

Comprehensive Analysis of Alternative Splicing in Gastric Cancer Identifies Epithelial–Mesenchymal Transition Subtypes Associated with Survival



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ABSTRACT

Alternatively spliced RNA isoforms are a hallmark of tumors, but their nature, prevalence, and clinical implications in gastric cancer have not been comprehensively characterized. We systematically profiled the splicing landscape of 83 gastric tumors and matched normal mucosa, identifying and experimentally validating eight splicing events that can classify all gastric cancers into three subtypes: epithelial-splicing (EpiS), mesenchymal-splicing (MesS), and hybrid-splicing. These subtypes were associated with distinct molecular signatures and epithelial–mesenchymal transition markers. Subtype-specific splicing events were enriched in motifs for splicing factors RBM24 and ESRP1, which were upregulated in

MesS and EpiS tumors, respectively. A simple classifier based only on RNA levels of *RBM24* and *ESRP1*, which can be readily implemented in the clinic, was sufficient to distinguish gastric cancer subtypes and predict patient survival in multiple independent patient cohorts. Overall, this study provides insights into alternative splicing in gastric cancer and the potential clinical utility of splicing-based patient classification.

Significance: This study presents a comprehensive analysis of alternative splicing in the context of patient classification, molecular mechanisms, and prognosis in gastric cancer.

Introduction

Alternative splicing (AS) is a major mechanism for increasing transcript diversity in eukaryotes, affecting approximately 95% of multi-exon genes that express multiple spliced isoforms in humans (1, 2). Dozens of aberrant splicing variants have been implicated in human disease, including cancer (3–5). These variants play roles in a broad spectrum of oncogenic processes, such as cell proliferation, apoptosis, hypoxia, angiogenesis, and immune escape (3–5). Aberrant

splicing has also been reported to promote epithelial–mesenchymal transition (EMT) and contribute to the invasion and metastasis of various tumor types, including breast, lung, and gastric cancer (6, 7).

Gastric cancer is the fifth most frequently diagnosed cancer and the third leading cause of cancer-related mortality worldwide (8). A large-scale project for genomic characterization of cancer, The Cancer Genome Atlas (TCGA) consortium, revealed that gastric cancer is highly heterogeneous and, based on mutations, copy number changes, and translocations, can be divided into four distinct tumor subtypes: Epstein-Barr virus (EBV)-associated, chromosomal instable (CIN), microsatellite instable (MSI), and genomically stable (GS; ref. 9). However, TCGA classification does not correlate well with oncologic outcome and thus is of limited prognostic utility (9). In a separate cohort, the Asian Cancer Research Group (ACRG), four gastric tumor subtypes were defined on the basis of gene expression patterns: MSI, Microsatellite Stable/EMT (MSS/EMT), MSS/TP53⁺, and MSS/TP53⁻. The ACRG subtypes are more closely associated with patient outcome, with the MSI and MSS/EMT patient subtypes having the best and worst prognosis, respectively (10). Neither of these classification schemes use AS profiles to stratify patients with respect to outcomes or treatment efficacy. AS-based classifications have been shown to accomplish these goals in other tumor types (11, 12) and therefore could be leveraged to improve patient stratification in gastric cancer.

Several transcriptomic studies have reported that AS of genes implicated in gastric cancer, such as skipping exon 2 or exons 18–19 of the receptor tyrosine kinase proto-oncogene *MET* (9), and isoform-specific expression of *ZAK* kinase (13). In addition, EBV-associated gastric cancers display a unique AS signature and changes in splicing factor levels compared with EBV-negative gastric cancers (14). Recent reanalysis of the TCGA database focusing on gastric cancer and gastrointestinal pan-cancer cohorts identified hundreds of tumor-associated AS events, several of which were associated with differential patient survival (15–18). Yet the role of AS in gastric cancer pathogenesis, and its potential clinical utility, has not been fully explored due to the lack of transcription-wide analysis on matched normal-tumor

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pair samples. A systematic analysis of AS events in matched normal-tumor gastric samples would allow identification of cancer-specific events, in addition to intertumor AS variation, and is currently lacking. Furthermore, deeper insights into AS events and their regulation are greatly needed to better understand the fundamental molecular features of, and potential causative mechanisms leading to, gastric cancer.

Here, we report an AS-based stratification of gastric cancer cases based on a transcriptome-wide analysis of a new cohort of 83 tumors and matched normal samples. We identify eight AS events and three RNA-binding protein (RBP) splicing factors, and demonstrate that either an AS- or RBP-based signature can be used to classify patients from multiple cohorts into subtypes with distinct clinical outcomes.

Materials and Methods

Samples

We selected 83 patients with gastric cancer with tumor and matched normal samples (Supplementary Table S1) from the fresh snap-frozen tissue repository at the Laboratory of Gastric Cancer Biology, Cancer Research Institute, Seoul National University, South Korea (from 2001 to 2015). All cases were staged according to the AJCC TNM 7th edition staging system. Pathological analysis classified papillary, well-differentiated and moderately differentiated types as the differentiated group, whereas poorly differentiated, mucinous, poorly cohesive cell types were classified as the undifferentiated group. For MSI, fragment analysis was used to compare tumor and normal tissue samples at 5 bp locations after PCR using Primer 1 consisted of BAT26 (116 bp) and BAT25 (148 bp), and primer 2 consisted of D5S346 (96–122 bp), D17S250 (146–165 bp) and D2S123 (144–174 bp). Study protocol was approved by the institutional review board (IRB H-2001-044-1092) and all methods were carried out in accordance with relevant guidelines and regulations.

RNA sequencing and data analysis

Total RNA was extracted from tumor and corresponding normal gastric mucosa samples and libraries were prepared using the TruSeq RNA Kit 2 per manufacturer's instructions. RNA sequencing (RNA-seq) was performed with Illumina HiSeq2000 (Illumina Inc.) for 101-mer paired-end reads. Reads were aligned against the human reference genome (GRCh38, Ensembl 94) using STAR (19) 2-pass mode v2.6.1. d and RSEM (20) v1.3.1 was used to quantify transcriptome abundance. Transcript abundance was analyzed by DESeq2 (21), and variance stabilizing transformation was used for downstream analysis. Gene set enrichment analysis (GSEA) was used to identify overrepresented biological functions using GSEA (22). We calculated enrichment scores using gene sets from MSigDB Hallmark ($n = 50$) and Kyoto Encyclopedia of Genes and Genomes (KEGG; $n = 186$) gene sets collection (23).

AS analysis

We used rMATS v4.0.2 (24) to identify differential AS events between two groups of samples. To keep high-rMATS confidence calls, we filtered out cases where the sum of inclusion junction counts (IJC) and skipping junction counts (SJC) was <25 in either set of samples. Differentially spliced exons were defined using a splicing ratio $|\text{PSI} (\text{Percent-Spliced-In})| > 0.2$ and $\text{FDR} < 0.05$ as thresholds. AS frequency for each patient was calculated as the total number of five major types of AS events.

Molecular subtype classification

To find variable AS events for patient subtyping, we performed hierarchical clustering using ComplexHeatmap v2.2.0 with ΔPSI values of 20 differential AS events between tumor and matched normal tissues, and focused on 5 variable AS events selected by visual

inspection of variability in the heatmap. Next, we defined two patient groups defined by principal component analysis using ΔPSI values of these 5 variable AS events. To expand the list of variable AS events with similar behavior, we compared AS events in group 1 versus group 2 using rMATS, yielding 28 variable AS events (Supplementary Fig. S2).

We applied the TCGA and ACRG subtype classifications to our gene expression data. We used approximately 800 signature gene classifiers (25) from the TCGA classification (9) to divide patients into four groups (EBV, MSI, GS, and CIN, Supplementary Fig. S22). Signature scores based on ACRG classification signature gene lists (e.g., MSI, EMT, and TP53 activity) were calculated as described previously in (Supplementary Fig. S23; ref. 10).

For the RBP-expression-based patient classification scheme, we used an average expression value of *RBM24* and *ESRP1* to divide patients into high and low expression groups. We defined the MesS (mesenchymal-splicing)-like subtype as *RBM24*_{high} and *ESRP1*_{low}, and EpiS (epithelial-splicing)-like subtype as *RBM24*_{low} and *ESRP1*_{high}. For the three RBP-based classifications, we used the average of both *RBM24* and *ESRP1* for the EpiS subtype.

Somatic mutation and copy number variation analysis

Exome enrichment, whole-exome sequencing, as well as copy number, mutation, and variant calling analysis are described previously in Supplementary Methods.

