CpG Island Methylator Phenotype Associates with Low-Degree Chromosomal Abnormalities in Colorectal Cancer

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Abstract

Purpose: Aberrant promoter methylation and genomic instability occur frequently during colorectal cancer development. CpG island methylator phenotype (CIMP) has been shown to associate with microsatellite instability, and BRAF mutation and is often found in the right-side colon. Nevertheless, the relative importance of CIMP and chromosomal instability (CIN) for tumorigenesis has yet to be thoroughly investigated in sporadic colorectal cancers.

Experimental Design: We determined CIMP in 161 primary colorectal cancers and 66 matched normal mucosae using a quantitative bisulfite/PCR/ligase detection reaction (LDR)/Universal Array assay. The validity of CIMP was confirmed in a subset of 60 primary tumors using MethyLight assay and five independent markers. In parallel, CIN was analyzed in the same study cohort using Affymetrix 50K Human Mapping arrays.

Results: The identified CIMP-positive cancers correlate with microsatellite instability (P = 0.075) and the BRAF mutation V600E (P = 0.00005). The array-based high-resolution analysis of chromosomal aberrations indicated that the degree of aneuploidy is spread over a wide spectrum among analyzed colorectal cancers. Whether CIN was defined by copy number variations in selected microsatellite loci (criterion 1) or considered as a continuous variable (criterion 2), CIMP-positive samples showed a strong correlation with low-degree chromosomal aberrations (P = 0.075 and P = 0.012, respectively). Similar correlations were observed when CIMP was determined by MethyLight assay (P = 0.001 and P = 0.013, respectively).

Conclusion: CIMP-positive tumors generally possess lower chromosomal aberrations, which may only be revealed using a genome-wide approach. The significant difference in the degree of chromosomal aberrations between CIMP-positive and the remainder of samples suggests that epigenetic (CIMP) and genetic (CIN) abnormalities may arise from independent molecular mechanisms of tumor progression.

Aberrant DNA methylation and genomic instability are hallmarks of colorectal cancer development. Two types of altered methylation patterns, gene-specific promoter hypermethylation and genome-wide hypomethylation, have been shown to interfere with gene expression in colorectal cancer. Whereas aberrant promoter hypermethylation at the 5′ regulatory regions or CpG islands impairs gene function by transcription silencing, which has been shown in numerous well-known tumor suppressor genes (1–3), global demethylation tends to occur at parasitic sequences or transposons resulting in the reactivation of this otherwise silenced genetic information (4–6). Changes in genome integrity during carcinogenesis are typically attributed to either chromosomal instability (CIN) or microsatellite instability (MSI). The characteristic feature of CIN tumors is aneuploidy, which presents as the change of an entire or a significant portion of a chromosomal arm, including, but not limited to, deletion, amplification, and loss of heterozygosity (LOH; refs. 7, 8). Alternatively, MSI tumors are known to have a different number of short microsatellite DNA repeats but retain a diploid genome. The relative

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Translational Relevance

This study stratifies a subset of colon cancers—CpG island methylator phenotype (CIMP) that possess multiple hypermethylated genomic loci and correlate with microsatellite instability (MSI) and BRAF mutations. The presence of significant low-degree of chromosomal aberrations in these tumors suggests that CIMP may arise from an independent mechanism of tumor progression compared with the majority of sporadic colon cancers. The different clinicopathologic characteristics of this subgroup may be beneficial for future development of cancer intervention therapy (e.g., demethylating drug treatments) and patient management. Additionally, because some CIMP tumors contain a high-degree of chromosomal instability (CIN), the mechanisms of CIMP and CIN do not seem to be mutually exclusive. Our report of a quantitative and high-resolution study of CIMP and CIN provides a more refined classification of CIMP tumors, including identifying those CIMP-positive/MSI-negative samples with high levels of CIN. The clinical characteristic or prognostic value of the refined CIMP classifications may be further established in a larger tumor set with known outcomes.

