Genomic and Epigenomic Integration Identifies a Prognostic Signature in Colon Cancer

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Abstract

**Purpose:** The importance of genetic and epigenetic alterations maybe in their aggregate role in altering core pathways in tumorigenesis.

**Experimental Design:** Merging genome-wide genomic and epigenomic alterations, we identify key genes and pathways altered in colorectal cancers (CRC). DNA methylation analysis was tested for predicting survival in CRC patients using Cox proportional hazard model.

**Results:** We identified 29 low frequency-mutated genes that are also inactivated by epigenetic mechanisms in CRC. Pathway analysis showed the extracellular matrix (ECM) remodeling pathway is silenced in CRC. Six ECM pathway genes were tested for their prognostic potential in large CRC cohorts (n = 777). DNA methylation of IGFBP3 and EVL predicted for poor survival (IGFBP3: HR = 2.58, 95% CI: 1.37–4.87, P = 0.004; EVL: HR = 2.48, 95% CI: 1.07–5.74, P = 0.034) and simultaneous methylation of multiple genes predicted significantly worse survival (HR = 8.61, 95% CI: 2.16–34.36, P < 0.001 for methylation of IGFBP3, EVL, CD109, and FLNC). DNA methylation of IGFBP3 and EVL was validated as a prognostic marker in an independent contemporary-matched cohort (IGFBP3 HR = 2.06, 95% CI: 1.04–4.09, P = 0.038; EVL HR = 2.23, 95% CI: 1.00–5.0, P = 0.05) and EVL DNA methylation remained significant in a secondary historical validation cohort (HR = 1.41, 95% CI: 1.05–1.89, P = 0.022). Moreover, DNA methylation of selected ECM genes helps to stratify the high-risk stage 2 colon cancers patients who would benefit from adjuvant chemotherapy (HR: 5.85, 95% CI: 2.03–16.83, P = 0.001 for simultaneous methylation of IGFBP3, EVL, and CD109).

**Conclusions:** CRC that have silenced genes in ECM pathway components show worse survival suggesting that our finding provides novel prognostic biomarkers for CRC and reflects the high importance of integrative analyses linking genetic and epigenetic abnormalities with pathway disruption in cancer. Clin Cancer Res; 17(6); 1535–45. ©2011 AACR.

Introduction

Recent efforts in genome-wide sequencing of multiple human cancers have identified a large number of mutations that cluster within a small number of intracellular pathways such as cellular senescence bypass, invasion, and metastasis (1, 2). Epigenetic abnormalities can also operate with genetic alterations to effect aberrant gene function in cancer (3). DNA hypermethylation of promoter-associated CpG islands is a frequent and early mechanism of tumor-suppressor gene inactivation in many cancers and may have great promise for identifying biomarkers (4–6). We have previously emphasized that DNA hypermethylation can affect many of the new genes found to be mutated in both colorectal (CRC) and breast cancer cells (7, 8). However, an understanding of such genome wide alterations, by itself, has not led to many clinically applicable biomarkers in predicting tumor behavior. We hypothesized that integration of the genome and hypermethylome may provide...
Translational Relevance

We have used an exciting integrative approach of multiple whole genome analysis (genetic and epigenetic) to identify an important pathway, the extracellular matrix (ECM) maintenance, and remodeling pathway, which is silenced in all colon cancers. Colon cancers that have silenced multiple genes in this pathway have a poor prognosis based on analyses of large cohort of patients.

DNA methylation of these genes is particularly useful in stratifying the low- versus high-risk stage 2 colon cancer patients. Colorectal cancers that have silenced multiple genes in the ECM pathway appear to show a significantly worse survival in adjusted multivariate analysis of a large cohort of colon cancer patients. Our study has important ramifications in the clinical management of colon cancer patients as 30% to 40% of early stage colon cancers will present with recurrence/metastases but current strategies are not helpful in stratifying this high-risk cohort. We show that our prognostic markers are the most efficacious in stratifying the high-risk subset of stage 2 colon cancers who may then benefit from adjuvant chemotherapy.

