

ARTICLE

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The Cancer Genome Atlas Research Network

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Integrated genomic and molecular characterization of cervical cancer

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Cervical cancer remains one of the leading causes of cancer-related deaths worldwide. Here we report the extensive molecular characterization of 228 primary cervical cancers, the largest comprehensive genomic study of cervical cancer to date. We observed striking APOBEC mutagenesis patterns and identified *SHKBPI*, *ERBB3*, *CASP8*, *HLA-A*, and *TGFBR2* as novel significantly mutated genes in cervical cancer. We also discovered novel amplifications in immune targets *CD274/PD-L1* and *PDCD1LG2/PD-L2*, and the *BCAR4* lncRNA that has been associated with response to lapatinib. HPV integration was observed in all HPV18-related cases and 76% of HPV16-related cases, and was associated with structural aberrations and increased target gene expression. We identified a unique set of endometrial-like cervical cancers, comprised predominantly of HPV-negative tumors with high frequencies of *KRAS*, *ARID1A*, and *PTEN* mutations. Integrative clustering of 178 samples identified Keratin-low Squamous, Keratin-high Squamous, and Adenocarcinoma-rich subgroups. These molecular analyses reveal new potential therapeutic targets for cervical cancers.

Cervical cancer accounts for 528,000 new cases and 266,000 deaths worldwide each year, more than any other gynecologic tumor¹. Ninety-five percent of cases are caused by persistent infections with carcinogenic human papillomaviruses (HPV)². Effective prophylactic vaccines against the most important carcinogenic HPV types are available, but uptake remains poor. Although early cervical cancer can be treated with surgery or radiation, metastatic cervical cancer is incurable and new therapeutic approaches are needed³.

While most HPV infections are cleared within months, some persist and express viral oncogenes that inactivate p53 and Rb, leading to increased genomic instability, accumulation of somatic mutations, and in some cases integration of HPV into the host genome⁴. The association with cancer risk and histological subtypes varies substantially among carcinogenic HPV types, but the reasons for these differences are poorly understood. Further, clinically relevant cervical cancer patient subgroups have yet to be identified. Presented here is a comprehensive study of invasive cervical cancer conducted as part of The Cancer Genome Atlas (TCGA) project, with a focus on identifying novel clinical and molecular associations as well as functionally altered signaling pathways that may drive tumorigenesis and serve as prognostic or therapeutic markers.

Samples and clinical data

Primary frozen tumor tissue and blood were obtained from women with cervical cancer without prior chemotherapy or radiotherapy (Supplemental Information S1 and Supplemental Tables 1 and 2). DNA, RNA, and protein were processed as previously described⁵ (Supplementary Information S1, S3, S5, and S8). Mutations were called for 192 samples (Extended Set), while all other platform (aside from protein) and integrated analyses were performed on a subset of 178 samples (Core Set). Protein levels were measured on 155 samples (119 samples from both the Core and Extended Sets plus 36 additional samples). The total number of non-overlapping samples in these three sets was 228 (Extended Data Fig. 1a). Of the 178 Core Set samples, surgery was the primary treatment in 121 cases, median follow-up was 17 months, and 145 patients were alive at the time of last follow-up. A committee of expert gynecologic pathologists reviewed most cases (Supplemental Information S1 and Extended Data Fig. 1b-g). The Core

Set included 144 squamous cell carcinomas, 31 adenocarcinomas, and 3 adenosquamous cancers.

Somatic genomic alterations

Whole exome sequencing was performed on 192 Extended Set tumor-blood normal pairs. All samples had at least 32 Mbp of target exons covered with a median of 49× (range: 7–351×) coverage for tumor samples and 47× (range: 9–341×) coverage for normal samples. Collectively, the samples harbored 43,324 somatic mutations, including 24,551 missense, 2,470 nonsense, 9,260 silent, 5,841 non-coding, 535 splice site, 74 nonstop, 475 frameshift indels, and 118 in-frame indels. Eleven tumors with outlier mutation frequencies (>600 per sample) were classified as “hypermutant.” The aggregate mutation density was 4.04 mutations per megabase across all tumors, and 2.53 when the hypermutant tumors were excluded.

Fourteen significantly mutated genes (SMGs) with false discovery rates (FDR) <0.1 were found using the MutSig2CV⁶ algorithm (Supplemental Table 4). We identified *SHKBPI*, *ERBB3*, *CASP8*, *HLA-A*, and *TGFBR2* as novel SMGs in cervical cancer, and confirmed *PIK3CA*, *EP300*, *FBXW7*, *HLA-B*, *PTEN*, *NFE2L2*, *ARID1A*, *KRAS*, and *MAPK1* which have been previously reported (Fig. 1, Extended Data Fig. 2a-g, and Supplemental Fig. S6)^{7,8}. Supplemental Table 4 shows the comparison of SMGs identified in the TCGA and Ojesina *et al.* datasets. Mutations in 7 of the 14 SMGs in the TCGA set were present in at least one squamous cell carcinoma and one adenocarcinoma; however, mutations in *HLA-A*, *HLA-B*, *NFE2L2*, *MAPK1*, *CASP8*, *SHKBPI*, and *TGFBR2* were found exclusively in squamous tumors.

PIK3CA harbored mostly activating helical domain mutations E542K and E545K, with a marked relative decrease in mutations elsewhere in the gene (Extended Data Fig. 2g). This observation resembles findings in bladder cancer⁹ and HPV-positive head and neck squamous cell cancers (HNSC)¹⁰, but it differs from observations in breast and most other cancers¹¹. The underlying nucleotide substitution pattern in the E542K and E545K mutations is associated with mutagenesis by a subclass of APOBEC cytidine deaminases^{8,12–15}, with 150 of 192 exomes displaying statistically significant ($q < 0.05$) enrichment (up to 6-fold) for the APOBEC signature. Further, the APOBEC mutation load correlated strongly with the total number of mutations per sample (Extended

*Lists of participants and their affiliations appear at the end of the paper.

Data Fig. 2h), suggesting that APOBEC mutagenesis is the predominant source of mutations in cervical cancers.

We found an average of 88 somatic copy number alterations (SCNAs) per tumor, fewer than in HNSC, ovarian, and serous endometrial carcinomas but more than in endometrioid endometrial carcinomas^{10,16,17}. GISTIC2.0 analysis (threshold $q < 0.25$) revealed 26 focal amplifications and 37 focal deletions along with 23 recurrently altered whole arms (Extended Data Fig. 3c and Supplemental Table 7). Novel recurrent focal amplification events were identified (in genomic order) at 7p11.2 (*EGFR*; 17%), 9p24.1 (*CD274*, *PDCD1LG2*; 21%), 13q22.1 (*KLF5*; 18%), and 16p13.13 (*BCAR4*; 20%). Other previously reported amplification events occurred at 3q26.31 (*TERC*, *MECOM*; 78%), 3q28 (*TP63*; 77%), 8q24.21 (*MYC*, *PVT1*; 42%), 11q22.1 (*YAP1*, *BIRC2/3*; 17%), and 17q12 (*ERBB2*; 17%). Novel recurrent deletions were identified at 3p24.1 (*TGFBR2*; 36%) and 18q21.2 (*SMAD4*; 28%), in addition to previously identified deletions at 4q35.2 (*FAT1*; 36%) and 10q23.31 (*PTEN*; 31%). A CN high cluster largely contained squamous tumors with amplification events involving 11q22 (*YAP1*, *BIRC2/3*) and 7p11.2 (*EGFR*), while the CN low cluster included most adenocarcinomas and was enriched for tumors with deletions in *TGFBR2* and *SMAD4* and gains in *ERBB2* and *KLF5* (Extended Data Fig. 3a, b). Notably, both groups had amplifications involving *CD274* (PD-L1) and *PDCD1LG2* (PD-L2) that correlated significantly ($p < 0.0001$) with expression of two key immune cytolytic effector genes, granzyme A and perforin¹⁸ (Extended Data Fig. 3d). This highlights the potential of immunotherapeutic strategies for a subset of cervical cancers.

Structural rearrangements were identified through analysis of RNA-seq (Core Set, $n = 178$) and whole genome sequencing (WGS) data with low-pass ($n = 50$) and deep ($n = 19$) coverage. Both RNA-seq and WGS detected 22 putative structural rearrangements in 14 patients (Supplemental Table 8). In total, 26 recurrent fusions were found (Supplemental Table 9, with examples in Extended Data Fig. 4d). RNA-seq analysis revealed 4 cases harboring 16p13 *ZC3H7A-BCAR4* gene fusions, with exon 1 of *ZC3H7A* linked to the last exon of *BCAR4*. Whole genome sequencing revealed tandem duplication and copy number gain of *BCAR4* on chromosome 16p13.13 (Extended Data Fig. 4c). *BCAR4* is a metastasis-promoting lncRNA that enhances cell proliferation in estrogen-resistant breast cancer by activating the HER2/3 pathway. Lapatinib, an EGFR/HER2 inhibitor, counteracts *BCAR4*-driven tumor growth *in vitro*, and warrants evaluation as a possible therapeutic agent in *BCAR4*-positive cervical cancer¹⁹.

Integrated analysis of molecular subgroups and pathways

Integration of copy number, methylation, mRNA, and miRNA data using iCluster²⁰ highlighted the molecular heterogeneity of cervical carcinomas. Three clusters were identified that largely corresponded to mRNA clusters (Supplemental Fig. S9): a squamous cluster with high expression of keratin gene family members (Keratin-high), another squamous cluster with lower expression of keratin genes (Keratin-low), and an Adenocarcinoma-rich cluster (Adenocarcinoma). Keratin-high and Keratin-low clusters included 133 of 144 squamous cell carcinomas and the Adenocarcinoma cluster contained 29 of 31 adenocarcinomas (Fig. 2). *KRAS* ($p = 9.7e-5$), *ERBB3* ($p = 2.6e-3$), and *HLA-A* ($p = 0.03$) mutations were significantly associated with iClusters, with *KRAS* mutations absent from the Keratin-high cluster and *HLA-A* mutations missing in the Adenocarcinoma cluster (Fig. 2). Members of the *SPRR* and *TMPRSS* cornification gene families and the SMGs *ARID1A* ($p = 0.02$), *NFE2L2* ($p = 6.9e-6$), and *PIK3CA* ($p = 0.01$) were differentially expressed between Keratin-low and Keratin-high clusters (Extended Data Fig. 4b).

Unsupervised hierarchical clustering of variable DNA methylation probes produced three groups (Extended Data Fig. 5a), including a small “CpG island hypermethylated” (CIMP-high) cluster, a CIMP-intermediate cluster, and a CIMP-low cluster that were associated with an epithelial-mesenchymal transition (EMT) mRNA score (Extended

Data Fig. 5b)^{10,21}. Most of the cases in the Adenocarcinoma iCluster were CIMP-high, while the other iClusters contained a mixture of CIMP-intermediate and CIMP-low samples (Fig. 2). Comparing all cervical carcinomas to 120 normal samples drawn from 12 TCGA projects, we identified 1026 epigenetically silenced genes that were methylated to a greater extent in cancers than in normal tissues, including several zinc-finger (*ZNF*), protease (*ADAM*, *ADAMTS*), and collagen (*COL*) genes (Supplemental Tables 11 and 12).

Unsupervised clustering resulted in 6 miRNA clusters associated with iClusters ($p = 1.7e-19$) (Extended Data Fig. 6a). Samples from the Adenocarcinoma iCluster almost exclusively overlapped with miRNA cluster 5, and were characterized by high expression of miR-375 and low expression of miR-205-5p and miR-944 (Supplemental Table 31). Expression levels of tumor suppressors miR-99a-5p and miR-203a were significantly higher in Keratin-high cluster samples than Keratin-low cluster samples (Supplemental Table 31; $p = 0.01$ and 0.008, respectively). Among miRNAs with significant and functionally validated gene and protein anti-correlations²², one large subnetwork involved miR-200-family and other miRNAs with expression patterns anti-correlated with those of the EMT-related transcription factors *ZEB1*, *ZEB2*, and *SNAI2*, the Hippo and p73 transcriptional co-factor *YAP1*, the receptor tyrosine kinases (RTKs) *ERBB2*, *ERBB3*, and *AXL*, and the hormone receptor *ESR1* (Extended Data Fig. 6b, Supplemental Fig. S17, Supplemental Fig. S18, and Supplemental Table 15).

Reverse Phase Protein Array (RPPA) analysis of 155 samples with 192 antibodies (Extended Data Fig. 1a and Supplemental Table 17) identified three clusters significantly associated with iClusters ($p = 1.8e-4$) and EMT mRNA score (Fig. 3a, c, d and Supplemental Table 16). EMT cluster samples were enriched in the Keratin-low iCluster, while PI3K/AKT and Hormone cluster samples were enriched in the Keratin-high and Adenocarcinoma iClusters, respectively, suggesting distinct pathway activation across integrated cervical cancer subtypes. Differential expression levels of Phospho-MAPK, Phospho-EGFR (Y1068), Phospho-Src (Y416), IGFBP2, and TIGAR between Keratin-high and Keratin-low iClusters suggest diverse activation patterns of RTK, MAPK, PI3K, and metabolic signaling pathways that may underlie the molecular diversity of cervical squamous cancers (Fig. 2).