importance of aberrant DNA methylation and genomic instability for tumorigenesis has been described in several areas. Evidence for biallelic inactivation of tumor suppressor genes resulting from methylation silencing and LOH has been established in human cancers (9–11). Genome-wide demethylation has been shown to correlate with CIN phenotype in primary colorectal cancers (12). CpG island methylator phenotype (CIMP), a subtype of colorectal cancers originally discovered using a set of hypermethylated genomic loci that were significantly presented in the tumors (also referred to as type “C” or “cancer-specific” markers; ref. 13), has been suggested to precede the “mutator phenotype” in MSI tumor progression. Most interestingly, a recent study proposed that sporadic colorectal cancers with no CIN and CIN/MSI phenotypes were associated with CIMP (14), suggesting that CIN and CIMP represent two independent mechanisms of genetic and epigenetic instability.

Although earlier reports suggested that the designation of a CIMP subgroup is merely an event with normal distribution of methylation rate and may be age or MSI dependent, the observation of such a phenotype has since been documented using various methylation markers (15–18). Recently, Weisenberger et al. (19) conducted an elegant, systematic screening of 195 CpG island methylation markers using a quantitative MethyLight assay, and a new panel of robust markers was proposed to stratify CIMP tumors. All these carefully designed studies have shown that CIMP correlates with DNA mutations and clinicopathologic features, including MSI, mutations in KRAS and BRAF, and presentation on the proximal side of colon (18–23). Traditionally, the study of chromosomal imbalance has relied on methods such as flow cytometry–based ploidy assay, fluorescence in situ hybridization, and LOH analysis, yet only selected genomic regions were typically studied in each case. Consequently, these assays may not examine sufficient loci to distinguish the near-diploid genome from those with low-degree aberrations. The recent advances in DNA microarray technology have made the analysis of global aneuploidy possible using comparative genomic hybridization (CGH) and single-nucleotide polymorphism (SNP) arrays (24, 25). In particular, the latter offers a higher-resolution mapping at the genome-wide scale and provides a means to study CIN based on the degree of chromosomal aberrations.

The current study aimed to thoroughly investigate the relative importance of CIMP and CIN in colorectal cancer. The DNA methylation status of each tumor sample was determined using a quantitative bisulfite/PCR/ligase detection reaction (LDR)/Universal Array assay in a panel of 10 tumor suppressor genes (26). Tumor-specific promoter hypermethylation emerged in the p16INK4a, p14ARF, TIMP3, MGMT, and RARβ genes. These five markers, also referred to as classic CIMP genes, were used for CIMP study, where a total of 161 primary colorectal tumors were analyzed. Our results have shown that the presence of CIMP in primary colorectal tumors correlated with MSI and the BRAF mutation V600E. Chromosomal aberrations were analyzed in 60 primary tumors from the same study cohort using Affymetrix 50K Human Mapping arrays. Aneuploidy was found in >80% of cases when LOH, gains, or losses of chromosomes were considered. Our data reveal a variable extent of CIN among colorectal cancer samples, and tumors with a high level of hypermethylated loci associate with low-degree chromosomal aberrations. Similar results were obtained when CIMP was evaluated using MethyLight assay and markers of CACNA1G, IGF2, NEUROG1, RUNX3, and SOCS1. We have shown that the degree of chromosomal aberrations in tumors bearing a high level of hypermethylated loci (“CIMP positive”) differs significantly from other tumors within the same study cohort. This work emphasizes the importance of using quantitative and genome-wide approaches to study CIMP and CIN phenotypes in colorectal cancer.

Materials and Methods

Study population. The study population consisted of 161 male and female patients who presented at Memorial Sloan-Kettering Cancer Center with a colonic neoplasm between 1992 and 2004. Following a review of clinical and pathologic data stored in institutional databases, cases of sporadic colorectal cancers were selected for further study. This sample cohort was representative of colonic neoplasms at different stages of development. Primary colon tumors were located in the proximal colon, distal colon, or the rectum. Stage of disease (American Joint Committee on Cancer criteria) also varied and included patients who presented to Memorial Sloan-Kettering Cancer Center with primary colon adenocarcinomas and/or adenomas, patients with primary adenocarcinomas and synchronous metastases to the liver and/or lung, and patients with recurrent disease (local and distant). The study cohort included White non-Hispanic, White Hispanic, Black non-Hispanic, Asian, and American Indian patients. Ages ranged from 17 to 86 y with a median of 66 y. Biological specimens used in this study included primary colon adenocarcinomas, adenomas, and corresponding normal mucosa. All 161 analyzed colorectal cancer specimens were thoroughly examined by Memorial Sloan-Kettering Cancer Center pathologists and microdissected. A subset of 60 tumors with >80% carcinoma cells were chosen for Affymetrix SNP array experiments. Tissues were procured at the time of surgical resection and stored for future use under Institutional Review Board–approved protocols.