DNA methylation and gene expression analyses

Primer pairs were preferentially designed near the putative transcriptional start site (TSS) in the 5′-CpG islands of the genes. Primer sequences for methylation-specific PCR (MSP) analysis were designed using MSP Primer 3. All primer sequences are listed in Supplementary Table S5. For expression studies using reverse transcription PCR (RT-PCR), primers were designed using the open access program Primer3.

For MSP analysis, DNA was extracted using the standard phenol–chloroform extraction method. Bisulfite modification of genomic DNA was carried out using the EZ DNA Methylation Kit (Zymo Research). DNA methylation analysis was performed using MSP primer pairs located close to the putative transcriptional start site in the 5′-CpG island with 2 μL of bisulfite-treated DNA as template and JumpStart Red Taq DNA Polymerase (Sigma) for amplification as previously described (7).

Total RNA was extracted from cell lines using the RNeasy Mini Kit (Qiagen), treated with DNase (Qiagen). For RT reaction, 1 μg of total RNA was subjected to the first strand cDNA synthesis using Superscript III first strand cDNA synthesis kit (Invitrogen) according to the manufacturer’s instructions. Expression analysis was performed by RT-PCR using 1 μL of cDNA as template and JumpStart Red Taq DNA Polymerase (Sigma) for amplification.

Quantitative methylation analysis

Quantitative MSP (qMSP) was performed after sodium bisulfite modification for selected genes including IGFBP3, EVL, CD109, FLNC, and FBN2 on a small series of normal colon and colon cancers (n = 5, each; Supplementary Fig. S2). qMSP analysis helped us to confirm whether our candidate genes showed cancer-specific DNA methylation.

Materials and Methods

Patients

Johns Hopkins (JHU) training and validation cohort samples for the colorectal cancer (CRC) studies were prepared from formalin-fixed and paraffin-embedded (FFPE) colon cancer tissue samples from the pathology archives of the JHU Hospital in accordance with the Institutional Review Board (IRB) and as per HIPAA compliance. The JHU training cohort consisted of 147 tissue samples from stage 1 to 4 colon cancer patients who underwent primary surgery from 1995 to 2005 (median follow-up of 6.4 years). The JHU validation cohort consisted of 72 tissue samples from stage 1 to 4 colon cancer patients who underwent primary surgery from 1995 through 2005 (median follow-up of 7.5 years). Patients with neoadjuvant chemotherapy, including all rectal cancers, were excluded from the current study. Patients in both JHU cohorts were similar with respect to age, sex, race, proportion of cases with lymphovascular invasion, pathologic grade, location, and proportion of cases with recurrence (Table 1).

Netherlands Cohort study on Diet and Cancer (NLCS) Validation cohort were prepared from 558 FFPE colon cancers from stages 1 to 4 (median follow-up of 7.2 years). The NLCS has been previously described in detail (13). A total of 925 incident CRC cases were identified from 1989 to 1994, and DNA was available for analyses on our current subset (Table 1).

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patterns consistent with the qualitative MSP analysis. For quantitative real-time analyses, the Power SYBR Green PCR kit (Applied Biosystems) was used and the amplification conditions consisted of an initial 10-minute denaturation step at 95°C, followed by 40 cycles of denaturation at 95°C for 15 s and annealing and extension for 30 s and 60 s, respectively. An ABI StepOnePlus Real-Time PCR System was used (Applied Biosystems), and for quantitation the comparative cycle threshold (Ct) method was used, normalizing the Ct values for the indicated gene to the Ct values of unmethylated reaction relative to a methylated reaction sample.

Cell culture and treatment
Cancer cell lines (CRC cell lines: HCT116, SW480, RKO, HT29, Caco-2, Lovo, COLO 320, COLO 205, DLD1, SW48, and SW620; breast cancer cell lines: MCF7, T47D, MDA-MB-231, and MDA-MB-468) were obtained from American Type Culture Collection (ATCC) and cultured in appropriate media and under conditions described by ATCC, with media obtained from Invitrogen, supplemented with 10% fetal bovine serum (Gemini Bio-Products) and 1% penicillin/streptomycin (Invitrogen). DKO cells (HCT116 cells with genetic disruption of DNMT1 and DNMT3b) were cultured as described previously (7).

