ovarian cancer<sup>14</sup> and IRF4 in chronic lymphocytic leukaemia<sup>15</sup> (Extended Data Figs. 7, 8). Most eQTLs (68.4%) involved associations with flanking non-coding mutation burdens (Extended Data Fig. 6e). Next, we considered eQTLs in flanking regions (n = 556) and tested for enrichment in cell-type-specific annotations from the Epigenetics Roadmap<sup>16</sup>. This identified 13 enriched annotations (FDR ≤ 10%) (Extended Data Fig. 9, Supplementary Table 6), including poised promoters, weak and active enhancers, and heterochromatin, but notably no enrichment for transcription-factor-binding sites (Supplementary Table 7). This enrichment in transcriptionally inactive regions may be due to an increased mutation rate in these regions (Extended Data Fig. 9), which has previously been reported in cancer<sup>17</sup>.

We also looked at the functional characterization of somatic eGenes and observed an enrichment for somatic eQTLs in bivalent promoters for cancer testis genes (P = 0.04, Fisher's exact test) such as  $TEKT5^{18}$  (Fig. 1c, Extended Data Fig. 8h). Furthermore, we found a global enrichment (FDR  $\leq$  10%) for Gene Ontology (GO) categories related to cell differentiation and developmental processes (Supplementary Table 8). Overall, somatic eQTL analysis identified mostly non-coding regions

associated with changes in local gene expression and, similar to cancerspecific germline eQTLs, showed enrichment for transcriptionally inactive regions such as heterochromatin.

## **Expression and mutational signatures**

Global variations in mutational patterns can be quantified using mutational signatures, which tag mutational processes specific to their tissue-of-origin and environmental exposures<sup>19</sup>. However, the extraction of mutational signatures is an intrinsically statistical process that requires a posteriori functional annotation. We performed a pan-cancer association analysis between genome-wide mutational signatures and gene expression levels to decipher the molecular processes that accompany the presence of mutational signatures.

We considered 28 mutational signatures derived using non-negative matrix factorization of context-specific mutation frequencies. We tested for association between signature prevalence in donors and total gene expression, accounting for total mutational burden, cancer type, and other technical and biological confounders. This identified 1,176 genes associated with at least one signature (FDR  $\leq$  10%) (Extended Data Fig. 10, Supplementary Table 19).

We considered 18 signatures with 20 or more associated genes for further annotation (Extended Data Fig. 11) and assessed enrichment using GO categories and Reactome pathways We found that 11 signatures were enriched for at least one category (FDR  $\leq$  10%) (Supplementary Table 19), revealing associations consistent with known and unknown aetiologies (Fig. 1d). For example, signature 38, which is correlated with the canonical UV signature 7 ( $r^2 = 0.375, P = 5 \times 10^{-40}$ ) (Extended Data Fig. 11c), was linked to melanin processes (Fig. 1d). The synthesis of melanin causes oxidative stress to melanocytes and we found signature 38 associated with the oxidative-stress-promoting gene  $TYR^{23}$  ( $P = 1.0 \times 10^{-4}$ ). A hallmark of signature 38 genes are C>A mutations, a typical product of reactive oxygen species Hat signature 38 may capture DNA damage that is indirectly caused by UV-induced oxidative damage after direct sun exposure, with TYR as a possible mediator of the effect.

#### Genomic basis of allelic expression

To analyse expression at the level of individual haplotypes, we tested for allelic expression imbalance (AEI) (FDR  $\leq$  5%, binomial test). We observed substantial differences in the fraction of genes with AEI between different types of cancer (Extended Data Fig. 12), and between cancer and the corresponding healthy tissues, with a high observed concordance between allelic imbalance at the DNA and RNA levels (Extended Data Fig. 13).

We used a logistic regression model to identify the determinants of AEI, accounting for known imprinting status<sup>26</sup>, the germline eQTL genotype, SCNAs and the weighted mutational burden of proximal somatic SNVs stratified into functional categories (Extended Data Fig. 2). In aggregate, SCNAs accounted for 84.3% of the total explained variation, which confirmed our findings from the somatic eQTL analysis, followed by germline eQTL lead variants (9.1%), somatic SNVs (4.9%) and imprinting status (1.7%) (Extended Data Fig. 14). Although cumulatively, non-coding variants were more relevant than coding variants, somatic protein-truncating variants ('stop-gained' variants) that triggered nonsense-mediated decay<sup>27</sup> were the most predictive individually. SNVs within splice regions, 5' untranslated regions (UTRs) and promoters were also strongly associated with the presence of AEI, and we observed a global trend of decreasing relevance of variants with increasing distance from the TSS (Fig. 1e, Extended Data Fig. 14).

Gene-centric attribution of AEI to individual sources of genetic variation (Supplementary Table 9) revealed an enrichment of somatically induced AEI in several known cancer-driver genes, as well as new candidates, such as the mismatch-repair-related gene *EXO1* that is associated

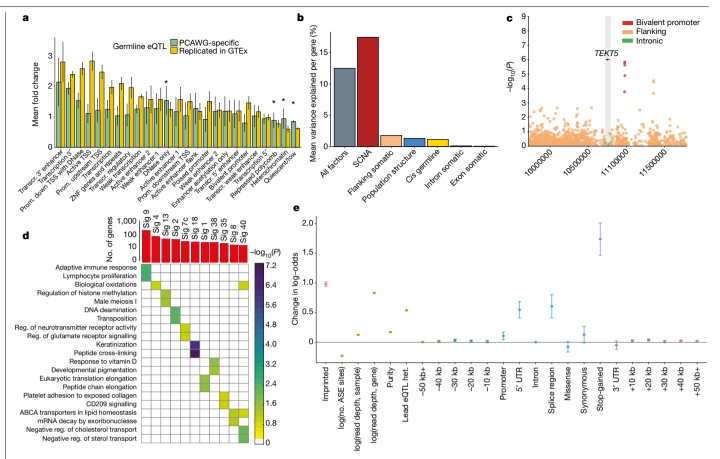


Fig. 1 | Germline and somatic SNVs associated with expression. a, Epigenetics Roadmap enrichment analysis, showing the average fold change in Roadmap factors across cell lines in PCAWG-specific eQTLs of the pan-analysis as well as eQTLs that replicate in GTEx tissues. \*P<0.05/25, one-sided Wilcoxon ranksum test in PCAWG-specific eQTLs corrected for the number of Roadmap factors used (that is, 25). Data are mean and s.d. b, Variance component analysis for gene expression levels, showing the average proportion of variance explained by different germline and somatic factors for different sets of genes including the mean effect across all factors: (1) all genetic factors (germline and somatic); (2) SCNAs; (3) somatic variants in flanking regions; (4) population structure; (5) cis-germline effects; and (6) somatic intron and exon mutation effects. c, Manhattan plot showing nominal P values of association for TEKT5 (highlighted in grey), considering flanking, intronic and exonic intervals.

The leading somatic burden is associated with increased TEKT5 expression  $(P=1.61\times10^{-6})$  and overlaps an upstream bivalent promoter (red dots; annotated in 81 Roadmap cell lines, including 8 embryonic stem cells, 9 embryonic-stem-cell-derived and 5 induced pluripotent stem-cell lines). d, Summary of significant associations between mutational signatures (Sig) and gene expression. Top, the total number of associated genes per signature (FDR≤10%). Bottom, enriched GO categories or Reactome pathways for genes associated with each signature (FDR≤10%, significance level encoded in colour, -log<sub>10</sub>-transformed adjusted P value). **e**, Standardized effect sizes on the presence of AEI, taking only SCNAs, germline eQTLs, coding and noncoding mutations into account. Data are the estimate and standard error of the estimate of the effect size.

with survival in colorectal adenocarcinoma (log-rank P = 0.022, hazard ratio = 0.57) (Supplementary Results). We further observed a strong enrichment in the AEI score of cancer testis genes based on somatic SNVs only ( $\chi^2$  test  $P = 6 \times 10^{-3}$ ). In summary, we identify somatic and germline genetic variation that is associated with allele-specific dysregulation of genes across cancer types.

## Mutations associated with promoter usage

We considered promoter activity<sup>28-30</sup> as another molecular phenotype to study the effect of promoter mutations. Although cancer-specific alternative promoter usage has previously been shown<sup>28</sup>, the association of underlying genomic alterations with promoter activities have not been broadly explored. To estimate the activity of individual gene promoters, we combined the expression of isoforms initiated in TSSs that are identical or nearby, assuming that these are transcribed from the same promoter (Extended Data Fig. 15a-c). We divided promoters into three categories: (1) inactive promoters (activity < 1 fragment per kilobase of transcript per million mapped reads (FPKM)), (2) major promoters (most active per gene) and (3) minor (all remaining) promoters,

and examined the rates of mutation across varying activity levels. We observed an increase in the number of mutations near the TSS of major promoters compared with minor or inactive promoters (Extended Data Fig. 15d). This pattern is most prominent in skin melanoma, in which it has been attributed to impaired nucleotide-excision repair (Extended Data Fig. 15e, f, k, l). The cancer type that shows the strongest deviation from this pattern is colorectal adenocarcinoma, which highlights the tissue-specificity of mutational patterns at promoters (Extended Data Fig. 15e, f, m, n). Only 171 promoters show mutations in more than 5 samples per tumour type in a 200-bp window upstream of the promoter (Extended Data Fig. 15g, h). Most mutations occur in skin melanoma and lymphoma, which is expected owing to reduced nucleotide-excision repair and activation-induced cytidine deaminase (Extended Data Fig. 15h). We did not find significant pan-cancer associations between promoter mutational burden and promoter activity (Extended Data Fig. 15i, j). However, TERT has the highest number of promoter mutations <sup>1,5,31</sup> (Extended Data Fig. 16a), and these mutations have previously been reported to be associated with TERT expression<sup>1</sup>; therefore, we investigated the TERT locus in more detail (Extended Data Fig. 16b). Although TERT does not show a significant association in the

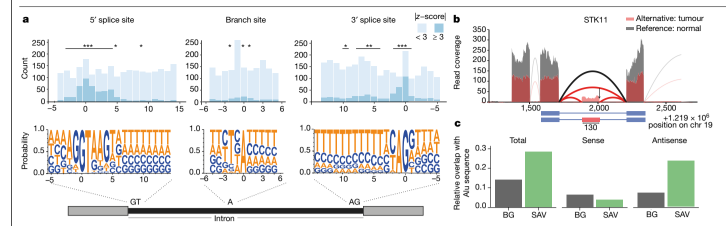


Fig. 2 | Position-specific effect of somatic mutations on alternative splicing. a, Top, proportion of mutations near exon–intron junctions and at branch sites that are associated with exon-skipping events. Mutations with associated splicing changes are those in which the percentage spliced in-derived |z-score| is  $\geq$  3 (dark blue). Asterisks denote intron positions significantly enriched for splicing changes relative to background based on a permutation test. \*P< 0.05, \*\*P< 0.01, \*\*\*P< 0.001. Bottom, sequence motifs of regions. **b**, Example of an

exonization event in the tumour-suppressor gene STK11. The RNA-seq read coverage for a part of the gene is shown in red for a donor carrying the alternative allele, and in grey for a random donor with reference allele. The cassette exon event is shown as a schematic below.  $\mathbf{c}$ , Enrichment of SINE elements in SAVs compared to sequence background (BG). Shown for SINE elements overlapping in sense (middle) and antisense (right) directions.

pan-cancer analysis, we found an association with increased promoter activity in individual types of cancer<sup>1</sup> (Extended Data Fig. 16c).

#### Mutations associated with splicing

Extending the classical hallmarks of cancer, alternative splicing is seen as increasingly relevant to explain cancer heterogeneity<sup>32</sup>. On the basis of our observations of a globally changing splicing landscape (Extended Data Fig. 17a-c), we sought to specifically understand the relationship between splicing changes and somatic mutations within introns. Focusing on cassette exon events, we integrated the quantification of splice events with somatic variants and identified 5,282 mutations near exon-intron boundaries, 1,800 (34%) of which were associated with a change in splicing (|z-score $| \ge 3$ ) (Supplementary Table 10). Consistent with previous findings using exome sequencing<sup>33,34</sup>, most mutations overlapping the essential dinucleotide motifs of the acceptor or donor site are associated with a splicing change-61% or 57%, respectively (Fig. 2a). Nearly one-third of all mutations (226) out of 469) in a 5-nucleotide window downstream of the 5' site were significantly enriched for splicing changes (Fig. 2a). Almost all changes significantly associated with somatic mutations had a negative effect on splicing (96%) (Extended Data Fig. 17d). For mutations in or near the poly-pyrimidine tract, we found a significant (permutation test, P < 0.05) enrichment for mutations linked to outlier splicing (Fig. 2a). We also found an enrichment (P < 0.05, fold change > 2) of splicing outliers at branch-site adenosines (Fig. 2a middle, Extended Data Fig. 17d, Supplementary Table 11). Together, these results suggest that somatic mutations in the extended splice site region, poly-pyrimidine tract and branch point can affect splicing.

We also identified 1,900 rare splicing-associated variants (SAVs) that appear in only a small number of samples using the SAVNet approach  $^{35}$  (Extended Data Fig. 17e; see 'Data availability' in the Methods). Notably, 862 SAVs affected canonical splice sites, whereas the other 1,038 disrupted non-canonical sites or created new splice sites. Notably, we find a twofold enrichment of cancer genes in SAVs (Extended Data Fig. 17f).

Although we find that those SAVs that create splice sites strongly concentrate near exon–intron boundaries (Extended Data Fig. 17g), 45.9% of SAVs are further than 100 bp away from the nearest annotated exon. Mutations at those sites generally changed the sequences towards the donor or acceptor motif consensus (Extended Data Fig. 17h). Focusing on novel splice sites deep in introns, we analysed the extent of exonizations—that is, the formation of new exons within an intron (Extended

Data Fig. 17j, Supplementary Tables 13, 14). More than one-fifth of these new exons (9 out of 43) occur in cancer-related genes, such as the well-known tumour-suppressor gene *STK11*. As expected, the exonization event would cause a frameshift in *STK11* (Fig. 2b, Extended Data Fig. 17k).

Alu elements that are inserted in an antisense direction have sequences that resemble consensus splice sites that, together with activating mutations, can lead to the formation of a new exon<sup>36</sup> (Extended Data Fig. 17l). We found a significant enrichment of splice-site-creating SAVs within annotated Alu sequences ( $P = 2.8 \times 10^{-9}$ ), particularly in the antisense direction ( $P = 2.6 \times 10^{-15}$ ) (Fig. 2c). Our results indicate that the exonization of Alu sequences, which has been extensively studied in the context of primate genome evolution, is also observed in cancer genome evolution.

### Patterns of gene fusions across cancer

Gene fusions are an important class of cancer-driving event with therapeutic and diagnostic value<sup>37</sup>. We identified a total of 925 known and 2,372 new cancer-specific gene fusions by combining the output of two fusion discovery methods as well as genomic rearrangement (structural variants) information and excluding artefacts or fusions in non-cancer samples<sup>38</sup> (Fig. 3a). For the 3,540 identified fusion events representing 3,297 unique gene fusions, we categorized them on the basis of novelty, recurrence and known oncogenic gene partners (Fig. 3a).

Only 149 (approximately 5%) of the fusions occur in more than one sample, among which 78 are novel. Most of these (46 out of 78) were found across several histotypes. Of the 27 most recurrent gene fusions (Extended Data Fig. 18a), 8 have previously been reported (for example, *CCDC6-RET*<sup>39</sup>, *FGFR3-TACC3*<sup>40</sup> and *PTPRK-RSPO3*) or independently detected in the TCGA cohort<sup>41</sup>, whereas 6 were new (such as *NUMB-HEATR4*, *ESR1-AKAP12* and *TRAF3IP2-FYN*). In total, 105 fusion transcripts involved the UTR region of one gene and the complete coding sequences of another gene, possibly resulting from structural variation in promoter regions.

Although most genes involved in fusions engaged with only one fusion partner, 35 genes had more than 5 partners. These 'promiscuous' genes tended to be selective in being either a 5' or a 3' partner with conserved break points and positions (3' or 5'), and were overrepresented in cancer census genes and the PCAWG cancer-driver genes (one-tailed Fisher's exact test, odds ratio = 8.66,  $P \le 1.1 \times 10^{-15}$ , and odds ratio = 12.27,  $P \le 2.2 \times 10^{-16}$ , respectively). Network analysis of promiscuous genes and their partners revealed several large gene clusters containing at least 10 genes (Extended Data Fig. 18b), enriched

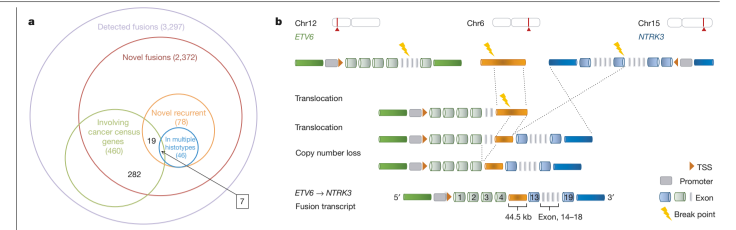


Fig. 3 | Structural rearrangements associated with RNA fusions. a, The number of all detected and new fusions and their overlap with the cancer census genes, **b**. Schematic of an example of bridged fusions. Bridged fusions

are those composite fusions formed by a third genomic segment that bridges two genes. Only one of the possible orders of genomic arrangement is depicted in each case, with break points highlighted as thunderbolts.

in cancer-related pathways (Benjamini-Hochberg corrected  $P \le 0.01$ ) and in protein-protein interactions ( $P \le 1.0 \times 10^{-7}$ ), which suggests a possible functional role in cancer.

Notably, a large number of fusions, including known fusions, could not be associated with only a single structural-variation event. For example, the ETV6-NTRK3 gene fusion 42 was present in a head and neck thyroid carcinoma sample, linking exon 4 of ETV6 to exon 12 of NTRK3. We found three separate structural variants in the same sample: (1) a translocation of ETV6 to chromosome 6; (2) a translocation of NTRK3 also to chromosome 6; and (3) an additional copy-number loss spanning from intron 5 of ETV6 to the exact structural variant break points, jointly bringing ETV6 within 45 kb upstream of NTRK3-a distance that would allow transcriptional read-through<sup>43</sup> or splicing<sup>44</sup> to yield the ETV6-NTRK3 fusion<sup>45</sup> (Fig. 3b). Thus, the short chromosome-6 segment appeared to function as a bridge, which linked two genomic locations to facilitate a gene fusion. We term such products 'bridged fusions'. This class of fusion is not uncommon. Out of a total of 436 gene fusions supported by 2 separate structural variants, 75 are bridged fusions (Supplementary Table 15).

On the basis of the nature of the underlying genomic rearrangements, we propose a unified fusion classification system (Extended Data Fig. 19a). Aside from bridged fusions, 344 additional fusions are linked to more than one structural variant in the same sample. These multi- structural variant fusions are collectively termed 'composite fusions' (Extended Data Fig. 19a, b). We find 284 intercomposite fusions (interchromosomal translocation) and 124 intracomposite fusions (intrachromosomal rearrangement), exemplified by ERC1-RET1 and NUMB-HEATR4 fusions, respectively (Extended Data Fig. 19b). Composite rearrangements bring the fusion partners significantly closer to each other, from the median natural distance of 6.8 Mb to the median of 7.9 kb (Wilcoxon rank-sum test,  $P \le 2.2 \times 10^{-16}$ ; Extended Data Fig. 19c) after translocation. For 18% of fusions, no evidence of structural variation was found. Given that 340 structural-variant-independent, intrachromosomal fusions had significantly closer break points than those with structural variation (Extended Data Fig. 19d), it is possible that they could result from RNA read-through events. The other possibility is that the underlying supporting structural variants escaped detection, as shown by the observation that known gene fusions that are driven by structural variation, such as TMPRSS2-ERG46, did not have consistent evidence for structural variation in matching samples.