The core members of each RPPA cluster with the highest silhouette width (> 0.02 , $n = 115$) were associated with five-year survival (Fig. 3b; $p = 6.1e-4$), with the EMT group exhibiting worse outcome. Interestingly, this was the only platform where clusters associated with outcomes (Supplemental Figs. S8, S9, S12, and S22; Supplemental Information S6). Samples in the EMT cluster exhibited high “reactive” pathway scores (Supplemental Fig. S20)¹¹, illustrating for the first time in cervical cancer the presence of a subset of stromal “reactive” tumors with high expression of Caveolin-1, MYH11, and Rab11, which also appear in other diseases (Supplemental Table 16)²³. YAP was the most significantly differentially expressed protein distinguishing EMT cluster samples from all others (Supplemental Table 18; $p = 1.7e-15$) and *YAP1* was significantly amplified in the EMT cluster samples compared with the Hormone ($p = 1.1e-5$) and PI3K/AKT cluster ($p = 6.4e-4$) samples. Regulation of the EMT-related molecules YAP and *ZEB1*²⁴⁻²⁶ may also be driven by significantly lower expression levels of miR-200a-3p in the EMT cluster samples compared with other RPPA cluster samples (Extended Data Fig. 6b and Extended Data Fig. 7a; $p = 3.8e-3$). These results highlight potential roles for YAP and reactive stroma in the context of EMT-regulated cervical cancer progression.

The Mutual Exclusivity Modules in cancer (MEMo) algorithm²⁷ uses somatic mutation and copy number data to identify oncogenic networks with mutually exclusive genomic alterations. Since miR-200a and miR-200b (miR-200a/b) expression were negatively correlated with EMT mRNA scores (Extended Data Fig. 7b, d), we used MEMo to examine alterations in miR-200a/b and EMT networks and found a potential link between TGF β pathway and miR-200a/b alterations in regulating EMT^{28,29}. Deletions and mutations affecting the receptor gene *TGFBR2*, the modulating genes *CREBBP* and *EP300*,

and the transcription factor *SMAD4* all likely impinge on growth suppressive and pro-apoptotic functions driven by TGF β (Fig. 4c) and were observed in 30% of squamous cell carcinomas (Fig. 4d). Tumors with both hypermethylation and downregulation of miR-200a/b (referred to as altered) were restricted to squamous cell carcinomas, were enriched in the Keratin-low iCluster (Fig. 4d and Extended Data Fig. 8; $p = 0.001$ for both miR-200a and miR-200b), showed significant upregulation of both *ZEB1* and *ZEB2* (Extended Data Fig. 9a-d), and were mutually exclusive with TGF β signaling pathway alterations (Fig. 4d). Importantly, samples with altered miR-200a/b exhibited higher EMT mRNA scores than unaltered samples, while there was no significant difference between samples with or without TGF β pathway alterations (Fig. 4d and Extended Data Fig. 7c, e). These findings highlight potential treatment approaches for this subgroup of cervical cancer patients, as targeting EMT may render tumors more sensitive to small molecule inhibitors and cytotoxic chemotherapy^{21,30,31}.

MEMo analysis also showed differences in therapeutically-relevant RTK, PI3K, and MAPK pathway alterations across cervical cancers. MEMo identified mutual exclusivity modules involving alterations within both the PI3K and MAPK pathways (Supplemental Table 27; adjusted $p = 0.06$); however, there was a strong tendency for co-occurrence of *ERBB2* and *ERBB3* alterations within adenocarcinomas ($p < 0.001$, log odds-ratio > 3), indicating that a subset of these tumors may exhibit aberrant HER3 signaling through interactions between mutant HER3 and activated HER2 and therefore could potentially benefit from HER2- and HER3-targeted therapies (Fig. 4a, b)³². Although not statistically significant, aberrations in *PIK3CA* also tended to co-occur with *PTEN* somatic mutations and deletions ($p = 0.078$, log-odds ratio = 0.71), which is similar to copy number-low endometrial tumors and suggests potential therapeutic benefit from PI3K pathway targeting agents¹⁷.

PARADIGM^{33,34}, which integrates copy number, RNA-seq, and pathway interaction data, showed markedly different pathway activation profiles between squamous carcinomas and adenocarcinomas (Extended Data Fig. 10 and Supplemental Fig. S48). PARADIGM identified higher inferred activation of p53, p63, p73, AP-1, MYC, HIF1A, FGFR3, and MAPK signaling as key distinguishing signaling features of squamous cell carcinomas, similar to other squamous cancers³⁵. In contrast, adenocarcinomas exhibited higher inferred activation of ER α , FOXA1, FOXA2, and FGFR1 pathways (Extended Data Fig. 10, Supplemental Fig. S25, Supplemental Fig. S48, and Supplemental Table 18). Possible underlying mechanisms for ER α upregulation may stem from the expression of miR-193b-3p, a direct regulator of *ESR1* that was significantly downregulated in adenocarcinomas compared with squamous carcinomas (Fig. 2, Extended Data Fig. 6, and Supplemental Table 14; $p = 0.04$), or from estrogen signaling in stromal cells³⁶.

Cross-cancer analysis

To evaluate the relationship of cervical cancer subtypes with endometrial cancer, an adjacent cancer site with hormone-related carcinogenesis, and HNSC, a subset of which is caused by HPV, hierarchical clustering of cervical, uterine corpus endometrial (UCEC)¹⁷, and HNSC¹⁰ mRNA expression data was performed. Three major groups were observed, with Cluster 1 including all UCEC samples and most cervical adenocarcinomas and characterized by overexpression of hormone receptor genes *ESR1* and *PGR* (Extended Data Fig. 4a). Cluster 2 included predominantly squamous cervical carcinomas and 23/27 HPV-positive HNSC samples. Cluster 3 included few cervical cancers and the remaining HNSC cancers, which were mostly HPV-negative. This highlights the similarity of HPV-related squamous cancers at different anatomical sites.

Since a subset of cervical cancers clustered with endometrial samples, a gene expression classifier was developed to predict whether carcinomas were cervical or endometrial (Supplemental Information S5). We classified 8 of 178 (4.5%) cervical cancer samples

as endometrial-like (UCEC-like) cancers, which were confirmed to be cervical cancers by study pathologists (Extended Data Fig. 1f, g). These tumors included 7 of 9 HPV-negative cancers and 5 of the 8 were adenocarcinomas. Six UCEC-like cancers were in the Adenocarcinoma iCluster and 2 were in the Keratin-low iCluster. Despite their low number, the UCEC-like tumors accounted for 33%, 27%, and 20% of mutations in *ARID1A*, *KRAS*, and *PTEN*, respectively. They were associated with the RPPA Hormone and miRNA C6 clusters, and all but one sample was CIMP-low and CN low (Supplemental Table 1).

HPV genotypes, variants, and integration

Of 178 Core Set tumors, 169 (95%) were HPV-positive, 120 (67%) had alpha-9 (A9) types (103 HPV16), 45 (25%) had alpha-7 (A7) types (27 HPV18), and 9 (5%) were HPV-negative (Supplemental Table 3). HPV variants were predominantly European (137 of 169, 81% A variants), and there was a significant association of non-European HPV16 variants with cervical adenocarcinomas (Supplemental Table 3; OR 5.3, $p = 3e-3$). All HPV-positive cancers had detectable expression of HPV E6 and E7 oncogene mRNAs, which encode proteins that inhibit p53 and Rb function, respectively^{37,38}. Interestingly, HPV18 cancers had significantly higher levels of unspliced/spliced transcripts encoding active E6 oncoprotein than the HPV16 cancers (Extended Data Fig. 11a; $p = 2e-10$), suggesting different functional implications of E6 and E7 in cancers associated with different HPV genotypes.

HPV A7 types were enriched in Keratin-low and Adenocarcinoma iClusters ($p = 5e-4$). Most HPV clade A7 tumors were CIMP-low, and HPV-negative tumors formed a distinct subgroup within the CIMP-low cluster with a significantly lower mean promoter methylation level than other samples in that cluster (Extended Data Fig. 5a; $p = 5e-3$). Samples with the highest rate of silencing were HPV-positive adenocarcinomas, particularly those related to A9 types (t-test p -values < 0.001). Functional Epigenetic Module (FEM; Supplemental Information S13) analysis³⁹, which integrates DNA methylation and gene expression data using protein-protein-interaction networks, identified inverse correlations between methylation and gene expression in HPV-positive vs. HPV-negative cervical cancers and HPV-positive ($n = 36$) vs. HPV-negative ($n = 243$) HNSCs. The analysis revealed 12 statistically significant subnetworks for cervical cancer and 11 for HNSCs, with one common subnetwork centered around Forkhead Box A2 (*FOXA2*) (Supplemental Table 19 and Supplemental Fig. S32). miR-944, miR-767-5p, and miR-105-5p were the most differentially expressed miRNAs between HPV-positive and HPV-negative samples (Supplemental Fig. S14e). miR-944 expression was also significantly higher while miR-375 expression was significantly lower in HPV16-positive squamous cancers compared with HPV18-positive squamous cancers (Supplemental Fig. S14d). Interestingly, HPV-negative cancers displayed a significantly higher EMT mRNA score and a lower frequency of the APOBEC mutagenesis signature compared with HPV-positive tumors (Extended Data Fig. 11b and Supplementary Figure S27; $p = 0.02$ and $p = 0.004$, respectively).

PARADIGM was used to evaluate molecular pathways differentially activated in squamous samples with A7 and A9 HPV infections. We observed higher inferred activation of p53 and p63 signaling and lower FOXA1 signaling in tumors infected with A9 types (Fig. 5a and Supplemental Fig. S23a). Higher *SFN* pathway activation was also observed for A9-positive tumors, which is consistent with the low methylation and high gene expression patterns of *SFN* revealed by FEM analysis (Fig. 5a and Supplemental Table 19). Interestingly, the *SFN*-encoded Stratifin/14-3-3 σ adapter protein has previously been associated with epithelial immortalization and squamous cell cancers^{40,41}, altered p53 pathway activation⁴², and Wnt-mediated β -catenin signaling⁴³.

Viral-cellular fusion transcripts indicating integration of HPV into the host genome were observed in 141 of 169 (83%) HPV-positive cancers, including all HPV18-positive cancers. Of these 141 cases, 90 (64%) had a single HPV integration event, 35 had two events,

and 16 had three or more events (totaling 220 unique integration events) (Supplemental Table 3). HPV integration events affected all chromosomes, including some previously described hotspots such as 3q28 and 8q24 (Fig. 5b)⁴⁴. Genomic loci affected by integration were characterized by increased SCNAs ($p = 6.9e-13$ for HPV16 and $p = 0.058$ for HPV18) and increased gene expression ($p = 1.6e-11$ for HPV16 and $p = 0.011$ for HPV18) (Extended Data Fig. 11c, d). One hundred fifty-three (70%) fusion transcripts included known or predicted genes, while the remainder included intergenic regions (Fig. 5b and Supplemental Table 3).

Conclusion

Through comprehensive molecular and integrative profiling, we identified novel genomic and proteomic characteristics that subclassify cervical cancers. Integrated clustering identified Keratin-low squamous, Keratin-high squamous, and Adenocarcinoma-rich clusters defined by different HPV and molecular features (Extended Data Fig. 8). *ERBB3*, *CASP8*, *HLA-A*, *SHKBP1*, and *TGFBR2* were identified as SMGs for the first time in cervical cancer, with *ERBB3* (HER3) immediately applicable as a therapeutic target. Notably, we report amplifications and fusion events involving the *BCAR4* gene for the first time in cancer, which can be targeted indirectly by lapatinib. Further, we identified amplifications in *CD274* and *PDCD1LG2*, two genes that encode for well-known immunotherapy targets. A set of endometrial-like cervical cancers comprised predominantly of HPV-negative tumors and characterized by mutations in *KRAS*, *ARID1A*, and *PTEN* was discovered, with *PTEN* and potentially *ARID1A* proteins serving as therapeutic targets. Importantly, over 70% of cervical cancers exhibited genomic alterations in either one or both of the PI3K/MAPK and TGF β signaling pathways (Extended Data Fig. 9e), illustrating the potential clinical significance of therapeutic agents targeting members of these pathways. For the first time, we report distinct molecular pathways activated in cervical carcinomas caused by different HPV types, highlighting the biologic diversity of HPV.

Together, these findings provide insight into the molecular subtypes of cervical cancers and rationales for developing clinical trials to treat populations of cervical cancer patients with distinct therapies.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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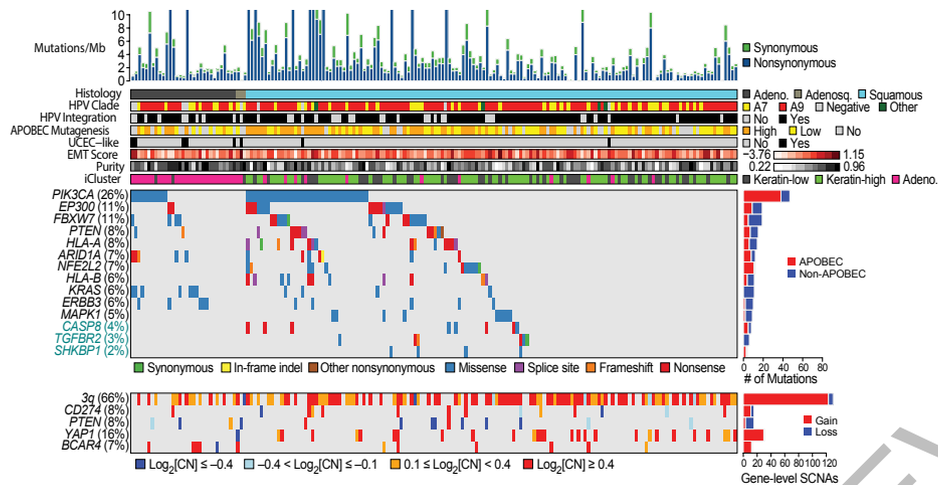


Figure 1 | Somatic alterations in cervical cancer and associations with molecular platform features. CESC samples are ordered by histology and mutation rate (top panel), clinical and molecular platform features (second panel), significantly mutated genes (SMGs; third panel), and select somatic copy number alterations (SCNAs; fourth panel) are presented. SMGs are ordered by the overall mutation frequency and

color-coded by mutation type. Novel SMGs identified in squamous cell carcinomas are labeled in turquoise text. The number of APOBEC signature mutations (red) and other mutations (blue) present in every SMG is plotted to the right of the SMG panel and the number of gene level SCNAs across all genes is plotted as gain (red) and loss (blue) to the right of the SCNA panel.