Signaling pathway analysis of methylated genes
We used a highly curated database (Metacore) that includes human protein–protein interactions, signal transduction, and metabolic pathways, and a variety of cellular functions and processes for signaling pathway analysis. Functional ontology enrichment selected the significantly altered canonical pathways, networks, and gene ontology terms. Analysis was run to detect deregulated interaction networks in CRC, using a subset of genes that are both mutated (1, 2) and hypermethylated in CRC tissue. Significant pathways were selected and intersected with those found by Wood and colleagues (2). While the article by Wood and colleagues (2) was focusing on those networks targeted by mutations alone, the combination of both approaches was designed to zoom in on those pathways that are important to CRC as a whole through alterations of these molecular interaction networks, be it by gene mutations or promoter hypermethylation.

Survival and statistical analysis
Univariate and multivariate Cox regression was used to identify genes that could significantly predict poor overall survival. All ties were handled by the Breslow method, and the proportional hazards assumption was verified by examination of residual plots. Adjusted Kaplan–Meier survival plots were created to demonstrate the ability of the genes to predict poor overall survival. Statistical significance in this study was set at 0.05. To control for multiple testing, Sidak adjustment method was used to correct for family-wise error rate. Results of all models are reported as odds ratios with 95% CIs. All data were analyzed using StataTM v9.2 statistical analyses software (Stata Corporation).

Table 1. Demographics of training and validation cohorts in this study

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<tr>
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<th>JHU Training cohort</th>
<th>Validation cohorts</th>
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<tr>
<td>N</td>
<td>n = 147</td>
<td>n = 72</td>
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<td>Median age, y</td>
<td>66</td>
<td>70</td>
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<tr>
<td>5-y survival</td>
<td>67.4%</td>
<td>65.3%</td>
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<td>Stage I</td>
<td>83.3%</td>
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<td>67.4%</td>
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<td>Male</td>
<td>84 (57%)</td>
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<td>Female</td>
<td>63 (43%)</td>
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<td>Right colon</td>
<td>79 (54%)</td>
<td>40 (56%)</td>
</tr>
<tr>
<td>Left colon</td>
<td>66 (45%)</td>
<td>32 (44%)</td>
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<td>Unknown</td>
<td>9 (1%)</td>
<td></td>
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<table>
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<td>Stage I</td>
<td>36 (24.4%)</td>
<td>18 (25.0%)</td>
</tr>
<tr>
<td>Stage II</td>
<td>49 (33.3%)</td>
<td>27 (37.5%)</td>
</tr>
<tr>
<td>Stage III</td>
<td>37 (25.2%)</td>
<td>20 (27.8%)</td>
</tr>
<tr>
<td>Stage IV</td>
<td>25 (17.1%)</td>
<td>7 (9.7%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>41 (9%)</td>
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Results

DNA methylation analysis of mutated genes

Using the recently compiled database of human cancer mutations in CRC and breast cancers, termed the candidate cancer genes (or CAN genes; refs 1, 2), we have now identified 119 genes which, in colon and breast cancer cell cultures, are also identified by our recently published gene discovery approaches (7, 8). As candidate genes for promoter DNA hypermethylation. As the majority of these genes were found to be hypermethylated in colon tumors, we focused the remainder of our studies on colon cancers. These 119 genes then represent the intersection of the consensus-mutated and DNA-hypermethylated genes in colon cancer. Seventy-five of these 119 genes contained promoter CpG islands (14) and were then first experimentally validated for (1) expression in normal colon tissues by RT-PCR (n = 65 genes); Of these, 45 genes showed (2) correlation between loss of gene expression by RT-PCR and promoter CpG island DNA methylation by MSP analysis in colon cancer cell lines (HCT116, RKO, SW480, and Colo320; Fig. 1A, Supplementary Table S1). To filter down to genes that only showed cancer-specific DNA methylation, we next looked at DNA methylation frequencies of these 45 genes in a series of primary colon cancers (n = 21, stages 2 and 3) as well as normal colon from cancer-free individuals (n = 20). (3) Thirty-four genes showed more than 5% frequency of DNA methylation in a series of primary colon cancers and (4) 29 of these genes showed cancer-specific DNA methylation with the absence of DNA methylation in normal colon (Supplementary Table S1 for schema and genes; Supplementary Table S2 for primary colon cancer sample information). Cancer-specific DNA methylation was also confirmed using qMSP for a subset of genes and showed identical results to the gel-based MSP assay (Supplementary Fig. S2).