KRAS and BRAF mutation detection. Briefly, KRAS and BRAF mutations were detected using PCR/LDR approaches (27). The PCR
Fig. 1. Identification of tumor-specific aberrant promoter hypermethylation. The cytosine methylation levels of a total of 86 primary colorectal tumors and 66 matched normal mucosa were determined using bisulfite/PCR/LDR/Universal Array assay. Six CpG sites at each promoter region were analyzed for 10 candidate tumor suppressor genes (except RAR) with only four CpG sites. The color scale represents the percentage of methylation levels determined from the standard curves at each CpG dinucleotide as described previously (26). Promoter methylation status was determined by averaging the methylation levels of all interrogated cytosines within a particular gene. Using matched normal tissues as controls, the statistical significance of tumor-specific hypermethylation was calculated for each promoter region (see Table 1).

primers used for K-RAS were forward primer 5'-TTTCATTTTATTATTATTAAACGCCTGCTGA-3' and reverse primer 5'-GAATGGCTCTGACCAGTAAATAAGC-3'. The PCR primers used for BRAF were exon 15 forward primer 5'-TTCTAATACCTGCTCTAGTAGA-3' and reverse primer 5'-GGCCAAAAATTTAATCGTGGGA-3'. Fluorescently labeled LDR primers were designed for the seven common KRAS (Val, Asp, Ala, Arg, Ser, and Cys at codon 12 and Asp at codon 13) and BRAF V600E mutations. The sequences of KRAS LDR primers were reported previously (27). The LDR primer sequences for detecting BRAF mutation were as follows: wild-type specific primer, 5'-TTTAGTATTTTTAGGTCCTGCACCAG-3' and reverse primer 5'-GAATGGTCCTGCACCAGA-3'; mutant specific primer, 5'-TTTAGTATTTTTAGGTCCTGCACAGT-3'; and common primer, Phos-5'-GAATGGTCCTGCACCAGA-3' and reverse primer -TTTAGTAAAAA-3'. The PCR primers used for K-RAS were designed previously (26). The raw methylation levels of cytosines at CpG dinucleotides were determined using a quantitative bisulfite/PCR/LDR/Universal Array approach. The reaction conditions and primer sequences were described previously (26). Briefly, oligonucleotide primers for BAT25, BAT26, D2S123, DSS346, and D17S250 were designed and fluorescently labeled. The microsatellite assays were done in a multiplex fashion and the PCR products were resolved on an ABI 3100 DNA Sequencer (Applied Biosystems). Every colorectal cancer specimen and its corresponding normal tissue were analyzed. Allele size instability of microsatellite markers in cancer specimens was detected and distinguished from PCR stutter using Genotyper 3.7 software (Applied Biosystems). Allele size instability was defined as the identification in a tumor of a novel-size allele (i.e., an allele with a length not seen in normal colonic mucosa from the same individual). MSI status was determined in 161 colorectal cancers and in all 161 matched normal tissues. MSI was scored as present when at least two of the five markers showed size instability. Microsatellite stability (MSS) was always scored when all markers were allele size stable.