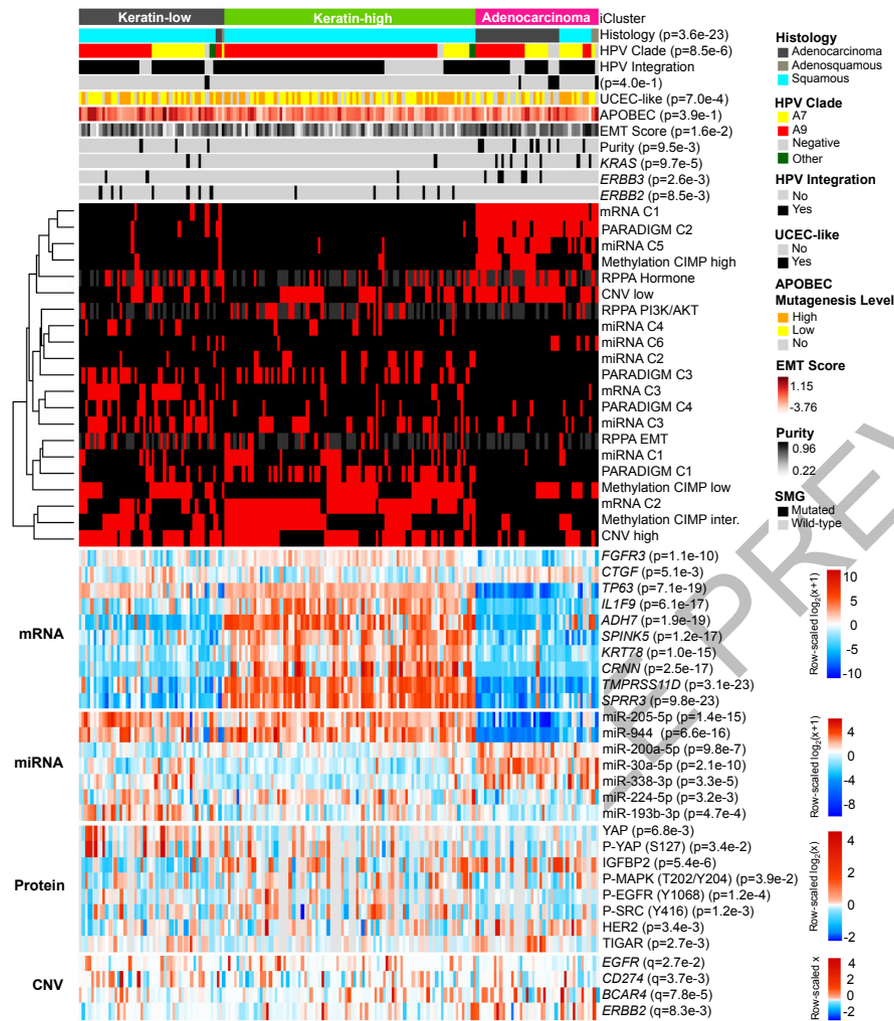


Figure 2 | Multiplatform integrative clustering of cervical cancers. Integrative clustering of 178 Core Set cervical cancer cases using mRNA, methylation, miRNA, and copy number (CNV) data identifies two squamous carcinoma-enriched groups (Keratin-low and Keratin-high) and one adenocarcinoma-enriched group as shown in the feature bars. Features presented include histology, HPV clade, HPV integration status, UCEC-like status, APOBEC mutagenesis level, mRNA EMT score, tumor purity, and three SMGs that are significantly associated across the three iClusters (*ERBB2* is presented for comparison purposes with its family member *ERBB3*). The cluster of cluster panel displays subtypes defined

independently by mRNA, miRNA, methylation, reverse phase protein array (RPPA), CNV, and PARADIGM data. Black indicates that the sample is not represented in the cluster, and gray represents data not available. The bottom heatmap panel shows select mRNAs, miRNAs, proteins, and CNVs that are either significantly associated with iCluster groups or identified as markers in other analyses. The heatmap color scale bar represents the scale for the features presented in the heatmap panel with a breakpoint of zero represented by white. APOBEC Mut., APOBEC Mutagenesis; inter., intermediate.

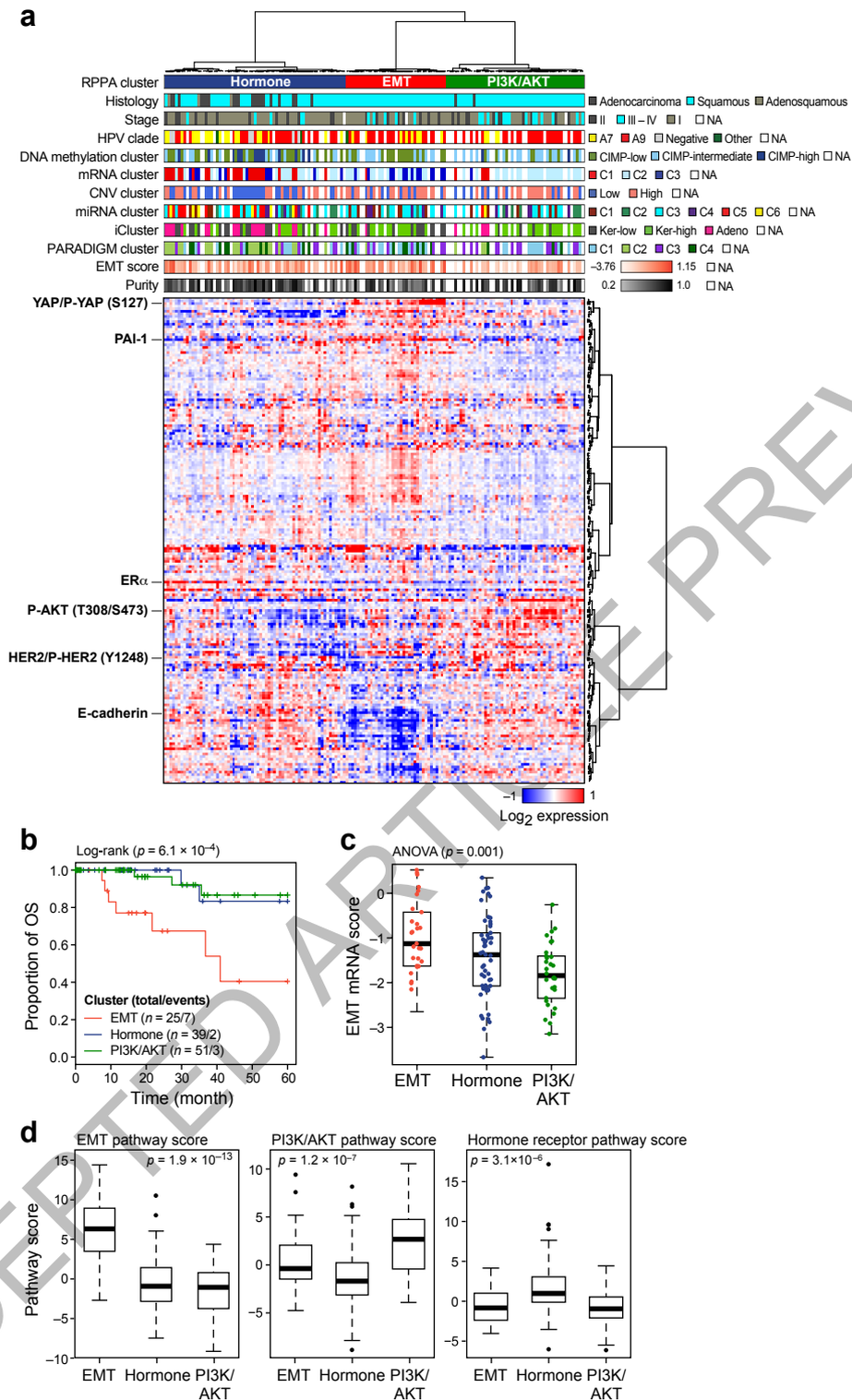


Figure 3 | Proteomic landscape of cervical cancer. a, Clustered heatmap of samples (columns) and 192 antibodies (rows) for 155 samples (112 overlap with the Core Set of 178; see Extended Data Fig. 1a). Clusters presented from left to right include Hormone (dark blue), EMT (red), and PI3K/AKT (green). A subset of proteins differentially expressed between the clusters is highlighted. Clinical and molecular feature tracks are shown for those features which were significantly associated with RPPA clusters ($p < 0.05$). Correlation between RPPA clusters and other categorical variables were detected by Chi-Squared test, while correlations with continuous variables were examined using the non-parametric Kruskal-Wallis test. In the heatmap blue color represents downregulated

expression, red represents upregulated expression, and white represents no change in expression. NA represents data not available. **b**, Five-year Kaplan-Meier survival curves and log-rank test's p-value comparing overall survival (OS) across all RPPA clusters using 115 Silhouette Width Core samples (Silhouette Core; see Supplemental Information S8). **c**, EMT mRNA score levels were calculated for all samples and compared across RPPA clusters. A significant p-value is presented for a one-way ANOVA analysis. **d**, Pathway scores for EMT, hormone receptor, and PI3K/AKT signaling pathways are presented for all RPPA clusters (x-axis), with significant pathway score differences between the clusters measured by Kruskal Wallis test.

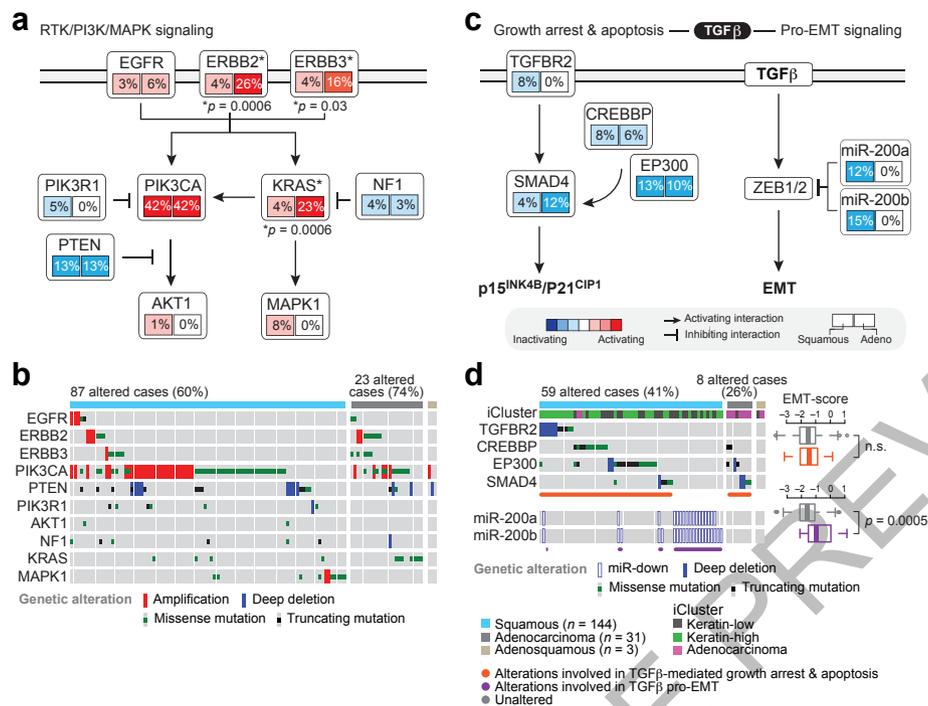


Figure 4 | Mutual exclusivity of somatic alterations within the PI3K/MAPK and TGFβR2 pathways. **a**, Multiple alterations affect receptor tyrosine kinase (RTK), AKT, and MAPK signaling in both squamous cell and adenocarcinoma cases. A schematic diagram of the pathways is shown for altered genes along with percentage of alteration in squamous cell and adenocarcinoma cases. Significant ($p < 0.05$) Student's *t*-test *p*-values for alteration frequency differences between squamous cell and adenocarcinomas are listed at the gene level, with genes marked with an asterisk (*). **b**, Distinct types of alterations (amplification, deletion, missense mutation, and truncating mutation) affect genes in these

pathways in each sample (columns). **c**, TGFβ signaling is frequently altered in cervical tumors. Alterations in this pathway are divided between those likely impinging on TGFβ tumor suppressive functions and those affecting the TGFβ-driven EMT program. Legend also corresponds to layout in panel **a**. **d**, Samples with alterations targeting TGFβ tumor suppressive functions do not show significantly different EMT scores compared with all other samples (*n.s.* = not significant); however, samples with low expression/high methylation of miR-200a/b have significantly higher EMT scores than all other samples. miR-down: met double-threshold of methylated and downregulated as described in Methods.