RBP motif enrichment analysis

Motif analysis for 114 RBPs in differential exon skipping events was performed using rMAPS (26), comparing differentially skipped exons of specific subtypes with control exons. 24,704 exons control exons, not differentially spliced, were defined as $\text{FDR} > 0.5$, and $\text{maxPSI} > 0.15$, and $\text{minPSI} < 0.85$. Regions for motif analysis included: 250 bp upstream or downstream into the intronic sequences and the first and last 50 bp of exonic sequences. To avoid enrichment of canonical splice sites, rMAPS excluded 20 bp within the 3' splice site and 6 bp within 5' splice site in introns. P value were calculated for each motif after counting the number of occurrences in skipped exons and control exons, and enriched motifs were identified as $P < 0.001$. To build a correlation network, we calculated Pearson correlation coefficients between tumor PSI and RBP expression for each pair of AS event and RBP. P values were converted into FDR for multiple test correction. Significant correlation pairs ($\text{FDR} < 0.05$) were used for network construction.

RNA extraction from tumors and normal tissues

50–100 mg of frozen gastric tumor or adjacent matched normal tissue was pulverized using a mortar with liquid nitrogen. Total RNA was extracted using the RNeasy kit (QIAGEN), including DNase I treatment per the manufacturer's instructions. RNA concentration and quality were assessed using a NanoDrop 2000 (Thermo Fisher Scientific) as well as an Agilent RNA 6000 Pico Kit with Agilent 2100 Bioanalyzer (Agilent). Samples with RNA Integration number >4.0 were used.

Cell cultures and RT-PCR for splicing events validations

AGS, MKN-74, and MKN-1 human gastric cancer cell lines were obtained from the Korean Cell Line Bank (KCLB, Seoul, Korea), and used in experiments after being certified by the KCLB. All cell lines were cultured in RPMI-1640 cell culture media (Gibco, Life Technologies) supplemented with 10% FBS (Gibco) and 1% penicillin-streptomycin (Gibco) and subcultured at intervals of 2–3 days. All cell lines were grown in 5% CO_2 humidified incubator at 37°C and *Mycoplasma* testing was routinely performed using e-Myco plus

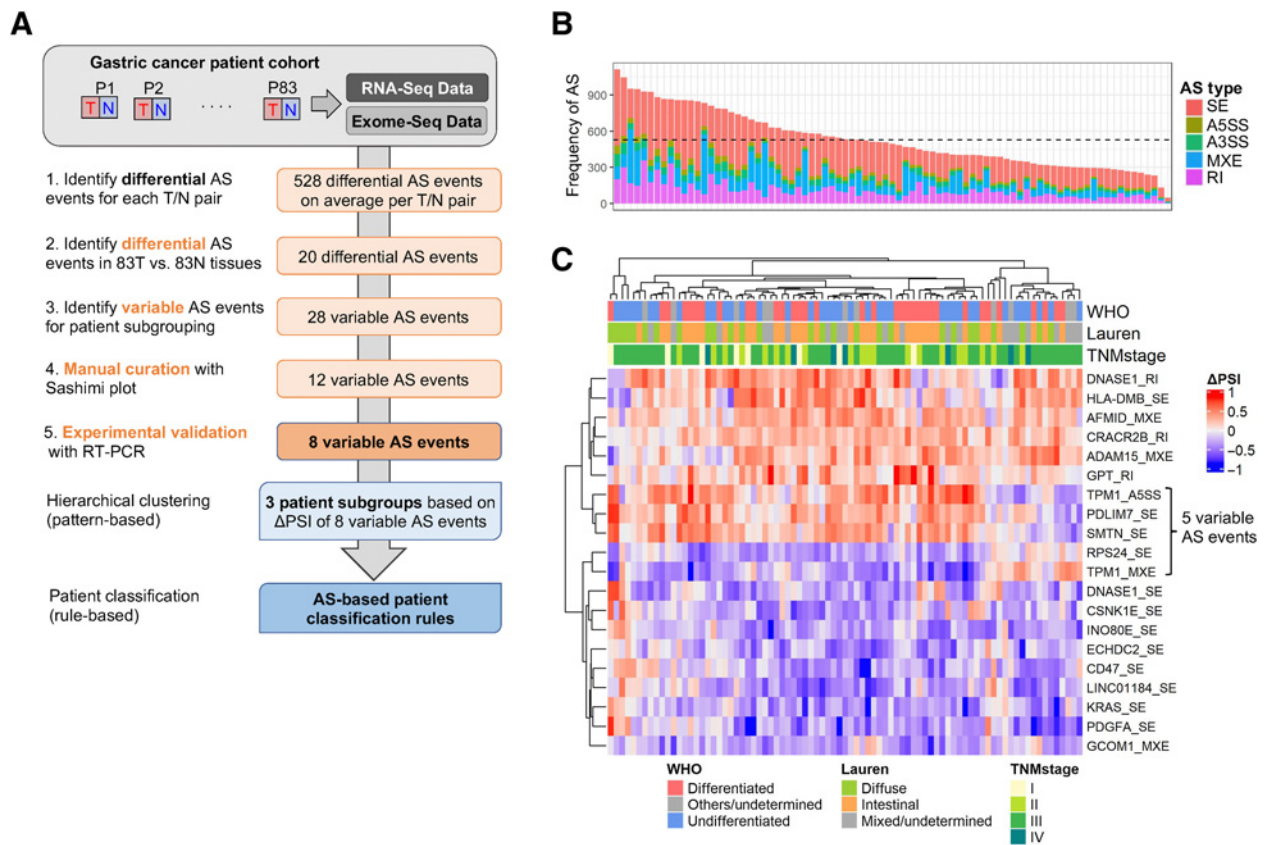


Figure 1.

Cancer-associated AS events in gastric tumors. **A**, Summary of AS-based patient classification scheme. N, normal tissue; P, patient; SNU, Seoul National University; T, tumor. **B**, Number of differential AS events in tumor versus normal pairs, colored by event type (SE, skipped exon; A5/3SS, alternative 5'/3' splice site; MXE, mutually exclusive exons; RI, retained intron). The mean number of AS events is shown as a horizontal dashed line. **C**, Differential AS events in 83 tumors versus normal tissues, plotted as Δ PSI values for each patient. WHO and Lauren classification, as well as tumor stage, are shown.

Mycoplasma PCR Detection Kit (Intron), verifying that the cells were *Mycoplasma* free. For the silencing of ESRP1 or RBM24 expression, cells were transfected with ESRP1 siRNA (siESRP1; Santa Cruz Biotechnology; sc-77526), RBM24 siRNA (sc-95527), or nontargeted siRNA (sc-37007) using Lipofectamine 2000 transfection reagent (Life Technologies) according to the manufacturer's protocol. Two microgram of total RNA was reverse-transcribed with Superscript III reverse transcriptase (Invitrogen) and cDNA from 10 ng of total RNA was subsequently used for each reaction. RT-PCR was performed with Solg Taq DNA Polymerase (SolGent co., Ltd.) using oligonucleotide primers detecting both the included and skipped isoforms (Supplementary Table S5). PCR products were separated on a 2% or 2.5% agarose gel stained with Dyne LoadingSTAR (DYNEBIO INC.). Bands corresponding to the included and skipped isoforms were quantified with the Image Laboratory 6.0 software (Bio-Rad) and for each splicing event exon inclusion was expressed as PSI as (intensity included)/(intensity included + skipped).

RT-PCR for splicing events validations

500 ng of total RNA was reverse-transcribed with Superscript III reverse transcriptase (Invitrogen) and cDNA from 5–10 ng of total RNA was used for each reaction. Semiquantitative PCR was performed with Hot-start Taq DNA polymerase (Thomas Scientific) using primers (IDT) detecting both included and skipped isoforms (Supplementary

Table S6). Optimal PCR conditions were defined for each primer pair to select semiquantitative conditions. PCR products were separated on a 2% agarose gel stained with SYBR Safe (Invitrogen). Bands corresponding to included and skipped isoforms were quantified with a ChemiDoc MP (Bio-Rad) and for each event exon inclusion was expressed as PSI as (intensity included)/(intensity included + skipped).