#### Landscape of RNA alterations in cancer

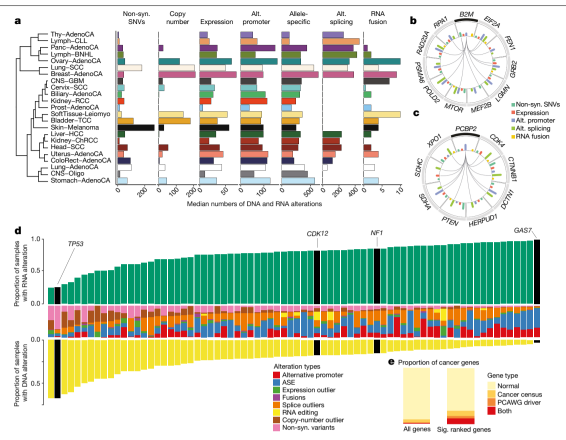
Given our comprehensive set of RNA alterations, we sought to characterize the heterogeneous mechanisms of cancer genome and transcriptome alterations. To enable joint analyses of RNA and DNA alterations, we created a gene-level table, which indicates the presence or absence of possible functional changes to RNA or DNA for each gene and donor. After stringent filtering, we identified 1,523,098 alteration events, in which an event is a gene-sample-alteration triplet (Extended Data Table 1, Supplementary Table 14). It should be noted that we chose to include only RNA alterations with potential functional effects or with the strongest quantitative affect, resembling similar strategies for filtering DNA alterations<sup>47</sup>. Recurrence analysis across several alteration types helped us to further enrich for functionally relevant genes. Building on the gene-centric table, we characterized gene alterations at the RNA level and contrasted these with DNA alterations (non-synonymous SNVs or SCNAs)<sup>5</sup>. On the basis of the calculated association between each RNA- and DNA-level alteration across all histotypes, we found that half of the RNA alterations significantly correlated with DNA alterations (likelihood ratio test, FDR  $< 1 \times 10^{-4}$ ) (Extended Data Fig. 20).

When comparing gene alteration frequencies across all histotypes (Fig. 4a), we note that different types of cancer contain distinct combinations of DNA- and RNA-level alterations (Fig. 4a, Supplementary Table 17). Although, as expected, skin melanoma significantly exceeds other cancers in the number of non-synonymous SNVs<sup>48</sup> (Wilcoxon rank-sum test, P < 0.012), lymphatic cancers have low numbers of SNVs (Wilcoxon rank-sum test,  $P = 5.3 \times 10^{-15}$ ), but high incidences of alternative splicing outliers (Wilcoxon rank-sum test,  $P = 4.9 \times 10^{-47}$ ), which suggests that transcriptomic alterations can be relatively more pronounced in certain cancer types.

To evaluate to which extent RNA changes provide additional mechanisms for cancer gene alterations, we examined DNA- and RNA-level alterations both in sets of genes in pathways (Extended Data Fig. 21) and in individual genes with known roles in cancer (Extended Data Fig. 22). We found that RNA alterations occur at a high proportion in many pathways, including the NOTCH and TGF-β pathways. In addition, KRAS exhibits more RNA alterations than DNA alterations in some types of cancer. Given the recent finding that alternative splicing of KRAS expanded the prognostic affect beyond mutation status in colorectal cancer<sup>49</sup>, our data further support several modes of alteration for KRAS in tumours.

#### Co-occurrence of RNA and DNA alterations

The diverse types of alteration in this study enabled us to investigate trans-associations between different genetic and expression characteristics involving cancer-related genes (FDR < 5%) (Supplementary Table 18). By investigating whether somatic mutations of known cancer



**Fig. 4** | **Global view of DNA and RNA alterations that affect tumours. a**, The median numbers of different alterations across histotypes. Histotypes are ordered by hierarchical clustering based on the pattern of different types of alteration. Only histotypes with more than 10 donors are shown. Alt., alternative; non-syn, non-synonymous. Cancer-type abbreviations are listed in Supplementary Table 23. **b, c**, Circular representations of the selected genes significantly co-occurred with  $B2M(\mathbf{b})$  and  $PCBP2(\mathbf{c})$ . Connecting lines indicate the specific types of co-occurrence of alteration pairs. The inner histograms

indicate the frequencies of incidences of different alteration types shown in different colours.  ${\bf d}$ , All 74 Catalogue of Somatic Mutations in Cancer (COSMIC) cancer census genes or PCAWG driver genes that are both frequently and heterogeneously altered across both RNA- and DNA-level alterations. Yellow bars indicate the proportion of samples that had DNA-level alterations, and green bars indicate the proportion of samples with RNA-level alterations. Middle column is the proportion of each alteration type observed for that gene.  ${\bf e}$ , The enrichment of cancer genes within our list of significantly recurrent genes.

genes are associated with the expression of other genes, we found *IDH1* and *NFKBIE* to be widely linked to the dysregulation of many genes (Extended Data Fig. 23a, b). Notable co-occurrences were present in several types of cancer. For example, *B2M* and *EIF4G2* alterations were simultaneously observed in both B-cell non-Hodgkin lymphoma and lung squamous cell carcinoma. Pathway enrichment analysis of the top 100 genes associated with all *B2M* alterations indicates that the most affected genes are involved in DNA repair (FDR  $\leq$  1%), and approximately two-thirds of those associations were significant in more than one cancer type (Fig. 4b, Extended Data Fig. 23c).

We also examined how cancer genes could be affected by other genes by co-occurrence analyses. Expression outliers of PCBP2 co-occurred with aberrant splicing of a large number of cancer-related genes, including CTNNB1 and CDK4 (Fig. 4c). PCBP2 has been reported to enhance the splicing of cassette exons 50. Our results thus further support the possible role of PCBP2 in regulating the splicing of cancer-related genes.

## Recurrent RNA alterations in driver genes

In our analyses of *cis*-acting mutations that are associated with these individual RNA phenotypes, the vast majority were observed rarely in the PCAWG cohort. Many cancer genes (such as *MET*<sup>51,52</sup>) are known to be somatically altered by heterogeneous mechanisms such as gene fusions, splicing mutations and non-synonymous mutations; therefore, examining genes that are altered by several *cis*-acting mechanisms may help to identify cancer genes in which an individual alteration

type is rare. A total of 5,413 genes were altered by gene expression, allele-specific expression (ASE), splicing and/or gene fusion, and had an associated DNA-level mutation in cis (Supplementary Table 20). PCAWG-defined driver genes<sup>8</sup> tended to have more diverse mechanisms of RNA-level alterations when compared to genes that have not previously been identified as a cancer gene (P < 0.001) (Extended Data Fig. 24a). We identified, for example, a somatic eQTL, a splicing-associated variant and fusions in the known tumour-suppressor NF1 in the MAPK pathway (Extended Data Fig. 24b).

Owing to the fact that most somatic mutations are rare<sup>5</sup>, it is difficult to statistically distinguish functionally relevant, potential driver alterations from passenger alterations. Therefore, we aimed to identify genes that are both recurrently and heterogeneously altered, under the hypothesis that these genes have increased functional relevance. This analysis identified 731 genes with significant recurrent aberrations (FDR < 5%) (Extended Data Fig. 25a), with the top-ranking genes carrying both RNA and DNA alterations. RNA alterations accounted for 0.05-99.14% (mean: 78.23%) of all identified alterations in each gene (Extended Data Fig. 25a, Supplementary Table 21). This ranking is enriched for the union of cancer census genes<sup>53</sup> (60 out of 603) and PCAWG-defined driver genes (33 out of 157, unioned: 74 out of 674  $P=4.6\times10^{-13}$ , enrichment: 2.45) (Fig. 4d, e).

Among the top 10% of our ranked genes is CDK12 (rank 55). We find 91 samples that have an alteration involving its protein kinase domain, which has been implicated in DNA repair dysregulation 54. Many of these samples have no DNA-level alterations in CDK12 (46%) (Extended Data

Fig. 26a). Furthermore, splicing, alternative promoter, SNV, RNA-editing and fusion alterations in this gene are mutually exclusive (adjusted  $P=4.8\times10^{-3}$ ) (Extended Data Fig. 26b, c). Upon further investigation, we find that somatic eQTL mutations in CDK12 are associated with a tandem duplicator phenotype<sup>55</sup>. Although this association was not replicated with other RNA alterations, it provides evidence that somatic CDK12 mutations may alter its function through gene expression changes. This example illustrates that performing a recurrence analysis over diverse RNA and DNA alterations can help to identify genes known to be important in tumorigenesis.

### Discussion

Here we present a comprehensive catalogue of RNA-level alterations in cancer, spanning 27 different tumour types, and provide a harmonized resource of matched transcriptome and whole-genome sequences. We identified 731 genes that were recurrently altered by several mechanisms, jointly enriched for known cancer census and PCAWG driver genes<sup>8</sup>. The list includes genes that are primarily altered at the DNA level (such as TP53), but also genes for which the alteration most frequently manifests in RNA (such as GAS7). Out of 87 samples from the PCAWG study that did not have a driver alteration at the DNA level<sup>5</sup>, and had RNA-sequencing (RNA-seq) data, every sample had an RNA-level alteration identified. Although cancer is thought to be driven by changes in DNA primarily, some driver alterations may manifest themselves via changes in RNA rather than DNA sequence mutations.

We identified germline eQTLs for around 20% of expressed genes. The number of eGenes found is generally low compared with some other studies, reflecting the heterogeneity of our samples. Only 422 genes appeared to be specific to cancer; this is likely to be an underestimate owing to the heterogeneity, small sample numbers and the rather conservative strategy chosen. We have also mapped linkages between genes and somatic aberrations in cis, in which 68.4% of associations were between non-coding somatic variants and gene expression. Allelic copy-number imbalance is a major determinant of ASE dysregulation in cancer. We found mutations associated with splicing changes including novel cancer-specific exons that can be partially explained by mutation-driven exonization. We systematically compared gene fusions with whole-genome rearrangements across many tumour types and found 82% of detected fusions were associated with specific genomic rearrangements. For the remaining fusions, it is possible that the relevant genomic rearrangements have not been detected, or that some fusions happen directly at the RNA level, as trans-splicing or read-through events. The availability of whole-genome sequences allowed us to develop a systematic classification of fusion events and to propose a new bridged fusion mechanism.

Because global differences in RNA expression phenotypes are largely tissue-specific, our ability to associate mutations in *cis* or *trans* are limited by the small and variable sample sizes within each histotype. Further work is needed to investigate other mechanisms of genome alteration that can lead to changes in RNA such as epigenetic changes<sup>56</sup> or enhancer hijacking<sup>57</sup>. Our work will help to prioritize further investigations.

Overall, our analyses show diverse modes of alteration of cancer genes and pathways at the DNA and RNA levels, and demonstrate that RNA analyses reveal cancer-associated pathway alterations that have not yet been detected via DNA-only approaches. These insights illustrate the power of integrated transcriptome and whole-genome sequencing analysis for cancer studies.

#### Online content

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- Weinhold, N., Jacobsen, A., Schultz, N., Sander, C. & Lee, W. Genome-wide analysis of noncoding regulatory mutations in cancer, Nat. Genet. 46, 1160-1165 (2014).
- Owens, M. A., Horten, B. C. & Da Silva, M. M. HER2 amplification ratios by fluorescence in situ hybridization and correlation with immunohistochemistry in a cohort of 6556 breast cancer tissues. Clin. Breast Cancer 5, 63-69 (2004).
- Climente-González, H., Porta-Pardo, E., Godzik, A. & Eyras, E. The functional impact of alternative splicing in cancer, Cell Reports 20, 2215-2226 (2017)
- Faderl, S. et al. The biology of chronic myeloid leukemia. N. Engl. J. Med. 341, 164–172 (1999).
- The ICGC/TCGA Pan-Cancer Analysis of Whole Genomes Network, Pan-cancer analysis of whole genomes. Nature https://doi.org/10.1038/s41586-020-1969-6 (2020)
- Gerstung M et al. The evolutionary history of 2 658 cancers. Nature https://doi. org/101038/s41586-019-1907-7 (2020)
- Li. Y. et al. Patterns of somatic structural variation in human cancer genomes. Nature https://doi.org/10.1038/s41586-019-1913-9 (2020).
- Rheinbay, E. et al. Analyses of non-coding somatic drivers in 2,693 cancer whole genomes. Nature https://doi.org/10.1038/s41586-020-1965-x (2020).
- Alexandrov, L. B. et al. The repertoire of mutational signatures in human cancer. Nature https://doi.org/10.1038/s41586-020-1943-3 (2020)
- 10. GTEx Consortium. Genetic effects on gene expression across human tissues. Nature 550, 204-213 (2017)
- Kilpinen, H. et al. Common genetic variation drives molecular heterogeneity in human iPSCs. Nature 546, 370-375 (2017).
  - Fredriksson, N. J., Ny, L., Nilsson, J. A. & Larsson, E. Systematic analysis of noncoding somatic mutations and gene expression alterations across 14 tumor types. Nat. Genet. 46, 1258-1263 (2014).
- Gong, J. et al. PancanQTL: systematic identification of cis-eQTLs and trans-eQTLs in 33 cancer types. Nucleic Acids Res. 46, D971-D976 (2018).
- 14. Bajrami, I. et al. Genome-wide profiling of genetic synthetic lethality identifies CDK12 as a novel determinant of PARP1/2 inhibitor sensitivity. Cancer Res. 74, 287-297 (2014).
- Havelange, V. et al. IRF4 mutations in chronic lymphocytic leukemia. Blood 118, 2827-2829 (2011)
- 16. Roadmap Epigenomics Consortium et al. Integrative analysis of 111 reference human epigenomes. Nature 518, 317-330 (2015).
- Zheng, C. L. et al. Transcription restores DNA repair to heterochromatin, determining regional mutation rates in cancer genomes. Cell Reports 9, 1228-1234 (2014).
- Hanafusa, T., Mohamed, A. E. A., Domae, S., Nakayama, E. & Ono, T. Serological identification of Tektin5 as a cancer/testis antigen and its immunogenicity. BMC Cancer 12. 520 (2012).
- Alexandrov, L. B. et al. Signatures of mutational processes in human cancer. Nature 500, 415-421 (2013) Milacic, M. et al. Annotating cancer variants and anti-cancer therapeutics in reactome.
- Cancers (Basel) 4, 1180-1211 (2012)
- Fabregat, A. et al. The Reactome pathway Knowledgebase. Nucleic Acids Res. 44, D481-D487 (2016)
- Kvam, E. & Tyrrell, R. M. The role of melanin in the induction of oxidative DNA base damage by ultraviolet A irradiation of DNA or melanoma cells. J. Invest. Dermatol. 113,
- Jimbow, K., Chen, H., Park, J. S. & Thomas, P. D. Increased sensitivity of melanocytes to oxidative stress and abnormal expression of tyrosinase-related protein in vitiligo. Br. J. Dermatol. 144, 55-65 (2001)
- Pilger, A. & Rüdiger, H. W. 8-Hydroxy-2'-deoxyguanosine as a marker of oxidative DNA damage related to occupational and environmental exposures. Int. Arch. Occup. Environ. Health 80, 1-15 (2006)
- Premi, S. & Brash, D. E. Unanticipated role of melanin in causing carcinogenic cyclobutane pyrimidine dimmers. Mol. Cell. Oncol. 3, e1033588 (2015).
- Morison, I. M., Ramsay, J. P. & Spencer, H. G. A census of mammalian imprinting, Trends Genet. 21, 457-465 (2005).
- Lindeboom, R. G. H., Supek, F. & Lehner, B. The rules and impact of nonsense-mediated mRNA decay in human cancers. Nat. Genet. 48, 1112-1118 (2016).
- Demircioğlu, D. et al. A pan-cancer transcriptome analysis reveals pervasive regulation through alternative promoters, Cell 178, 1465-1477,e17 (2019).
- Reves, A. & Huber, W. Alternative start and termination sites of transcription drive most transcript isoform differences across human tissues. Nucleic Acids Res. 46, 582-592 (2018)
- 30. Feng, G. et al. Ubiquitously expressed genes participate in cell-specific functions via alternative promoter usage. EMBO Rep. 17, 1304-1313 (2016).
- Huang, F. W. et al. Highly recurrent TERT promoter mutations in human melanoma Science 339, 957-959 (2013)
- 32. Oltean, S. & Bates, D. O. Hallmarks of alternative splicing in cancer. Oncogene 33, 5311-5318 (2014).
- Jung, H. et al. Intron retention is a widespread mechanism of tumor-suppressor inactivation. Nat. Genet. 47, 1242-1248 (2015).
- Kahles, A. et al. Comprehensive analysis of alternative splicing across tumors from 8.705 patients. Cancer Cell 34, 211-224.e6 (2018).
- 35. Shiraishi, Y. et al. A comprehensive characterization of cis-acting splicing-associated variants in human cancer. Genome Res. 28, 1111-1125 (2018).
- Sorek, R. The birth of new exons: mechanisms and evolutionary consequences. RNA 13, Mertens, F., Johansson, B., Fioretos, T. & Mitelman, F. The emerging complexity of gene
- fusions in cancer. Nat. Rev. Cancer 15, 371-381 (2015).
- Melé, M. et al. Human genomics. The human transcriptome across tissues and individuals, Science 348, 660-665 (2015).

- Matsubara, D. et al. Identification of CCDC6-RET fusion in the human lung adenocarcinoma cell line, LC-2/ad. J. Thorac. Oncol. 7, 1872-1876 (2012).
- Carneiro, B. A. et al. FGFR3-TACC3: a novel gene fusion in cervical cancer. Gynecol Oncol Rep 13, 53-56 (2015).
- Lee, M. et al. ChimerDB 3.0: an enhanced database for fusion genes from cancer transcriptome and literature data mining. Nucleic Acids Res. 45 (D1), D784-D789 (2017).
- Knezevich, S. R., McFadden, D. E., Tao, W., Lim, J. F. & Sorensen, P. H. A novel ETV6-NTRK3 gene fusion in congenital fibrosarcoma. Nat. Genet. 18, 184-187 (1998).
- Nacu, S. et al. Deep RNA sequencing analysis of readthrough gene fusions in human prostate adenocarcinoma and reference samples. BMC Med. Genomics 4, 11 (2011).
- Jia, Y., Xie, Z. & Li, H. Intergenically spliced chimeric RNAs in cancer. Trends Cancer 2, 475-484 (2016)
- Greger, L. et al. Tandem RNA chimeras contribute to transcriptome diversity in human 45. population and are associated with intronic genetic variants. PLoS ONE 9, e104567
- 46. Tomlins, S. A. et al. Recurrent fusion of TMPRSS2 and ETS transcription factor genes in prostate cancer, Science 310, 644-648 (2005).
- 47. Koboldt, D. C. et al. VarScan 2: somatic mutation and copy number alteration discovery in cancer by exome sequencing. Genome Res. 22, 568-576 (2012).
- 48. Lawrence, M. S. et al. Mutational heterogeneity in cancer and the search for new cancerassociated genes, Nature 499, 214-218 (2013).
- 49. Eilertsen, I. A. et al. Alternative splicing expands the prognostic impact of KRAS in microsatellite stable primary colorectal cancer. Int. J. Cancer 144, 841-847 (2019).
- Ji, X. et al. aCP binding to a cytosine-rich subset of polypyrimidine tracts drives a novel pathway of cassette exon splicing in the mammalian transcriptome. Nucleic Acids Res. 44, 2283-2297 (2016).
- Stransky, N., Cerami, E., Schalm, S., Kim, J. L. & Lengauer, C. The landscape of kinase fusions in cancer. Nat. Commun. 5, 4846 (2014).
- Cancer Genome Atlas Research Network. Comprehensive molecular profiling of lung adenocarcinoma. Nature 511, 543-550 (2014).
- Forbes, S. A. et al. COSMIC: somatic cancer genetics at high-resolution. Nucleic Acids Res. 45 (D1), D777-D783 (2017).
- 54. Blazek, D. et al. The Cyclin K/Cdk12 complex maintains genomic stability via regulation of expression of DNA damage response genes. Genes Dev. 25, 2158-2172
- Menghi, F. et al. The tandem duplicator phenotype is a prevalent genome-wide cancer configuration driven by distinct gene mutations. Cancer Cell 34, 197-210.e5 (2018).
- Dawson, M. A. & Kouzarides, T. Cancer epigenetics: from mechanism to therapy. Cell 150, 56 12-27 (2012).
- Zhang, X. et al. Identification of focally amplified lineage-specific super-enhancers in 57 human epithelial cancers, Nat. Genet, 48, 176-182 (2016).

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## Methods

#### RNA-seq alignment and quality-control analysis

Tumour and healthy ICGC RNA-seg data, included in the PCAWG cohort<sup>5</sup>, was aligned to the human reference genome (GRCh37.p13) using two read aligners: STAR<sup>58</sup> (v.2.4.0i, two-pass), performed at MSKCC and ETH Zürich, and TopHat2<sup>59</sup> (v.2.0.12), performed at the European Bioinformatics Institute. Both tools used Gencode (release 19)60 as the reference gene annotation. For the STAR two-pass alignment, an initial alignment run was performed on each sample to generate a list of splice junctions derived from the RNA-seg data. These junctions were then used to build an augmented index of the reference genome per sample. In a second pass, the augmented index was used for a more sensitive alignment. Alignment parameters have been fixed to the values reported in https://github.com/ICGC-TCGA-PanCancer/ pcawg3-rnaseq-align-star. The TopHat2 alignment strategies also followed the two-pass alignment principle, but was performed in a single alignment step with the respective parameter set. For the TopHat2 alignments, the irap analysis suite<sup>61</sup> was used. The full set of parameters is available along with the alignment code in https:// hub.docker.com/r/nunofonseca/irap\_pcawg/. For both aligners, the resulting files in BAM format were sorted by alignment position, indexed and are available for download in the GDC portal (https:// portal.gdc.cancer.gov/) and the ICGC Data Portal (https://dcc.icgc. org/). The individual accession numbers and download links can be found in the PCAWG data release table: http://pancancer.info/data releases/may2016/release may2016.v1.4.tsv. Cancer-type abbreviations are listed in Supplementary Table 23. Histology was derived from an older version released by the PCAWG Pathology and Clinical Correlates Working Group. Assignments of donor to histology used in this study can be found in the file rnaseq.extended.metadata. aliquot\_id.V4.tsv.gz at https://dcc.icgc.org/releases/PCAWG/ transcriptome/metadata/.