Detection of promoter methylation status using bisulfite/PCR/LDR/Universal Array. The methylation levels of cytosines at CpG dinucleotides were determined using a quantitative bisulfite/PCR/LDR/Universal Array approach. The reaction conditions and primer sequences were described previously (26). The raw methylation levels of the interrogated cytosines were calculated as the fluorescence ratios of Cy3/Cy5. By measuring the fluorescence intensity of an equal molar amount of Cy3- and Cy5-labeled LDR primers, a fluorescence intensity ratio (W = I_{Cy3}/I_{Cy5}) was used to normalize the label bias. To determine the percentage of methylation, calibration curves of interrogated cytosines were established by mixing the genomic DNAs of tumor cell lines (or in vitro methylated DNAs) and normal human lymphocytes in appropriate ratios. The methylation status of selected CpG sites present in the promoters of 10 tumor suppressor genes was initially determined in normal human lymphocytes using bisulfite sequencing. The majority of analyzed CpG cytosines did not show detectable hypermethylation, except for a few sites at the
Statistical analysis. All correlations were done by computing likelihood ratio $\chi^2$ statistics and associated $P$ values from the $n \times m$ contingency tables. When computing the $\chi^2$ statistics, the neighboring columns were merged into a single-cell count if the expected number of observations was smaller than 1. The odds ratio and 95% confidence interval were computed from $2 \times 2$ contingency tables extracted from the $n \times m$ tables (SYSTAT software). Multiple testing correction was done using the Benjamini-Hochberg method to find those tests with a false discovery rate of, at most, 5%.

Results

Quantitative bisulfite/PCR/LDR/Universal Array to determine CIMP. To select differentially hypermethylated candidate promoters in which their presence in tumors is statistically significant, a total of 86 primary colorectal tumors and 66 matched normal tissues were initially analyzed (Fig. 1). We used a recently developed quantitative bisulfite/PCR/LDR/Universal Array assay, which simultaneously detects 58 CpG loci methylation status in a panel of 10 tumor suppressor genes, p16NK4a, p14ARF, TIMP3, APC, RASSF1, CDH1, MGMT, p16NK4a and RARβ promoters. These few cytosines were either eliminated from the subsequent LDR assay or replaced with other neighboring unmethylated cytosines. The CpG cytosines with no detectable hypermethylation were considered as “zero” methylation for all the calibration curves. A promoter methylation status was determined by averaging the methylation levels of all six interrogated CpG sites (at the exception of RARβ with only four sites). In general, hypermethylation was scored when the averaged methylation of a promoter region was $\geq$10%. Using this cutoff point, we have confirmed that MGMT promoter hypermethylation correlated with a marked decrease of RNA expression (data not shown).

Determination of promoter methylation status using MethyLight. Real-time PCR to measure DNA methylation (MethyLight) was done as previously reported (19). Briefly, five CIMP-specific markers, including CACNA1C, IC2, NEUROG1, RUNX1, and SOCS1, were analyzed in each tumor sample using ABI 7500 (Applied Biosystems). ALL (Alu repeats) was used to normalize the amount of input bisulfite-treated genomic DNA. All the primer and probe sequences were published previously (19). The percentage of methylated reference (i.e., degree of methylation) at each locus was calculated by dividing the ratio of GENE/ALL in a sample by the ratio of GENE/ALL in SssI-treated human genomic DNA (presumably fully methylated) and multiplying this value by 100. Positive methylation in each locus was defined as percentage of methylated reference $>10$ as previously suggested (19). Tumor samples were scored as CIMP positive when three or more of five markers were hypermethylated.

Copy number and LOH analysis. The Affymetrix GeneChip Human Mapping 50K Xba 240 arrays (“SNP array”) were used to study overall chromosomal abnormalities. A total of 58,960 SNPs were detected on each array and the detailed protocol is described in the manufacturer’s manual and summarized by Tsafrir et al. (29). Briefly, genomic DNA was digested with XbaI and ligated onto adapters. PCR conditions were optimized to amplify fragments in the range of 250 to 2,000 bp in a GeneAmp PCR system 9700 (Applied Biosystems). PCR products were purified with a Qiagen MiniElute 96 UF PCR Purification Kit (Qiagen) and visualized on a 4% Tris-borate EDTA agarose gel to confirm that the average size was $\leq$180 bp. The fragmented DNA was then biotin labeled and hybridized to the array as described in the Affymetrix protocol. Following a staining and washing step, the array was scanned using GeneChip Scanner 3000 to generate image (DAT) and cell intensity (CEL) files. CEL files were imported into GeneChip DNA Analysis Software 4.0 (GDAS 4.0, Affymetrix) to generate the SNP calls. The Chromosomal Copy Number Analysis Tool 3.0 software (CNAT, Affymetrix) uses the probe intensity data as well as the SNP calls to generate both the single point analysis and genomic smoothed analysis copy number estimates and the corresponding $P$ values. The degree of chromosomal aberrations in a given sample was determined by counting the number of abnormal chromosomal arms with deletion, amplification, or LOH. The concepts and algorithms of CNAT were previously described (30). In addition, CNAT generates LOH calls based on the SNP calls. A default value of 0.5 Mb was used as the window size for averaging genomic smoothed analysis values.