Quantitative PCR analysis of RBPs

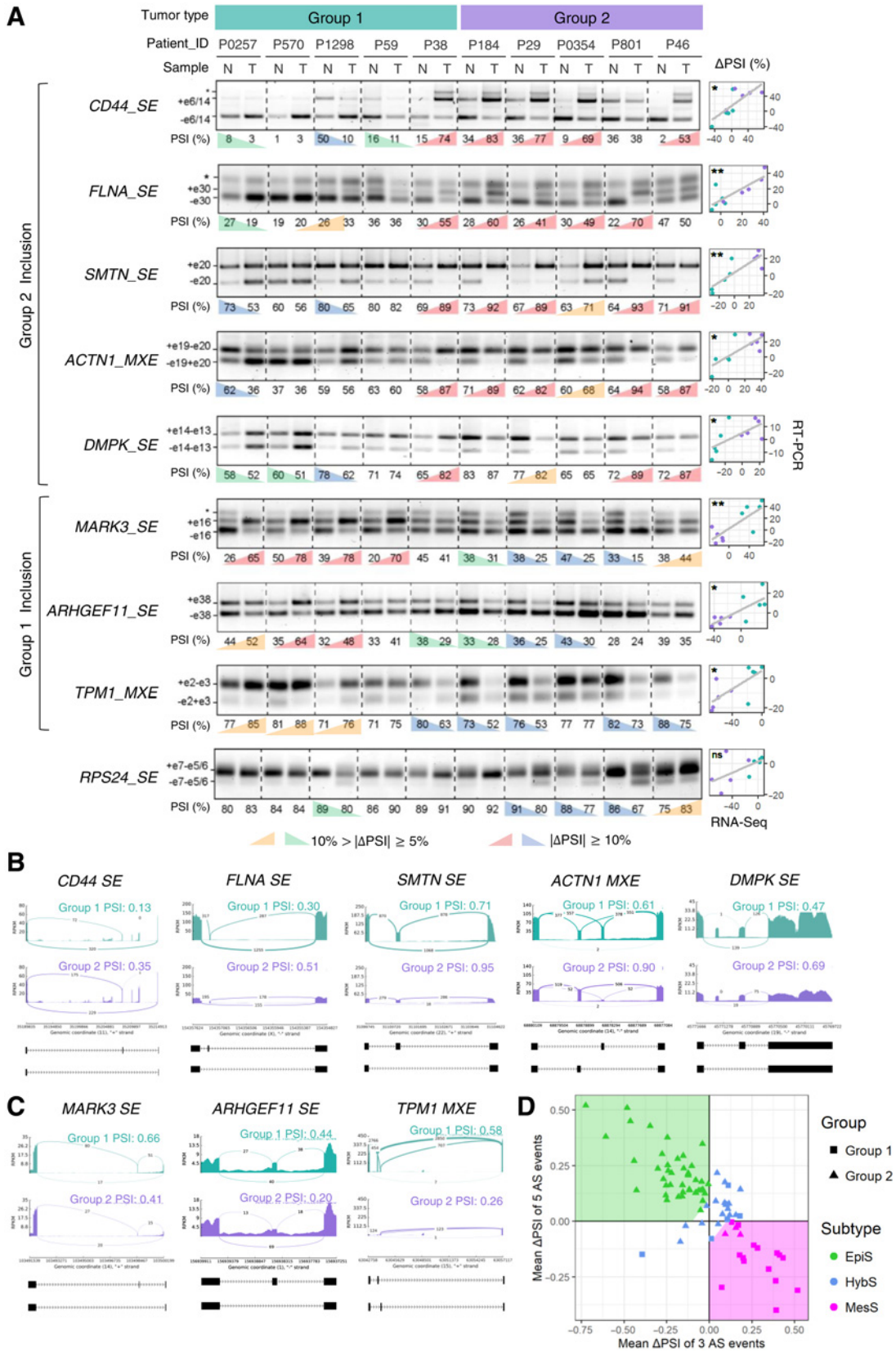
qPCR was performed using cDNA as above with iTaq Universal SYBR Green Supermix (Bio-Rad) using gene-specific oligonucleotide primers (IDT; Supplementary Table S7) per the manufacturer's instructions using a ViiA7 (Life Technologies). Expression for each target gene was normalized to GAPDH, and then normalized across all samples.

Validation in independent patient cohorts

RNA-seq data of paired normal and tumor samples for 80 early-onset gastric cancers from Mun and colleagues (SRP172499; ref. 27) were processed as described above. Clinical data and gene expression profiles of 295 TCGA gastric cancers were downloaded from cBioportal (28, 29). ACRG microarray data (GSE62254; ref. 10) were analyzed using RMA normalization.

Statistical analysis

Fisher exact and χ^2 tests were performed to evaluate differences between subgroups. Log-rank and cox regression analysis were used to



test survival differences. Kaplan–Meier survival analysis was performed using the survival package and Cox proportional hazards regression for multivariable survival analysis was applied using the *coxph* function. All analyses were performed using R v3.6.0.

Results

Comprehensive genomic, transcriptomic, and splicing profiles of matched gastric tumor and normal tissue samples

We performed RNA-seq of paired tumor and adjacent normal mucosa tissues from 83 patients with gastric cancer. On the basis of the Lauren classification, the patient cohort included 35 intestinal, 27 diffuse, and 21 mixed or undetermined types of gastric cancer (Supplementary Table S1). We generated 9.5 billion paired-end reads, with an average of 64.2 million read pairs per tumor, and 49.8 million read pairs for the corresponding normal mucosal samples (Supplementary Fig. S1). In addition, we performed whole-exome sequencing at an average read depth of 129X for the tumor samples and an average read depth of 85X for the matched normal samples.

Taking advantage of the paired sampling that was available for all of our patients (Fig. 1A), we first looked for differential AS events between tumor and normal tissue. We used rMATS, a computational tool for splicing analysis that uses both exon body and exon junction reads from RNA-seq data, to quantify the “Percent Spliced-In” (PSI) value for each AS event (24). We determined AS profile differences between each tumor and paired normal tissue using a stringent cutoff value of at least 20% change in PSI (Δ PSI). We detected on average 528 AS events in each tumor versus matched normal pair, with skipped exon (SE) events being the most frequent (Fig. 1B). We further identified 20 AS events in 18 genes that are differentially spliced between tumors and normal tissues, including 12 SE events, four mutually exclusive exons (MXE), three retained introns (RI), and one alternative 5' splice site (A5SS; Fig. 1C; Supplementary Table S2). Spliced genes were associated with cell adhesion and the cytoskeleton (e.g., *SMTN*, *TPM1*, *PDLIM7*, and *ADAM15*), translational control (e.g., *RPS24*), cell proliferation (e.g., *PDGFA*, *KRAS*), cell metabolism (e.g., *AFMID*, *ECHDC2*, *INOE80*, and *GPT*), or immune cell signaling (e.g., *CD47*). Our analysis also detected a skipped exon in the *KRAS* oncogene that was previously identified in a smaller cohort of patients with gastric cancer (30), as well as mutually exclusive exons in *AFMID* detected in hepatocellular carcinoma (31).

Patient subtype classification based on AS events

We next sought to identify AS events that could be used for patient classification. These were defined as AS events that exhibit variable profiles across tumors (hereafter referred to as *variable* AS events). Five such variable AS events were identified upon hierarchical clustering of the 20 differential AS events (Fig. 1C). We then divided the patients with cancer into two groups, Group 1 and Group 2, based on the patterns of these five variable AS events. By comparing Groups 1 and 2, 23 additional variable AS events were identified to facilitate the further

dichotomization of patients (See Materials and Methods). This analysis yielded a total of 28 variable AS events in 13 genes (Supplementary Fig. S2A and S2B). Because many of these events were partially overlapping in the same gene (e.g., *TPM1* contains eight AS events) and rMATS reports each AS event separately, all events were manually inspected, and merged to produce a group of 12 nonredundant variable AS events in nine genes (Supplementary Table S3).

We experimentally validated the splicing patterns of these nine genes from Groups 1 and 2, using an adjacent section of tumor and matched normal tissues from 10 of the patients originally used for RNA-seq analysis. Differential splicing patterns detected by our transcriptome analysis between tumor and matched normal samples were measured using semiquantitative RT-PCR for AS events in nine genes (Fig. 2A). The Δ PSI values were determined for each tumor-normal pair, and eight out of nine AS events showed a significant positive correlation ($P < 0.04$) of Δ PSI values between RNA-seq data and the RT-PCR results (Fig. 2A). We also observed that these eight validated AS events could be divided into two classes where the “Group 1 inclusion” consisted of 5 AS events (i.e., SE events in *CD44*, *FLNA*, *SMTN*, *DMPK*, and an MXE event in *ACTN1*) and the “Group 2 inclusion” consisted of 3 AS events (i.e., SE events in *MARK3*, *ARHGEF11*, and an MXE event in *TPM1*). Inclusion patterns of these 8 AS events in the two patient groups were visualized using Sashimi plots (Fig. 2B and C).

Hierarchical clustering of patients based on Δ PSI values of these 8 events showed that patients can be divided into two distinct subgroups plus one subgroup with an intermediate pattern (Supplementary Fig. S3). Plotting the average Δ PSI values of the Group 1 inclusion events versus Group 2 inclusion events showed a significant negative correlation with a mutually exclusive pattern between these two Δ PSI variables ($r = -0.85$, $P < 2.2e-16$, Fig. 2D), and suggested a simple AS-based patient classification scheme.

According to this novel AS event-based classification scheme, we divided patients with gastric cancer into three subtypes (Fig. 2D): EpiS (top left region), MesS (bottom right region), and Hybrid-Epithelial/MesS (HybS, intermediate region), which were named according to the results of GSEA, described below. This new classification scheme can be applied to any single patient without the need for complex clustering algorithms.