Quality control of all datasets was performed at three main levels: (1) assessment of initial raw data using FastQC<sup>62</sup> (v.0.11.3) (Supplementary Fig. 4); (2) assessment of aligned data (percentage of mapped and unmapped reads for both alignment approaches); and (3) quantification (by correlating the expression values produced by the STAR and TopHat2 based expression pipelines) (Supplementary Fig. 2). In total, we defined six quality-control criteria to assess the quality of the samples. We marked a sample as a candidate for exclusion if: (1) 3 out of 5 main FastOC measures (base-wise quality, k-mer overrepresentation, guanine-cytosine content, content of N bases and sequence quality) did not pass: (2) more than 50% of reads were unmapped or fewer than 1 million reads could be mapped in total using the STAR pipeline; (3) more than 50% of reads were unmapped or fewer than 1 million reads could be mapped in total using the TopHat2 pipeline; (4) we measured a degradation score<sup>63</sup> greater than 10; (5) the fragment count in the aligned sample (averaged over STAR and TopHat2) was <5 million; and (6) the correlation between the expression counts of both pipelines was < 0.95. If a sample did not pass one of these six criteria it was marked as problematic and placed on a greylist. If more than two criteria were not passed, we excluded the sample.

A subset of 722 libraries from the projects ESAD-UK, OV-AU, PACA-AU and STAD-US were identified as technical replicates generated from the same sample aliquot. These libraries were integrated postalignment for both the STAR and the TopHat2 pipelines using samtools<sup>64</sup> into combined alignment files. Further analysis was based on these files. Read counts of the individual libraries were integrated to a sample-level count by adding the read counts of the technical replicates.

Initially, a total of 2,217 RNA-seq libraries were fully processed by the pipeline. Quality-control filtering and integration of technical replicates (722 libraries) gave a final number of 1,359 fully processed RNA-seq sample aliquots from 1,188 donors.

#### GTEx data analysis

For a panel of RNA-seq data from a variety of healthy tissues, data from 3,274 samples from GTEx (phs000424.v4.p1) were used and analysed with the same pipeline as PCAWG data for quantifying gene expression. A list of GTEx identifiers are provided at https://dcc.icgc.org/releases/PCAWG/transcriptome/metadata.

# $\label{eq:Quantification} Quantification and normalization of transcript and gene expression$

STAR and TopHat2 alignments were used as input for HTSeq<sup>65</sup> (v.0.6.1p1) to produce gene expression counts. Gencode v.19<sup>60</sup> was used as the gene annotation reference. Quantification on a per-transcript level was performed with Kallisto<sup>66</sup> (v.0.42.1). This implementation is available as a Docker container at https://hub.docker.com/r/nunofonseca/irap\_pcawg. The implementation of the STAR and TopHat2 quantification is available as docker containers in: https://github.com/ICGC-TCGA-PanCancer/pcawg3-rnaseq-align-star and https://hub.docker.com/r/nunofonseca/irap\_pcawg/, respectively. Quantification of consensus expression was performed by taking the average expression based on STAR and TopHat2 alignments. Gene counts were normalized by adjusting the counts to FPKM<sup>67</sup> as well as FPKM with upper quartile normalization (FPKM-UQ) in which the total read counts in the FPKM definition has been replaced by the upper quartile of the read count distribution multiplied by the total number of protein-coding genes.

The FPKM and FPKM-UQ calculations were as follows. FPKM =  $(C \times 10^9)/(NL)$ , in which N denotes the total fragment count to protein-coding genes, L denotes the length of the gene and C denotes the fragment count. FPKM-UQ =  $(C \times 10^9)/(ULG)$ , in which U denotes the upper quartile of fragment counts to protein-coding genes on autosomes unequal to zero, and G denotes the number of protein-coding genes on autosomes.

#### t-Distributed stochastic neighbour embedding analysis

The *t*-distributed stochastic neighbour embedding (*t*-SNE) plots in Supplementary Figs. 5 and 6 were produced using the RTsne package<sup>68</sup> (with a perplexity value of 3) based on the Pearson correlation of the aggregated expression (log + 1) of the 1,500 most variable genes. FPKM expression values per gene were aggregated (median) by tissue (GTEx) and study (PCAWG). Coefficient of variation for each gene was also computed per tissue (GTEx) and study (PCAWG) to determine the 1,500 most variable genes. Purity values were previously described<sup>69</sup>.

The t-SNE plot in Extended Data Fig. 17c is based on all exon-skipping events in protein-coding genes confirmed by SplAdder 70. Each event was quantified in both the PCAWG and GTEx cohort. All events with more than 1% of missing percentage spliced in (PSI) values across the concatenated PCAWG and GTEx samples were removed. The remaining missing values were imputed as the mean over the non-missing samples. The centred data were then visualized using the TSNE package from the Scikit Learn toolkit 71 with a perplexity value of 100, random state 0 and an initialization with PCA.

## Associations between genetic variation and gene expression: patient cohort

To associate genetic variation with gene expression, we analysed whole-genome sequencing (WGS) of the 1,188 donors with matched whitelisted RNA-seq data from the PCAWG cohort. Germline genotypes, SNV calls and segmented allele-specific SCNA calls were previously reported<sup>5</sup>. We matched 1,188 tumour RNA-seq IDs<sup>5</sup> to WGS whitelist tumour IDs (synapse entry syn10389164). For patients with multiple WGS IDs (2 out of 1,188) or RNA-seq aliquot IDs (17 out of 1,188), we resolved the matching by pairing samples with the same 'tumor\_wgs\_submitter\_specimen\_id' (Supplementary Table 1). The 1,188 patients are spread across 27 types of cancer and 29 project codes and include

899 carcinomas; 34 patients are metastatic and 13 recurrent with the remaining patients being primary tumours (Supplementary Table 1).

We used the data of these 1,188 patients for performing somatic and germline eQTL mapping, ASE analysis and association studies between gene expression and mutational signatures.

### Gene expression filtering

Gene expression values (measured in FPKM; https://dcc.icgc.org/releases/PCAWG/transcriptome/gene\_expression) from consensus expression quantification as described above were used for this analysis.

Genes with FPKM  $\geq$  0.1 in at least 1% of the patients (12 patients) were retained, resulting in 47,730 genes. Only 18,898 protein-coding genes (according to the 'gene type' biotype reported in Gencode v.19<sup>60</sup>) were used for the subsequent OTL analyses. The log<sub>2</sub>-transformed expression values (FPKM + 1) were subjected to peer analysis 72 to account for hidden covariates (syn7850427; https://dcc.icgc.org/releases/PCAWG/ transcriptome/eQTL/phenotype). To balance the number of covariates, statistical power and available sample sizes per cancer type, we followed the GTEx protocol and estimated 15, 30 and 35 hidden covariates to be used depending on sample size<sup>73</sup> ( $n < 150, 150 \le n < 250, n \ge 250$ ). Peer residuals were then rank-standardized across patients. The FPKM cut-off values and peer correction were also applied to the subset of 899 patients with carcinoma, yielding 18,837 protein-coding genes after filtering. Furthermore, we used ordinary least-squares regression to correlate each of the 35 peer factors with per-sample covariates, including cancer project codes, gender, tumour purity, somatic burden and several sequence metrics (Supplementary Notes), to understand the proportion of variance explained by known biological and technical covariates.

#### **Covariates**

In all linear models, we accounted for known confounding factors by modelling them as fixed effects. In all association studies, we accounted for sex, project code (describing cancer type and country of origin) and per-gene copy-number status (Supplementary Table 1 for the list of per patient covariates; syn7253568 and syn7253569 for sex and project codes; syn9661460 for per gene copy number). Per-gene copy-number alterations were derived as the average copy number across all copy-number aberrations called within the annotated gene boundaries based on syn8042988.

The somatic eQTL, ASE and mutational signature analyses also accounted for total somatic mutation burden (number of SNVs and short insertions and deletions (indels)) and sample purity (Supplementary Table 1). Purity was estimated based on copy-number segmentation. In addition, the somatic eQTL and ASE analyses accounted for local SNV burden calculated in a 1-Mb window from the gene coordinates (https://dcc.icgc.org/api/v1/download?fn=/PCAWG/transcriptome/eQTL/covariates/pergene.somatic.snv.cis.burden.1188.wl.donors.tsv.gz).

The germline eQTL analysis also modelled the population structure as random effect. The population structure was assessed by a kinship matrix that was calculated based on every twentieth germline variant, processed as described below (see 'Germline eQTL variants'). The kinship matrix was then calculated as an empirical patient-by-patient covariance matrix.

Different covariates were accounted for per-analysis method (Supplementary Table 1). The project code describes cancer type and country-of-origin. Somatic burden is the total number of SNVs and indels. Purity was estimated based on copy-number segmentation. Local somatic burden is the number of SNVs in a 1-Mb window around the gene coordinates. Local copy number was defined as the average copy-number state across all SCNAs called within the annotated gene boundaries.

## GO and Reactome pathway enrichment

We performed GO<sup>74,75</sup> and Reactome pathway<sup>20,21</sup> enrichment with the Bioconductor packages biomaRt<sup>76,77</sup>, clusterProfiler<sup>78</sup> and ReactomePA<sup>79</sup>

(FDR  $\leq$  10%). The number of genes used as background set is described per analysis method.

#### Germline eOTL variants

PCAWG variant calls v.0.1<sup>5</sup> were downloaded from GNOS and processed following the PCAWG-8 protocol: (1) VCF files were indexed and merged using bcftools<sup>80</sup>. (2) All variants were filtered for 'PASS' flag. (3) All variants were filtered for quality larger than 20. (4) Only bi-allelic sites were considered.

HDF5 files for each 100-kb chunk of the VCF files were generated, assuming additivity that was numerically encoded as 0, 1 or 2 for homozygous reference, heterozygous or homozygous alternative state, respectively. For indels, we encoded the presence or absence of the variant as 0 or 1, respectively. Each variant was normalized to mean 0 and standard deviation 1. Missing variants were mean-imputed. To create our eQTL release set v.1.0, the resulting HDF5 files were subsequently merged into a global HDF5 file and all variants which follow any of the following conditions were removed: (1) minor allele frequency  $\leq 1\%$ ; and (2) missing values  $\geq 5\%$ 

#### Germline eQTL analysis

In the germline eQTL analyses, we used the processed gene expression dataset from 1,178 patients for which germline variant calls (eQTL release set v.1.0, see 'Germline eQTL variants') were available. Linear mixed models were used to model the correlation between germline variants (within 100 kb of gene boundaries) and gene expression values (see 'Gene expression filtering') using the limix package<sup>81</sup>. Known covariates were modelled as fixed effects and population structure as random effect (see 'Covariates').

A two-step approach was used to adjust for multiple testing. First, for each gene, we adjusted for the number of independent tests estimated based on local linkage disequilibrium  $^{82}$ . Second, we performed a global correction across the lead variants, that is, the most significant SNPs, per eQTL. Germline eGenes were defined as genes with an eQTL with global FDR  $\leq 5\%$ .

#### **GTEx comparative analysis**

The GTEx comparative eQTL analysis was based on the eQTL maps v.6p<sup>10</sup>. We mapped the positions and alleles of our PCAWG-specific eQTL to the eQTL in all GTEx tissues. To determine whether a lead eOTL variant is replicated in a given GTEx tissue, we followed the previously described strategy<sup>10</sup>. For each eGene, we considered the eQTL lead variant and assessed the replicability of the signal in the GTEx cohort based on marginal association statistics using 42 GTEx tissues without cell lines (P < 0.00024 = 0.01/42, corrected for the number of GTEx tissues—that is, 42)). If the lead variant did not replicate or was not tested, we determined replication based on the variant with the smallest P value within the linkage disequilibrium block ( $r^2 \ge 0.8$ estimated based on UK10K project) of the lead variant across 25 (or 42) tissue-matched GTEx analyses. If neither lead nor any variant within the linkage disequilibrium block was tested, we determined replication based on the smallest P value of any variant within the 100-kb window tested within the GTEx cohort. We also derived less stringent sets of PCAWG-specific eGenes by allowing replication in up to 1, 5 or 10 GTEx tissues.

## Tissue sharing of germline eGenes between histotypes

Using the R package qvalue (https://github.com/StoreyLab/qvalue, v.2.14.0), we generated  $\pi_1$  statistics comparing the lead variants of one histotype against their P value distribution in the other histotypes. Because  $\pi_1$  statistics are known to be confounded by sample size and number of eQTL found, we subsampled the eQTL lead variants to a randomly selected set of 100 variants. After 20 rounds of subsampling, we derived the same  $\pi_1$  statistics as mentioned earlier and reported the average.

#### Roadmap enrichment of germline eGenes

For each lead variant, we generated a matching background set of 1,000 variants using SNPsnap<sup>83</sup>. Each variant (background and foreground) was intersected with the location of 25 Roadmap factors<sup>16</sup> in 127 cell types. From this we derived fold change and P values. Significant changes of fold change between PCAWG-specific and unspecific eQTLs is based on a one-sided Wilcoxon rank-sum test.

#### **Enrichment analysis**

Enrichment of Reactome pathways of PCAWG-specific eGenes was performed using the Bioconductor package ReactomePA<sup>79</sup>.

#### Somatic calls and mutational burden

We used the set of consensus SNVs somatic calls provided by PCAWG (syn7357330) based on three core caller pipelines and MuSE<sup>84</sup>. On average, we counted 22,144 somatic SNVs per patient, with different median numbers of SNVs per cancer type, ranging from 1,139 in thyroid adenocarcinoma to 72,804 SNVs in skin melanoma (Extended Data Fig. 5a). Owing to the low frequency of somatic SNVs across the cohort (Extended Data Fig. 5b), we collapsed the variants by genomic regions defined by gene annotations (Gencode v.19<sup>60</sup>). Specifically, we generated a set of disjoint gene exons by collapsing overlapping exon annotations into single features using bedtools<sup>85</sup>. The set of disjoint introns was generated using bedtools by subtracting the collapsed exonic regions from the gene regions. To map local effects of somatic mutations in flanking features outside the gene body, we binned the surrounding regions (plus and minus 1 Mb from the gene boundaries) into 2-kb windows (flanking) overlapping by 1 kb.

We defined three different types of aggregated somatic burden to assess differences in power in detecting somatic eGenes and P value calibration. The burden in a genomic region was defined as (1) a binary value that indicates presence or absence of SNVs; (2) the aggregated burden as sum of SNVs; or as (3) weighted burden, that is, sum of variant allele frequencies of the SNVs (Supplementary Fig. 10a) to take into account their clonality (https://dcc.icgc.org/releases/PCAWG/transcriptome/eQTL/genotypes). We assessed calibration of all three analyses with Q-Q plots of nominal and permuted P values (permutation of the patients in the gene expression matrix) (Supplementary Fig. 10b-d). Moreover, for the linear regression analysis, genotypes were standardized across patients (to mean zero and standard deviation one) and standardized effect sizes are provided in Supplementary Table 5.

Overall, somatic burden within flanking regions was the most prevalent type of burden tested per gene (Extended Data Fig. 6a). We found similar average relative mutation density per type of genomic region (flanking = 0.008 mutations per kb; introns = 0.007 mutations per kb; exons = 0.006 mutations per kb) (Extended Data Fig. 6b) and average recurrence of the same mutated region across the cohort was rather low (flanking = 1.4%; exons = 1.7%; introns = 4%) (Extended Data Fig. 6c).

## Somatic eQTL analysis

Linear models were used to model the correlation between recurrent somatic burden and gene expression of up to 18,898 protein-coding genes, using the limix package<sup>81</sup> (see 'Gene expression filtering'). Gene expression was corrected for 35 hidden Peer factors. Known covariates were modelled as fixed effects (see 'Covariates'). We considered only somatic burdens with frequency greater than 1%, including exonic and intronic burdens, as well as flanking burdens, within 1 Mb from gene boundaries.

The somatic eQTL analysis was performed on all 1,188 patients and on the subset of 899 patients with carcinoma (representing 20 of the 27 types of cancer) to replicate the analysis on a more homogeneous set of tumours. A cis window of 1 Mb from the gene boundaries was used to find mutated genomic intervals with a burden frequency  $\geq 1\%$  in the cohort (at least 12 patients in the full cohort and 9 patients in

the carcinoma cohort). Together, 18,708 of the genes had at least one mutated interval at that frequency and were included in the analysis and 1,049,102 regions showed a burden frequency  $\geq$  1%

Bonferroni correction was applied to correct for multiple *cis* windows tested within the same gene. Then, Benjamini–Hochberg correction was applied to adjust the P values of the lead genomic regions across genes. Somatic eGenes were defined as genes with an eQTL at a FDR  $\leq$  5%.

#### Somatic cis-eQTL comparative analysis

We compared our 649 somatic eQTL set with three previous cancer studies<sup>86–88</sup> to identify independent evidence of interaction between our eGenes and the associated cis-genomic regions with somatic burden. Studies were chosen if they provided lists of cancer regulatory elements linked to genes or regulatory elements with somatic mutations linked to gene expression deregulation in cancer. All the three studies examined were based on TCGA cancers. For this, we checked perfect overlaps with both the somatic burden location and the eGene. Moreover, we looked at the overlap between somatic eQTL and 72,987 GeneHancer89 enhancers-to-genes interactions, with at least two independent supporting methods (called 'double-elite'), downloaded from the UCSC hg19 GeneHancer track90. We then compared this overlap with a set of nulls generated by 1,000 random permutations of the GeneHancer regulatory elements with nearby genes located within 1Mb. We then retrieved an empirical P value of enrichment by counting the number of random nulls (N) showing greater number of overlaps than those found between the somatic eQTL set and the GeneHancer set (P = (N+1)/(1,000+1)).

#### Functional enrichment in somatic cis-eQTL

To identify putative regulatory sites enriched for somatic eQTL, we retrieved functional annotations of the lead genomic flanking intervals of the somatic eQTL (556 intervals linked to 638 somatic eQTL). Therefore, we mapped somatic eQTL to 25 Roadmap Epigenomics chromatin marks of 127 different cell types<sup>16</sup> and ENCODE transcriptionfactor binding site annotations in 9 cell types (including 8 cancer and one embryonic stem-cell lines<sup>91</sup>) (Supplementary Tables 6 and 7). We compared annotations in the significant set of eQTLs with a null distribution based on 1,000 random samplings of a matched set of genomic intervals. To define the matched sets of genomic intervals, we selected flanking genomic intervals from the whole set of tested genes that showed a similar distance from the gene start (exact distance  $\pm 2 \text{ kb}$ ) and that matched the exact burden frequency of the corresponding interval in the significant associations. We then overlapped the 1,000 matched sets with Roadmap Epigenomics and ENCODE annotations. To avoid ambiguous overlaps (with multiple annotations), we retained only genomic intervals showing a minimum overlap of 10% of their length.

We retrieved an empirical P value of enrichment for each annotation by counting the number of randomly sampled flanking intervals (N) showing greater number of overlaps compared to the eQTL set (P=(N+1)/(1,000+1)). Benjamini–Hochberg correction was applied to the empirical P values (over 25 marks in 127 cell lines for Roadmap Epigenomics annotations and over 149 transcription-factor-binding sites for 9 ENCODE cell lines). We then computed the fold change per annotation and cell line as a ratio of annotated lead flanking intervals and mean number of annotated matched random flanking intervals over the 1,000 samplings.

Furthermore, we performed GO<sup>74,75</sup> and Reactome pathway<sup>20,21</sup> enrichment with the Bioconductor packages biomaRt<sup>76,77</sup>, clusterProfiler<sup>78</sup> and ReactomePA<sup>79</sup> (FDR  $\leq$  10%) and also looked at enrichment within high-confidence cancer testis genes previously described<sup>92</sup>, using 18,708 genes with at least one mutated interval as background.