### Table 1. The significance of tumor-specific promoter hypermethylation

<table>
<thead>
<tr>
<th>Gene</th>
<th>$\chi^2$</th>
<th>$P$</th>
<th>Gene</th>
<th>$\chi^2$</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MGMT</td>
<td>44.692</td>
<td>0.000</td>
<td>DAPK1</td>
<td>3.463</td>
<td>0.063</td>
</tr>
<tr>
<td>p14ARF</td>
<td>42.924</td>
<td>0.000</td>
<td>GSTP1</td>
<td>3.463</td>
<td>0.063</td>
</tr>
<tr>
<td>RARβ</td>
<td>35.183</td>
<td>0.000</td>
<td>APC</td>
<td>3.352</td>
<td>0.067</td>
</tr>
<tr>
<td>TIMP3</td>
<td>11.929</td>
<td>0.001</td>
<td>RASSF1</td>
<td>0.228</td>
<td>0.633</td>
</tr>
<tr>
<td>p16NK4a</td>
<td>10.106</td>
<td>0.001</td>
<td>CDH1</td>
<td>0.005</td>
<td>0.942</td>
</tr>
</tbody>
</table>

NOTE: $P$ values in boldface remain significant at 5% level after multiple testing correction.
DAPK1, GSTP1, and RARβ. PCR primers were designed to target sequences that are essentially free of CpG dinucleotides, allowing equal amplification of both methylated and unmethylated DNAs in multiple genomic regions. The methylation level of each interrogated cytosine was determined using LDR and displayed on a Universal Array (26). Six CpG dinucleotide sites per promoter region (except for RARβ, where four sites were used) were analyzed. Promoter methylation status was scored by averaging the methylation levels of all interrogated CpGs of that gene. This criterion effectively eliminated background signals and low-level methylation that may not be relevant to the control of gene expression. An example is shown in the RARβ gene, where the fourth CpG site was frequently methylated from low to medium levels in the matched normal tissues (Fig. 1). By averaging the methylation levels of all four sites in RARβ, the effect of the sporadic methylation site was reduced and the scoring of false-positive hypermethylation was minimized. Five promoter regions (p16INK4a, p14ARF, MGMT, RARβ, and TIMP3) showed statistically significant increases of methylation in tumors compared with normal tissues (P < 0.001; Table 1). Using these five markers in a later validation study, the methylation status of 161 colorectal cancer samples was profiled and the number of tumor-specific, hypermethylated promoters in each sample was considered in a continuous fashion (Fig. 2; Supplementary Fig. S1). The lack of clear bimodal distribution of the hypermethylated loci in colorectal cancers disfavored a simple dichotomization of the analyzed samples; instead, CIMP was regarded as a unimodal distribution in all subsequent studies.

**CIMP correlates with MSI and BRAF mutation.** A characteristic feature of the previously reported CIMP subgroup of colorectal cancers is its substantial enrichment in MSI cases and BRAF mutations. To evaluate whether CIMP-positive samples identified in our approach had similar genetic alterations, we established the MSI and BRAF and KRAS mutation profiles in 161 colorectal cancer samples (Fig. 2). Our data show that CIMP correlates with MSI, suggesting that MSI tumors tend to have a higher proportion of methylated loci than their MSS counterparts (P = 0.075; Table 2). A significant difference was also observed between BRAF V600E mutations and the number of methylated genes (Table 2). Individuals with BRAF mutations had ~18-fold risk of being CIMP positive than individuals without BRAF mutations [P = 0.0001; odds ratio, 18.45 (95% confidence interval, 4.51-75.11)]. The presence of higher percentage of BRAF mutants in the CIMP subgroup indicates that BRAF mutations significantly correlate with the hypermethylation phenotype (P = 0.00005).