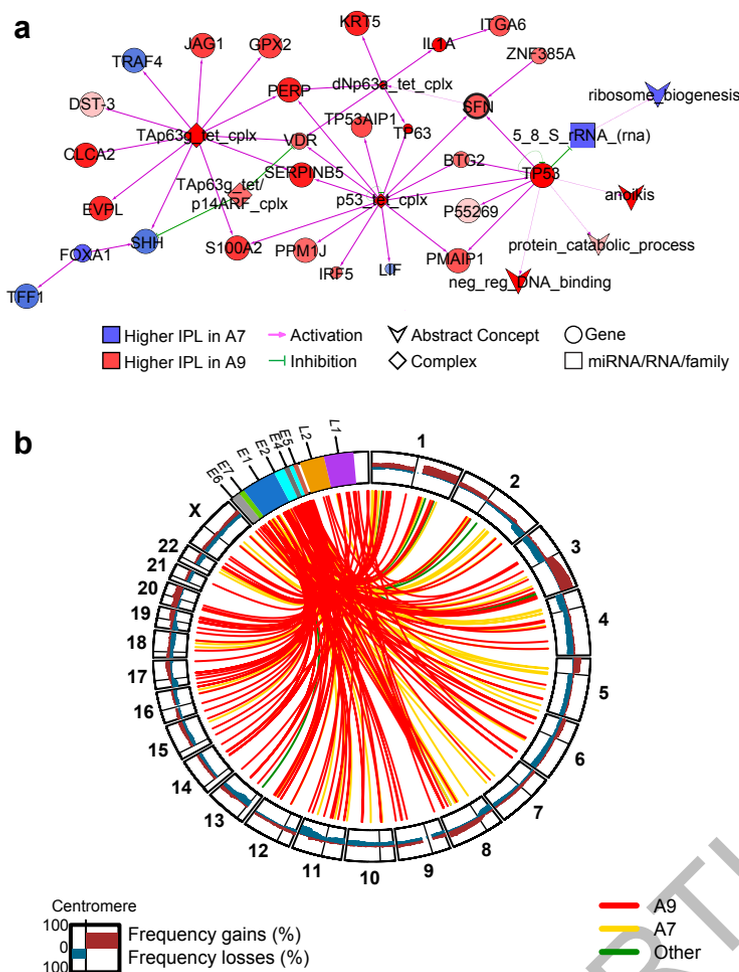


Figure 5 | HPV integration and differential pathway activation between HPV subtypes. a, Cytoscape display of the largest interconnected regulatory network of PARADIGM integrated pathway level (IPL) features showing differential inferred activation between HPV A9 and A7 squamous carcinomas ($n = 101$ and $n = 35$, respectively). Node color and intensity reflect the level of differential activation. Node size represents level of significance. Regulatory nodes with at least 5 downstream targets are highlighted in bold text. *SFN* is within a subnetwork identified by Functional Epigenetic Module (FEM) analysis (Supplemental Information S13) as disrupted between HPV A9 and A7 squamous cell carcinomas, and is highlighted using a bold black outline. **b,** Circos plot showing frequency (0-100%) of gains and losses for regions of each chromosome (outer circle). Lines within inner circle indicate integration breakpoints from the HPV genome to the human genome as defined in Methods, Supplemental Information S2, and Supplemental Table 3. Lines are color coded by HPV clade.

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METHODS

Samples and data freeze. The Core Data Freeze (Core Set) included 178 cases from cervical carcinoma (CESC) batches 88, 114, 127, 148, 169, 179, 200, 217, 236, 256, 280, 297, 335, and 350 (Supplemental Table 1). Samples in the Core Set had mRNA-seq, whole exome DNA-seq (WES), miRNA-seq, methylation, SNP6 copy number, and clinical data available. Additional cases having multicenter mutation calls and/or RPPA data included 67 cases from CESC batches 88, 114, 127, 148, 169, 179, 200, 217, 236, 256, 280, 297, 335, 350, 361, 373, 380, 394, and 420 (Supplemental Table 2). Of these cases, 14 had mutations called and 60 had RPPA data available; however, RPPA data for 17 cases was excluded due to low protein content within samples (Supplemental Table 2). Mutations were called for 192 samples (Extended Set), while all other platform and integrated analyses (aside from protein) were performed on the subset of 178 Core Set samples. Protein levels were measured on 155 samples, which included 119 total samples from both the Core and Extended Sets as well as 36 samples outside of these sets. The total number of non-overlapping samples across Core, Extended, and RPPA datasets is 228 (Extended Data Fig. 1a).

HPV detection, variant calling, and transcript analysis. HPV status was determined using consensus results from MassArray and RNA-seq (Supplemental Information S2). MassArray uses real-time competitive polymerase chain reaction and matrix-assisted laser desorption/ionization-time of flight mass spectroscopy with separation of products on a matrix-loaded silicon chip array, similar to the work described in Tang *et al.*⁴⁵. Two approaches for pathogen detection from RNA-seq data were used. The first used the microbial detection pipeline at the British Columbia Cancer Agency's Genome Sciences Centre (BC), which is based on BioBloom Tools (BBT, v1.2.4b1)⁴⁶. The second used the PathSeq algorithm⁴⁷ at the Broad Institute (BI) to perform computational subtraction of human reads followed by alignment of residual reads to a combined database of human reference genomes and microbial reference genomes including HPV. In 97% of samples, complete agreement between MassArray and both RNA-seq approaches was observed. The remaining discrepant samples were resolved by majority decision, assigning the genotype called by at least two of the methods. RNA-seq data in FASTA format was used to identify HPV variants (Supplemental Fig. S1). Unaligned reads were taken from the PathSeq analysis and aligned to HPV reference genomes using TopHat⁴⁸ with default parameters⁴⁹. The HPV variant lineages/sublineages were assigned based on the phylogenetic topology and confirmed visually using the SNP patterns⁵⁰. HPV splice junctions from RNA-seq were determined using TopHat. Two transcript types were distinguished for HPV16 and HPV18: (a) transcripts that included evidence of an unspliced sequence of E6, and (b) a transcript spliced at the E6 splice donor site (position 226 for HPV16 and position 233 for HPV18) (Supplemental Fig. S2). Read counts for unspliced, spliced, as well as the ratio of unspliced/spliced transcripts were categorized into quartiles separately for HPV16 and HPV18.

HPV integration analysis. Using RNA-seq data, concordance of integration events based on alignments of contigs from *de novo* transcriptome assembly (BC) and read alignments (BI) was evaluated (Supplemental Fig. S3). We identified method-specific integration events by assigning all sites within a 500-kb sliding window to a single integration event located at the median coordinate of that event's assigned sites. An integration event was labeled as 'confident' when the total read support for each of its supporting integration sites passed center-specific read evidence thresholds. To take advantage of differences between the two integration methods (i.e. contig and read), for the concordance analysis we used all method-specific integration events (both confident and non-confident events). We labeled an integration event as 'concordant' when both methods reported an integration event within 500 kb in the same patient. For some concordant events, both methods reported a confident event. An integration event was labeled as 'discordant' when only one center reported a confident integration event within 500 kb (Supplemental Figs. S4 and S5). For both intragenic and intergenic concordant events, we reported a range of coordinates that extends from the most proximal to the most distal supporting integration site. We assessed gene-level expression relative to somatic copy number and structural variant data for genes into which we had mapped viral-human junctions from RNA sequencing data and for genes that were associated with enhancers into which we had mapped RNA junctions.

DNA sequencing and mutation calling. Detailed methods for library hybrid capture, read alignments, and somatic variant calling are documented in Supplemental Information S3. MutSig2CV⁶ was utilized to identify significantly mutated genes (SMGs) within the cervical cancer exome sequencing data. Mutations were analyzed for the Core Set plus 14 samples to total 192 Extended Set samples. Eleven samples were identified to exhibit greater than average mutations rates and were termed "hypermutants" (somatic mutations > 600). These 11 samples were excluded from the analysis for identifying SMGs. All 3 sample subsets (all samples, squamous carcinomas only, adenocarcinomas only) without "hypermutants" (Supplemental Table 4) were analyzed using an FDR cutoff of

0.1. FDR values are shown in Supplemental Table 4. SMG analysis using the entire sample cohort in Ojesina *et al.* was performed as described previously⁸.

Copy number analysis. DNA from each tumor or germline sample was hybridized to Affymetrix SNP 6.0 arrays using protocols at the Genome Analysis Platform of the Broad Institute as previously described⁵¹. Briefly, Birdseed was used to infer a preliminary copy number at each probe locus from raw.CEL files⁵². For each tumor, genome-wide copy number estimates were refined using tangent normalization, in which tumor signal intensities are divided by signal intensities from the linear combination of all normal samples that are most similar to the tumor¹⁶. Individual copy number estimates then underwent segmentation using Circular Binary Segmentation⁵³, and segmented copy number profiles for tumor and matched control DNAs were analyzed using Ziggurat Deconstruction⁵⁴. Significance of copy number alterations were assessed from the segmented data using GISTIC2.0 (Version 2.0.22)⁵⁴. For the purpose of this analysis, an arm-level event was defined as any event spanning more than 50% of a chromosome arm. For copy number based clustering, tumors were clustered based on log2 copy number at regions revealed by GISTIC analysis. Clustering was done in R based on Euclidean distance using Ward's method. Allelic and integer copy number, tumor purity, and tumor ploidy were calculated using the ABSOLUTE algorithm⁵⁵.

Detecting structural variants from RNA-seq and WGS data. Integrative analysis was performed to identify putative driver fusions using both WGS (low-pass and high-coverage) and RNA-seq data. RNA-seq data for 178 Core Set cases were analyzed using the TopHat-Fusion and BreakFusion, PRADA, and MapSplice algorithms. To identify structural variations in WGS data, 50 low-pass WGS and 19 high-pass WGS samples were analyzed. Detection of structural variations in low-pass WGS data was performed using two algorithms, BreakDancer⁵⁶ and Meerkat⁵⁷, with a requirement for at least two discordant read pairs supporting each event and at least one read covering the breakpoint junction. High-pass WGS data were analyzed to detect somatic structural variations using two runs of BreakDancer and one run of SquareDancer (<https://github.com/ding-lab/squaredancer>). The gene fusion lists generated by all methods and platforms were integrated (See Supplemental Tables 8-10).

APOBEC mutagenesis analysis. Analysis is based on previous findings that APOBECs deaminate cytidines predominantly in a tCw motif and that the APOBEC mutagenesis signature is composed of approximately equal numbers of two kinds of changes in this motif: tCw→tTw and tCw→tGw mutations (flanking nucleotides are shown in small letters; w = A or T). Using mutation data from all 192 Extended Set samples, we calculated on a per sample basis the enrichment of the APOBEC mutation signature among all mutated cytosines in comparison to the fraction of cytosines that occur in the tCw motif among the +/- 20 nucleotides surrounding each mutated cytosine ("APOBEC_enrich" column in data files). The minimum estimate of the number of APOBEC-induced mutations in a sample (APOBEC_MutLoad_MinEstimate) was calculated using the formula: $[(tCw \rightarrow G + tCw \rightarrow T)^x] / [(APOBEC_enrich - 1) / (APOBEC_enrich)]$, which allows estimating the number of APOBEC signature mutations in excess of what would be expected by random mutagenesis. "APOBEC_MutLoad_MinEstimate" was calculated only for samples passing 0.05 FDR threshold for APOBEC enrichment ($[BH_Fisher_p_value_tCw] < 0.05$). Samples with "BH_Fisher_p-value_tCw" value greater than 0.05 received a value of 0. The "APOBEC_MutLoad_MinEstimate" value shows high correlation (0.9-0.95) with all other parameters used to characterize the APOBEC mutagenesis pattern, such as APOBEC enrichment as well as absolute and relative APOBEC mutation loads. For some analyses and figures, the "APOBEC_MutLoad_MinEstimate" parameter was converted into categorical values as follows:

1. "no": "APOBEC_MutLoad_MinEstimate" = 0
2. "low": $0 < \text{"APOBEC_MutLoad_MinEstimate"} \leq \text{median of non-zero values}$
3. "high": "APOBEC_MutLoad_MinEstimate" > median of non-zero values

The median of non-zero values in the Extended Set = 33.

Methylation analysis. The Illumina Infinium HM450 array⁵⁸ was used to evaluate DNA methylation in the Core Set of cervical cancer samples. Unsupervised consensus clustering was performed with Euclidean distance and partitioning around medoids (PAM) using the most variable 1% of CpG island promoter probes. Epigenetically silenced genes were identified as previously described⁵⁹. A total of 120 normal samples were used for this analysis by selecting 10 samples at random from the 12 TCGA projects that included normal samples.

RNA-seq analysis. RNA was extracted, converted into mRNA libraries, and paired-end sequenced (paired 50 nt reads) on Illumina HiSeq 2000 Genome Analyzers as previously described⁵. RNA reads were aligned to the hg19 genome assembly using MapSplice v12_07⁶⁰. Gene expression was quantified for the transcript models corresponding to the TCGA GAF2.1 (<https://tcga-data.nci.nih.gov/docs/GAF/GAF.hg19.June2011.bundle/outputs/TCGA.hg19.June2011.gaf>) using RSEM4⁶¹ and normalized within a sample to a fixed upper quartile.