## Variance component analysis

Limix was used to perform variance decomposition using the same covariates as in the somatic variant analyses except for local

copy-number state (see 'Covariates'). The random effects were based on the following common germline variants and somatic burden (frequency > 1%) (see 'Somatic calls and mutational burden' for detailed description of burden): (1) *cis*-somatic intronic: weighted burden in introns; (2) *cis*-somatic exonic: weighted burden in exons; (3) *cis*-somatic flanking: weighted burden in 1-kb-overlapping regions of 2 kb within 1 Mb from gene boundaries; (4) somatic intergenic: weighted burden in 1-kb-overlapping regions of 2 kb outside the 1 Mb window; (5) *cis*-germline: germline variants within 100 kb from gene boundaries; (6) *trans*-germline: genome-wide population structure (see 'Covariates'); and (7) local copy-number variation (see 'Covariates').

All the data was mean-centred and standardized. For each of the random effects, a linear kernel was computed and used as covariance matrix. The resulting variance components were normalized to add up to one.

#### Mutational signature associations

We obtained 39 mutational signatures from PCAWG-7 beta 2 release and used linear models to associate the mutational signatures with gene expression of up to 18,898 protein-coding genes across 1,159 patients while accounting for known covariates (see 'Covariates') (quality control) (Extended Data Fig. 10a–e). The 1,159 patients were a subset of the total 1,188 patients, for whom mutational signature profiles were available. Gene expression was corrected for 35 hidden peer factors (see 'Gene expression filtering').

We retained 18,888 genes that showed a minimum FPKM of 0.1 in at least 1% of 1,159 the patients (see 'Gene expression filtering'). Signatures with zero variance and a prevalence below 1% were filtered, and we obtained 28 signatures. We applied linear models to associate expression of these genes with the signatures across all 1,159 patients, a subset of 877 patients with carcinoma or a subset of 891 European patients to assess consistency of the associations (Extended Data Fig. 10f, g).

Across all patients, we found 1,176 significantly associated genes after Benjamini–Hochberg correction (we used an FDR  $\leq$  10% for enrichment analyses, multiple testing was applied across all signature–gene pairs) (Supplementary Tables 19a–c). We performed gene enrichment analyses of the significant genes per signature (see 'GO and Reactome pathway enrichment') (here 18,831 background genes, multiple testing correction across all ontologies per signature FDR  $\leq$  10%) (Supplementary Table 19d). Whereas most signatures were associated with only few genes, 18 showed recurrent *trans* effects and affected expression of over 20 genes (Extended Data Fig. 11d, Supplementary Table 19e). We further found that the vast majority of genes (85.8%) were associated with only one signature (1,009 genes); 129 genes were associated with two, 32 with three, 5 with four and 1 with five signatures.

To assess how tissue-specific both mutational signatures and their associations with gene expression are, we analysed the occurrence of each signature in each of the types of cancer. We assessed the presence (at least one SNV of a signature in at least one patient with a specific cancer type) and mean prevalence (mean number of SNVs of a certain signature across all patients of a specific cancer type) of the signatures in the types of cancer (Extended Data Fig. 13c, d). We defined cancer-typespecific signatures to occur in up to four types of cancer (signatures 4, 7, 9, 12, 16, 38 and 39) and common signatures to be missing in up to five types of cancer (signatures 2, 13 and 18). For each of these signatures, we performed cancer-type-specific analyses, that is, we assessed the association between the respective signature and gene expression in just the patients who are of a cancer type that shows mutations of the respective signature (Extended Data Fig. 13c, left heat map). We then correlated the P values of these cancer-type-specific analyses with the P values of the analysis across all patients and calculated the Pearson correlation coefficients (Supplementary Fig. 24a-e). We show that the correlation between cancer-type-specific and whole-cohort *P* values is dependent on the sample size of the respective analysis ( $r^2 = 0.671$ ) (Supplementary Fig. 1f).

We further performed PCA on the signatures across both, patients (PCA on signature-specific SNVs per patient) and genes (PCA on adjusted *P* values of signature-gene expression associations) (Extended Data Fig. 11a, b).

To assess significance of the functional annotation of SNVs by mutational signatures, we also associated gene expression with the total number of SNVs and correlated the P values ( $-\log_{10}(P)$ ) of the associations with the respective signature-specific P values. The absolute Pearson correlation coefficients remain below 0.1 (Supplementary Table 19f).

To establish causality of signature–gene expression associations, we included the germline eQTL into the analysis using linear mixed models; 197 of our 1,176 signature-associated genes were also germline eGenes. These 197 associations involved 26 of the 28 mutational signatures. We associated the lead variants of these eGenes with the rank-standardized signature SNVs across 2,507 patients. We used the subset of the 2,818 WGS patients for which mutational signature profiles and all known covariates were available. We accounted for the same fixed covariates as in the mutational signature–gene expression association studies and, in addition, for kinship as a random effect (see 'Covariates').

We then performed proportional colocalization analysis with Bayesian model averaging using the R package  $\operatorname{coloc}^{93}$  to test whether gene expression and mutational signatures share common causal genetic variants in a given gene region. A proportional colocalization analysis tests the null hypothesis of colocalization by assuming that two phenotypes that share causal variants will have proportional regression coefficients for either phenotype with any variant selection in the vicinity of the causal variant. We applied the Bayesian model averaging approach, with each tested model consisting of a selection of two variants. The P values are then averaged over all models to generate posterior predictive P values  $^{93}$ . We filtered variants so that no pair of variants showed  $r^2 > 0.95$  and each variant's marginal posterior probability of inclusion with one of the phenotypes was greater than 0.01. The nominal P values of rejecting the null hypothesis of colocalization are listed in Supplementary Table 19e.

We then performed mediation analysis <sup>94,95</sup> to assess directionality of the effect between germline eQTL, gene expression and mutational signature. First, causal mediation analysis was applied to each of the triples of eQTL lead variant, gene and mutational signature using a structural equation model from the R package lavaan <sup>96</sup>. Then, we used the R package mediation <sup>97</sup> to assess significance of mediation and estimate the proportion of mediated effect by non-parametric bootstrap confidence intervals (1,000 simulations).

## ASE analysis: assembling phased germline and somatic variants

To understand the precise effect of somatic variations in their genomic context and for subsequent allele-specific analyses, both germline and somatic variants were phased. For assembling phased germline genotypes, we used the Sanger 1000G callset<sup>6</sup>, and applied IMPUTE2<sup>98</sup> for phasing of heterozygous germline variants. The IMPUTE2 output was corrected using results from the Battenberg CN calling algorithm<sup>99</sup> to ascertain that no haplotype switches occur within regions of consecutive copy-number gain. The resulting phased germline genotypes were arranged such that haplotype 1 always corresponded to the amplified alleles in regions with SCNAs (major allele). In cases in which both cooccur on the same NGS read (approximately 10 million variants, 20% of all SNVs), we phased individual somatic variants to the nearest germline heterozygous site. For downstream analyses, we considered only SNVs that were phased by at least three reads to the respective germline variant (approximately 6 million out of 10 million SNVs).

All phased SNVs were aggregated into functional categories based on their genomic regions defined by gene annotations (upstream, downstream, promoter, 5' UTR, intron, synonymous, missense, stop gain and 3' UTR) and mapped to the nearest gene within a *cis* window of 100 kb using the Variant Effect Predictor (VEP) tool<sup>100</sup>. Promoter

variants were defined as 1-kb upstream of the TSS. We included flanking regions by using the VEP 'UpDownDistance' plugin with a maximum range parameter of 100 kb. We divided the upstream and downstream variant categories into disjoint categories using 10-kb windows from 10 to 100 kb. We integrated 'splice donor' and 'splice acceptor' variants into the general 'splice region' variant category and mapped 'stop retained' variants to the 'synonymous' variant category. We averaged transcript-level annotations to gene-level annotations to retrieve the expected functional effect of a variant for a given gene. We analysed the relationship between SNV variant allele frequency and SCNAs at the same locus to determine whether variants occurred before ('early') or after ('late') the corresponding SCNA (PCAWG-11). We computed a weighted *cis*-mutational burden per category by estimating the cancer cell fraction of each SNV and aggregating SNVs to a total localized burden weighted by their respective cancer cell fraction.

#### **ASE read counts**

The positional information of the heterozygous germline variants was used together with the RNA-seq BAM files as input to the GATK ASEReadCounter<sup>101</sup> algorithms for counting ASE reads. We considered reads with a minimum mapping quality of 20 and a minimum base quality of 10. Only heterozygous variants with a minimum coverage of eight RNA-seq reads were considered for all further analyses.

The raw ASE read counts were post-processed as follows: (1) ASE sites were converted to BED files and aligned against the ENCODE 50-mer mappability track (wgEncodeCrgMapabilityAlign50mer.bigWig) to extract mappability scores for all sites. All sites with mappability scores unequal to 1 were removed. (2) All sites with allelic read counts less or equal to 1 were removed to prevent genotyping error to influence ASE quantification. (3) All sex chromosomes were dropped for further analysis. (4) We estimated sequencing error per patient as the sum of non-reference and non-alternative bases over the total number of bases. We assessed statistical mono-allelicity through a binomial test using the estimated sequencing error probabilities, corrected using the Benjamini-Hochberg step down procedure. All sites that appeared to be statistically mono-allelic were removed. (5) For each ASE site, copynumber states were retrieved from the Sanger copy-number consensus callset (PCAWG-11). Purity estimates for each patients were retrieved from the accompanying purity tables.

To aggregate site-level ASE to a gene-level readout and to allow for estimation of effect directionality, we used the phased germline genotypes. Gene mapping was performed against ENSEMBL release 75 using the pyEnsembl Python library. We retrieved all genes at each ASE site and summed up the read counts on the respective haplotypes to genelevel haplotype-specific read counts. We further averaged haplotypespecific copy-number states to a mean haplotype-specific copy-number state per gene and computed the gene-level copy-number ratio as the major over total ratio of those averages. To allow for a robust assessment of gene-level ASE, we considered only genes with at least 15 reads total, yielding 4,379,378 gene-patient pairs of 1,120 patients and 17,009 unique genes across 12,441,502 accessible sites in total. Every remaining gene was tested for AEI using a binomial test against an expected read ratio of 0.5 to derive nominal P values, and a binomial test against the expected copy-number ratio modified by tumour purity to derive copy-number-corrected Pvalues. Nominal and copy-number-corrected P values were adjusted separately for multiple testing using the Benjamini-Hochberg procedure. Significant AEI was called at FDR ≤ 5%. We further annotated each gene with the number of ASE sites used for aggregation. For all downstream analyses, we considered only genes annotated as protein coding (ENSEMBL biotype = 'protein\_coding').

#### **Generalized linear models**

Across all 4,379,378 gene–patient pairs, we trained multivariate linear models using (i) logistic regression against a binary indicator of AEI absence or presence in a gene, or (ii) standard linear regression against

the phased ASE ratio of a gene to assess the directionality of the regulatory change. For (i), haplotype-specific mutations were summed up to a total burden per category, whereas for (ii) we used the difference in burden between the haplotypes 1 and 2. The consistency of the phasing map between somatic variants and ASE sites ensured that model coefficients kept their directionality independent of the arbitrary labelling of haplotypes as 1 or 2. The full set of considered factors is as follows: (1) copy-number ratio at the gene locus  $(0.5 \le x \le 1)$ ; (2) sample purity (0 < x < 1); (3) natural logarithm of total gene length (x > 0); (4) natural logarithm of the length of the canonical transcript (x > 0); (5) heterozygosity of the lead eQTL variant (x = 0) if homozygous, x = 1 if not homozygous); (6) all mutational burden categories as determined by VEP annotations (upstream in 10-kb windows, downstream in 10-kb windows, promoter, 5' UTR, intron, synonymous, missense, stop gain and 3' UTR;  $x \ge 0$  for logistic model,  $x \in \mathbb{R}$  for directed model).

To compare global effects and different contributions of SCNA, germline eQTL, coding and non-coding SNVs, a simplified logistic model was trained after accumulating all coding and non-coding variants to separate categories and reporting standardised effect sizes (Fig. 1e).

#### Cancer gene enrichment

Cancer gene enrichment was conducted on the COSMIC census<sup>53</sup> using Fisher's exact test and gene set enrichment analysis as previously described<sup>102</sup>. For enrichment, the average score of a gene was computed across the cohort and only genes with at least five replicates in the cohort were kept, yielding a total of 16,078 genes.

#### Chromosomal distribution of ASE

We calculated the recurrence of ASE genes in each tumour type. To examine the chromosomal distribution of ASE genes, we calculated the average recurrence of all genes for every 200-gene window with a 10-gene step, and then subtracted the average ASE occurrence in each tumour type to obtain the peaks of ASE surplus across all chromosomes. The recurrence of copy-number genes was calculated in an analogous manner.

#### Estimation of alternative promoter activity

We estimated promoter activities using RNA-seq data and Gencode (release 19) annotations for 70,937 promoters in 20,738 genes. We grouped transcripts with overlapping first exons under the assumption that they are regulated by the same promoter<sup>103</sup>. TSSs that are located within internal exons, or which overlap with splice acceptor sites, were removed from this analysis as these promoters are difficult to estimate from RNA-seq data<sup>28</sup>. Promoter activity can be estimated using exon usage<sup>29</sup>, spliced reads<sup>28</sup> or isoform-based estimates<sup>30</sup>. Here we used an isoform-based approach to quantify promoter activity. We quantified the expression of each transcript from the RNA-seq data using Kallisto<sup>66</sup> and calculated the sum of expression of the transcripts initiated at each promoter to obtain an estimate of promoter activity. To obtain the relative activity for each promoter, we normalized each promoter's activity by the overall gene's expression. We divided the promoters of each gene into three categories based on their average pan-cancer promoter activity. The promoters with <1 FPKM average activity are called inactive promoters, and the most active promoter of each gene is called the major promoter. The remaining active promoters of the gene are called minor promoters.

The association between promoter activities and promoter mutation burden was estimated using the same framework as the somatic eQTL analysis. We examined associations for the promoters of expressed multi-promoter genes with a burden frequency  $\geq 1\%$  in the cohort (at least 12 patients in the full cohort). The weighted burden of the region 1-kb upstream of the TSS—that is, the sum of variant allele frequencies of the SNVs for each gene—was used as the genotype for the promoters of the respective genes. We used linear models to study the associations between the recurrent somatic burden and the promoter activity (both

for the relative activity and the  $\log_2$ -transformed absolute activity). Similar to the somatic eQTL analysis, the known covariates and the 35 hidden peer factors were provided as cofactors to the linear models. We adjusted the *P* values using Benjamini–Hochberg correction method and looked for associations with FDR  $\leq 5\%$ .

#### Identification of alternative splicing

We used the alignments based on the STAR pipeline to collect and quantify alternative splicing events with SplAdder<sup>70</sup>. The software has been run with its default parameters with confidence level 3. We generated individual splicing graphs for each RNA-seq sample for both tumour samples as well as matched healthy samples (when available). All graphs were then integrated into a merged graph to comprehensively reflect all splice junctions observed in all samples together. On the basis of this combined graph, SplAdder was used to extract alternative splicing events of the following types: alternative 3' splice site, alternative 5' splice site, cassette exon, intron retention, mutually exclusive exons, coordinated exon skip (see supplementary figure 3 in ref. 70). Each identified event was then quantified in all samples by counting split alignments for each splice junction in any previously identified event and the average read coverage of each exonic segment involved in the event was determined. We then computed a PSI value for each event that was then used for further analysis. We further generated different subsets of events, filtered at different levels of confidence, in which confidence is defined by the SplAdder confidence level (generally 2), the number of aligned reads supporting each event, the number of samples that were found to support the event by SplAdder, and the number of samples that passed the minimum aligned read threshold.

# Enrichment of outlier splicing associated with splice sites and branchpoint motifs

We assessed the significance of mutational enrichment for 5' and 3' splice sites, and branch-point  $^{104,105}$  intronic regions using a permutation-based approach. Impactful mutations were defined as mutations overlapping exons and introns involved in cassette exon events, in which the PSI-derived z-score was  $\geq 3$  or  $\leq -3$ . For each intronic site, we compared the frequency of observed impactful mutations against frequencies of randomly sampled intronic regions (number of iterations = 1,000). For exonic sites, the null distribution was established from randomly sampled exonic sites. Randomly sampled sites were within a 100-bp window around the 5' and 3' splice site. For branch-point regions, sampled sites were within a 50-bp window around the branch-point sequence. The P value was computed as the number of randomly sampled frequencies greater or equal to the observed frequency.

#### SAVNet analysis for identifying rare SAVs

The SAVNet approach  $^{35}$  was designed for identifying somatic variants associated with local aberrant splicing alterations from matched genome and transcriptome sequencing data. It uses permutations to calculate an FDR and by restricting to two classes of relationships between somatic mutations and splicing alterations to focus: (1) splice site disruption, in which exon skipping, alternative 5' or 3' splice site, or intron retention is associated with a mutation in a splice site motif; and (2) splice site creation, in which alternative 5' or 3' splice sites are associated with mutations that create a novel splice motif (FDR  $\leq$  10%) (Extended Data Fig. 17e).

#### Identification of RNA fusions

Gene fusions between any two genes were identified based on two gene fusions detection pipelines: FusionMap (v.2015-03-31) pipeline<sup>106</sup> and FusionCatcher (v.0.99.6a)/STAR-Fusion (v.0.8.0) pipeline<sup>107</sup>. ChimerDB 3.0 was used as a reference of previously reported gene fusions. The database contains 32,949 fusion genes split into three groups: (1) KB: 1,067 fusion genes manually curated based on public resources of fusion genes with experimental evidences; (2) Pub: 2,770 fusion genes

obtained from text mining of PubMed abstracts; and (3) Seq: archive with 30,001 fusion gene candidates from deep-sequencing data. This set includes fusions found by re-analysing the RNA-seq data of the TCGA project encompassing 4,569 patients from 23 types of cancer.

In brief, FusionMap was applied to all unaligned reads from the PCAWG aligned TopHat2 RNA-seq BAM files for each aliquot to detect gene fusions. In the FusionCatcher/STAR-Fusion pipeline, for each aliquot with paired-end RNA-seq reads FusionCatcher was applied to the raw reads, with the genome reference. Specifically, for each aliquot with paired-end RNA-seq reads FusionCatcher was applied to the raw reads. The '-U True; -V True' runtime options were used. For each aliquot with single-end RNA-seq reads, STAR-Fusion was applied to the raw reads, with the same reference genome and gene models as FusionCatcher and with default settings. In parallel, FusionMap was applied to all unaligned reads from the PCAWG aligned TopHat2 RNA-seq BAM files for each aliquot to detect gene fusions with the following non-default options values: MinimalHit = 4; OutputFusionReads = True; RnaMode = True; FileFormat = BAM.

To reduce the number of false-positive fusions, the two sets of fusions were filtered to exclude fusions based on the number of supporting junction reads, sequence homology, and occurrence in normal samples (from the GTEx and the PCAWG cohort). To get a high-confident consensus fusion call set from these two pipelines, a fusion to be included in the final set of fusions had to: (i) be detected by both fusion detection tools in at least one sample; and/or (ii) be detected by one of the methods and have a matched structural variant in at least one sample. The consensus WGS-based somatic structural variants (v.1.6) were obtained from the PCAWG repository in https://dcc.icgc.org/releases/PCAWG.

For integration with matched structural variant evidence, a fusion was considered to match a structural variant if the absolute distance between the fusion break points and structural variant break points did not exceed 500 kb (the distance was considered infinite when the chromosomes of the fusion and structural variant break point differ). When there was no evidence for a direct structural variant fusion, the search was expanded to look for composite fusions. In this case, an exhaustive search was performed to look for two structural variants with break points close to the fusion break points and with an effective distance smaller than 250 kb.

Finally, 3,540 fusion events were included as the consensus fusion call set, from these 2,268 were detected by both FusionCatcher/STAR-Fusion and FusionMap (from these, 1,821 had matched structural variant evidence) and 1,112 were detected by only one method and had matched structural variance evidence.

In total, approximately 36% of all detected fusion transcripts were predicted to be in-frame, several UTR-mediated fusion transcripts preserve complete coding sequences of one fusion partner. These include a known fusion *TBL1XR1-PIK3CA* in a breast tumour and a notable new example *CTBP2-CTNNB1* in a gastric tumour.

All fusions are available in Synapse: https://dcc.icgc.org/releases/PCAWG/transcriptome/fusion.