A subset of these genes had been previously identified by us during genome-wide screens (7, 8) and were included in the current integrative approaches to provide a more comprehensive view of frequent epigenetic alterations in colon cancer. Among the 29 genes, 15 genes are newly identified by this integrative approach as potential targets of both mutation and DNA methylation (FLNC, SH3TC1, ZNF569, SLCL22A15, HAPLN1, PRKD1, IGFBP3, TCECG1L, LAMA1, FBN2, CPAMD8, NTNG1, EYA4, GRID1, and PPM1E) in this study with 5% to 100% DNA methylation frequencies in CRC (Fig. 1B), whereas the remaining 14 genes were also previously identified (7). A direct comparison of DNA methylation and known mutation frequencies of these 29 genes reveals that they are more often epigenetically than genetically altered in colon cancer (Supplementary Fig. S1). DNA methylation frequencies of the 29 genes show variable frequencies in the primary colon cancers ranging from 5% to 100% (Supplementary Fig. S1). 17 out of 29 genes (GUCY1A2, NRCAM, PTPRD, PPME1, STARD8, GRID1, EVL, EYA4, NTNG1, CPAMD8, FBN2, MMP2, SYNE1, LAMA1, TCECG1L, APC2, and GPNMB) displayed high level (≥50%) hypermethylation in colon cancers, whereas 12 genes (SLCL22A15, ICAM5, HAPLN1, CHD5, RET, LGi6, PRKD1, IGFBP3, FLNC, CD109, ZNF569, and SH3TC1) showed lower frequency of DNA methylation (<50%) in colon cancers (Fig. 1B).

Pathway analysis identifies global disruption of the ECM pathway in CRC

A growing body of research is defining that, particularly with respect to new genes being discovered in cancer resequencing efforts and those genes which are genetically altered with low frequency, alterations of pathways rather than of individual genes may be the important issue for affecting the course of tumorigenesis (1, 2). These mutated genes, then, belong to a handful of important signaling pathways in breast, colorectal, and other cancer types (2, 15). We queried an extensive gene ontology set (MetaCore; GeneGo) and nearly half of the genes that we identified to be both mutated and hypermethylated (13 of 29, 44.8%) mapped to the ECM pathway in CRC (Fig. 2). ECM remodeling is critical for many developmental processes, and recently, abnormally remodeled ECM has been shown to contribute to the neoplastic process (9).

We, therefore, focused the remainder of our studies to understanding the functional implications of the dysregulation of the ECM pathway in CRC. In our analyses of these ECM pathway genes, 8 of the genes (NRCAM, EVL, NTNG1, CPAMD8, FBN2, MMP2, LAMA1, and GPNMB) showed high frequency (>50%) of DNA methylation in CRC and are amongst those mapping to the ECM pathway (refs. 10, 16, 17; Fig. 2), and the remainder show lower frequency of DNA methylation in CRC (ICAM5, HAPLN1, IGFBP3, FLNC, and CD109; refs 18–20).