To predict whether a cancer sample was from the cervix or the uterus, the data matrix of normalized gene-level RSEM values from 170 UCEC samples was merged with the data matrix from the Core Set ($n = 178$) of cervical cancers. This merged dataset was then randomly split into a training set (87 CESC samples; 86 UCEC samples) and a test set (91 CESC samples; 84 UCEC samples). A sample was predicted to be CESC if the t-statistic vs. UCEC was significant ($p < 0.05$), but was not significantly different from the CESC mean (and vice versa for the UCEC prediction). A data matrix of RSEM values from 178 CESC, 170 UCEC, and 279 HNSC samples was used to identify expression patterns across the 3 cancer types. The gene expression matrix was further filtered to only include the top 25% most variable genes by mean absolute deviation ($n = 4,039$ genes).

EMT mRNA score analysis. The EMT score was computed as previously described^{10,21}. Briefly, the EMT score was the value resulting from the difference between the average expression of mesenchymal (M) genes minus the average expression of epithelial (E) genes. All NA values were removed from the calculation. Two-sample t-test and ANOVA were applied to each comparison accordingly.

miRNA sequencing and analysis. MicroRNA sequence (miRNA-seq) data was generated for the Core Set of tumor samples using methods described previously¹¹. We identified miRNAs that have been associated with EMT^{62–66} and then calculated Spearman correlations between the EMT scores and RPMs for 5p and 3p mature strands for each of these miRNAs using MatrixEQTL and filtering by $FDR < 0.05$. An miRNA was considered to be epigenetically controlled if BH-corrected p-values were less than 0.01 for both a) a Spearman correlation of miRNA abundance (RPM) to beta for probes in promoter regions associated with the miRNAs, and for b) a t-test of RPM between unmethylated ($\beta < 0.1$) and methylated ($\beta > 0.3$) samples (an “epigenetically-controlled pattern”). We assessed potential miRNA targeting for all 178 samples and then separately for the 144 squamous samples by calculating miR-mRNA and miR-protein (RPPA) Spearman correlations with MatrixEQTL v2.1.1 using gene-level normalized abundance RNA-seq (RSEM) data and normalized RPPA data. Correlations were calculated with a p-value threshold of 0.05, and then the anti-correlations were filtered at $FDR < 0.05$. We extracted miR-gene pairs that corresponded to functional validation publications reported by miRTarBase v4.5²². For miR-RPPA anti-correlations, all gene names that were associated with each antibody were used. Results were displayed with Cytoscape v2.8.3.

PARADIGM analysis. Integration of copy number, RNA-seq, and pathway interaction data was performed on the Core Set of samples using PARADIGM^{33,34}. Briefly, PARADIGM infers integrated pathway levels (IPLs) for genes, complexes, and processes using pathway interactions, genomic, and functional genomic data from each patient sample. One was added to all expression values, which were then \log_2 -transformed and median-centered across samples for each gene. The \log_2 -transformed, median-centered mRNA data were rank-transformed based on the global ranking across all samples and all genes and discretized (+1 for values with ranks in the highest tertile, -1 for values with ranks in the lowest tertile, and 0 otherwise) prior to PARADIGM analysis.

Pathways were obtained in BioPax Level 3 format, and included the NCIPID and BioCarta databases from <http://pid.nci.nih.gov> and the Reactome database from <http://reactome.org>. Gene identifiers were unified by UniProt ID and then converted to Human Genome Nomenclature Committee’s HUGO symbols using mappings provided by HGNC (<http://www.genenames.org/>). Altogether, 1524 pathways were obtained. Interactions from all of these sources were then combined into a merged Superimposed Pathway (SuperPathway). Genes, complexes, and abstract processes (e.g. “cell cycle” and “apoptosis”) were retained and henceforth referred to collectively as pathway features. The resulting pathway structure contained a total of 19504 features, representing 7369 protein-coding genes, 9354 complexes, 2092 families, 82 RNAs, 15 miRNAs, and 592 abstract processes.

The PARADIGM algorithm infers an IPL for each pathway element that reflects the log likelihood contrasting the probability of activity against inactivity. An initial minimum variation filter (at least 1 sample with absolute activity > 0.05) was applied, resulting in 15502 concepts (5898 protein-coding genes, 7307 complexes, 1916 families, 12 RNAs, 15 miRNAs, and 354 abstract processes) with relative activities showing distinguishable variation across tumors.

iCluster analysis. Integrative clustering of RNA-seq, methylation, copy number, and miRNA data was performed using R package “iCluster²⁰”. The Core Set of samples was used since all samples in this Set had data available across these four platforms. RNA-seq, methylation, copy number, and mature-strand miRNA datasets had 20531, 395552, 23109, and 1213 features, respectively. The 500 most variable features based on the standard deviation from each dataset were selected for the integrative clustering analyses. For analysis involving the RNA-seq and miRNA datasets, a $\log(x+1)$ transformation was used in order to deal with skewness in the data⁶⁷. Methylation data was logit transformed to make it closer to normal distribution. The copy number variation data included the regions

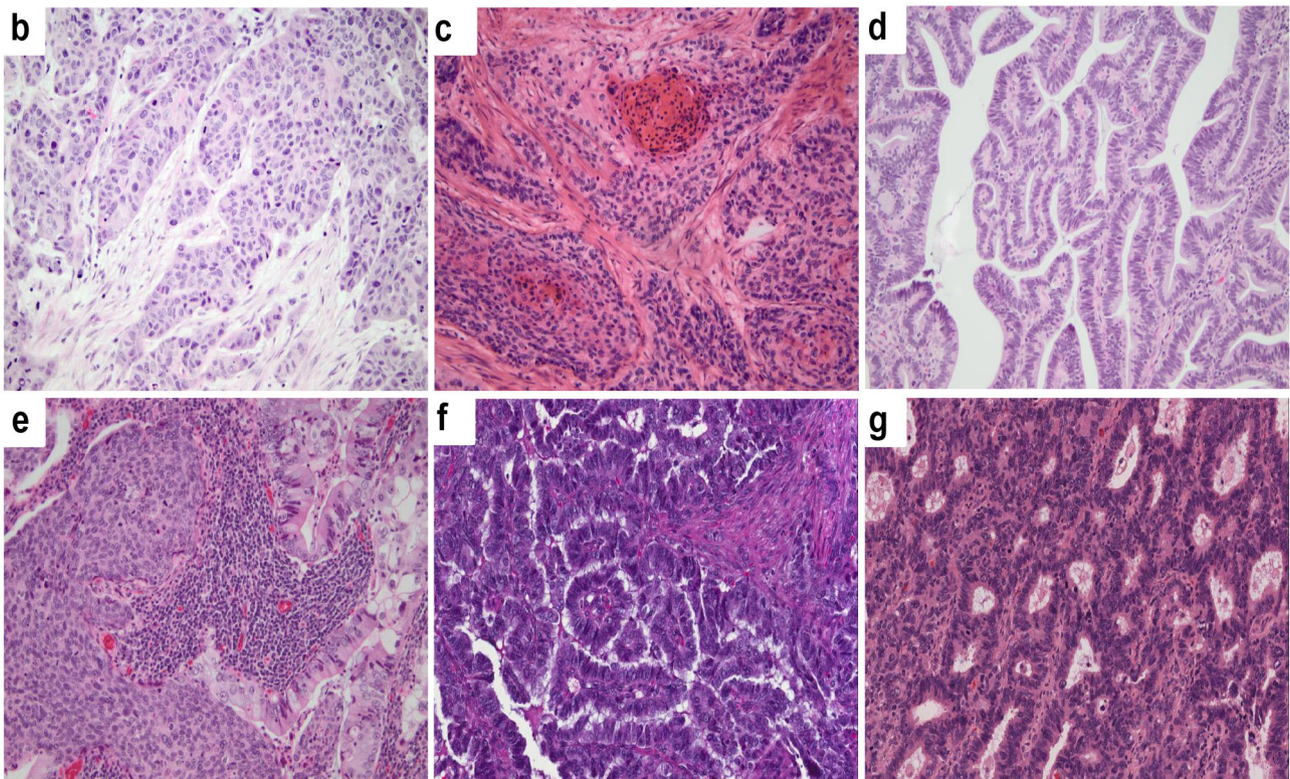
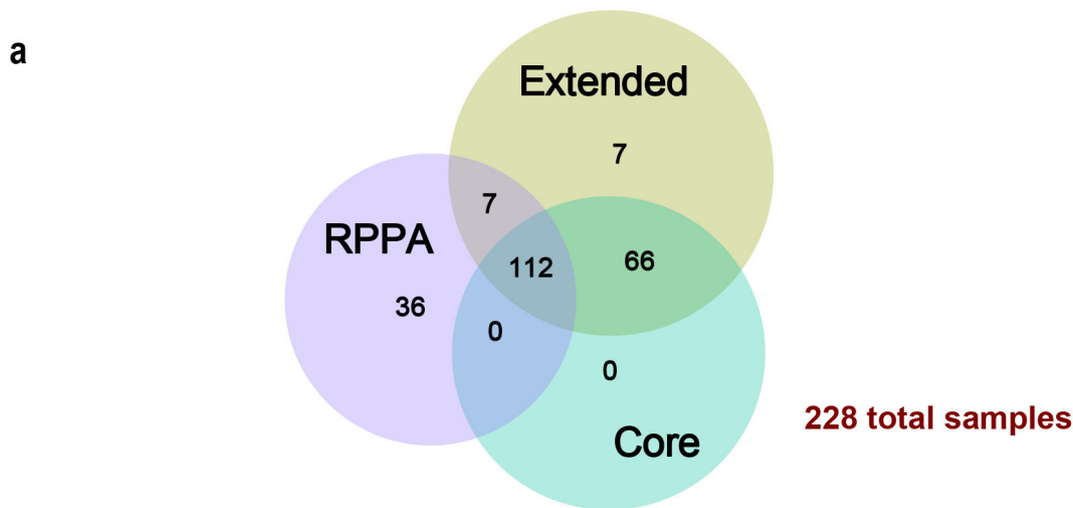
determined from GISTIC2.0, with copy number variation treated as a continuous measurement based on the segmentation mean value for the region.

MEMo analysis. High DNA methylation levels upstream of miR-200a and miR-200b corresponded to transcriptional downregulation of the miRs (Extended Data Fig. 9a). For a sample to be called altered for either miR-200a or miR-200b (or both), we required both high DNA methylation level upstream of the miR (β -value > 0.3) and low miR expression ($\log_2(\text{RPM}) < 9.3$ for miR-200a and $\log_2(\text{RPM}) < 9$ for miR-200b). Binary calls were given to altered and unaltered samples based on this double threshold (1 = altered, 0 = unaltered).

The Mutual Exclusivity Modules in cancer (MEMo) algorithm²⁷ was run on all Core Set samples. MEMo was initially run on 27 regions of recurrent copy number gain, 36 of copy number loss, and 22 recurrently mutated genes. In order to include alterations for miR-200a and miR-200b in the MEMo analysis, a custom network was designed where each miR was connected to its known and validated targets (see above). Second, this network was merged with the comprehensive pathway network used by MEMo to search for modules of altered genes that include at least one of the miRs. Extracted modules were tested for mutual exclusivity using MEMo’s statistical framework (Supplemental Table 27). Student’s t-test was performed for comparing EMT mRNA scores between groups.

Data Availability. The primary and processed data used in analyses can be downloaded by registered users from <https://gdc-portal.nci.nih.gov/> and the TCGA publication page (https://tcga-data.nci.nih.gov/docs/publications/cesc_2016/).