#### Identification of RNA-editing events

We used an RNA-editing events calling pipeline, which is an improved version of that previously published <sup>108</sup>. First, we summarized the base calls of pre-processed aligned RNA reads to the human reference in pileup format. Second, the initially identified editing sites were then filtered by the following quality-aware steps: (1) the depth of candidate editing site, base quality, mapping quality and the frequency of variation were taken into account to do a basic filter: the candidate variant sites should be with base-quality  $\geq$  20, mapping quality  $\geq$  50, mapped reads  $\geq$  4, variant-supporting reads  $\geq$  3, and mismatch frequencies (variant-supporting-reads/mapped-reads)  $\geq$  0.1. (2) Statistical tests based on the binomial distribution B(n,p) were used to distinguish true variants from sequencing errors on every mismatch site<sup>109</sup>, in which p denotes the background mismatch rate of each transcriptome

sequencing, and n denotes sequencing depth on this site. (3) Discard the sites present in combined DNA SNP datasets (dbSNP v.138, 1000) Genome SNP phase 3, human Dutch populations<sup>110</sup>, and BGI in-house data; combined datasets deposited at: ftp://ftp.genomics.org.cn/ pub/icgc-pcawg3). (4) Estimate strand bias and filter out variants with strand bias based on two-tailed Fisher's exact test. (5) Estimate and filter out variants with position bias, such as sites only found at the 3' end or at 5' end of a read. (6) Discard the variation site in simple repeat region or homopolymer region or <5 bp from splicing site. (7) To reduce false positives introduced by misalignment of reads to highly similar regions of the reference genome, we performed a realignment filtering. Specifically, we extracted variant-supporting reads on candidate variant sites and realign them against a combination reference (hg19 genome plus Ensembl transcript reference v.75) by bwa0.5.9-r16. We retain a candidate variant site if at least 90% of its variant-supporting reads are realigned to this site. Finally, all high confident RNA-editing sites were annotated by ANNOVAR<sup>111</sup>. (8) To remove the possibility of an RNA-editing variant being a somatic variant, the variant sites are positionally filtered against PCAWG WGS somatic variant calls (9). The final two steps of filtering are designed to enrich the number of functional RNA editing sites. First, we keep only events that occur more than two times in at least one cancer type. Second, we keep only events that occur in exonic regions with a predicted function of missense, nonsense or stop-loss. The final step of filtering within exonic regions with a specific predicted function induces the largest difference in observed frequencies of RNA-editing events between our analysis and the published one 108. A comparative depiction of the frequencies of RNA-editing events identified in our analysis (Supplementary Table 24) and the previously published analysis <sup>108</sup> is seen in Supplementary Fig. 23.

#### Gene-centric table creation

To perform joint analysis across RNA and DNA alterations, each alteration type was condensed into a binary gene-centric format. Because alterations occur at many different scales (nucleotide, exonic, gene or transcript), to make them comparable we projected each alteration type onto the gene body. We summarized each alteration type by its presence or absence within a single gene, yielding a binary value per type for each gene-sample pair.

The events we included in this analysis were: RNA editing, nonsynonymous variants, expression, splicing alterations, copy-number alterations, fusions and alternative promoters. Each alteration type was summarized differently owing to their inherent differences.

RNA-editing events and non-synonymous variants can occur several times within a single gene body, so these events were denoted as 1 if they occurred at least once within a gene–sample pair.

For copy number, to obtain a single numerical value per gene-sample pair, the copy-number alteration was averaged over the gene body. Because we do not have matched normal samples against which to compare, we instead consider outlying events within each histotype as significant. Thus, a value of 1 was given to average copy-number alterations larger than 6 or smaller than 1.

Similar to non-synonymous variants, multiple splice events can occur within a gene body. The event with the most extreme PSI value within the gene body is selected as the candidate event for the gene. The candidate's PSI value for a gene is compared over all samples within a histotype and it is set to 1 (that is, significant) only if it the absolute value of its z-score is larger than 6 and the standard deviation is larger than 0.01 within that histotype.

Similar to expression outliers, we calculate a *z*-score using the log-transformed upper-quartile normalized FPKM values with a pseudo-count of 1. All genes within a histotype with a standard deviation larger than zero and an absolute value larger than three were identified as an outlier. Alternative promoter outliers were calculated based on relative promoter activity within each cancer type. To binarize the promoter

activity, a z-score cut-off of two over the relative expression distribution within each cancer type was used.

For ASE outliers, only genes with significant allelic imbalance (FDR  $\leq$  5% and allelic imbalance > 0.2, binomial test) were denoted as 1. All ASE events that were identified were further filtered to keep only genes that have not been identified as imprinted  $^{26}$ .

In addition to the z-score-filtering mentioned above, we further filtered non-synonymous SNVs, RNA-editing events and splicing events such that they either induce a frameshift or the alternative region contains an HGMD variant <sup>112</sup> of the category 'damaging'.

It must be noted that in many cases, the z-score calculated is not from a Gaussian distribution, so some events may be missed or falsely included. Through our choice of very stringent z-score thresholds and functional filters, we hope that spurious outlier events are minimized.

#### Pathway analysis

For our pathway analysis, we used the TCGA pathway definitions to examine genes and pathways that have several alterations at both the DNA and RNA level<sup>113</sup>.

#### Co-occurrence analysis

The co-occurrence analysis was also performed on the aforementioned binarized gene-centric table, but only including variants, expression outliers, alternative promoters, alternative splicing and fusions. SCNA and ASE are excluded owing to a large number of anticipated co-occurrence. In this analysis, we required at least one gene of a given alteration pair to be a COSMIC gene. For each alteration pair, based on the number of donors with both alterations, one alteration only and neither alterations in a set of cancer samples, we performed Fisher's exact test to determine whether the alteration pair was independent of each other. Such tests were followed by Benjamini-Hochberg multiple testing correction to obtain the FDR (or q values). To rule out the potential false-positive association caused by tissue-specific alterations, we performed the same analysis for each of the tumour types with at least 50 patients, and retained only those alteration pairs that were significantly associated in both the pan-cancer analysis and in at least one specific cancer indication. Among the significantly associated alteration pairs, the co-occurred pairs were those with odds ratio greater than 1. Pathway enrichment and visualization<sup>21,114</sup> were conducted using the R package ReactomePA<sup>79</sup>. The circos plots were generated using the R package circlize<sup>115</sup>. The splicing related genes were derived from the genes annotated as 'REACTOME MRNA SPLICING' or 'REACTOME MRNA SPLICING MINOR PATHWAY' in the Molecular Signatures Database (MSigDB)<sup>116</sup>.

# Identifying genes with heterogeneous mechanisms of alterations in *cis*

Genes with multiple heterogeneous mechanisms of RNA alteration were identified from associations of *cis* variants with gene expression, ASE, fusions and splicing. For gene expression, genes associated with somatic eQTL with FDR < 5% were selected. For ASE, the top 5% of genes ranked by the predicted contribution of somatic variants on ASE. For fusions, all RNA fusions with structural variant support were selected. For splicing, genes having somatic mutations within 10 bp of an annotated splice site or 3 bp of a branch point and associated splicing were selected. These associated splicing events also had to have a |z-score| greater than or equal to 3 and the difference of percent spliced in the outlier event was greater than or equal to 10%.

#### Recurrence analysis

The recurrence analysis was performed on the binarized gene-centric table for all nine alteration types. The recurrence analysis was performed in three main steps: (1) Aggregate within each alteration type across all samples. This results in a sum for each gene-alteration pair. (2) Convert the counts to ranks within each alteration. The smallest rank

goes to the most frequently altered genes. Ranks are split evenly across ties. (3) To generate a single score for each gene, the second smallest rank across alterations is used as the score. To identify a score cut-off value for significantly altered genes, a null distribution was generated through permutation. The permutations were performed over the samples within each gene-alteration pair, this was done over all genes and samples 1,000 times, concatenating together all observations, results in 16.8 million permuted scores. P < 0.05 as derived from the null distribution was defined as significant, resulting in a score greater than or equal to 774 considered as significant.

WExT<sup>117</sup> was used to test the significance of mutually exclusivity of RNA and DNA alterations. As further evidence that *CDK12* alterations may have a functional affect, we find evidence of the previously detected link<sup>55</sup> between a large tandem duplicator phenotype (here defined as more than 10 tandem duplications of size greater than 100 kb) and *CDK12* somatic eQTL mutation (7 out of 18 somatic eQTL carriers are also among the 215 large tandem duplicator cases, P = 0.032, hypergeometric test).

#### Statistical tests

All common statistical tests are two-sided unless otherwise specified. No statistical methods were used to predetermine sample size. The experiments were not randomized and investigators were not blinded to allocation during experiments and outcome assessment.

#### **Reporting summary**

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

### **Data availability**

Somatic and germline variant calls, mutational signatures, subclonal reconstructions, and other core data generated by the ICGC and TCGA PCAWG Consortium are described in an accompanying Article<sup>5</sup> and are available for download at https://dcc.icgc.org/releases/PCAWG. Additional information on accessing the data, including raw read files, can be found at https://docs.icgc.org/pcawg/data/. In accordance with the data access policies of the ICGC and TCGA projects, most molecular, clinical and specimen data are in an open tier that does not require access approval. To access potentially identification information, such as germline alleles and underlying sequencing data, researchers will need to apply to the TCGA data access committee via dbGaP (https:// dbgap.ncbi.nlm.nih.gov/aa/wga.cgi?page=login) for access to the TCGA portion of the dataset, and to the ICGC data access compliance office (http://icgc.org/daco) for the ICGC portion of the dataset. In addition, to access somatic SNVs derived from TCGA donors, researchers will also need to obtain dbGaP authorization. Data derived specifically from RNA-seq analysis can be found at https://dcc.icgc.org/releases/PCAWG/ transcriptome. Subfolders contain identification and quantification of alternative promoter usage, alternative splicing, RNA fusions, gene expression, transcript-level expression and RNA editing. Identified eQTLs are in https://dcc.icgc.org/releases/PCAWG/transcriptome/eQTL and a binarized table indicating all RNA and DNA alterations for each gene can be found in the subfolder https://dcc.icgc.org/releases/PCAWG/transcriptome/recurrence\_analyses/. In addition, quality-control metrics and metadata are also included. Some datasets are denoted with synXXXXX accession numbers and available at Synapse (https://www.synapse.org/).

#### **Code availability**

The core computational pipelines used by the PCAWG Consortium for alignment, quality control and variant calling are available to the public at https://dockstore.org/search?search=pcawg under the GNU General Public License v.3.0, which allows for reuse and distribution. Further details on code availability are in the Supplementary Information.

- 58. Dobin, A. et al. STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29, 15-21 (2013).
- Kim, D. et al. TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions. Genome Biol. 14, R36 (2013).
- Harrow, J. et al. GENCODE: the reference human genome annotation for The ENCODE Project. Genome Res. 22, 1760–1774 (2012).
- Fonseca, N. A., Petryszak, R., Marioni, J. & Brazma, A. iRAP an integrated RNA-seq analysis pipeline. Preprint at https://www.bioRxiv.org/content/10.1101/005991v1 (2014).
- Bioinformatics, B. FastQC: a quality control tool for high throughput sequence data; http://www.bioinformatics.babraham.ac.uk/projects/fastqc (2011).
- Cancer Genome Atlas Research Network. The molecular taxonomy of primary prostate cancer. Cell 163, 1011–1025 (2015).
- Li, H. et al. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25, 2078–2079 (2009).
- Anders, S., Pyl, P. T. & Huber, W. HTSeq—a Python framework to work with highthroughput sequencing data. Bioinformatics 31, 166–169 (2015).
- Bray, N. L., Pimentel, H., Melsted, P. & Pachter, L. Near-optimal probabilistic RNA-seq quantification. Nat. Biotechnol. 34, 525–527 (2016).
- Mortazavi, A., Williams, B. A., McCue, K., Schaeffer, L. & Wold, B. Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat. Methods 5, 621–628 (2008).
- Krijthe, J. H. Rtsne: t-distributed stochastic neighbor embedding using barnes-hut implementation; https://github.com/jkrijthe/Rtsne (2015).
- Dentro, S. C. et al. Portraits of genetic intra-tumour heterogeneity and subclonal selection across cancer types. Preprint at https://www.biorxiv.org/content/10.1101/312041v4 (2018)
- Kahles, A., Ong, C. S., Zhong, Y. & Rätsch, G. SplAdder: identification, quantification and testing of alternative splicing events from RNA-Seq data. *Bioinformatics* 32, 1840–1847 (2016).
- Pedregosa, F. et al. Scikit-learn: machine learning in python. J. Mach. Learn. Res. 12, 2825–2830 (2011).
- Stegle, O., Parts, L., Piipari, M., Winn, J. & Durbin, R. Using probabilistic estimation of expression residuals (PEER) to obtain increased power and interpretability of gene expression analyses. *Nat. Protocols* 7, 500–507 (2012).
- GTEx Consortium. The Genotype-Tissue Expression (GTEx) project. Nat. Genet. 45, 580–585 (2013)
- The Gene Ontology Consortium. Gene Ontology: tool for the unification of biology. Nat. Genet. 25, 25–29 (2000).
- The Gene Ontology Consortium. Gene Ontology Consortium: going forward. Nucleic Acids Res. 43, D1049–D1056 (2015).
- Durinck, S., Spellman, P. T., Birney, E. & Huber, W. Mapping identifiers for the integration of genomic datasets with the R/Bioconductor package biomaRt. *Nat. Protocols* 4, 1184–1191 (2009).
- Durinck, S. et al. BioMart and Bioconductor: a powerful link between biological databases and microarray data analysis. *Bioinformatics* 21, 3439–3440 (2005).
- Yu, G., Wang, L.-G., Han, Y. & He, Q.-Y. clusterProfiler: an R package for comparing biological themes among gene clusters. OMICS 16, 284–287 (2012).
- Yu, G. & He, Q.-Y. ReactomePA: an R/Bioconductor package for reactome pathway analysis and visualization. Mol. Biosyst. 12. 477–479 (2016).
- Li, H. A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. *Bioinformatics* 27, 2987–2993 (2011).
- Lippert, C., Casale, F. P., Rakitsch, B. & Stegle, O. LIMIX: genetic analysis of multiple traits. Preprint at https://www.bioRxiv.org/content/10.1101/003905v2 (2014).
- Davis, J. R. et al. An efficient multiple-testing adjustment for eQTL studies that accounts for linkage disequilibrium between variants. Am. J. Hum. Genet. 98, 216–224 (2016).
- Pers, T. H., Timshel, P. & Hirschhorn, J. N. SNPsnap: a Web-based tool for identification and annotation of matched SNPs. *Bioinformatics* 31, 418–420 (2015).
- Fan, Y. et al. MuSE: accounting for tumor heterogeneity using a sample-specific error model improves sensitivity and specificity in mutation calling from sequencing data. Genome Biol. 17, 178 (2016).
- Quinlan, A. R. & Hall, I. M. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26, 841–842 (2010).
- Corces, M. R. et al. The chromatin accessibility landscape of primary human cancers. Science 362, 362 (2018).
- Zhang, W. et al. A global transcriptional network connecting noncoding mutations to changes in tumor gene expression. *Nat. Genet.* 50, 613–620 (2018).
- Smith, K. S. et al. Signatures of accelerated somatic evolution in gene promoters in multiple cancer types. *Nucleic Acids Res.* 43, 5307–5317 (2015).
- Fishilevich, S. et al. GeneHancer: genome-wide integration of enhancers and target genes in GeneCards. Database 2017, 2017 (2017).
- Haeussler, M. et al. The UCSC Genome Browser database: 2019 update. Nucleic Acids Res. 47, D853–D858 (2019).
- ENCODE Project Consortium. An integrated encyclopedia of DNA elements in the human genome. Nature 489, 57-74 (2012).
- Wang, C. et al. Systematic identification of genes with a cancer-testis expression pattern in 19 cancer types. Nat. Commun. 7, 10499 (2016).
- Wallace, C. Statistical testing of shared genetic control for potentially related traits. Genet. Epidemiol. 37, 802–813 (2013).
- Baron, R. M. & Kenny, D. A. The moderator-mediator variable distinction in social psychological research: conceptual, strategic, and statistical considerations. J. Pers. Soc. Psychol. 51, 1173–1182 (1986).
- Preacher, K. J. & Hayes, A. F. SPSS and SAS procedures for estimating indirect effects in simple mediation models. Behav. Res. Methods Instrum. Comput. 36, 717–731 (2004).
- Rosseel, Y. lavaan: AnRPackage for structural equation modeling. J. Stat. Softw. 48, 2 (2012).
- Tingley, D., Yamamoto, T., Hirose, K., Keele, L. & Imai, K. mediation: RPackage for causal mediation analysis. J. Stat. Softw. 59, 5 (2014).

- Howie, B. N., Donnelly, P. & Marchini, J. A flexible and accurate genotype imputation method for the next generation of genome-wide association studies. PLoS Genet. 5, e1000529 (2009).
- 99. Nik-Zainal, S. et al. The life history of 21 breast cancers. Cell 149, 994-1007 (2012).
- 100. McLaren, W. et al. The Ensembl variant effect predictor. Genome Biol. 17, 122 (2016).
- Castel, S. E., Levy-Moonshine, A., Mohammadi, P., Banks, E. & Lappalainen, T. Tools and best practices for data processing in allelic expression analysis. Genome Biol. 16, 195 (2015)
- Subramanian, A. et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc. Natl Acad. Sci. USA 102, 15545– 15550 (2005)
- Frith, M. C. et al. A code for transcription initiation in mammalian genomes. Genome Res. 18. 1–12 (2008).
- Signal, B., Gloss, B. S., Dinger, M. E. & Mercer, T. R. Machine learning annotation of human branchpoints. *Bioinformatics* 34, 920–927 (2018).
- Mercer, T. R. et al. Genome-wide discovery of human splicing branchpoints. Genome Res. 25, 290–303 (2015).
- 106. Ge, H. et al. FusionMap: detecting fusion genes from next-generation sequencing data at base-pair resolution. *Bioinformatics* **27**, 1922–1928 (2011).
- Nicorici, D. et al. FusionCatcher a tool for finding somatic fusion genes in paired-end RNA-sequencing data. Preprint at https://www.bioRxiv.org/content/10.1101/011650v1 (2014).
- Han, L. et al. The Genomic Landscape and Clinical Relevance of A-to-I RNA Editing in Human Cancers. Cancer Cell 28, 515–528 (2015).
- Li, Q. et al. Caste-specific RNA editomes in the leaf-cutting ant Acromyrmex echinatior. Nat. Commun. 5, 4943 (2014).
- Genome of the Netherlands Consortium. Whole-genome sequence variation, population structure and demographic history of the Dutch population. Nat. Genet. 46, 818–825 (2014).
- Wang, K., Li, M. & Hakonarson, H. ANNOVAR: functional annotation of genetic variants from high-throughput sequencing data. *Nucleic Acids Res.* 38, e164 (2010).
- Stenson, P. D. et al. The Human Gene Mutation Database: towards a comprehensive repository of inherited mutation data for medical research, genetic diagnosis and nextgeneration sequencing studies. Hum. Genet. 136, 665–677 (2017).
- Sanchez-Vega, F. et al. Oncogenic signaling pathways in the cancer genome atlas. Cell 173, 321–337,e10 (2018).
- Merico, D., Isserlin, R., Stueker, O., Emili, A. & Bader, G. D. Enrichment map: a networkbased method for gene-set enrichment visualization and interpretation. PLoS ONE 5, e13984 (2010).
- Gu, Z., Gu, L., Eils, R., Schlesner, M. & Brors, B. circlize implements and enhances circular visualization in R. Bioinformatics 30, 2811–2812 (2014).
- Liberzon, A. et al. Molecular signatures database (MSigDB) 3.0. Bioinformatics 27, 1739–1740 (2011).
- Leiserson, M. D. M., Reyna, M. A. & Raphael, B. J. A weighted exact test for mutually exclusive mutations in cancer. *Bioinformatics* 32, 1736–1745 (2016).
- Rafnar, T. et al. Sequence variants at the TERT-CLPTM1L locus associate with many cancer types. Nat. Genet. 41, 221–227(2009).
- Bojesen, S. E. et al. Multiple independent variants at the TERT locus are associated with telomere length and risks of breast and ovarian cancer. Nat. Genet. 45, 371–384 (2013).
- Ye, K. et al. Systematic discovery of complex insertions and deletions in human cancers. Nat. Med. 22, 97–104 (2016).