In addition, we also noted that genes such as TIMP2 and TIMP3 that have previously been identified by ourselves and others as being hypermethylated in colorectal and other cancers also belong to the ECM pathways (21, 22). Based on this observation, we also investigated, for potential epigenetic silencing in CRC, other genes in the ECM remodeling pathway which have previously been identified to show hypermethylation in other cancers by us and others. Six other genes, in addition to TIMP2 and TIMP3, were identified, including PLAU, Nidogen (NID1), Osteonectin, CD44, MMP9, and Laminin 5 (LAM5; refs 23–26) and these 8 genes were then investigated for potential DNA methylation in our current set of primary CRC. Six of these 8 ECM genes, including TIMP2, TIMP3, MMP9, PLAU, Nidogen, and Osteonectin were found to be methylated in CRC with variable DNA methylation frequencies, ranging from 5% to 95% (Supplementary Table S3). All 8 additional ECM genes also showed no evidence of DNA methylation in normal colon (data not shown; n = 5). The TIMPs (TIMP2 and TIMP3) and MMPs (MMP9) have long been thought to be an integral components of the ECM pathway and their expression has been critical in primary tumor growth, angiogenesis, invasion, and metastasis in different type of cancers (18). Nidogen forms a complex with laminins and...
collagen IV and helps to stabilize the basement membrane structure as well as being important for cell adhesion by establishing contact with integrins (24). Osteonectin (also known as SPARC) and PLAU are involved in wound repair (25), cell migration, and differentiation but their role in tumorigenesis has not yet been defined.

Together, all our data show that at least 19 genes in the ECM pathway are hypermethylated in CRC (Fig. 2 and Supplementary Table S3), and strongly suggests that the ECM pathway is globally and universally altered in CRC by mutation or epigenetic silencing but much more commonly by the latter abnormality.
<table>
<thead>
<tr>
<th>Gene name</th>
<th>RefSeq ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPA0D9</td>
<td>NM_015602</td>
<td>Complement 3 D9; transmembrane protein that binds heparin</td>
</tr>
<tr>
<td>CD109</td>
<td>NM_133493</td>
<td>GPI anchored member of a2 macroglobulin C3/4/5 family; involved in RET signaling, linked to TGFβ signaling and cytoskeleton maintenance</td>
</tr>
<tr>
<td>EVL</td>
<td>NM_016337</td>
<td>Eno/Ena/Vasp-like; links protein with actin cytoskeleton via GTPase Rap1; involved in cell spreading and lamellipodia formation</td>
</tr>
<tr>
<td>FBN2</td>
<td>NM_001999</td>
<td>Filibrin, colocalizes with laminin, collagen, fibronectin, versican, and others</td>
</tr>
<tr>
<td>FLNC</td>
<td>NM_001458</td>
<td>Filamin c; regulates actin cytoskeleton via interaction with fibronectin</td>
</tr>
<tr>
<td>GN1MB</td>
<td>NM_001005340</td>
<td>Osteoacticin; regulates MMP3 activity</td>
</tr>
<tr>
<td>HAPLN1</td>
<td>NM_001894</td>
<td>Hyaluronan and proteoglycan link protein 1</td>
</tr>
<tr>
<td>ICAM5</td>
<td>NM_003269</td>
<td>Binds ERM (ezrin/moesin/ radixin) proteins to regulate actin cytoskeleton</td>
</tr>
<tr>
<td>IGF1BP3</td>
<td>NM_006098</td>
<td>Regulates IGF1 activity through direct interaction</td>
</tr>
<tr>
<td>LAMA1</td>
<td>NM_0055659</td>
<td>Laminin alpha 1; basement membrane protein</td>
</tr>
<tr>
<td>MMP2</td>
<td>NM_004530</td>
<td>Matrix metalloproteinase 2; enhances cell–cell adhesion cleavage of ECM proteins fibronectin and vitronectin into fragments and facilitates binding to p-integrin receptors</td>
</tr>
<tr>
<td>NTNG1</td>
<td>NM_00113226</td>
<td>Netrin G1 transmembrane receptor mediates cell matrix adhesion</td>
</tr>
<tr>
<td>NRCAM</td>
<td>NM_001037132</td>
<td>Neuron related cell adhesion molecule; p-catenin target</td>
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<td>TIMP3</td>
<td>NM_000362</td>
<td>Tissue inhibitor of metalloproteinase 3; inhibitory role against metalloproteinases</td>
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<td>NM_003255</td>
<td>TIMP metalloproteinase inhibitor 2 precursor; inhibitory role against metalloproteinases</td>
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<td>Nidogen</td>
<td>NM_002608</td>
<td>Basement membrane glycoproteins; play a role in cell interactions with the extracellular matrix</td>
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<tr>
<td>Osteonectin</td>
<td>NM_003119</td>
<td>Secreted protein acidic and rich in cysteine/osteonectin/GM40, or SPARC</td>
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<tr>
<td>MMP9</td>
<td>NM_004994</td>
<td>Matrix metalloproteinase 9 proprotein; secreted as inactive proproteins which are activated when cleaved by extracellular proteinases</td>
</tr>
<tr>
<td>PLAUI</td>
<td>NM_002659</td>
<td>Urokinase plasminogen activator proprotein; involved in degradation of the extracellular matrix and possibly tumor cell migration and proliferation</td>
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</table>
CRC with ECM dysregulation demonstrate worse survival