Molecular characterization of AS event-based gastric cancer subtypes

The AS-based classification divided 83 patients into 43 EpiS, 22 HybS, and 18 MesS subtypes (Fig. 3A). At the molecular level, the frequency of AS events, mutation rate, and copy number alterations were significantly higher in the EpiS subtype compared with the MesS subtype (Fig. 3B–D). The average number of AS events, as well as the total number of SE events, per tumor sample was significantly higher than in EpiS tumors compared with MesS tumors (Fig. 3B). Five genes (i.e., *LRP1B*, *ZFH4*, *ROBO1*, *PLEC*, and *AHNAK*) were frequently mutated (>20%) in EpiS but not in MesS tumors and *MUC16* similarly showed a higher mutation rate in EpiS compared with MesS

Figure 2.

Eight AS events display distinct patterns in two patient groups. **A**, RT-PCR validation of AS events in gastric tumors (T) and matched normal (N) tissues from 10 patients, using primers that amplify both included and skipped isoforms. For each AS event, a representative gel image is shown along with PSI values, calculated as the percentage of included/(%included + %skipped); significant Δ PSI (T–N) values ($n = 3$, Δ PSI >|10%| and Δ PSI >|5%|) are colored. Exons included (+) or skipped (–) are annotated (*, bands of no interest). Correlations of Δ PSI values between RT-PCR and RNA-seq experiments are shown on the right. *, $P < 0.05$; **, $P < 0.01$; ns, not significant. **B** and **C**, Read coverage and junction reads showing inclusion of AS events that define “Group 2” (**B**) and “Group 1” (**C**) in gastric tumors. Gene name and AS type are indicated, along with average PSI for tumors. **D**, Scatter plot of patients using mean(Δ PSI; Group 2 vs. 1). The areas for three subtypes are shown in different background colors—EpiS, green; MesS, magenta; and HybS, white, where the x and y intercepts of linear boundaries are 0.1.

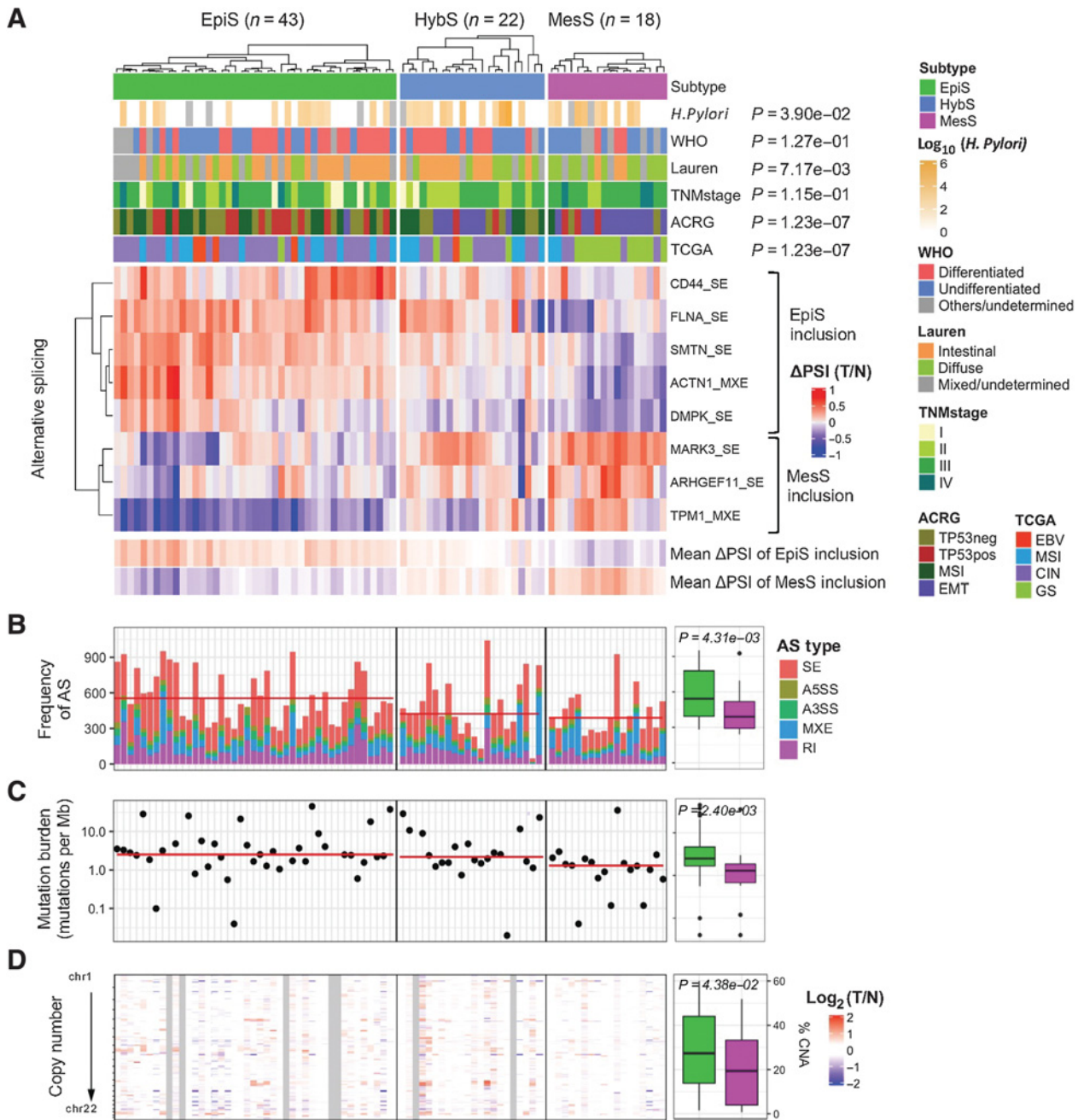


Figure 3.

AS event-based patient classification defines the EpiS and MesS subtypes of gastric cancer. **A**, Clinical information, including *H. pylori* status, and ACRG and TCGA classifications for EpiS, MesS, and HybS subtypes defined on the basis of AS events are shown. *P* values from χ^2 enrichment or Wilcoxon rank-sum tests are shown. For each event, Δ PSI for each tumor-normal pair are shown. **B**, The frequency of AS events for each patient is the sum of differential AS events in matched tumor versus normal samples. **C** and **D**, Difference in mutation burden (**C**) and copy number alteration (**D**) between EpiS and MesS subtypes are shown, along with average values, and Wilcoxon rank-sum test *P* values.

(Supplementary Fig. S4A). No mutations in splicing regulatory genes were found to be significantly associated with our subtypes (Supplementary Table S4; Supplementary Fig. S4B). The EpiS subtype also exhibits copy number gains in several regions, including regions overlapping the known cancer-associated genes *EGFR*, *ERBB2*, *MYC*, and *CCNE1*, which are involved in cancer cell proliferation (Supplementary Fig. S4C). Copy number alterations were relatively rare in

MesS patients, similar to the TCGA GS subtype. Furthermore, the copy number alterations observed in the MesS subtype were also present in the EpiS subtype, whereas most copy number amplifications in driver oncogenes were not present in the MesS subtype (Supplementary Fig. S4C).

To further characterize the patient subtypes defined by eight AS events, we performed GSEA using total gene expression data