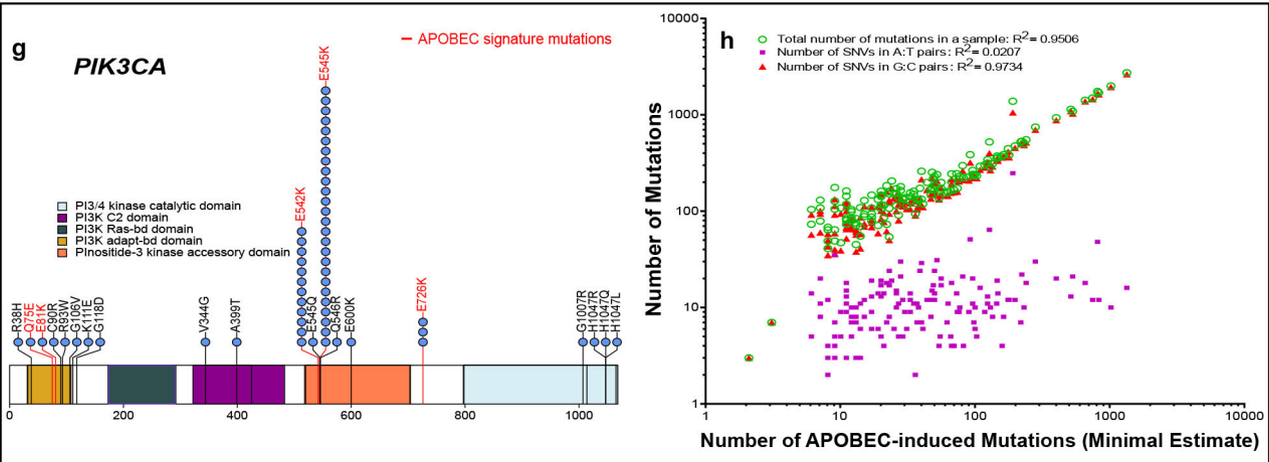
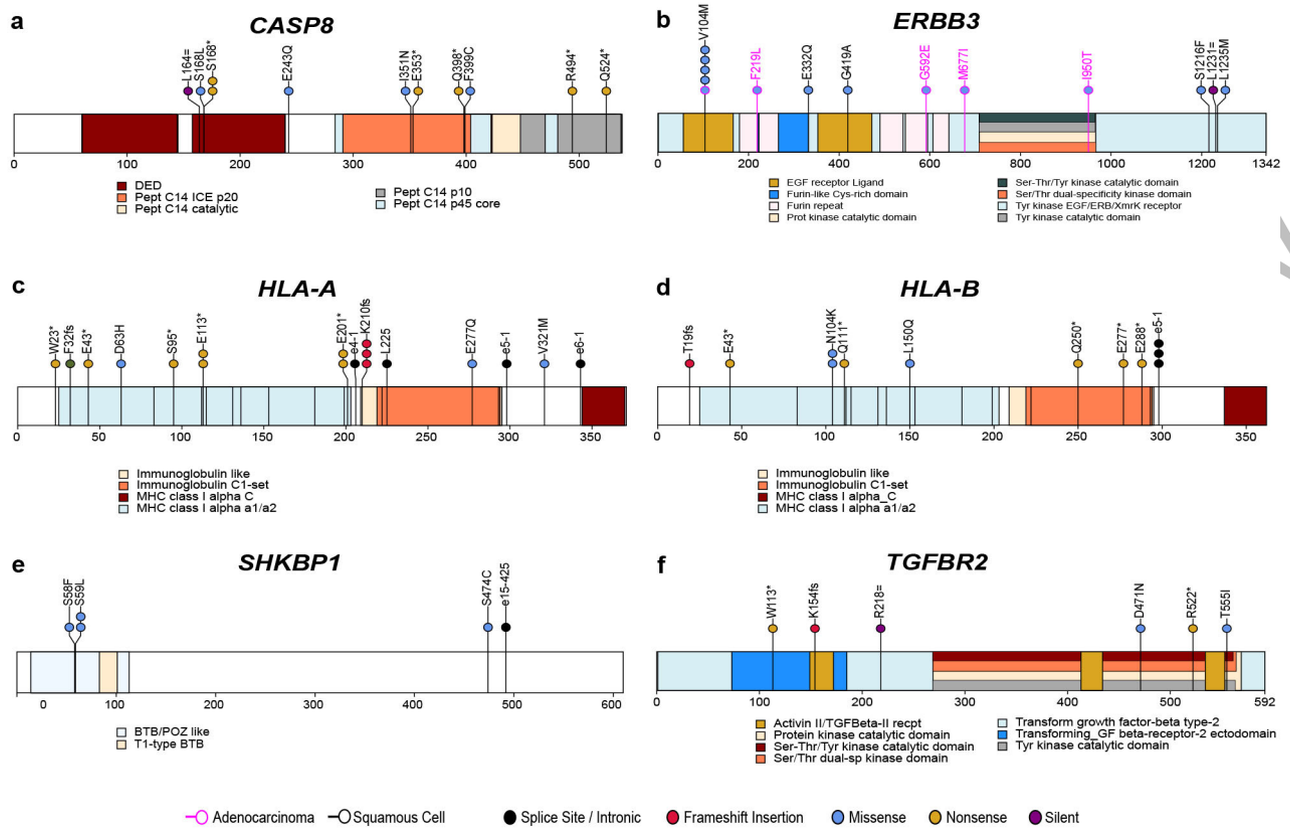
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Extended Data Figure 1 | Sample sets and histologic patterns of cervical cancer.

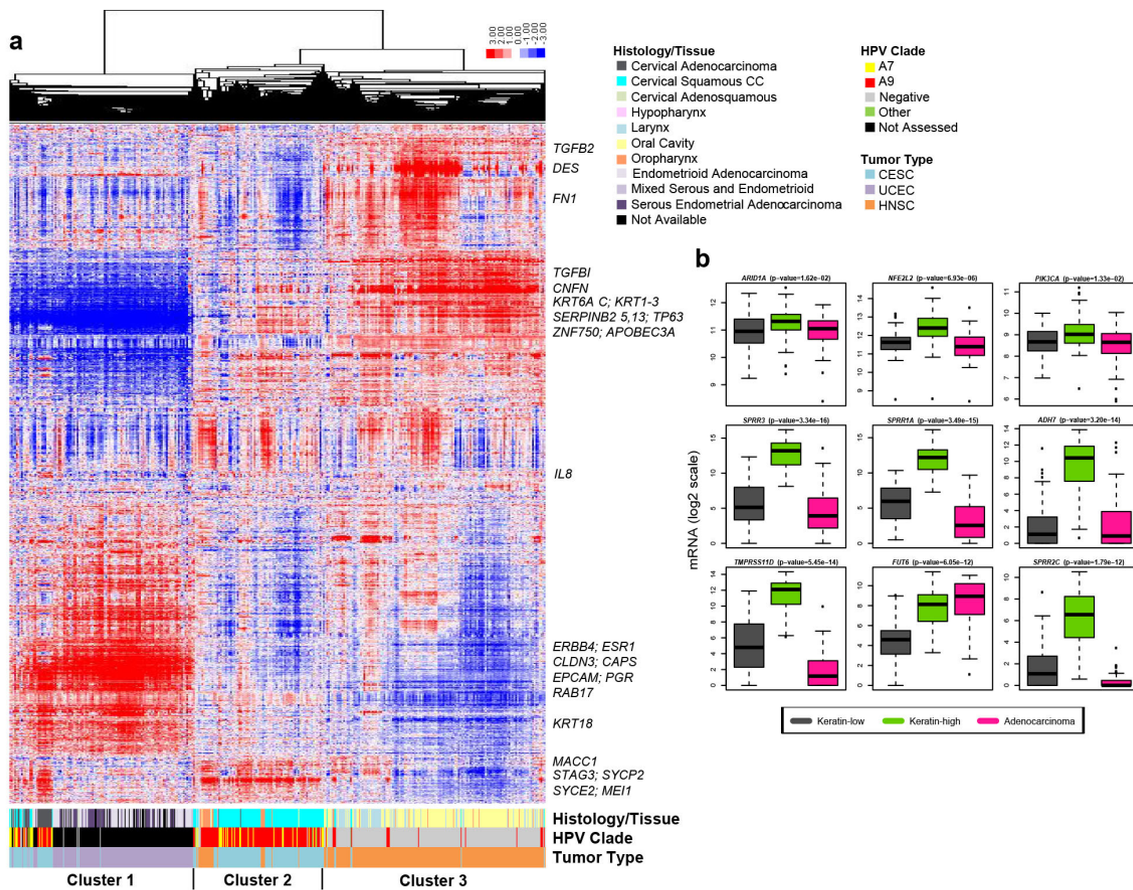
a, Summary of sample numbers and degree of overlap between the Core, Extended, and RPPA datasets. **b**, Squamous cell carcinoma of the large cell non-keratinizing type. Tongues of highly atypical polygonal neoplastic squamous cells infiltrate through a fibrotic stroma. The cells show abundant eosinophilic cytoplasm with pleomorphic nuclei and prominent mitotic figures. Although the tumor cells contain abundant cytokeratin filaments, this tumor has traditionally been termed “non-keratinizing” because of the absence of characteristic keratin pearls. **c**, Squamous cell carcinoma of the large cell keratinizing type. Nests of atypical squamous cells infiltrate through a fibrotic stroma. In addition, this tumor shows highly eosinophilic keratin pearls with small, inky dark nuclei that imperfectly mimic the normal keratinization that is found in the epidermis. This differentiation pattern is aberrant in the cervix in which the squamous epithelium is normally a non-keratinizing squamous mucosa. **d**, Adenocarcinoma of endocervical type (well-differentiated). Closely set, atypical glands with enlarged nuclei and scattered mitotic figures infiltrate through the connective tissue of the cervix. The tall

columnar tumor cells show basally-placed, crowded, enlarged nuclei that show frequent mitotic figures. Compared with normal endocervical cells, the tumor cells show relative loss of intra-cytoplasmic mucin and are frequently called “mucin-depleted,” although most, but not all endocervical adenocarcinomas show varying amounts of intracytoplasmic mucin at least focally. **e**, Adenosquamous carcinoma of cervix. This tumor shows both nests of non-keratinizing squamous cell carcinoma and glands composed of tall columnar adenocarcinoma reflecting the origin of most cervical cancers in the transformation zone of the cervix in which both squamous and glandular cells normally differentiate. Despite this biphasic differentiation potential, adenosquamous carcinomas are relatively uncommon in the cervix. **f**, UCEC-like HPV negative adenocarcinoma of endocervical type from a radical hysterectomy specimen. The endometrium in the uterus was benign. **g**, UCEC-like HPV positive adenocarcinoma of endocervical type from a radical hysterectomy specimen. The endometrium in the uterus was benign. All samples were stained with hematoxylin and eosin (20x).



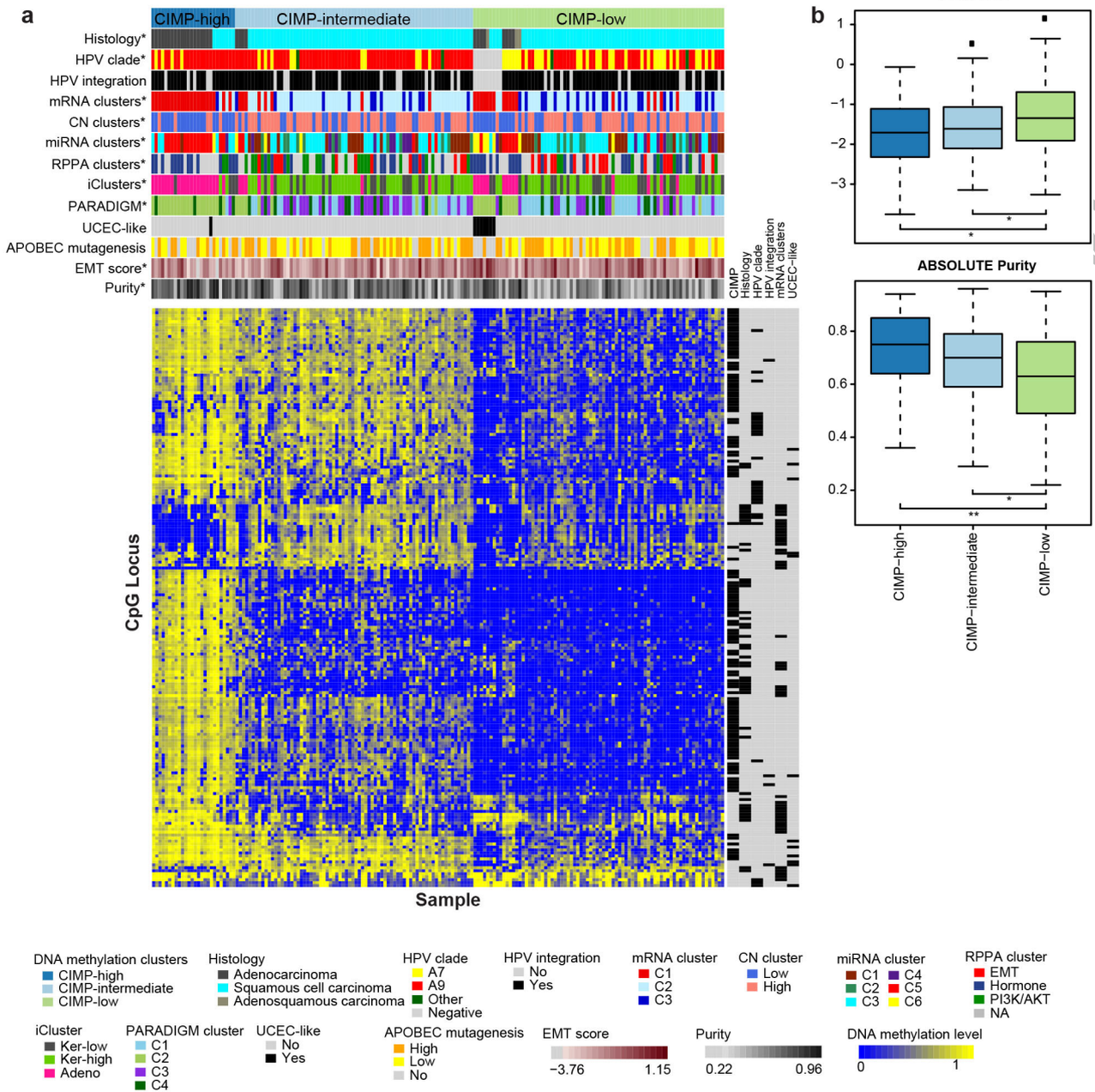
Extended Data Figure 2 | Significantly mutated genes and the role of APOBEC in cervical cancer mutagenesis. a-f, High-confidence somatic mutations in significantly mutated genes (SMGs) among 192 exome-sequenced samples in the Extended case set are shown. Domains are labeled in accordance with Gencode 19 corresponding to Ensembl 74. Mutations at canonical intronic splice acceptor (e-1 and e-2) are labeled based on proximity to the nearest coding exon. Panels display somatic mutations detected in novel cervical cancer SMGs, with *HLA-B* included for comparison with its family member *HLA-A*. Each axis is the protein-coding portion of a gene and each highlighted section represents the UniProt functional domain. Vertical lines indicate the boundaries of multiple annotation sources within common domain annotations as outlined in Supplemental Table 5. Horizontal lines distinguish overlapping domains. Circles represent a single mutation and are colored based on mutation type. Mutations present in squamous cell carcinomas are

outlined in black while those present in adenocarcinomas are outlined in pink. g, *PIK3CA* mutations and recurrence are shown in a stacked circle plot, as above. Additionally, lollipop sticks are colored red if the mutation type coincides with patterns of APOBEC mutagenesis. h, The minimal estimated number of APOBEC-induced mutations (“APOBEC_MutLoad_MinEstimate” column in Supplemental Table 1) strongly correlates with total number of mutations in a sample, as well as with the number of single nucleotide variants (SNVs) in G:C pairs which are the exclusive substrate for mutagenesis by APOBEC cytidine deaminases. While correlation with mutagenesis in A:T base pairs, which cannot be mutated by APOBEC enzymes is statistically significant (two-tailed P = 0.047), it is very weak. Pearson correlation and R² were calculated for all 192 exome-sequenced samples, including samples with zero values. Only samples with non-zero values of “APOBEC_MutLoad_MinEstimate” are presented.