We next wanted to study the clinical correlates of our gene silencing data in terms of the previously observed juxtaposition of embryonic gene expression patterns and EMT with such tumors having the most primitive and aggressive phenotypes (27). In this respect, an exploratory clinical outcome analysis in our initial small cohort of colon cancer patients (n = 21 patients) was performed for hypothesis-generation and showed that DNA methylation of selected ECM genes (IGFBP3 and EVL) was associated with significantly worse outcomes on unadjusted analysis (P = 0.02 and P = 0.03, respectively), whereas DNA methylation of other genes (CD109, NR CAM, NTNG, and FLNC) showed a trend toward poor survival which was not statistically significant. To explore the potential clinical significance of these data, we tested these above 6 ECM pathway genes for their prognostic potential in a large training cohort of patients with colon cancers and well-annotated clinical data that could be correlated with survival and gene methylation status. Strikingly, in analyses of tumors from 147 patients with stage 1 to 4 colon cancers (JHU training cohort; Table 1), we observed a statistically significant increased risk for mortality, when either individual genes or combinations of genes were methylated after adjusting for other prognostic variables such as age and stage of cancers (Figs. 3A and 4A). For example, adjusted Kaplan–Meier survival curves for 2 of the individual ECM genes, IGFBP3 and EVL, show that DNA methylation of each is associated with decreased survival (adjusted for age and stage; Fig. 3A). Multivariate HRs (Fig. 4A) for survival depicted in a forest plot show the HRs for DNA methylation of these 2 genes predicted risk for poor survival (IGFBP3 HR = 2.58, 95% CI, 1.37–4.87, P = 0.004; EVL HR = 2.48, 95% CI, 1.07–5.74, P = 0.034), whereas another gene CD109 shows a trend toward statistical significance (HR = 1.81, 95% CI, 0.94–3.50, P = NS). Moreover, in addition, in patients with simultaneous DNA methylation of multiple genes in the ECM pathway, significantly worse survival is predicted in adjusted analysis. For example, for DNA methylation of IGFBP3, EVL, CD109, and FLNC, the odds of dying are increased to greater than eight-fold after adjusting for age and stage of cancer compared with patients who lack DNA methylation of these genes (HR: 8.61, 95% CI: 2.16–34.36, P < 0.001) (Fig. 4A).

We further examined DNA methylation status of IGFBP3, EVL, CD109, NR CAM, and FLNC in an independent validation cohort of 72 patients with stage 1 to 4 colon cancers (Validation cohort JHU; Table 1 and Supplementary Table S4). In multivariate-adjusted analysis, both IGFBP3 (HR: 2.06, 95% CI: 1.04–4.09, P = 0.038) and EVL (HR: 2.23, 95% CI: 1.00–5.06, P = 0.050) remained statistically significant as markers associated with worse survival. CD109 showed a trend toward worse survival outcome that was not statistically significant (HR: 1.49, 95% CI: 0.72–3.08, P = NS), whereas FLNC and NR CAM had no prognostic value. Adjusted Kaplan–Meier survival curves (adjusted for age and stage) for 2 of the individual ECM genes, IGFBP3 and EVL, again show that DNA methylation of each is associated with decreased survival in our validation cohort (Fig. 3A). Multivariate HRs (Fig. 4B) for survival depicted in a forest plot of both the training and validation cohort shows that simultaneous DNA methylation of multiple genes in the ECM pathway is associated with significantly worse survival. These findings support our hypothesis that DNA methylation of ECM pathway genes may have an important role during cancer progression and metastasis.