Because BRAF mutations are known to occur frequently in MSI tumors (31), we further investigated whether the highly significant correlation between BRAF mutation and CIMP may be MSI dependent. Tumors samples were split into MSI and MSS subgroups according to their BRAF mutation status (Table 2). In the MSI category, BRAF mutations were only identified in samples with a high level of methylated loci but were not present in others (P = 0.022). In contrast, in the MSS subgroup, samples with no methylated loci harbored a BRAF mutation, which accounted for nearly two thirds of all BRAF mutants in MSS (P = 0.03). These findings indicate a higher frequency of BRAF mutations in the CIMP-positive subgroup independent

**Table 2. The correlation between CIMP and gender, location, MSI, and KRAS and BRAF mutations**

<table>
<thead>
<tr>
<th>No. methylated loci (CIMP)</th>
<th>Total no., n (%)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Total (primary CRC)</td>
<td>49</td>
<td>45</td>
</tr>
<tr>
<td>Men</td>
<td>26</td>
<td>23</td>
</tr>
<tr>
<td>Women</td>
<td>23</td>
<td>22</td>
</tr>
<tr>
<td>Distal</td>
<td>39</td>
<td>29</td>
</tr>
<tr>
<td>Proximal</td>
<td>10</td>
<td>16</td>
</tr>
<tr>
<td>KRAS_WT</td>
<td>32</td>
<td>28</td>
</tr>
<tr>
<td>KRAS_Mut</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>No info</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>MSI-H</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>MSS</td>
<td>40</td>
<td>32</td>
</tr>
<tr>
<td>No info</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>BRAF_WT</td>
<td>42</td>
<td>44</td>
</tr>
<tr>
<td>BRAF_Mut</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>No info</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>MSI-H BRAF_WT</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>MSI-H BRAF_Mut</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>MSS BRAF_WT</td>
<td>36</td>
<td>32</td>
</tr>
<tr>
<td>MSS BRAF_Mut</td>
<td>4</td>
<td>0</td>
</tr>
</tbody>
</table>

Abbreviation: CRC, colorectal cancer.
of the tumor MSI status. Thus, our results strongly suggest that the highly significant correlation between BRAF mutation and CIMP does not simply reflect a common feature of MSI.

**CIMP associates with low-degree chromosomal aberrations.** The chromosomal aberration profiles of 60 primary colorectal tumors from the same study cohort were established. Data obtained from the matched normal tissues, in conjunction with Affymetrix reference controls, were used as scoring criteria to minimize the effect of individual germ-line copy number variations and random noise of allele intensity at each SNP site (data not shown). To validate SNP array results, the overall DNA fold change in tumor samples was assessed by CGH arrays (29). A strong genome-wide correlation between SNP-derived and CGH-derived data was observed in all five tested carcinomas (an example is shown in Supplementary Fig. S2). In addition, copy number gains and losses at individual loci were determined using conventional assays. Frequent DNA amplification of chromosomal region 20q13.2 was confirmed in all six tested adenocarcinomas by fluorescence in situ hybridization, with up to 10- to 20-fold copy number increases (29). The common chromosomal deletion at 5q21-22 was corroborated using the analysis of microsatellite markers (Supplementary Fig. S3); allelic imbalance was detected by comparing tumors with the matched adjacent normal tissues.