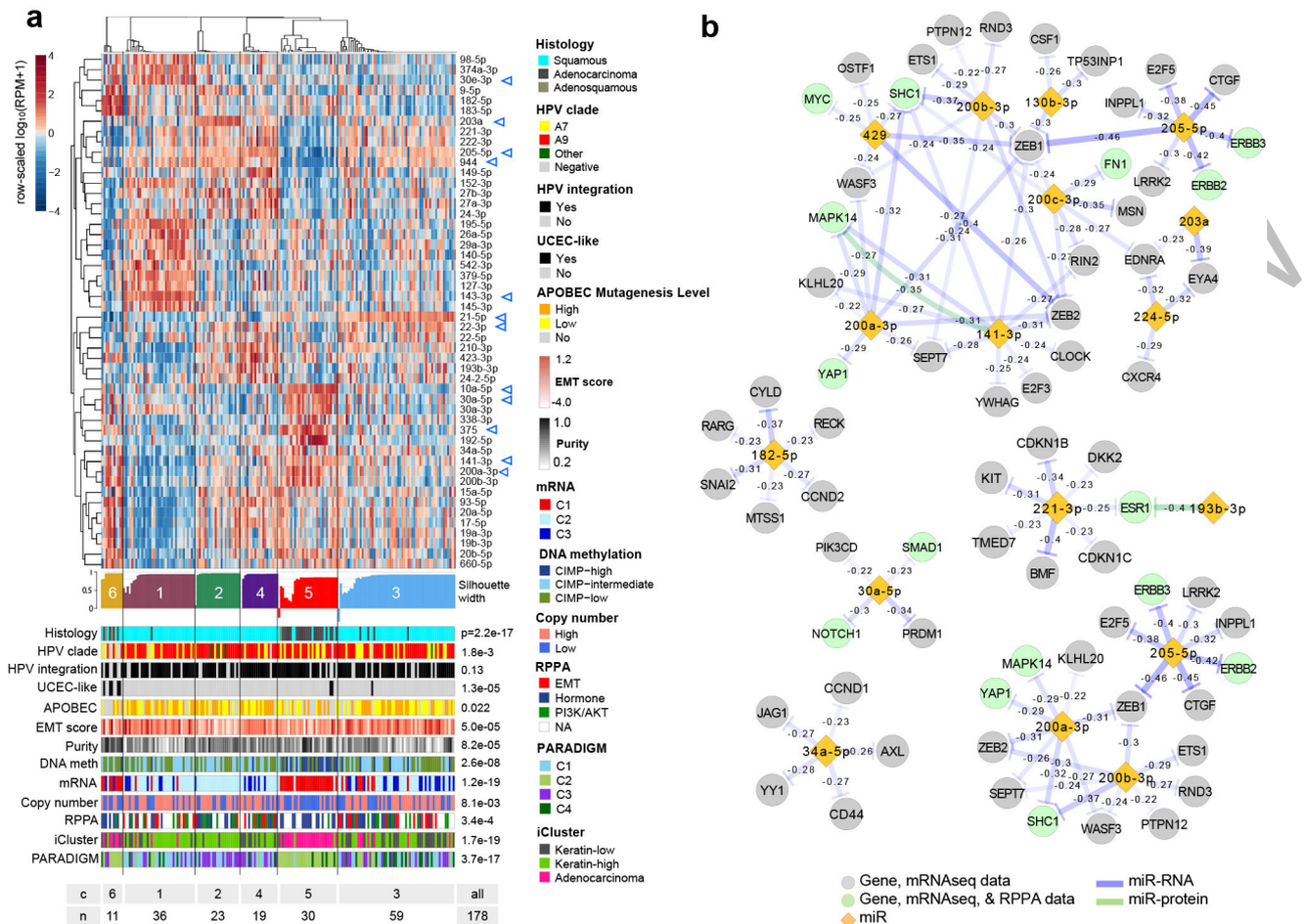
Extended Data Figure 4 | Gene expression patterns and fusion genes found in cervical cancer. **a**, Hierarchical clustering (uncentered correlation with centroid linkage as the clustering method) was performed on 4,039 expressed and highly variable genes across 178 cervical, 170 endometrial, and 279 head and neck cancer samples. Normalized gene-level RSEM values were median-centered prior to clustering and relative increased expression values are indicated by red color while relative decreased expression values are indicated by blue color. Cervical, endometrial, and head and neck cancer samples are indicated by different colors as noted in the figure at the bottom of the heatmap. Also included are indications of HPV status, histology of cervical and endometrial cancers, and tissue site for head and neck cancer samples. Select genes are noted to the right of their locations on the heatmap. **b**, Boxplots of the three differentially expressed SMGs and top six significantly differentially

expressed non-SMGs across the iCluster groups using Kruskal Wallis test. All genes are significantly different across the Keratin-low and Keratin-high clusters. Significant p-values across Keratin-low and Keratin-high clusters are presented. **c**, A schematic of *BCAR4* tandem duplication in one case (C5-A3HF), detected by analysis of somatic copy number (top) and structural variation (middle). Split reads and genomic breakpoints indicating the tandem duplication are shown. At the RNA level (bottom) the last exon of *BCAR4* forms a fusion gene with the first exon of *ZC3H7A* (red bars indicate location of mRNA breakpoints; NR_024049 shown as *BCAR4* representative transcript). **d**, Schematic of recurrent fusions (*CPSF6-C9orf3*, *ARL8B-ITPR1*, and *MYH9-TXN2*) or fusions with known occurrences in other cancer types (*FGFR3-TACC3*), detected by at least two RNA-seq fusion callers in 178 samples. Red bars indicate the mRNA breakpoints.



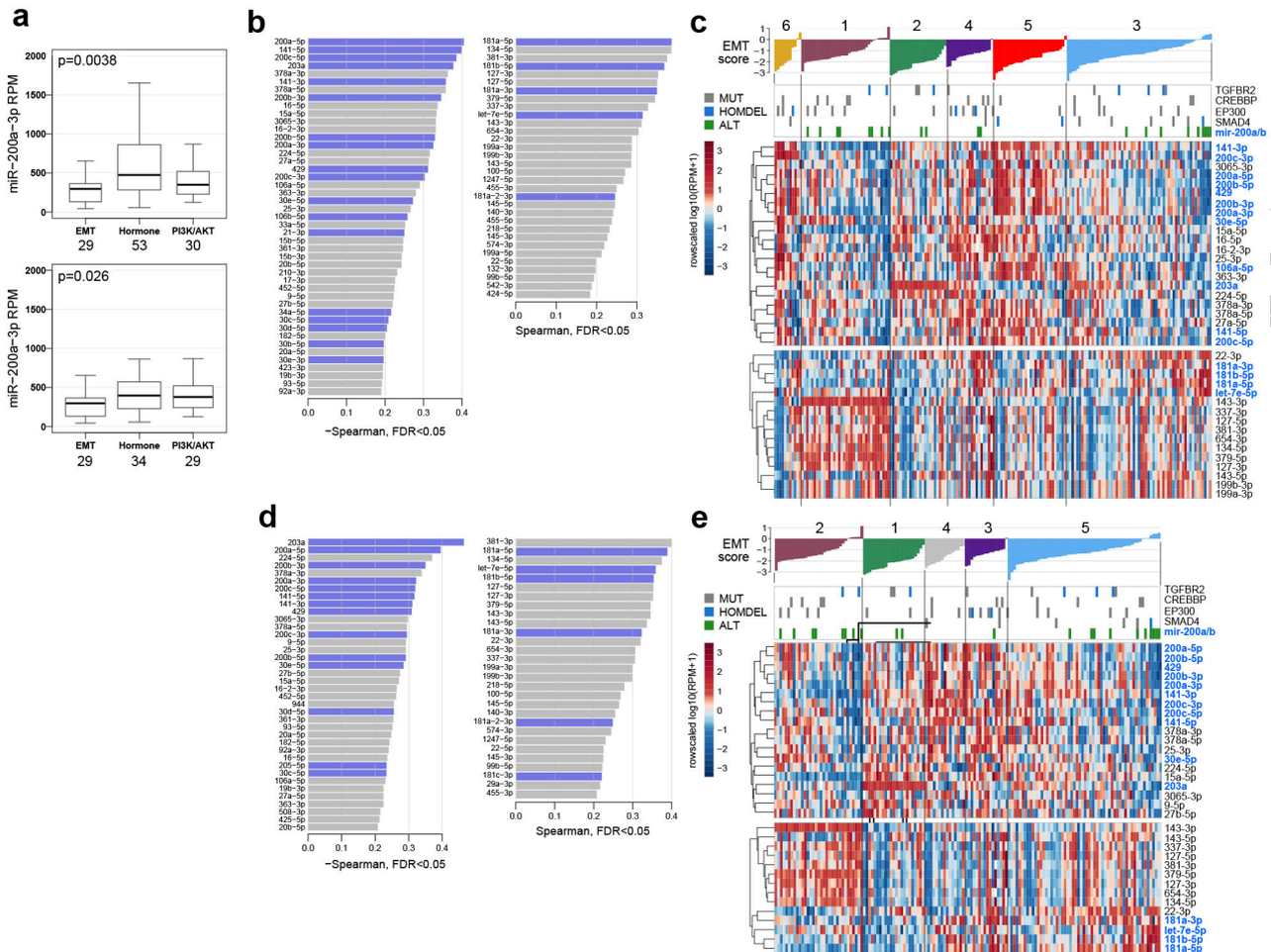
Extended Data Figure 5 | Unsupervised clusters of DNA methylation data. **a**, Heatmap showing beta values of 178 Core Set samples ordered by CIMP clusters. Samples are presented in columns and the CpG island promoter CpG loci are presented in rows. An annotation panel on the right of the heatmap indicates CpG loci that are differentially methylated within a particular feature (see Supplemental Table 13). All features (marked

with *) are statistically significantly associated with DNA methylation clusters (Fisher's Exact test p -value < 0.01) except APOBEC mutagenesis level, UCEC-like status, and HPV integration status. **b**, Box plots of the EMT mRNA score and tumor purity by CIMP clusters. Student's t -test p -value < 0.01 (**) and < 0.05 (*) are reported.



Extended Data Figure 6 | miRNA clusters and miR-gene/protein anti-correlations in cervical cancer. **a**, Unsupervised clustering for miR profiles across 178 Core Set tumor samples. Top to bottom: a normalized abundance heatmap for the fifty 5p or 3p strands that were highly ranked as differentially abundant by a SAMseq multiclass analysis, silhouette width profile calculated from the consensus membership matrix, a heatmap of tumor sample purity, covariates with association p-values, and a summary table of the number of samples in each cluster. The scale bar shows row-scaled $\log_{10}(\text{RPM}+1)$ normalized abundances. **b**, Subnetworks