We also looked at the prognostic significance of these ECM genes (IGFBP3, EVL, CD109, NR CAM, and FLNC) in a secondary validation cohort NLCS (13) of 558 patients with stage 1 to 4 colon cancers (Table 1). Of note, the NLCS cohort differs significantly from the JHU training and validation in being a historical cohort (study period 1986–1991) with marked differences in survival of colon cancers, a preponderance of distal colon cancers and markedly worse outcomes (see Table 1). EVL (HR: 1.41, 95% CI: 1.05–1.89, P = 0.022) again is statistically significant as a marker associated with worse survival and CD109 again shows a trend toward statistical significance (HR: 1.19; 95% CI: 0.95–1.48, P = NS), whereas IGFBP3 methylation was not significant in this historical cohort.

ECM biomarkers in stage 2 colon cancers

In the clinical management of colon cancer patients, there is an urgent need for identification of biomarkers that may stratify the high risk stage 2 colon cancers patients who are at the highest risk of recurrence. Previous large randomized trials have not shown a consistent benefit of adjuvant chemotherapy (28) but approximately 30% to 40% of these patients will nevertheless recur/metastasize (29). Identification of this high-risk cohort of stage 2 colon cancer patients would be useful as this subset may derive a benefit from adjuvant chemotherapy. Therefore, we investigated the biomarker potential of the ECM gene methylation in the stage 2 colon cancers in the combined JHU training and validation cohort (n = 219; Supplementary Table S6). Both IGFBP3 and CD109 are associated with worse survival in these stage 2 colon cancers (IGFBP3 HR: 3.02; 95% CI: 1.42–6.4;
$$P = 0.004$$ and $$CD109$$ HR: 2.41; 95% CI: 1.14–5.10; $$P = 0.021$$, respectively. Similarly, simultaneous DNA methylation of multiple genes in the ECM pathway is again associated with significantly worse survival for the stage 2 colon cancers (HR: 5.85, 95% CI: 2.03–16.83, $$P = 0.001$$ for DNA methylation of both $$IGFBP3$$ and $$CD109$$; Fig. 3B).

Discussion

Our data highlight the important role that integrative approaches may have in understanding cancer biology and implications of such in the translational arena. Our current approach of analyzing the genomic and epigenomic changes in colon cancer identified the role of the ECM pathway in colorectal carcinogenesis. These findings also are potentially useful in finding novel prognostic biomarkers as highlighted by our finding of DNA methylation of $$IGFBP3$$, $$EVL$$, and $$CD109$$ being associated with worse survival. As mentioned in the results, this may have direct clinical implications in stratifying the high-risk stage 2 colon cancers patients who have worse outcomes and who may benefit from adjuvant chemotherapy, although our findings would need to be validated in other retrospective studies of carefully clinically annotated tissues or a well-designed prospective study. Our findings were validated in an internal validation cohort (JHU validation) where DNA methylation of both $$IGFBP3$$ and $$EVL$$ again showed the association with worse survival, whereas $$CD109$$ showed a trend toward worse survival. We then validated our findings in another large external NLCs for DNA methylation of...
EVL and CD109 associated with worse outcomes. However, IGFBP3 methylation was not associated with worse survival in this cohort. Although at present, it is unclear why IGFBP3 methylation was not validated in the NLCS cohort, there are multiple possible explanations for this. This may simply reflect the different biology in the two cohorts. In fact, epigenetic alterations have been shown to vary in different populations and races and potentially affect outcomes (30, 31). Netherlands also has a lack of colon cancer screening program and this is reflected in the markedly different 5-year survival of this cohort (54.8% NLCS vs. 67.4% JHU training cohort or 65.3% JHU validation cohort) and the biases associated with screening (32). Moreover, the NLCS cohort has a preponderance of left-sided cancers, whereas...
the JHU cohorts reflect a distribution and survival similar to what is seen in the Surveillance, Epidemiology, and End Results database (a large national cancer database in the United States; ref. 33). The location of colon cancer has been shown to affect both genetic and epigenetic alterations (34–36) and survival (33, 37). In addition, adjuvant chemotherapy was not offered routinely for high risk patients in the NLCS cohort. Thus lack of validation of IGFBP3 may also reflect the fact that this is a predictive rather than a prognostic marker. Further validation studies in comparable modern-day cohorts similar to the JHU cohort may clarify this. In particular, our data on IGFBP3 and survival is intriguing as previous studies have shown that IGFBP3 induces apoptosis and inhibits DNA synthesis in breast, prostate, and non-small–cell lung cancer (38). Moreover, prior studies have also reported that high levels of circulating IGFBP3 decrease the risk of recurrent colorectal adenoma formation (39). Further studies are needed to confirm these findings particularly the risk stratification in stage 2 colon cancers.