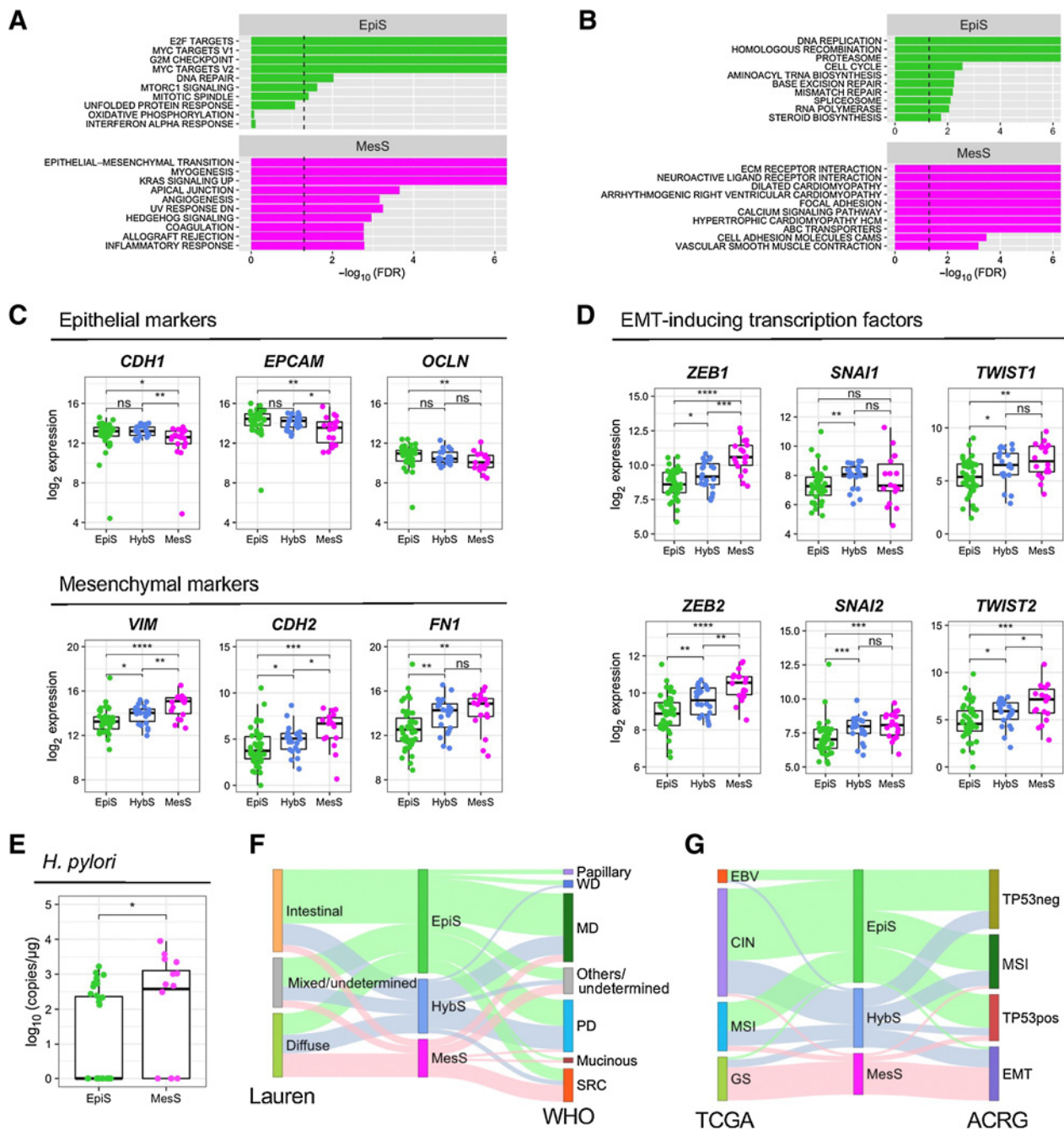


Figure 4. Functional characterization of AS subtypes. **A** and **B**, Statistically enriched gene sets in EpiS and MesS subtypes identified using GSEA 50 Hallmark gene set (**A**) and MSigDB KEGG pathways (**B**). **C**, Gene expression of epithelial *CDH1*, *EPCAM* and *OCLN*, and mesenchymal *VIM*, *CDH2* and *FN1* markers in each patient subtype. **D**, Gene expression of EMT-inducing transcription factors *ZEB1/2*, *SNAI1/2*, and *TWIST1/2* in each patient subtype. **E**, Absolute quantification of *H. pylori* copies in each patient subtype. **F**, Associations between AS event-based patient subtype and Lauren or WHO clinical subtypes. MD, moderately differentiated; PD, poorly differentiated; SRC, signet ring cell carcinoma; WD, well differentiated. **G**, Associations between AS event-based patient subtype and TCGA or ACRG molecular subtypes. Wilcoxon rank-sum test, *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; ****, $P < 0.0001$; ns, nonsignificant.

between the EpiS and MesS subtypes (Fig. 4A and B; Supplementary Fig. S5; ref. 23). The EpiS subtype showed an enrichment of terms related to E2 factor (E2F) targets, MYC targets, and cell-cycle regulation, which are typical proliferation signals for epithelial cancer cells (Fig. 4A), as well as DNA damage response pathways

(Fig. 4B). Higher mutation rates and copy number amplifications detected in the EpiS subtype (Fig. 3C and D) are consistent with the activation of DNA damage response pathways detected by GSEA. The MesS subtype showed various terms related to EMT (Fig. 4A), as well as extracellular matrix-receptor interaction, focal adhesion,

and cell adhesion molecules (Fig. 4B). Thus, the characteristic biological signature associated with our AS event-based patient classification is the EMT process, which we used to assign the tumor subtypes defined above.

To confirm the involvement of EMT in our patient subtypes, we investigated the expression of well-established EMT marker genes. In accordance with our GSEA, epithelial marker genes *CDH1*, *EPCAM* and *OCLN* showed higher expression in the EpiS subtype than in the other two subtypes, whereas mesenchymal marker genes *VIM*, *CDH2* and *FN1* showed higher expression in the MesS subtype (Fig. 4C). Of note, among the 8 variable AS events, aberrant splicing of *ARHGGEF11* and *CD44* is known to play important roles during EMT in diverse cancers (32, 33). We further examined the expression of known EMT-inducing transcription factors (TF), such as *ZEB1* and *ZEB2*, *SNAI1* and *SNAI2* (also known as Snail and Slug), as well as *TWIST1* and *TWIST2* (34). Gene expression of all these EMT TFs, except *SNAI1*, gradually increased across subtypes in the order EpiS-HyS-MesS (Fig. 4D), highlighting differences in EMT between our three patient subtypes. Expression of other TFs associated with EMT, such as *TCF4*, *KLF8*, *FOXO2* increased between EpiS and MesS, whereas *TCF3*, *GRHL2*, *ELF3*, and *H2AFX* decreased (Supplementary Fig. S6), consistent with the overall enrichment of mesenchymal markers in the MesS group.

Finally, we compared our AS event-based patient classification scheme with established clinical and molecular subtype classifications. MesS tumors exhibit higher *Helicobacter pylori* (*H. pylori*) copy numbers compared with EpiS tumors ($P = 3.9e-02$; Fig. 4E; Supplementary Table S1), in line with previous findings suggesting that *H. pylori* promotes EMT in gastric cancer (35, 36). In terms of the Lauren classification, we observed a significant association between our AS event-based MesS subtype and the diffuse subtype, whereas patients classified as EpiS were enriched in the intestinal subtype ($P = 7.17e-03$; Fig. 4F). Compared with the TCGA and ACRG molecular classifications, our AS event-based MesS subtype, which rarely contains copy number alterations, was significantly associated with the TCGA GS subtype ($P = 1.23e-07$) as well as with the ACRG EMT subtype ($P = 1.23e-07$; Fig. 4G).

Splicing factors RBM24, RBMS1, and ESRP1 regulate AS events in gastric cancer subtypes

RBPs regulate AS in a dose-dependent manner by binding specific sequences on the pre-mRNA and promoting or blocking exon inclusion. To investigate the molecular mechanisms regulating differential AS events between the EpiS and MesS subtypes, we performed an RBP-binding motif enrichment analysis using rMAPS (26). For the five AS events included in EpiS tumors, we identified binding sites for nine RBPs specifically enriched in exonic or intronic regions near the spliced exons (Fig. 5A; Supplementary Table S5; Supplementary Fig. S7). Similarly, we found binding motifs for four RBPs enriched in the three AS events included in MesS tumors. For example, the ESRP1-binding motif was enriched in upstream introns of AS events included in MesS tumors ($P = 2.18e-4$; Fig. 5A), whereas the RBM24-binding motif was enriched in downstream introns of AS events included in EpiS tumors ($P = 6.61e-07$; Fig. 5A). Finally, AS events from both subtypes contained motifs for two RBPs, PCBP1, and PCBP2, which are known oncogenic-splicing factors that are also involved in translation coactivation and RNA stability (37).

Next, we investigated whether these RBPs were differentially expressed between MesS and EpiS patient subtypes. Among the 13 RBPs that exhibit enriched binding sites, *RBM24* and *RBMS1* were upregulated in the MesS subtype, whereas *ESRP1* and *PTBPI* were

upregulated in the EpiS subtype (Fig. 5B; Supplementary Fig. S8A). We also examined the data for any evidence of a regulatory relationship between RBP level and the Δ PSI values of all eight AS events. Gene expressions of *RBM24*, *RBMS1*, and *ESRP1* were significantly correlated with the level of exon inclusion for most AS events (Supplementary Fig. S8B and S8C), supporting a regulatory role for *RBM24*, *RBMS1*, and *ESRP1* in modulating these eight AS events. A correlation network was built using significantly correlated RBP-AS event pairs (FDR < 0.05), and revealed *RBM24*, *RBMS1*, and *ESRP1*, but also *PTBPI*, to be major nodes connecting all eight variable AS events (Fig. 5C). A few additional RBPs (i.e., *CELF5*, *PCBP2*, *CELF4*, *FUS*, *SRSF3*, and *RBM38*) seem to play auxiliary roles in regulation of these specific AS events, connecting only a subset.