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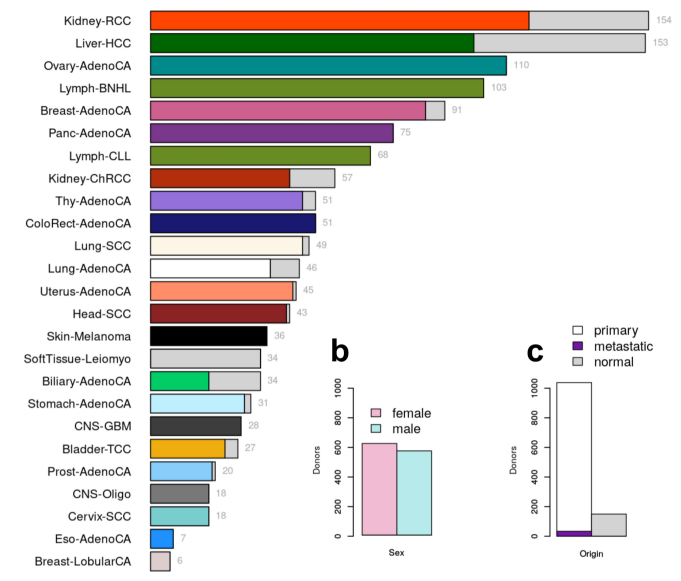
#### Additional information

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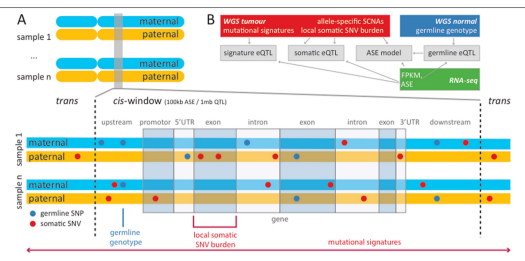
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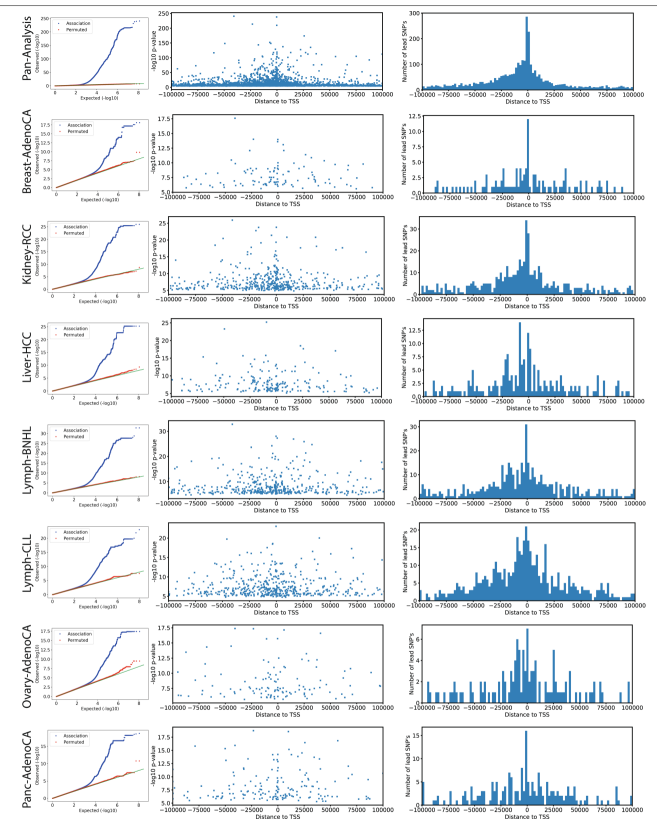
**Extended Data Fig. 1** | **Pan-cancer expression profiling of 1,188 PCAWG donors. a**, Tumour and normal RNA-seq data from 27 histotypes. The total number of samples is shown to the right of the bars. Grey bars denote matched

healthy samples. **b**, Number of female versus male donors. **c**, Total number of tumour and matched healthy samples from the PCAWG study. A subset of tumours (dark violet) was metastatic.



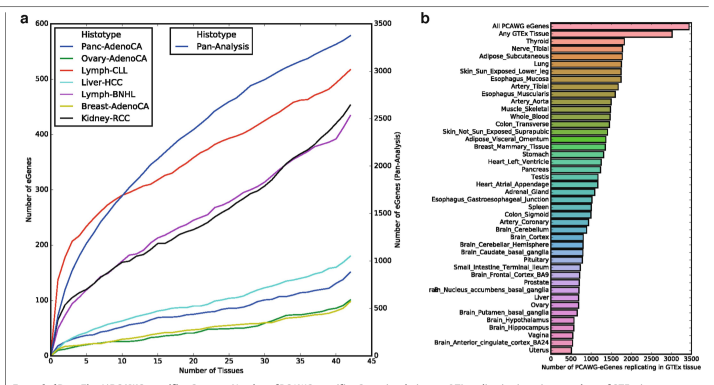
**Extended Data Fig. 2** | **Overview of the different sources of genetic variation considered in the analysis. a**, For analyses of cis regulation, mono-allelic single-nucleotide germline variants (single nucleotide polymorphisms (SNPs), blue) were individually tested for association with total gene expression using standard eQTL approaches. Owing to their low recurrence in the cohort, somatic SNVs were aggregated in burden categories depending on their position relative to the gene tested (for example, promoter, S' UTR or intron). Local SNV burdens were then tested for association with ASE globally across all genes, as well as with total expression on a per-gene level using eQTL approaches. *Trans* effects were estimated by testing total gene expression for

association with mutational and epigenetic signatures. Window sizes were 1 Mb for all somatic *cis*-eQTL analyses, and 100 kb for ASE and germline *cis*-eQTL. **b**, Overview of the different datasets and their contributions to the analyses described in **a**. Germline genotypes were derived from the matched healthy whole-genome sequencing (WGS) samples. Allele-specific SCNAs, mutational signatures and local SNV burdens were derived from the tumour WGS in comparison to the unaffected WGS samples. ASE and total expression (FPKM) were derived from the tumour and normal RNA-seq data. Arrows indicate dependencies between individual analyses carried out.

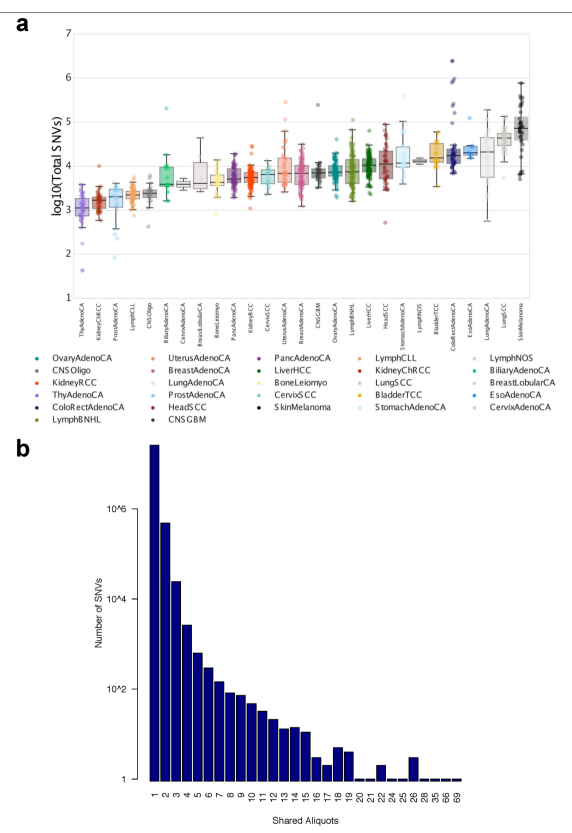


**Extended Data Fig. 3** | **Germline eQTL lead variants.** Left, quantile—quantile (Q-Q) plot of P values of germline eQTL lead variants in the pan-cancer and histotype-specific analysis (FDR  $\leq$  5%, blue) and P values of the same analysis

after permutation (random permutation of patients, red). Middle and right, distributions of distance to the respective TSS of all germline eQTL lead variants in the pan-cancer and histotype-specific analysis.

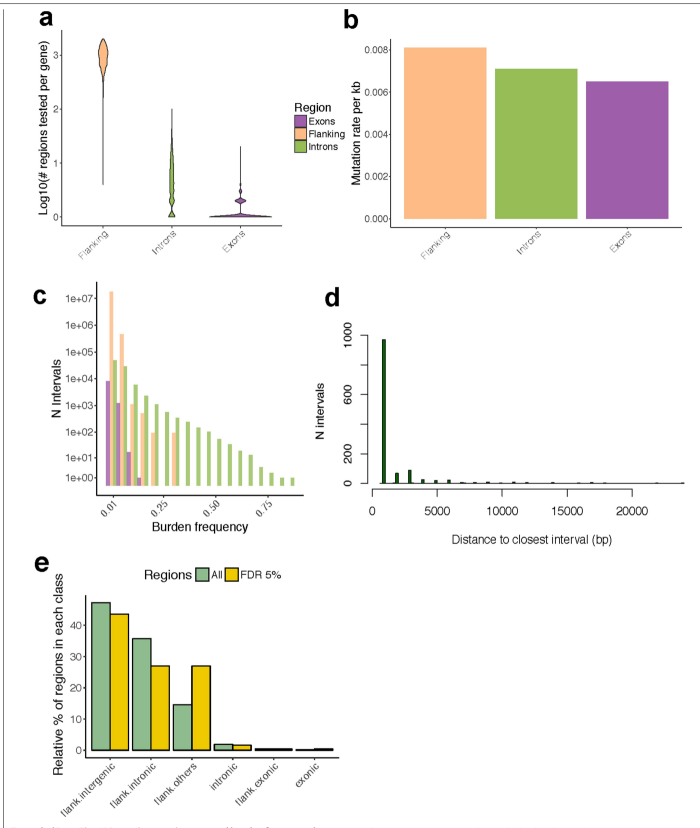


 $\textbf{Extended Data Fig. 4} | \textbf{PCAWG-specific eGenes. a}, \textbf{Number of PCAWG-specific eGenes in relation to eQTL replication in various numbers of GTEx tissues.} \\ \textbf{b}, \textbf{Number of eGenes of the PCAWG pan-analysis replicating in corresponding GTEx tissues.} \\ \textbf{c}, \textbf{c},$ 



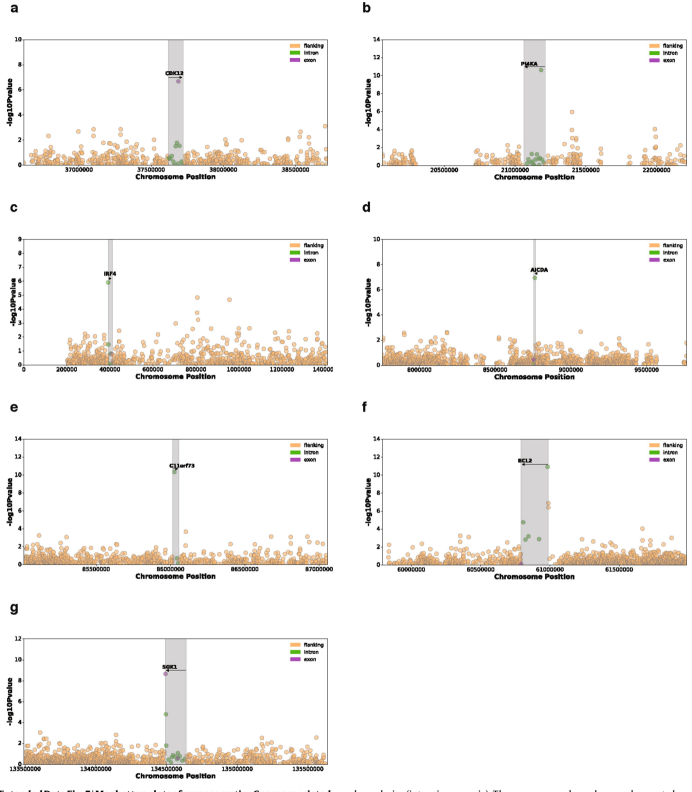
**Extended Data Fig. 5** | *Cis*-mutational somatic burden. a, Total number of somatic mutational load per cancer type. Median numbers of SNVs range from 1,139 in thyroid adenocarcinoma to 72,804 in skin melanoma. b, Number of

recurrent somatic SNVs shared by increasing numbers of patients. A small fraction of 86 SNVs is detected in more than 1% of the cohort (12 patients).



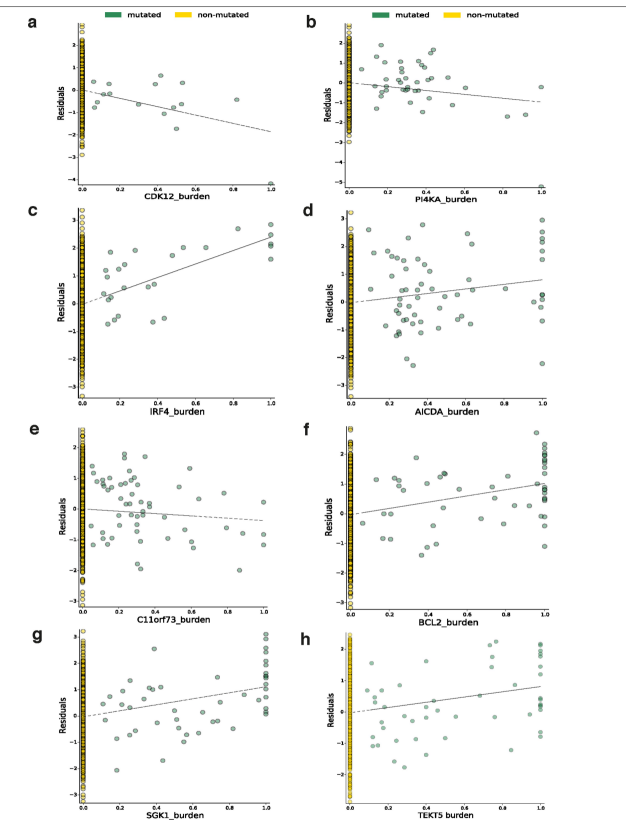
**Extended Data Fig. 6** | **Somatic mutation rate and burden frequency by type of region tested. a**, Number of mutated regions tested per gene with somatic burden frequency  $\geq 1\%$ . **b**, Mutation rate per kilobase. **c**, Burden frequency, stratified by the type of interval tested (flanking, exonic or intronic). **d**, Distribution of distances (bp) of the leading intervals (FDR  $\leq 5\%$ ) to the closest (left and right) interval such that the association *P* value decreases by at least one order of magnitude (99% of the distribution is shown). **e**, Breakdown of all genomic regions tested (n = 1,049,102 with burden frequency  $\geq 1\%$ ) and of

the 567 genomic regions that underlie the observed somatic cis-eQTL at a FDR of 5% (intronic denotes eGene intron; exonic denotes eGene exon; flank. denotes 2-kb flanking region within 1 Mb distance to the eGene start and end; flank.intergenic denotes flanking region in a genomic location without gene annotations; flank.intronic denotes flanking region overlapping an intron of a nearby gene; and flank.others denotes flanking region partially overlapping several annotations of a nearby gene).

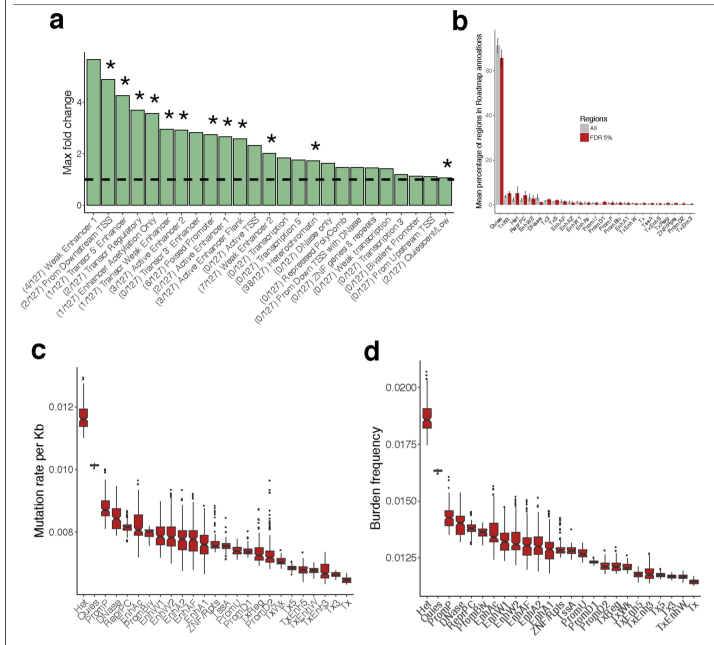


 $\label{lem:extended} \textbf{Extended Data Fig. 7} | \textbf{Manhattan plots of seven somatic eGenes associated} \\ \textbf{with genic lead burden.} & \textbf{Altogether, 11} genic somatic eQTLs showed significant} \\ \textbf{changes in gene expression associated with somatic burdens within the gene} \\ \textbf{Altogether, 12} | \textbf{Manhattan plots of seven somatic eGenes associated} \\ \textbf{Manhattan plots of seven somatic eGenes eGenes associated} \\ \textbf{Manhattan plots of eGenes eG$ 

boundaries (intronic or exonic). The seven genes shown here are known to be important in the pathogenesis of specific cancers. **a**, CDK12. **b**, PI4KA. **c**, IRF4. **d**, AICDA. **e**, CIIorf73 (also known as HIKESHI). **f**, BCL2. **g**, SGK1.

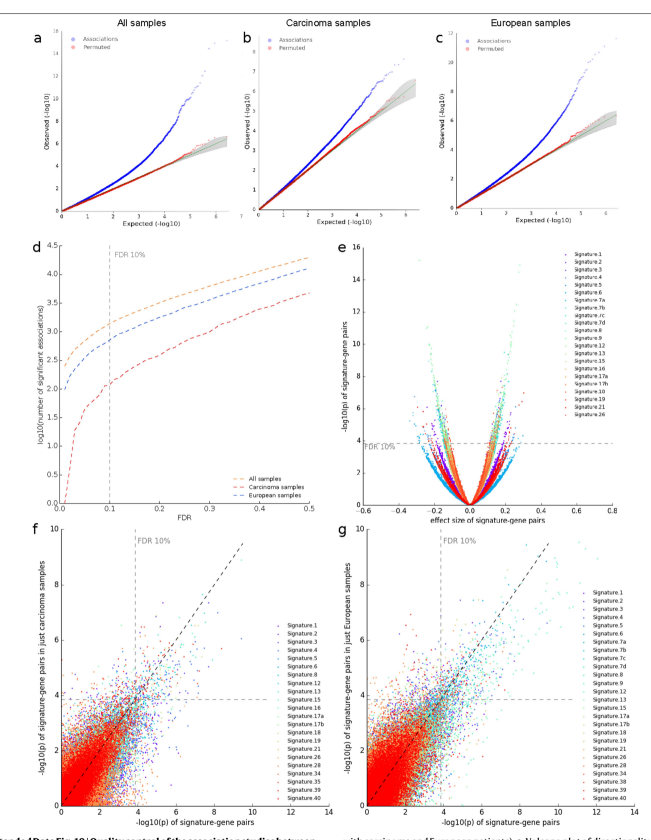


 $\textbf{Extended Data Fig. 8} | \textbf{Scatter plots of eight somatic e Genes.} | \textbf{Plots show the effect of the lead weighted burden on the gene expression residuals (obtained as described in the Methods) of these genes. \textbf{a}, \textit{CDK12.} \textbf{b}, \textit{Pl4KA.} \textbf{c}, \textit{IRF4.} \textbf{d}, \textit{AICDA.} \textbf{e}, \textit{C11orf73.} \textbf{f}, \textit{BCL2.} \textbf{g}, \textit{SGK1.} \textbf{h}, \textit{TEKT5.}$ 



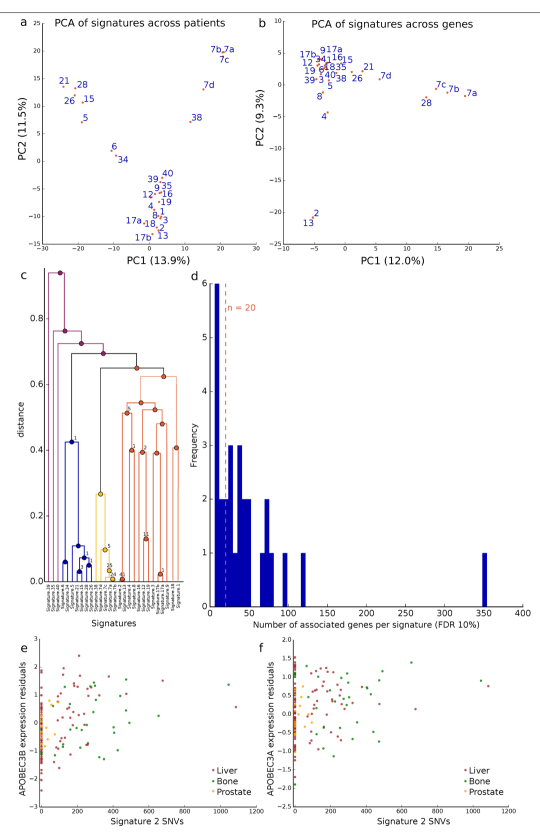
Extended Data Fig. 9 | Roadmap epigenome marks overlapping flanking intervals with somatic burden. a, Maximum fold enrichment of epigenetic marks from the Roadmap Epigenomics Project across 127 cell lines. The number of cell lines with significant enrichments is indicated in parentheses (FDR  $\leq$  10%); asterisks denote significant enrichments in at least one cell line. b, Mean percentages (over the 127 cell lines) of regions overlapping (by at least 10% of their length) Roadmap epigenome marks, calculated using all genomic flanking regions (n = 1,637,638) and the subset of 556 flanking intervals associated with somatic eQTL (FDR  $\leq$  5%). c, Mutation rate per kilobase. d, Burden frequency (across the 127 cell lines) of the 556 flanking intervals in

somatic eQTLs (FDR  $\leq$  5%), overlapping 25 Roadmap epigenome marks. DNase, DNase only; EnhA, active enhancer; EnhAc, enhancer acetylation only; EnhAF, active enhancer flank; EnhW, weak enhancer; Het, heterochromatin; PromBiv, bivalent promoters; PromD, promoter downstream; PromP, poised promoters; PromU, promoter upstream; Quies, quiescent/low; ReprPC, repressed PolyComb; TssA, active TSS; TxReg, transcription regulatory; ZNF/Rpts, ZNF genes and repeats; Tx, transcription; Tx3, transcription 3′, Tx5, transcription 5′; TxEnh3, transcription 3′ enhancer; TxEnh5, transcription 5′ enhancer; TxEnhW, transcription weak enhancer; TxWk, weak transcription.