Previous studies have also alluded to the potential of epigenetic changes, especially DNA hypermethylation of promoter-associated CpG islands, in serving as prognostic biomarkers. Brock and colleagues showed that hypermethylation of 2 genes, p16 and H-cad, in the primary lung tumor and the mediastinal lymph nodes could predict recurrence for early stage lung cancers (5). In another study, DNA methylation profiles of multiple methylated-in-tumor (MINT) loci predicted distal recurrence in a small cohort of rectal cancer patients (40), while ID4 methylation has also been associated with worse outcomes (41). However most of these studies used candidate genes from the literature and tested them for prognostic significance in a cohort of patients. In our study, we have not only identified a unique pathway, the ECM pathway, that is altered by DNA methylation of many genes in CRC, but then show that dysregulation of this core pathway has prognostic significance as well. However, our current studies do not clearly distinguish between the DNA hypermethylation arising from the tumorigenic cells or from the non-tumorigenic stromal cells. This will need to be investigated in future studies.

What are the functional consequences of loss of function for the above ECM genes in CRC? We also considered, from several perspectives, our findings in another context—namely, that they might represent an example of properties which reflect the primitive cell status increasingly attributed to multiple tumor types (42). First, multiple previous studies (3) have stressed that some 50% of genes that become DNA hypermethylated in colon cancers are marked by a chromatin pattern, termed bivalent chromatin, that keeps these same genes in a low expression pattern in ESC and embryonic mesenchymal (EM) progenitor cells (43). Indeed, a key chromatin component of bivalent chromatin, the presence of polycomb group (PcG) constituents around gene start sites, is present in ESC and EM cells in the promoter regions of several of the ECM genes (IGFBP3, NRCAM, CPAMD8, FBN2, MMP2, TIMP3, MMP9), based on our database analyses (unpublished data).

In summary, our results suggest that CRC may harbor defects in key ECM genes, induced infrequently by gene mutations, and even more frequently by gene promoter DNA hypermethylation. This loss of function for such genes appears to correlate with loss of cellular differentiation and to be associated with aspects of primitive, stem/progenitor cell, features of CRC (Supplementary Fig. S3). Loss of ECM gene function could foster metastatic behavior by altering events which require ECM remodeling such as allowing cells to invade through the basement membrane, migrate into the lymphovascular space, and establish metastatic foci in a distant organ. Our study suggests that this ECM alteration by hypermethylated genes may contribute to carcinogenesis, to some degree in virtually every CRC, through silencing of selected ECM pathway genes by both genetic and epigenetic alterations. Finally, our data indicate that detection of DNA hypermethylation of selected of the genes we have studied may provide a biomarker strategy for predicting the clinical behavior of CRC. Our data emphasize the importance of integrating cancer gene mutations and DNA hypermethylation changes to uncover the molecular events disrupting cell signaling in CRC and other cancers.

Disclosure of Potential Conflicts of Interest

S.B. Baylin and J.G. Herman are consultants of MDx Health and receive research support from MDx Health. The other authors disclosed no potential conflicts of interest.

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