The correlation between the expression of regulatory RBPs *RBM24*, *RBMS1*, and *ESRP1*, and exon inclusion of all eight MesS and EpiS AS events was further validated experimentally (Fig. 5D and E; Supplementary Fig. S8B and S8C). Differential RBP expression was measured with quantitative PCR and mirrored RNA-seq data almost perfectly. Notably, *RBM24* expression showed a significant correlation ($r > |0.5|$) with all five AS events from EpiS tumors, whereas *RBMS1*'s correlation was significant only for *CD44* and *ACTN1* AS events (Fig. 5D and E). *ESRP1* expression correlated with all three AS events in *MARK3*, *ARHGGEF11*, and *TPM1* from MesS tumors (Fig. 5E). Of note, *ESRP1*-regulated AS of *ARHGGEF11* has been shown to be essential for epithelial tight junction integrity (38). Thus, *RBM24* and *ESRP1* may be critical regulatory RBPs in gastric cancer, acting oppositely in the EpiS and MesS subtypes. Our working model is that, in the EpiS subtype, patients with low expression of *RBM24* and *RBMS1* and high expression of *ESRP1* show inclusion of "EpiS exons" and skipping of "MesS exons" (Fig. 5F), whereas we propose the converse to be the case in MesS subtype patients.

We performed siRNA-mediated *ESRP1* knockdown in AGS and MKN-74 epithelial-like gastric cancer cell lines, and *RBM24* knockdown in MKN-1 mesenchymal-like gastric cancer cells to demonstrate the causal role of these RBPs in the regulation of the splicing events that constitute our splicing-based signature of gastric tumors (39). At baseline, the epithelial-like gastric cancer cell lines expressed high levels of *ESRP1* and low levels of *RBM24*, whereas mesenchymal-like gastric cancer cells showed opposite expression patterns (Supplementary Fig. S9A). Similarly, at baseline, epithelial-like and mesenchymal-like gastric cancer cells often show opposite splicing patterns (Supplementary Fig. S9B). Knockdown of *ESRP1* significantly decreased inclusion of exons in *CD44* (in AGS and MKN-74) and *ACTN1* (in AGS), two EpiS-splicing events, and increased exon inclusion in *ARHGGEF11* (in AGS and MKN-74), a MesS-splicing event, compared with scrambled siRNA control (Supplementary Fig. S9B). Similarly, knockdown of *RBM24* in MKN-1 cells significantly increased exon inclusion in *ACTN1*, an EpiS-splicing event, and decreased exon inclusion in *TPM1*, a MesS-splicing event. Exons of *MARK3* in MKN-1 were either fully included in either cell line or fully skipped and were not affected by RBP knockdowns.

Validation of the AS event signature using independent cohorts

To ascertain the robustness of our findings, we verified whether the AS event-based classification and molecular findings were reproducible in other patient cohorts. We looked for public datasets that included (i) reasonably large number of patients; (ii) matched tumor and normal samples; and (iii) raw RNA-seq data available for analyzing splicing variants. The proteogenomic study by Mun and colleagues (27) met these conditions with 80 early-onset patients with gastric cancer even though most of them were of the Lauren diffuse subtype. Analysis of this cohort reproduced our salient findings, including (i) classification using

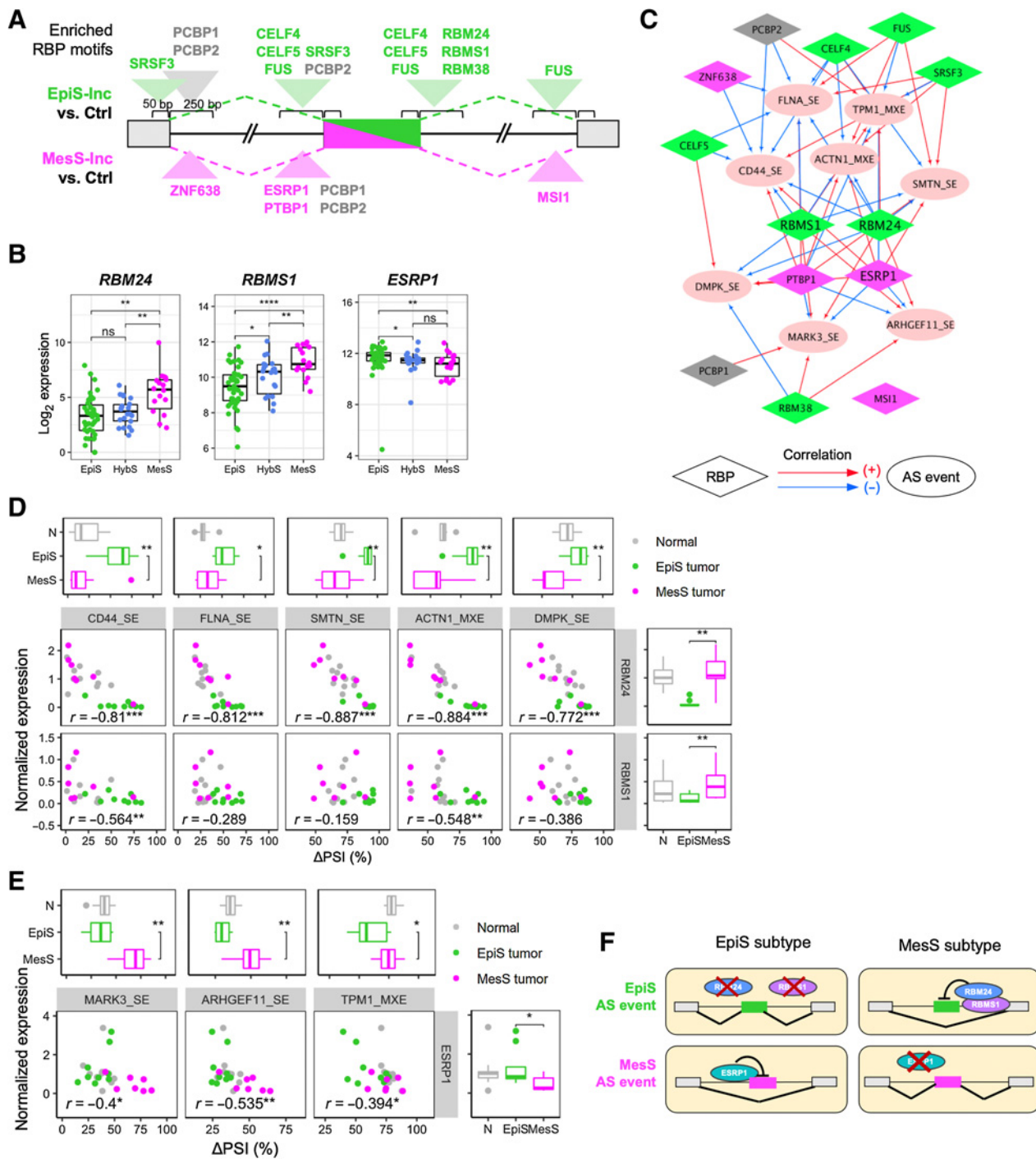


Figure 5.

RBPs as regulators of AS events in EpiS and MesS tumor subtypes. **A**, RBP motifs enriched near differentially included exons in EpiS and MesS subtypes. See also Supplementary Table S5. **B**, Normalized RBP gene expression in each patient subtype. **C**, Correlation network between RBP expression and Δ PSI values of each AS event. Positive and negative correlations are shown with red and blue arrows, respectively. Colors for RBP nodes indicate the subtypes of RBP motifs enriched in **A**. **D** and **E**, Correlation between RBP expression and AS for each RBP-AS pair, for EpiS (**D**) and MesS (**E**) AS events, colored by sample type. Box plots indicate subtype differences in AS event (top) and RBP expression (right). **F**, Model of AS regulation by RBP expression in gastric cancer. Wilcoxon rank-sum test, *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; ****, $P < 0.0001$; ns, nonsignificant.

the eight AS events grouped patients into three distinct subtypes (Fig. 6A and B); (ii) subtypes displayed differential expression of epithelial (*CDH1*, *EPCAM*, and *OCN*) and mesenchymal (*VIM*, *CDH2*, and *FN1*) markers; and (iii) *RBM24* and *RBMS1* were upregulated and *ESRP1* downregulated in the MesS subtype (Supplementary Fig. S10). Interestingly, the MesS subtype defined in our study was associated with the so-called integrated subtype 4 by Mun and colleagues ($P = 2.6e-07$; Fig. 6B; ref. 27), in which the key enriched biological process is RhoA-mediated tumor cell migration via actin reprogramming and focal adhesion assembly. It suggests that our MesS and EpiS AS events, including exon inclusion events in *ARHGEF11*

and *ACTN1*, might play similar roles in regulating cytoskeletal rearrangements via the RhoA signaling pathway (40).