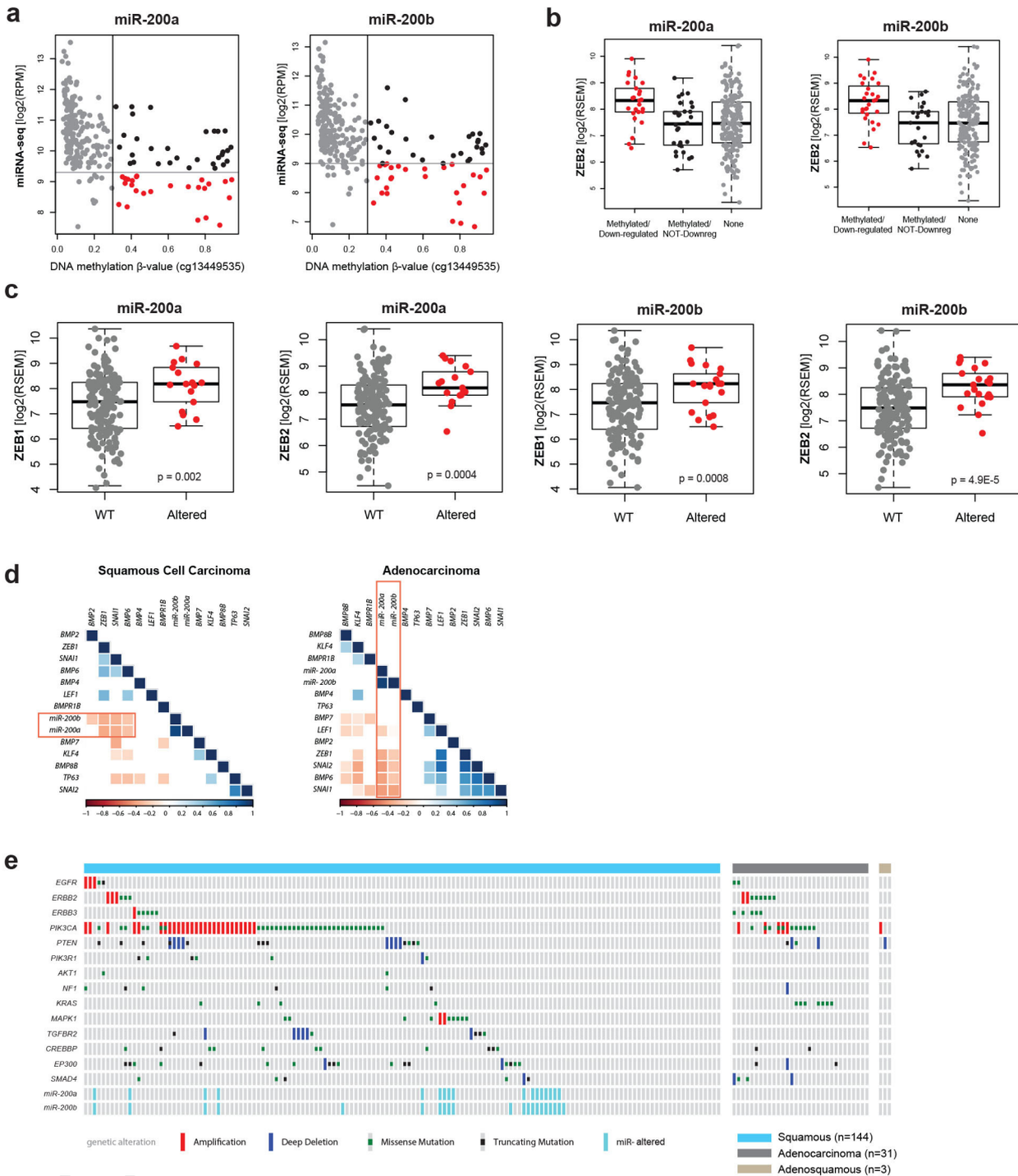
of potential targeting relationships for a subset of miRs, as significance-thresholded ($\text{FDR} < 0.05$) miR-mRNA and miR-RPPA anti-correlations that are supported by functional validation publications. For genes (nodes), color distinguishes those that are only present in mRNA data (grey) from those that are present in both mRNA and RPPA data (green). Edges represent anti-correlations, and color distinguishes anti-correlations between a miR and mRNA (purple) and a miR and an unphosphorylated protein (green). In the $n = 178$ Core Set cohort, no correlations satisfying $\text{FDR} < 0.05$ were reported between a miR and a phosphorylated protein.



Extended Data Figure 7 | EMT-associated miRs and their relationship to miR clusters and TGF β 2 somatic alterations. **a**, Normalized miR-200a-3p abundance (RPM) across RPPA clusters for all 112 (top) and 92 squamous (bottom) samples of the Core Set for which RPPA data is available. P-values presented are from two-sided Kolmogorov-Smirnov tests for RPPA-based EMT cluster vs non-EMT cluster samples. For $n = 112$ samples, median miR-200a-3p RPM = 296.4 within the EMT cluster ($n = 29$) and 410.0 ($n = 83$) in non-EMT cluster samples. For squamous samples, median miR-200a-3p RPM = 296.4 ($n = 29$) within the EMT cluster and 393.4 ($n = 63$) in non-EMT cluster samples. EK-A2R7, which is in the Hormone RPPA cluster, has an RPM value of 4267 and is not shown. Results are not presented for adenocarcinoma samples separately due to limiting sample numbers ($n = 18$ from the Core Set

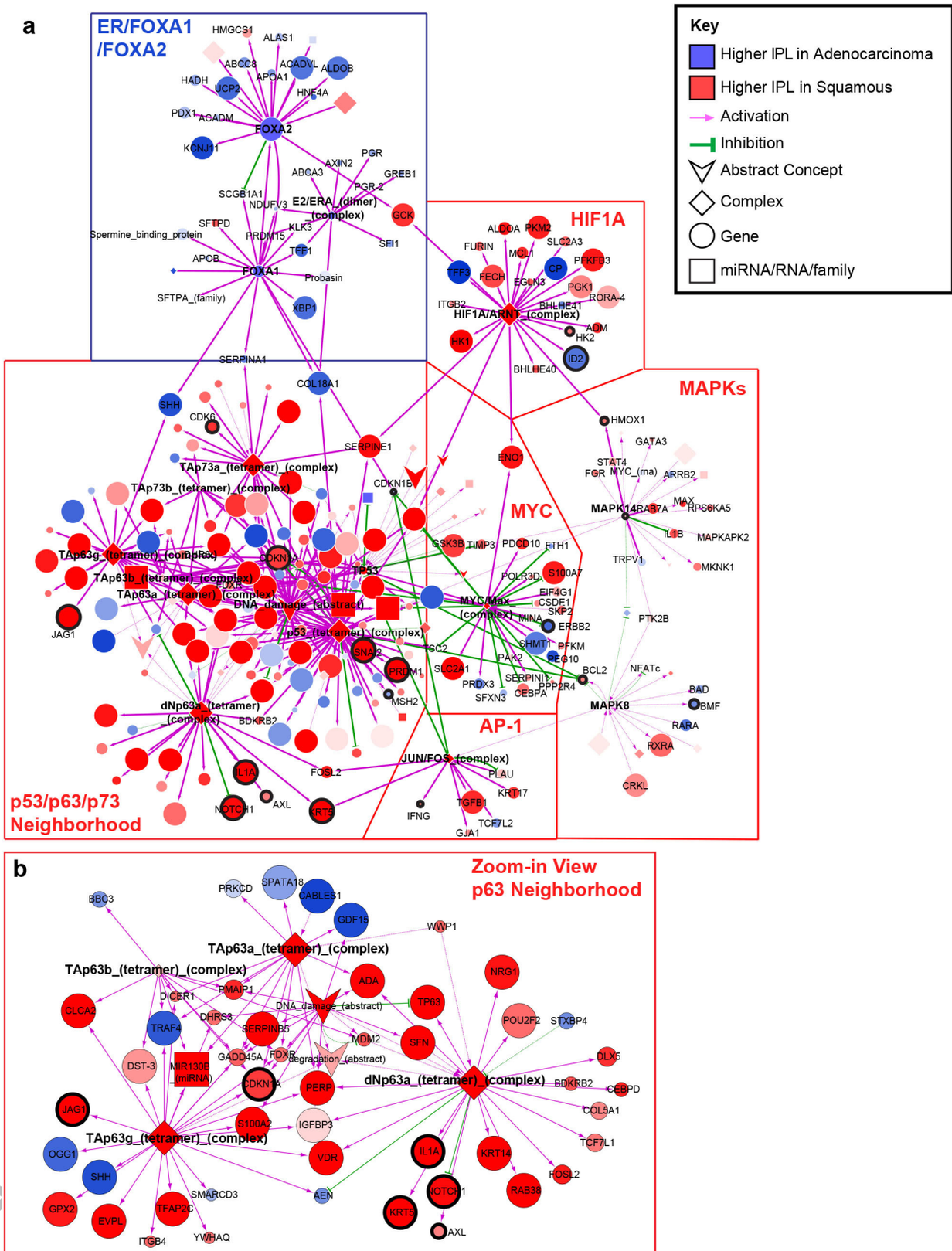
with RPPA data available). **b**, Negative and positive Spearman correlation coefficients (FDR < 0.05) between EMT mRNA score and normalized abundance (RPM) for miRNA mature strands ($n = 178$). miRNAs that have been reported as associated with EMT (see Methods) are highlighted by purple bars. **c**, Normalized abundance heatmap of miRNAs most strongly negatively and positively correlated with EMT mRNA scores, with samples grouped by miRNA cluster and sorted by EMT score within each cluster. Somatic mutations (MUT) and deletions (HOMDEL) are shown for *TGFB2*, *CREBBP*, *EP300*, and *SMAD4*. Methylation and concomitant downregulated expression alterations (ALT) as defined in Methods for miR-200a/b are also shown. miRNAs in blue text represent those highlighted by purple bars in **b**. **d-e**, Same as **b-c**, but for the $n = 144$ squamous tumor samples.





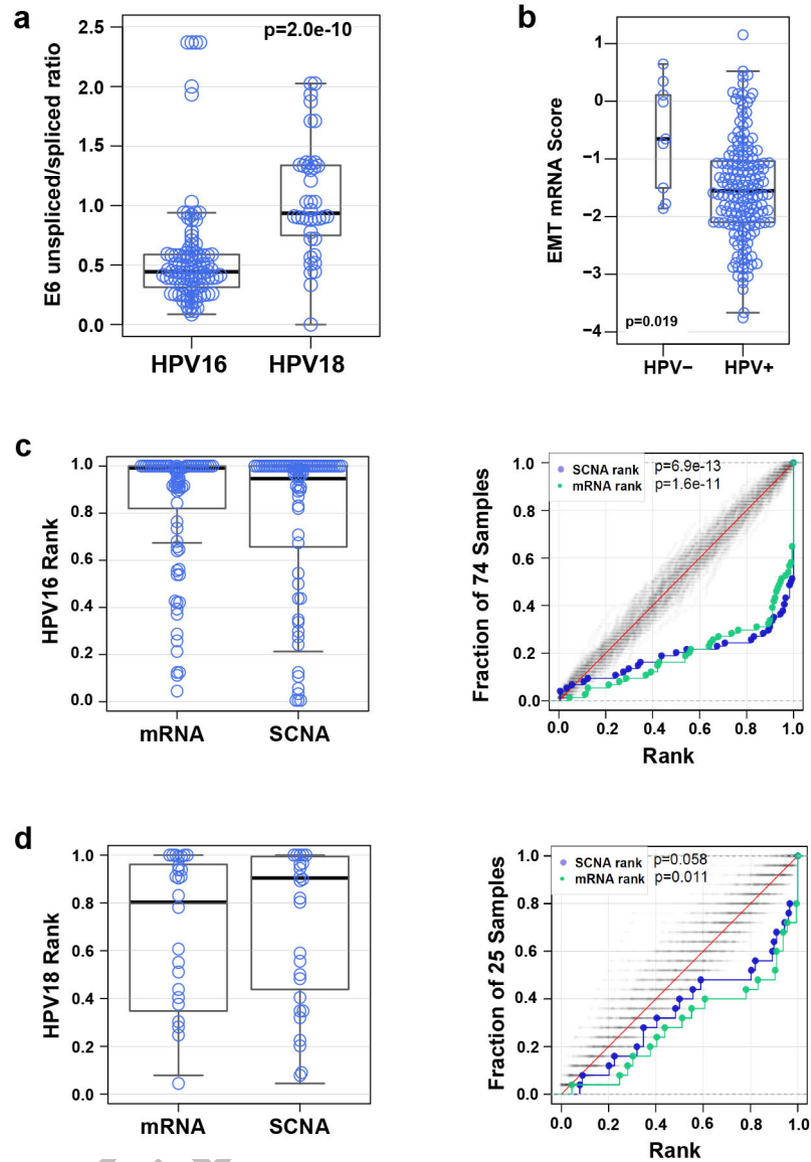
Extended Data Figure 9 | miR-200a/b associations with EMT-regulating genes and somatic alterations within RTK, PI3K, MAPK, and TGF β R2 pathways in cervical cancer. **a**, Expression levels for miR-200a and miR-200b compared to DNA methylation level at their promoter. Samples were called altered if the miRs were concurrently hypermethylated ($\beta > 0.3$) and downregulated (red cases). **b**, mRNA expression levels for ZEB2, a target of both miR-200a and miR-200b, in subsets of miR-200a/b altered samples. ZEB2 is upregulated in cases with concurrent hypermethylation

and downregulation of the miRs. **c**, mRNA expression levels of both ZEB1 and ZEB2 in miR-200a/b hypermethylated/downregulated (Altered) and all other (WT) samples. **d**, Correlations of miR-200a and miR-200b expression with multiple genes involved in EMT signaling across squamous cell carcinomas and adenocarcinomas. **e**, Extent of genetic alterations and miR downregulation in the RTK, PI3K, MAPK, and TGF β pathways across all cervical tumors.



Extended Data Figure 10 | Pathway biomarkers differentiating squamous cell carcinomas and adenocarcinomas. a, Cytoscape display of the largest interconnected regulatory network of PARADIGM pathway features differentially activated between squamous cell carcinomas and adenocarcinomas connected through hubs with ≥ 10 downstream targets. Hubs with ≥ 10 downstream targets are labeled. Genes showing

mRNA-miRNA expression anti-correlation with strong evidence support are highlighted with thicker black outline and labeled. Top differentially expressed genes relating to immune function are also labeled. Node size is proportional to significance of differential activation. b, Zoom-in display of the p63 sub-network neighborhood. First neighbors (upstream or downstream) of four p63 complexes (bold text) are displayed in this view.



Extended Data Figure 11 | HPV integration and molecular characteristics in cervical cancer. **a**, E6 unspliced/spliced ratio for HPV16 and HPV18 intragenic, enhancer, and intergenic sites. HPV16: median = 0.44 ($n = 102$), HPV18: median = 0.93 ($n = 40$). The p-value is from a two-sided Kolmogorov-Smirnov test. **b**, Distribution of RNAseq-based EMT score for HPV-negative (HPV-) and HPV-positive (HPV+) samples ($n = 178$). **c**, Distributions of SCNA and mRNA abundance ranks

(left panel) and distribution functions for SCNA and mRNA abundance ranks with 100 random expectation samples close to the diagonals (grey) (right panel) for genomic loci integrated with HPV16. **d**, Distributions described in **c** for genomic loci integrated with HPV18. BH-corrected p-values for the SCNA and mRNA abundance ranks (median p-values) are reported.