Extended Data Fig. 10 | Quality control of the association studies between gene expression and mutational signatures. a–c, Q–Q plots of the P values of the linear model to associate expression of 18,831 genes with 28 mutational signatures across all 1,159 patients (a), 877 patients with carcinoma (b), or 891 European patients (c). d, Number of significant associations ( $\log_{10}$ -transformed) at different FDR thresholds (across all patients, patients

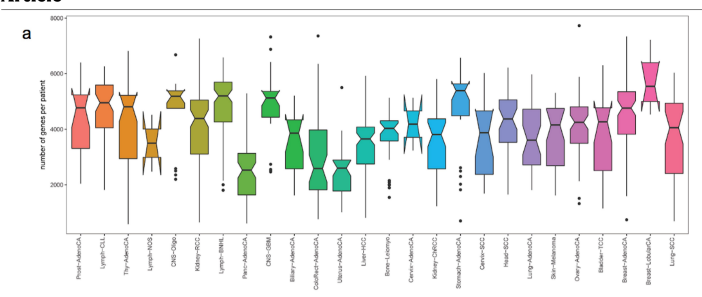
with carcinoma and European patients). **e**, Volcano plot of directionality of effects in the analysis of all patients. **f**, **g**, Comparison of analyses between all patients and patients with carcinoma (**f**) and between all patients and European patients (**g**). The  $-\log_{10}(P \text{ values})$  per signature–gene pair are correlated (r=0.763 (**f**) and r=0.789 (**g**), Pearson correlation coefficient), especially above an FDR threshold of 10%.



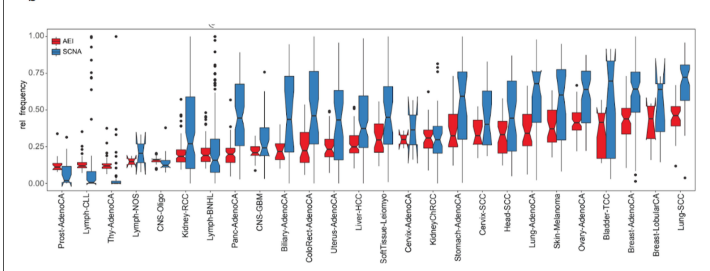
**Extended Data Fig. 11**|See next page for caption.

Extended Data Fig. 11 | Relationship between mutational signatures and gene expression patterns. a, b, Principal component analysis (PCA) of signatures across 1,159 patients (PCA on signature-specific SNVs per patient) (a) and signature-gene expression associations across 18,831 genes (PCA on adjusted P values of signature-gene expression associations) (b). The PCA on the SNVs recapitulates known interdependencies, for example, between signatures 7, whereas the PCA on the signature-gene association studies also emphasizes functional relatedness, for example, between signatures 2 and 13. c, Hierarchical clustering of signatures. The numbers at the nodes indicate the number of genes commonly associated with two to four respective signatures. The dendrogram shows genes that are associated with more than one signature mostly owing to similar SNV patterns of these signatures across patients.

**d**, Frequency of number of significantly associated genes per signature (FDR ≤ 10%). Although many signatures are significantly associated with a few genes, 18 signatures are associated with more than 20 genes. Signature 9 is associated with more than 350 genes. Vice versa, 1,009 genes are associated with only one signature, 129 with two, 32 with three, 5 with four and 1 with five signatures. **e**, **f**, Mutational signature–gene associations, depicting positive associations between the expression of the canonical APOBEC pathway genes *APOBEC3B* (**e**) and *APOBEC3A* (**f**) and signature 2. The associations within the three cancer type with the strongest correlation between signature and gene expression (hepatocellular carcinoma (Liver–HCC), bone leiomyosarcoma (Bone–Leiomyo) and prostate adenocarcinoma (Prost–AdenoCA)) are shown.

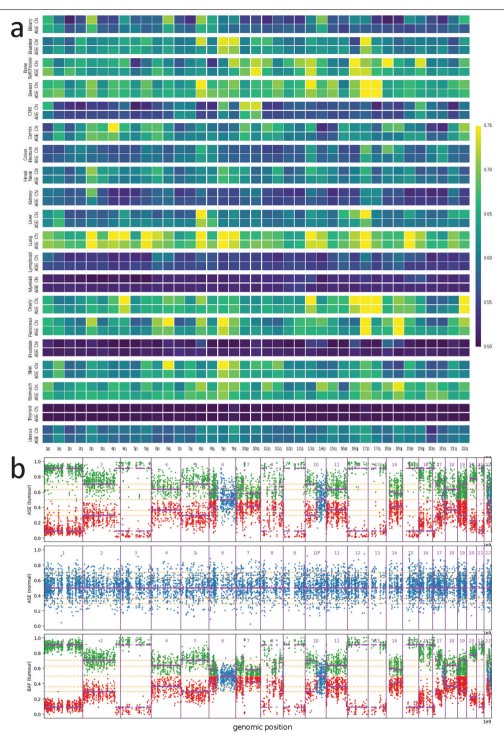






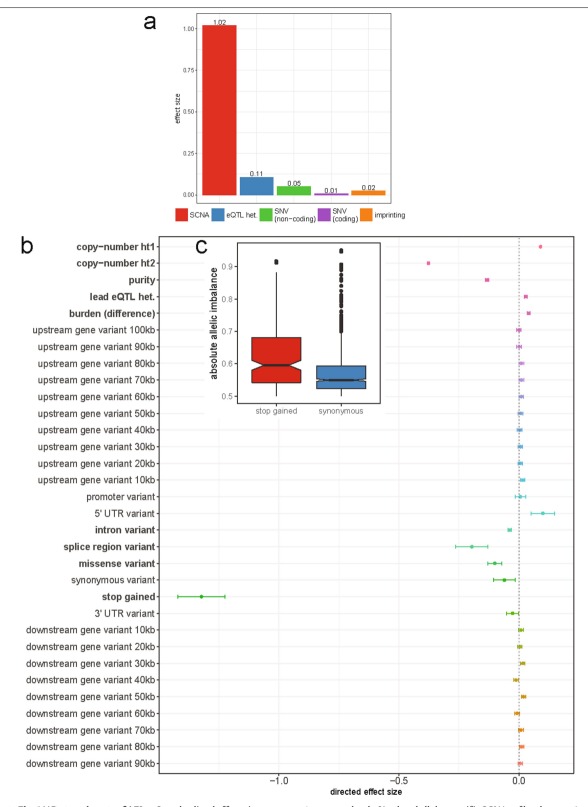
**Extended Data Fig. 12** | **ASE analysis. a**, All types of cancer are ordered by the average AEI frequency. The numbers of genes per patient for which ASE could be quantified are shown, stratified according to cancer type, resulting in between 588 and 7,728 genes per patient. **b**, Distribution of the fraction of

genes with AEI (red) and SCNAs (blue) over the number of measurable genes for each patient across the cohort. Cancer types with high chromosomal instability also exhibit highest amounts of AEI.



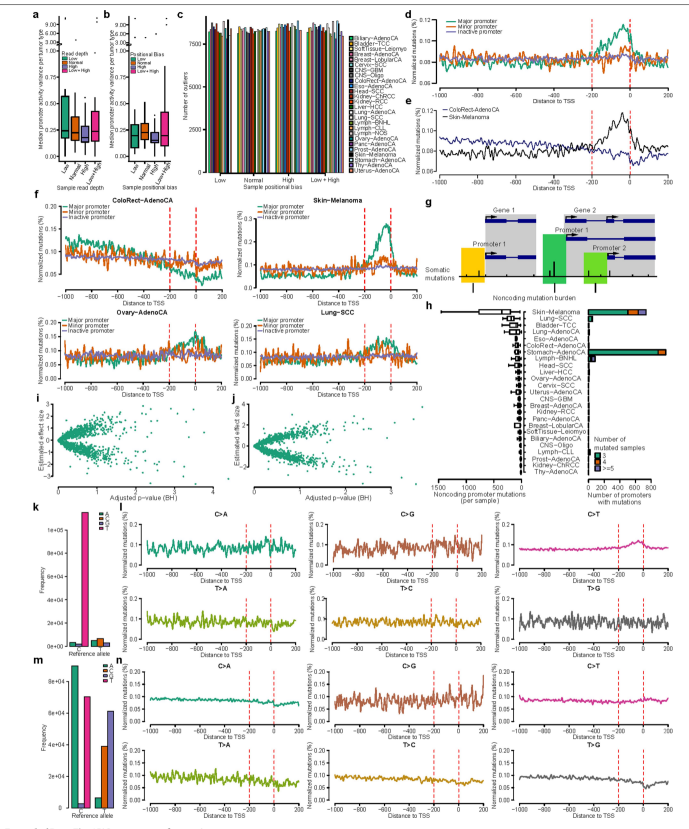
Extended Data Fig. 13 | SCNAs as major driver for allelic dysregulation in cancer. a, Absolute allelic expression imbalance closely follows allelic imbalance at the genomic level. Values of 0.5 (blue) denote equal number of reads from both alleles. Values of 1 (yellow) reflect mono-allelic expression or

regions with loss of heterozygosity. **b**, Comparison between B-allele frequency (BAF) and ASE ratios from a single patient with lung cancer (LUAD-US) with profound chromosomal instability shows strong correlation between allelic imbalance on expression and genomic levels.



 $\label{lem:extended Data Fig. 14 | Determinants of AEI. a}, Standardized effect sizes on the presence of AEI, taking only SCNAs, germline eQTLs, coding and non-coding mutations into account. In summary, SCNAs accounted for 86.1% of the total effect size, followed by germline eQTLs (9.0%) and somatic SNVs (4.8%).$ **b**, Relevance of individual somatic mutation types ('copy-number ht1' and

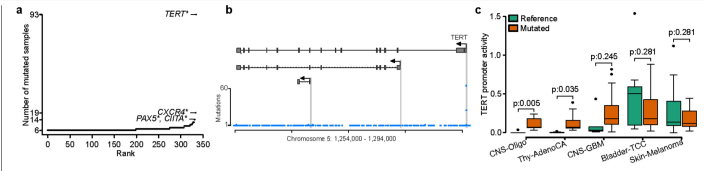
'copy-number ht2' as local allele-specific SCNAs of haplotypes 1 and 2, respectively), germline eQTLs and other covariates for the ASE ratio. Significant covariates (FDR  $\leq$  5%) are highlighted in bold.  $\boldsymbol{c}$ , Comparison of the effect of protein-truncating variants (stop-gained) and synonymous variants on the ASE ratio.



 $\textbf{Extended Data Fig. 15} \, | \, \textbf{See next page for caption}.$ 

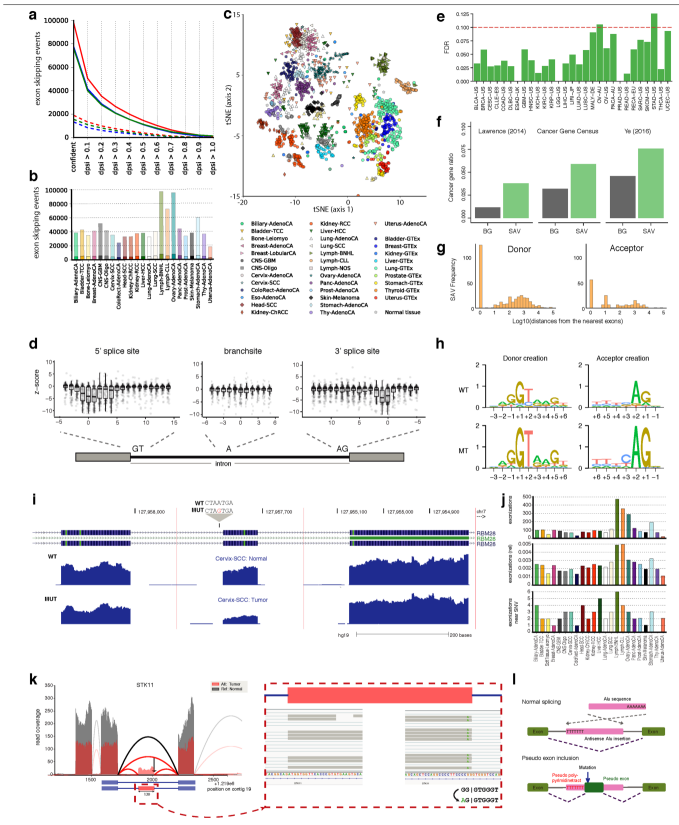
Extended Data Fig. 15 | Overview of estimations of promoter activity and non-coding promoter mutations associations and patterns. a, b, The technical variation of the promoter activity estimates across varying library depth (a) and positional bias (b). c, The number of outlier promoters per tumour type according to promoter activity variance (variance larger than  $1.5 \times 1.5 \times 1.5$ 

TSS, in which major promoters show an enrichment of mutations whereas minor and inactive promoters do not.  $\mathbf{g}$ , Schematic of the calculation of noncoding promoter mutational burden.  $\mathbf{h}$ , Overview of non-coding promoter mutations per sample and the number of mutated promoters per tumour type for promoters with at least three mutated samples.  $\mathbf{i}$ ,  $\mathbf{j}$ , Association of absolute ( $\mathbf{i}$ ) and relative ( $\mathbf{j}$ ) promoter activity with promoter mutations across all samples.  $\mathbf{k}$ ,  $\mathbf{l}$ , Overview of promoter mutations for skin melanoma tumours.  $\mathbf{k}$ , Most promoter mutations are C>T, which indicates UV-induced DNA damage.  $\mathbf{l}$ , Distribution of promoter mutations for each mutation class reveals the enrichment of C>T mutations around the 200-bp window upstream.  $\mathbf{m}$ ,  $\mathbf{n}$ , Overview of promoter mutations for colorectal adenocarcinoma tumours.  $\mathbf{m}$ , Most promoter mutations are C>A and C>T.  $\mathbf{n}$ , Distribution of promoter mutations for each mutation class does not display an enrichment of mutations around the 200-bp window upstream, differing from the mutation pattern of skin melanoma tumours.



**Extended Data Fig. 16** | *TERT* **promoter mutations. a**, Promoters ranked by the number of mutated samples across all types of cancer in a 200-bp window. Asterisk indicates cancer census genes. **b**, The *TERT* locus and number of mutations observed at each position. The first promoter shows a highly

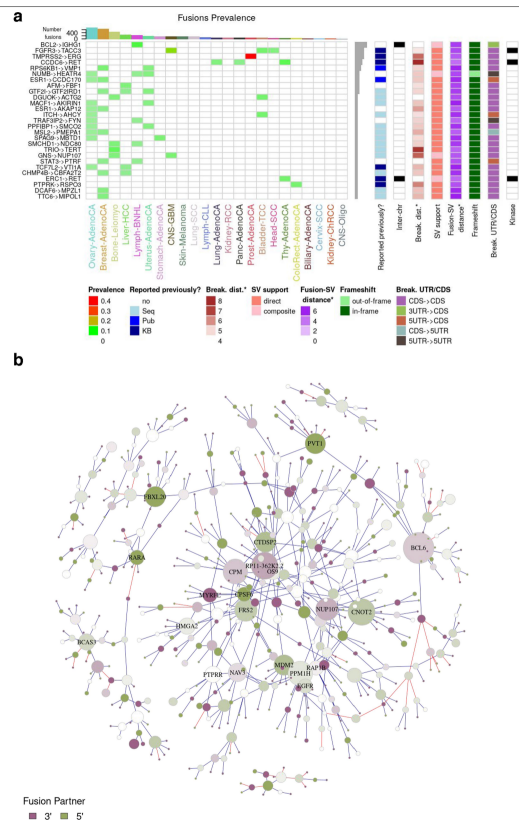
recurrent non-coding mutation reported previously ^118,119.  $\bf c$ , Comparison of TERT promoter activity for mutated and non-mutated samples per tumour type.



Extended Data Fig. 17 | See next page for caption.

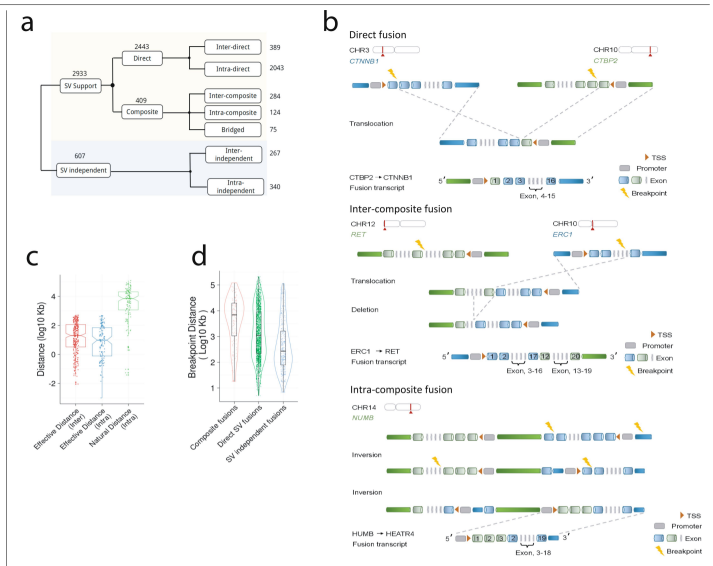
Extended Data Fig. 17 | Alternative splicing and association with somatic mutations. a, Number of exon-skipping events confirmed at different ΔPSI thresholds in tumour (red), matched healthy (green) and GTEx (blue) samples for liver tissue. Dashed lines show the subset of exon-skipping events that only contain annotated introns. b, Number of exon-skipping events confirmed at a ΔPSI level of greater than 0.3 for the individual histotypes. Transparent section of bars represents the fraction of novel events, containing at least one unannotated intron. c, Splicing landscape for exon-skipping events. t-SNE analysis based on exon-skipping PSI values for all ICGC tumour and healthy samples together with tissue-matched GTEx samples. d, Position-specific effect of somatic mutations on alternative splicing. Magnitude and direction of mutation-associated splicing alterations. e, Permutation-based FDR values for SAV detection based on the different types of cancer. f, Cancer gene set enrichment for SAV sets, shown for cancer census gene set (middle) and sets determined in ref. 48 (left) and ref. 120 (right). g, Positional distributions (logarithms of distance from the nearest exons) of somatic variant creating  $novel splicing donors \, and \, acceptors. \, \boldsymbol{h}, Sequence \, motiflogos \, around \, somatic$ mutation creating novel splicing motifs. i, Example splicing effect of a branchpoint mutation. UCSC genome browser RNA-seq coverage plots of cassette exon event in RBM28 between mutant and wild type. Mutant (bottom track)

contains an A>G mutation 29 nucleotides upstream from the acceptor site of an affected exon. i, Distribution of new cassette exon events detected only within the PCAWG cohort. Top, number of events per histology type. Middle, events normalized to the total number of cassette exons detected in the histology types. Bottom, the number of exonization events per histotype for the subset with the novel cassette exons colocated to a somatic alteration near the acceptor or donor of the exon.  $\boldsymbol{k},$  Example of an exonization event in the tumour-suppressor gene STK11. RNA-seq read coverage for a part of the gene is shown in red for a donor carrying the alternate allele and in grey for a random donor with reference allele. The cassette exon event is shown as a schematic below, with blue (red) boxes denoting constitutive (alternative) exons and blue  $solid \, lines \, denoting \, introns. \, Magnified \, panels \, at \, the \, bottom \, show \, details \, from \,$ Integrative Genomics Viewer visualization, highlighting a somatic mutation at the 3' end of the cassette exon. The associated sequencing change is illustrated on the bottom right corner, in which the vertical bar denotes the exon-intron boundary. I, Alu-based exonization mechanism. Top, the presence of an Alu element in an intron in antisense alone will still result in normal splicing. $Bottom, specific \, mutations \, of \, the \, Alu \, sequence \, creates \, new \, splice \, sites \, and \,$ results in exonization.