Validation of the RBP signature using independent cohorts

Given that high-depth RNA-seq and splicing analyses are expensive and not routinely performed in the clinic, we devised a classification scheme that is more readily implementable and only focuses on the differential expression of splicing regulatory RBPs. We first examined RBP expression in the TCGA and ACRG cohorts. *RBM24* and *RBMS1* were upregulated, and *ESRP1* was downregulated in both the TCGA GS subtype and the ACRG EMT subtype (Supplementary Fig. S11),

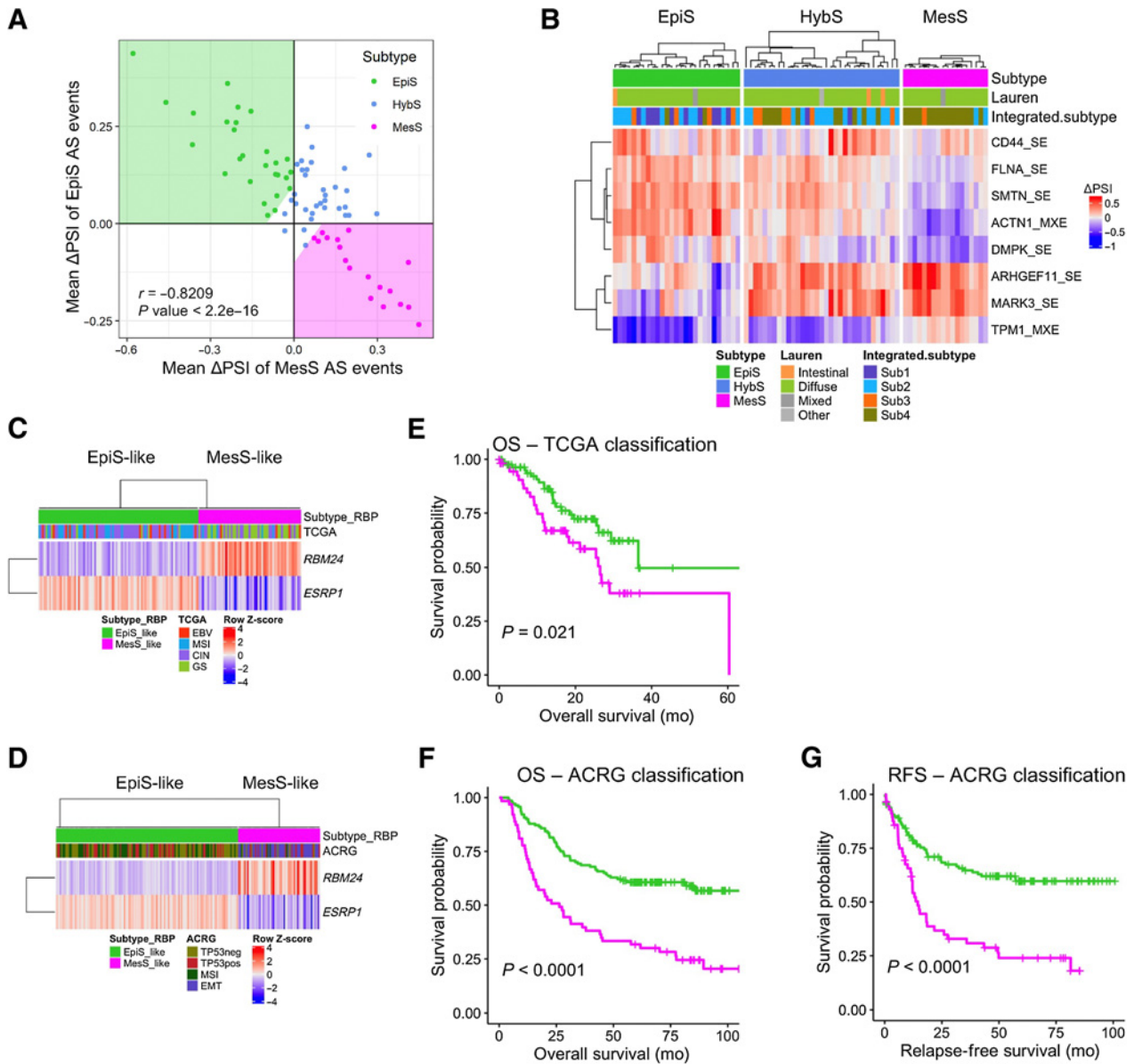


Figure 6. Validation of AS-based subtype classification in independent datasets. **A**, Scatter plot of patients from Hwang’s cohort using meanΔPSI of AS event-based EpiS and MesS tumor subtypes, colored by subtype. **B**, AS event-based patient classification applied to Hwang’s cohort. Patient clinical classification by Lauren and their integrated subtype are shown. **C** and **D**, RBP-based classification of TCGA (**C**) and ACRG (**D**) patients according to *RBM24* and *ESRP1* expression. **E** and **F**, Kaplan-Meier plot for OS between EpiS-like and MesS-like RBP-subtypes in TCGA (**E**) and ACRG (**F**) cohorts (log-rank test *P* values). **G**, Kaplan-Meier plot for relapse-free survival between EpiS-like and MesS-like RBP-subtypes in the ACRG cohort (log-rank test *P* values).

consistent with the overall association of the MesS subtype with these subtype classifications (Fig. 4F). Because our MesS subtype was characterized by high expression of *RBM24* and low expression of *ESRP1* (Fig. 5B), we thus defined MesS-like ($RBM24_{high}/ESRP1_{low}$) and EpiS-like ($RBM24_{low}/ESRP1_{high}$) subtypes that simply use the expression level of these two RBPs. We then evaluated this RBP-based classification in the TCGA and ACRG cohorts and confirmed that the gene expression levels of *ESRP1* and *RBM24* separated patients into two subtypes properly (Fig. 6C and D).

We further examined the prognostic value of our subtypes. Relapse-free survival showed a marginal difference between the EpiS and MesS subtypes (Supplementary Fig. S12). In addition, we discovered that TCGA patients with the MesS-like subtype had a significantly worse prognosis than those with the EpiS-like subtype (log-rank $P = 2.10 \times 10^{-2}$; Fig. 6E), whereas the TCGA subtypes did not show any difference in the overall survival (OS; Supplementary Fig. S13A). Similarly, ACRG patients with the MesS-like subtype, which was highly associated with the ACRG EMT subtype, showed worse survival than the EpiS-like subtype, both for OS and relapse-free survival (log-rank $P < 0.0001$ and $P < 0.0001$, respectively; Fig. 6F and G). The ACRG subtypes showed similar trends (Supplementary Fig. S13B and S13C). Results from a similar analysis, but using three RBPs additionally, including *RBMS1*, were virtually identical to findings using two RBPs (Supplementary Fig. S14).

For a further assessment of the association of our two RBP-based subtypes and patient survival, we developed a multivariate Cox proportional hazard model that included age, sex, pathologic stage, Lauren classification, and our subtypes (Supplementary Fig. S15). In the TCGA cohort, the RBP-based classification was an independent prognosis factor for OS (HR, 3.74; 95% confidence interval (CI), 1.43–9.80; $P = 0.007$). In the ACRG cohort, the RBP subtype was also an independent prognostic factor for both OS (HR, 1.82; 95% CI, 1.20–2.80; $P = 0.005$) and recurrence-free survival (HR, 1.61; 95% CI, 1.00–2.60; $P = 0.005$). Of note, another hazard model, including both splicing-based subtypes and TCGA subtypes for the TCGA cohort, revealed that our RBP subtype outperformed TCGA subtypes to predict OS (HR, 3.26; 95% CI, 1.137–9.4; $P = 0.028$; Supplementary Fig. S16A). Equivalent analysis with the ACRG cohort showed that RBP-based subtypes performed better in the OS only (HR, 1.92; 95% CI, 1.07–3.44; $P = 0.029$; Supplementary Fig. S16B and S16C). Collectively, survival analyses using multiple independent cohorts strongly indicate poor prognosis of the RBP-based MesS-like subtype.

Discussion

Although AS is a key regulatory mechanism that affects virtually all cellular processes, few studies have explored the role of AS in gastric cancer (30, 41–43), and even fewer studies have investigated the utility of AS-based patient classification (11). Given this, we (i) systematically profiled the AS landscape in gastric cancer, (ii) identified RBPs that regulate tumor subtype-specific AS events, and (iii) devised a patient classification scheme based on variable AS events or their regulatory RBPs. Unlike other signature-based classification methods that typically require whole transcriptome data and complex clustering algorithms, our first classification scheme is based on differential AS of only eight genes from tumor and paired normal tissue. Our second classifier, derived from the AS data, is even more straightforward and is based on expression of two RBPs, *RBM24*, and *ESRP1*, in tumor tissue only. The two splicing classification schemes, one based on AS events, the other on expression of RBP-splicing regulators, are closely correlated ($P = 5 \times 10^{-4}$) if we compare the subtypes (Supplementary Table S8), but are based on different molecular phenomena. Compared with the

TCGA gastric cancer patient classification relying on a complex clustering method using genome, RNA-seq, and microbiome data, or to the ACRG classification using a 675-gene signature after normalization of the RNA-seq data (9, 10), our classification based on expression level of two RBPs is analytically much more efficient for distinguishing gastric cancer subtypes and their associated prognosis, and therefore easier to implement in the clinic on gastric tumor biopsies.

On the basis of the eight AS events signature, we defined two tumor gastric subtypes, EpiS and MesS. One of the major molecular mechanisms behind our AS-based patient subgrouping was the EMT process. EMT plays essential roles in tumor initiation, progression, and metastasis in cancer, but its contribution is highly dependent on the cancer type (44). Tumors from the EpiS and MesS subtypes represent the two sides of the EMT spectrum. In the MesS subtype, EMT-TFs were upregulated and mesenchymal-epithelial transition (MET)-TFs were downregulated relative to the EpiS subtype (Fig. 2D; Supplementary Fig. S6). Furthermore, activation of the KRAS–MAPK and Hedgehog pathways in the MesS subtype (Fig. 3A) can in turn activate EMT-TFs *SNAIL1/2* and *ZEB1/2* indirectly via TGF β –SMAD and Notch signaling (34, 45). The enrichment of several gene sets related to the DNA-damage response pathways is observed in the EpiS subtype (Fig. 3B), whereas the upregulation of *H2AX* relative is observed in the MesS subtype (Supplementary Fig. S6). Indeed, Weyemi and colleagues (46) showed that histone variant *H2AX*, a sensitive marker of double-strand breaks, is a critical regulator of EMT, whose loss induces EMT and reintroduction partially reverses EMT. Oh and colleagues (47) reported epithelial and mesenchymal phenotypes by analyzing genomic and proteomic data from several cohorts. Even though AS was not included in their analysis, EMT markers were found to be a critical factor in patient subtypes associated with clinical outcomes such as patient survival and resistance to chemotherapy.

As for the interplay between regulation of AS and expression of EMT master regulators, several studies reported that EMT-TFs altered expression of splicing regulators. In particular, *ZEB1* and *SNAIL1* were shown to repress *ESRP1* expression and lead to EMT in various cancer cell lines (48, 49). Similarly, in our dataset, we observed significant correlations between the expression of EMT-TFs and RBPs levels, with *ZEB1* being positively correlated with both *RBM24* and *RBMS1* ($r > 0.5$), whereas *ESRP1* levels negatively correlated with *ZEB1/2*, *SNAIL2*, and *TWIST2* expression ($r < -0.5$; Supplementary Fig. S17). Furthermore, to determine whether RBPs could regulate downstream EMT-TFs, we analyzed the expression of six TFs in RNA-seq data from breast and lung cancer cell lines with *ESRP1* knockdown. Our analysis reveals that these six EMT-TFs did not change their expression upon *ESRP1* knockdown (Supplementary Fig. S18), thus suggesting EMT-activating TFs are presumably the upstream regulators of RBPs affecting AS.

Several of the eight AS events and three regulatory RBPs in gastric cancer identified herein have previously been associated with EMT in other tumor types (30, 38, 49, 50). For example, *ESRP1* is downregulated during EMT in breast cancer models, and drives AS of mesenchymal isoforms of *ARHGEF11*, *CD44*, and *TPM1* (38, 40). In addition, *ESRP1*-regulated AS events in *MARK3*, a kinase that regulates phosphorylation of microtubule-associated proteins, and *TPM1*, a protein involved in stabilizing cytoskeleton actin filaments are identified in this study. The function of these isoforms in EMT in MesS gastric tumors remains to be determined. The role of the other two RBPs, *RBM24*, and *RBMS1*, in EMT is not well characterized, and further studies of their splicing targets may provide valuable insights into the functional associations between splicing and EMT.

We observed several gene mutations and copy number variants (CNV) associated with AS subtype (Supplementary Fig. S4A and S4C).

The underlying molecular mechanisms linking mutations or CNVs in cancer genes with changes in AS patterns remain poorly understood in gastric cancer. One possible explanation would be that mutations or CNVs in cancer genes impact downstream signaling pathways and lead to changes in RBP levels, which in turn impact splicing of downstream targets. To test this hypothesis, we examined the association of mutations and CNVs with RBP expression. Wild-type and mutant patient groups were compared for differential RBP expression, which showed significant differences for *RBM24* expression and *ROBO1* or *PLEC* mutations ($P = 0.032$ or 0.013), as well as *ESRP1* expression and *LRP1B* or *AHNAK* mutations ($P = 0.018$ or 0.040 ; Supplementary Fig. S19A). However, this observation should be interpreted carefully because most mutations were not recurrent and occurred at different sites, therefore leading potentially to distinct functional consequences, and making it difficult to assess their effect on AS. In parallel, we examined the correlation between copy numbers and RBP expressions, which showed that *ESRP1* expression was positively correlated with *MYC* copy numbers ($r = 0.312$; Supplementary Fig. S19B). In breast cancer and lymphoma, *MYC* has been previously reported to regulate directly the transcription of several splicing regulators, including SRSF1, TRA2 β or PRMT5, by binding to their promoter regions (51, 52).

For nine cases out of 10 RT-PCR validations showed concordant results with the RNA-seq subtyping; however, we observed that one of our patient samples (P38) was predicted to be the MesS subtype in RNA-seq data and later determined to be the EpiS subtype based on RT-PCR validation (Fig. 2A; Supplementary Fig. S20A). To investigate the cause of this discrepancy, we repeated RNA-seq using the same RNA sample as in the RT-PCR validation experiment and observed that the new RNA-seq results also showed an EpiS subtype (Supplementary Fig. S20B). We confirmed that both RNA-seq datasets were from the same individual using NGSCheckMate (53). The source of discrepancy in the AS-based subtypes was therefore likely due to intratumor heterogeneity between the RNA samples used to generate the RNA-seq data, as are often the cases in cancer pathology studies (39). Hence, tumor heterogeneity introduces additional complexity to the evaluation of how robustly any approach classifies patient subtypes.

Finally, to explore whether our findings from primary tumors are recapitulated in cell models, we profiled splicing patterns as described above using available RNA-seq data from 37 gastric cancer cell lines from CCLE (54). We then applied the AS-based signature to gastric cancer cell lines and classified 11 cell lines as MesS, 7 as HybS, and 19 as EpiS subtypes (Supplementary Fig. S21A). Among the eight AS events that constituted our AS event-based signature derived from primary gastric tumors, AS events in *CD44*, *MARK3*, *ARHGEF11* showed variable splicing patterns across cell lines (Supplementary Fig. S21A), whereas the remaining five AS events exhibited an epithelial AS pattern across all cell lines, suggesting that they might be specific to primary tumors and lost during cell culture. Splicing inclusion ratios of *CD44*, *MARK3*, and *ARHGEF11* AS events were significantly different between EpiS and MesS cell lines (Supplementary Fig. S21B). Consistent with the observations in human primary tumors, *CD44* exon was more included in EpiS compared with MesS cell lines, whereas exons in *MARK3* and *ARHGEF11* were more skipped (Supplementary Fig. S21B). Finally, when

examining RBP levels, *ESRP1* showed a higher expression in EpiS than MesS cell lines, similarly to primary tumors (Supplementary Fig. S21C).

Future studies aimed at defining the molecular drivers of the transition from EpiS to MesS AS-based subtypes are needed. These changes could be triggered by alterations in genomic, epigenetic, or transcriptional regulators, which have been shown to drive splicing factor changes in other tumor types. Understanding these mechanisms will allow us to identify AS-based targets for development of gastric cancer therapies and to move toward harnessing the full potential of AS in precision medicine.

In summary, we systematically investigated the landscape and roles of AS in gastric cancer using RNA-seq data from matched tumor and normal samples. Association of AS with the EMT program was firmly established and suggested AS-based patient stratification schemes, which highlighted the potential of AS analysis as a tool for precision medicine. To our knowledge, this study presents the most comprehensive analysis to date of AS in the context of patient classification, molecular mechanisms, and prognosis in gastric cancer.

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Authors' Contributions

Y. Jun: Conceptualization, data curation, formal analysis, visualization, writing—original draft. **Y.-S. Suh:** Conceptualization, resources, writing—original draft. **S.H. Park:** Validation. **J. Lee:** Data curation, validation. **J.-I. Kim:** Data curation, validation. **S. Lee:** Conceptualization, data curation, formal analysis. **W.-P. Lee:** Conceptualization, formal analysis, supervision, validation, writing—review and editing. **O. Anczuków:** Conceptualization, resources, supervision, validation, writing—review and editing. **H.-K. Yang:** Conceptualization, resources, supervision, funding acquisition, writing—review and editing. **C. Lee:** Conceptualization, data curation, formal analysis, supervision, funding acquisition, writing—review and editing.

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