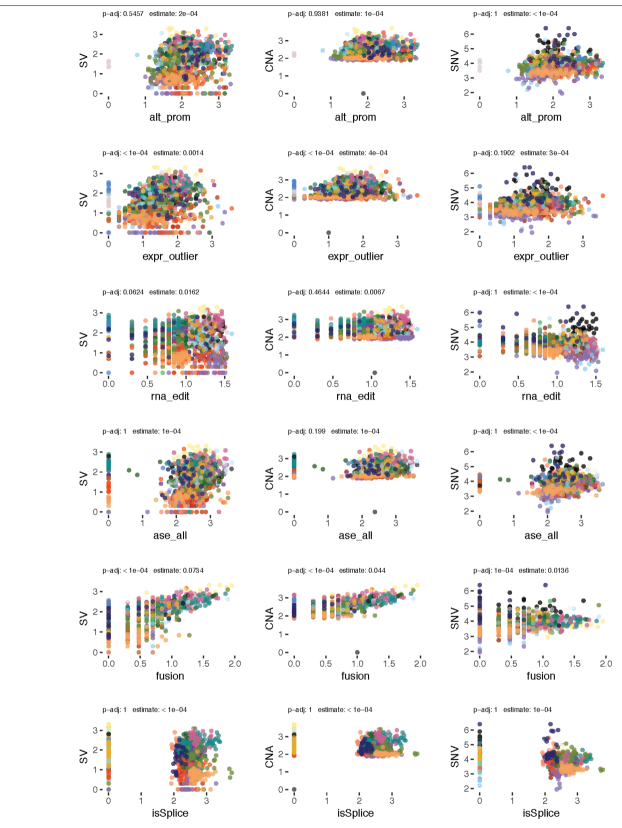
 $\label{lem:extended Data Fig. 18} Recurrent and promiscuous RNA fusions. a, \textit{Features of the 27 most recurrent in-frame or open-reading-frame-retaining fusions.} \\ Kinase column indicates whether one of the gene partners is a kinase gene b, Network with connected clusters of at least 10 genes. Genes are represented as nodes, and the size of a node is proportional to the number of gene-fusion partners. Two nodes are connected if one fusion was detected involving the$ 

two genes: an edge is coloured blue if the fusion has evidence for matched structural rearrangements and is coloured red otherwise. Nodes and connections are shown only between promiscuous genes. The colour intensity indicates whether a gene is involved more often in a fusion as a  $3^\prime$  (purple) or  $5^\prime$  (green) gene or both (white).



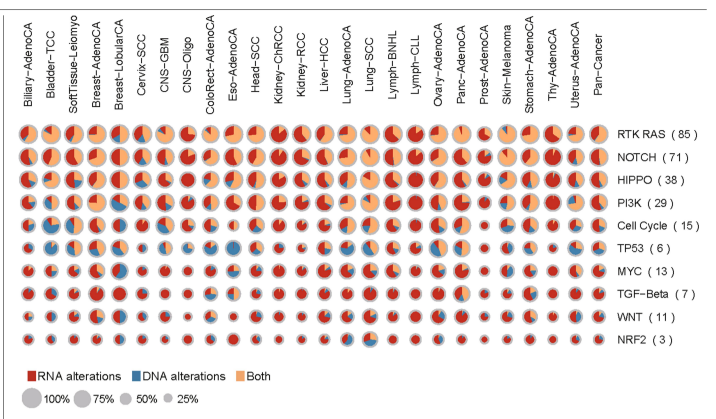
**Extended Data Fig. 19** | **Structural rearrangements associated with RNA fusions. a**, Systematic classification scheme of all gene fusions based on underlying structural variants (SVs). Numbers of fusion events of different classes are shown to the right. **b**, Schematic of examples of different types of structural-variant-supported fusions: (1) direct fusions; (2) intercomposite fusions; and (3) intracomposite fusions. Bridged fusions are shown in Fig. 3b. Only one of the possible orders of genomic arrangement is depicted in each case, with break points highlighted by thunderbolts. **c**, Supported rearrangements for composite fusions bring the fused segments of two genes

significantly closer. Natural distance indicates the native distance between two related structural variant break points. Effective distance indicates the distance between the final two break points of the intra- and intercomposite fusions.  ${\bf d}$ , The break points of structural-variant-independent fusions are typically closer than those for other interchromosomal fusions, which indicates that at least some of the structural-variant-independent fusions may occur directly at the RNA level, mediated either by trans-splicing or read-through events.



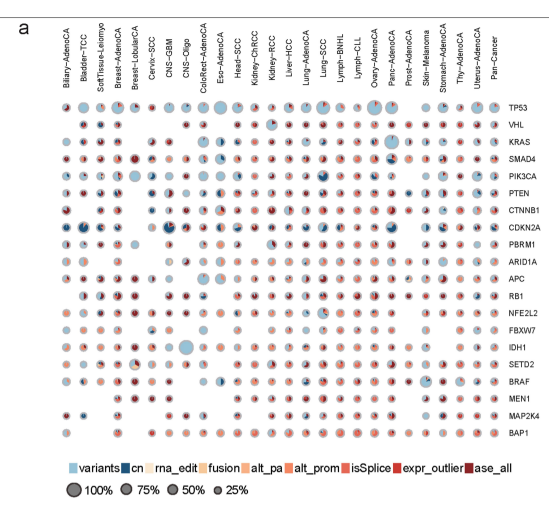
**Extended Data Fig. 20 | Correlation of the number of somatic genomic alterations with RNA alterations.** Scatter plots of  $\log_{10}$ -transformed frequency of DNA alterations versus  $\log_{10}$ -transformed frequency of RNA alterations, in which each row is a DNA alteration in the following order: structural variants, copy-number aberrations and non-synonymous variants. Each row is an RNA alteration in the following order: expression outliers, RNA

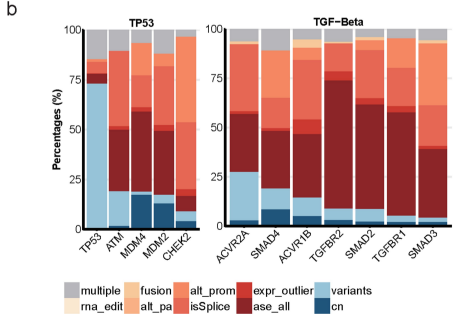
editing, ASE, fusions and splicing. Each point is a sample coloured by histotype, and its position is the log-transformed number of aberrations found in each sample. The Benjamini–Hochberg-adjusted P values are calculated from a likelihood ratio test assuming negative binomial distribution; histotype is used as a confounder.



 $\label{lem:continuous} \textbf{Extended Data Fig. 21} | \textbf{Global view of DNA} \ and \ \textbf{RNA} \ alterations \ affecting} \ \textbf{cancer pathways}. \ \textbf{Composite pie charts showing the percentages of RNA} \ alterations, \ \textbf{DNA} \ alterations \ or \ both, \ affecting sets \ of genes \ in \ well-characterized \ cancer \ pathways \ and \ known \ to \ be \ functionally \ altered \ in \ cancer.$ 

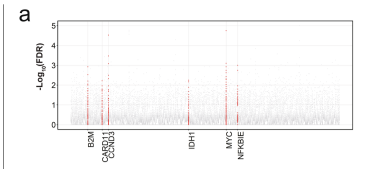
The sizes of circles represent the percentages of patients affected based on the given gene set. The columns indicate different types of cancer. The numbers in parenthesis indicate the number of genes analysed for the specific pathway.

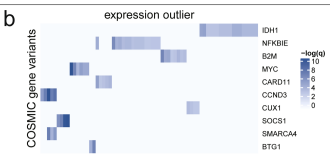


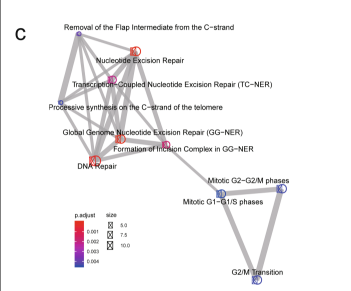


 $\label{lem:extended Data Fig. 22} Breakdown of DNA and RNA alterations of cancer genes. a, Composite pie charts showing percentages of DNA and RNA alterations for top cancer-driver genes. The 20 most significant cancer-driver genes identified by the PCAWG group in pan-cancer level are depicted, with the sizes of the pie charts indicating the percentages of patients carrying alterations in the given driver gene. The areas represent the relative$ 

percentages of patients exhibiting different alterations depicted by corresponding colours. When several types of alteration in one pathway affect the same patient, only a fraction is counted towards each type of alteration. **b**, Proportional bar plots showing the distribution of gene alterations for genes in the *TP53* and *TGFB* pathways.





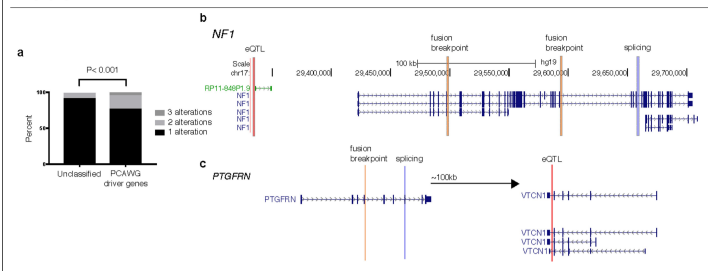


#### $\textbf{Extended Data Fig. 23} \,|\, \textbf{\textit{Trans}-} associations found by co-occurrence analyses.$

**a**, Scatter plot for association of gene expression outliers with cancer gene variants. Each dot represents an alteration pair. The *x* axis shows all COSMIC genes ordered alphabetically and the *y* axis represents the FDR-adjusted *P* values (*q* values) based on Fisher's exact tests. COSMIC genes with more than five significant associations (FDR < 5%) are coloured in red and labelled. **b**, Heat map showing the extent of associations between COSMIC gene somatic mutations and expression outliers of all genes. Each row indicates one gene,

and the colour intensity shows the significance of *trans*-association. COSMIC genes labelled to the right are ordered by the number of significant associations. Only the top 10 genes are shown. **c**, Enrichment map showing the significant (FDR  $\leq$  0.01) pathways based on the top 100 significant genes associated with B2M alterations. Colour intensity represents enrichment significance, node sizes the number of analysed genes belonging to the given pathway and edge sizes the degree of overlap between two gene sets. Only the top 10 enriched terms are shown.

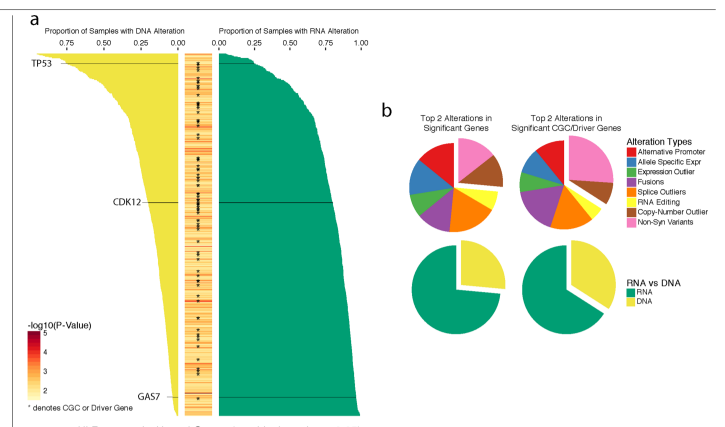
## **Article**



#### Extended Data Fig. 24 | Genes can be altered in cis by several mechanisms.

**a**, Genes with at least one type of RNA alteration that also has an associated change at the DNA-level in *cis*. Genes are either classified as a PCAWG driver

gene or not classified as a driver gene or a cancer gene from the cancer gene census.  $\mathbf{b}$ ,  $\mathbf{c}$ , Examples of a known cancer gene, NFI ( $\mathbf{b}$ ), and an unclassified gene, PTGFRN( $\mathbf{c}$ ), having heterogeneous mechanisms of alterations.



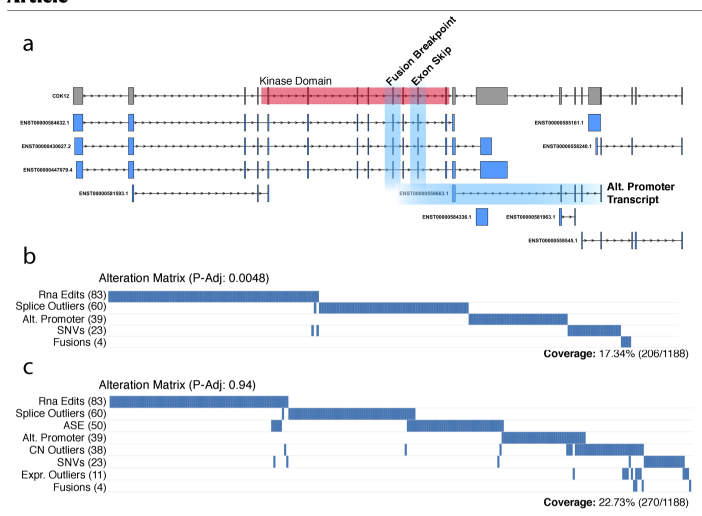
All Recurrently Altered Genes (empirical p-value < 0.05)

**Extended Data Fig. 25 | Proportion of genes with DNA or RNA alterations. a**, Full list of 731 genes that are both frequently and heterogeneously altered across both RNA- and DNA-level alterations. Yellow bars to the left indicate the proportion of samples that had DNA-level alterations, whereas green bars to the right indicate the proportion of samples with RNA-level alterations. Middle

 $column \ is \ a heat \ map \ corresponding \ to \ the -log_{10}(Pvalue). \ Asterisks \ indicate \ a \ COSMIC \ Cancer \ Gene \ Census \ (CGC) \ gene \ or \ PCAWG \ driver \ genes.$ 

 $\begin{tabular}{ll} \textbf{b}, Distribution of alteration types among all significant genes or just CGC or PCAWG driver genes. \end{tabular}$ 

## **Article**



**Extended Data Fig. 26** | **Outlier events in** *CDK12.* **a**, Fusion, splicing and alternative promoter outlier events of the RNA alterations that lead to either partial or full removal of the kinase domain in *CDK12.* **b**, All outlier events in *CDK12.* including those not contained directly within the kinase domain, across all 1,188 samples. Each column is a sample and each row is the alteration type.

Although not directly searching for mutually exclusive events across all genes, we find that CDK12 is marginally mutually exclusive in RNA editing, splicing outliers, alternative promoters, non-synonymous variants and fusions  $(4.810^{-3}, \text{unweighted WExT})$ . **c**, All alteration events that occur within CDK12 across all 1,188 samples, which is not mutually exclusive.

## Extended Data Table 1 | RNA alteration data

RNA data	Total number of gene alterations found	Mean number of gene alterations per donor
Gene expression (PCAWG)	93,481	78.69
RNA fusions	5,900	4.97
Alternative promoters	246,224	207.26
Alternative splicing	345,115	290.50
Allele-specific expression	544,664	458.47
RNA editing (with a non-synonymous change)	14486	12.19
Combined gene-centric table (DNA and RNA alterations)	1,523,098	1,282.07



Corresponding author(s): Alvis Brazma, Gunnar Raetsch, Angela N Brooks

# **Reporting Summary**

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## Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

n/a	Con	firmed
	$\boxtimes$	The $\underline{\text{exact sample size}}$ (n) for each experimental group/condition, given as a discrete number and unit of measurement
	$\boxtimes$	An indication of whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
		The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
	$\boxtimes$	A description of all covariates tested
	$\boxtimes$	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	$\boxtimes$	A full description of the statistics including <u>central tendency</u> (e.g. means) or other basic estimates (e.g. regression coefficient) AND <u>variation</u> (e.g. standard deviation) or associated <u>estimates of uncertainty</u> (e.g. confidence intervals)
		For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i> ) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable.</i>
	$\boxtimes$	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
$\boxtimes$		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
	$\boxtimes$	Estimates of effect sizes (e.g. Cohen's $d$ , Pearson's $r$ ), indicating how they were calculated
		Clearly defined error bars State explicitly what error bars represent (e.g. SD, SE, CI)

Our web collection on statistics for biologists may be useful.

#### Software and code

Policy information about availability of computer code

Data collection

Core data was collected through Pan-cancer Analysis of Whole Genomes Data Coordination center through https://dcc.icgc.org/releases/ PCAWG/

Data analysis

Core RNA-Seq alignment pipelines are available through Github/Docker: https://github.com/akahles/icgc\_rnaseq\_align, https:// hub.docker.com/r/nunofonseca/irap\_pcawg/. STAR, TopHat2, HTSeq, Kallisto, Limix, PLINK, Bedtools, Vcftools, Bcftools, Samtools, Tabix, GATK, ASEReadCounter, Lavaan, Mediate, SplAdder, SAVNet, Sv2gf

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

#### Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data

Human research participants

- A description of any restrictions on data availability

Somatic and germline variant calls, mutational signatures, subclonal reconstructions, transcript abundance, splice calls and other core data generated by the ICGC/ TCGA Pan-cancer Analysis of Whole Genomes Consortium is described here58 and available for download at https://dcc.icgc.org/releases/PCAWG. Additional information on accessing the data, including raw read files, can be found at https://docs.icgc.org/pcawg/data/. In accordance with the data access policies of the ICGC and TCGA projects, most molecular, clinical and specimen data are in an open tier which does not require access approval. To access potentially identification information, such as germline alleles and underlying sequencing data, researchers will need to apply to the TCGA Data Access Committee (DAC) via dbGaP (https:// dbgap.ncbi.nlm.nih.gov/aa/wga.cgi?page=login) for access to the TCGA portion of the dataset, and to the ICGC Data Access Compliance Office (DACO; http:// icgc.org/daco) for the ICGC portion. In addition, to access somatic single nucleotide variants derived from TCGA donors, researchers will also need to obtain dbGaP authorisation. Data derived specifically from RNA-Seq analysis can be found at https://dcc.icgc.org/releases/PCAWG/transcriptome. Subfolders contain identification and quantification of alternative promoter usage, alternative splicing, RNA fusions, gene expression, transcript-level expression, and RNA editing. Identified eQTLs are in https://dcc.icgc.org/releases/PCAWG/transcriptome/eQTL and a binarized table indicating all RNA and DNA alterations for each gene can be found in the subfolder https://dcc.icgc.org/releases/PCAWG/transcriptome/recurrence\_analyses/. Additionally, QC metrics and metadata are also included.

Field-specific reporting					
Please select the best fit for your research. If you are not sure, read the appropriate sections before making your selection.					
Life sciences	Behavioural & social sciences				
For a reference copy of t	the document with all sections, see <u>nature.com/authors/policies/ReportingSummary-flat.pdf</u>				
Life scier	nces				
Study design	1				
All studies must dis	sclose on these points even when the disclosure is negative.				
Sample size	This study represents the analysis of 1,188 donors from the Pan-Cancer Analysis of Whole Genomes project. The sample size was limited to the availability of RNA-Seq data from matched donors with WGS data from the PCAWG project.				
Data exclusions	A larger set of 2,217 RNA-Seq libraries were initially collected and data were excluded after QC analysis. The QC criteria was standard and preestablished before excluding data.				
Replication	Reproducibility of the analysis is ensured through data-sharing and code-sharing. Unfortunately, at the time of the analysis there were no approrpriate datasets to use for replication studies of associations, since this is one of the largest collections of pan-cancer whole genomes and matched transcriptomes. For the somatic eQTL analysis, there were some related studies that came to similar conclusions and are noted in the Supplementary Information.				
Randomization	Cancer histotypes were defined by the PCAWG Pathology and Clinical Correlates Working Group based on tumor histology. These tumor subtypes were accounted for as covariates in all applicable analyses of association.				
Blinding	Blinding was not relevant to our study as it was essential to understand underlying confounding variables in our associations, such as tumor subtype, sex, etc.				
Materials &	experimental systems				
Policy information	about <u>availability of materials</u>				
n/a Involved in t	he study				
Unique m	naterials				
Antibodies Antibodies					
	ic cell lines				
Research	animals				

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| reporting summary

March 2018

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N/IAthad	-specific re	norting
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n/a Involved in the study

ChIP-seq

Flow cytometry

Magnetic resonance imaging