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**Fig. 3.** A genome-wide profiling of chromosomal aberrations in 60 primary colorectal tumors. Chromosomal gains and losses are depicted in a semiquantitative heat map diagram. The color scale represents normalized copy number changes: loss of alleles (green), gain of one allele (red), gain of more than one allele (dark red), and no change of copy number (white). The yellow bars across multiple chromosomal loci represent the copy number neutral LOH events. X and Y axes, chromosome number and sample ID of each tumor, respectively. Tumor samples are organized and displayed based on their degree of aneuploidy. The dotted lines divide each chromosome into p and q arms. Four color-coded columns on the right depict the tumor CIN, MSI, and CIMP status. The degree of CIN is presented in a continuous fashion, with green to red colors representing the lowest to highest levels of overall chromosomal aberrations. Codes on the MSI column: red, MSI-high; orange, MSI-low; green, MSS. Two methods (LDR/Universal Array and MethyLight) were used to classify CIMP. Codes on the CIMP (LDR/array) column: red, tumors with four to five hypermethylated CIMP loci; orange, tumors with one to three hypermethylated CIMP loci; green, tumors with no hypermethylated marker. Codes on the CIMP (MethyLight) column: red, CIMP-positive; orange, CIMP-negative contains one to two hypermethylated CIMP loci; green, tumors with no hypermethylated marker.

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7 M. D. Bacolod et al., in preparation.
Consistent with the SNP data, samples with no change in copy number were correctly identified as such using the microsatellite marker based approach.

As shown in Fig. 3, sample nos. 1 to 8 showed relatively stable genomes with no apparent aneuploidy, whereas >80% of the analyzed colorectal cancers displayed characteristic CIN features of copy number alterations and allelic imbalances. DNA copy number gains occurred most frequently in chromosomes 7p, 7q, 8q, 13q, and 20q, whereas copy number losses were most often detected in chromosomes 1p, 4p, 4q, 8p, 14q, 15q, 17p, 18p, and 18q. We noticed that the majority of samples with MSI were not diploid and possessed low-degree chromosomal aberrations. Although this observation seems to contradict the conventional viewpoint of MSI (typically considered as diploid), it is consistent with the results of a recent study, which compared the extent of CIN in 23 MSI and 23 MSS sporadic colorectal cancers using array CGH (32). This study showed that chromosomal aberrations are present in both MSI and MSS sporadic colorectal cancers, but the degree of CIN differs significantly between the two groups.

Interestingly, genomic instability varied among CIMP-positive samples, with the most stable genome seen in sample no. 2 and the least stable one in no. 57. Sample nos. 10, 11, 12, and 14 were clearly not exempt from chromosomal aberrations; however, their abnormalities were less substantial than those in nos. 40 and 57. For instance, significant chromosomal losses and LOH events at chromosome 21 were identified in sample nos. 10, 11, and 12, whereas gains at chromosomes 8 and 9 were found in sample nos. 12, 13, and 14. To study the relationship between CIMP and CIN, we classified the CIN phenotype using two criteria. When defining CIN as aberrations in chromosomes 2p, 3p, 5q, 17p, and 18q (criterion 1) as previously proposed (14), sample nos. 17 to 60 were scored as CIN positive and inversely correlated with CIMP \( (P = 0.075) \). Alternatively, when considering CIN as the presence of significant gains, losses, or LOH in any one or more chromosomal regions (criterion 2), the degree of chromosomal aberrations varied in a wide spectrum across all tumors. This is of particular interest because samples scored as “CIN negative” (e.g., nos. 9-16) by criterion 1 were revealed as having low-degree aberrations. Consequently, under the latter criterion, which considers CIN as a continuous variable, only sample nos. 1 to 8 were qualified as true CIMP negative, and tumors bearing high levels of hypermethylated loci (i.e., CIMP positive) exhibited a significant correlation with low-degree CIN \( (P = 0.012) \).

To ensure that the relationship between CIN and CIMP was independent of the method of scoring CIMP, we used the five robust methylation markers and MethyLight assay to identify CIMP as proposed by Weisenberger et al. All experiments were conducted and the 60 tumors were analyzed based on the published protocol (Supplementary Fig. S4). As shown in Fig. 3, samples identified as CIMP positive showed a significant correlation with MSI \( (P = 0.0008, \text{two-tailed Fisher’s exact test}) \). Furthermore, under criterion 1, we found an inverse relationship between CIMP and CIN \( (P = 0.001, \text{two-tailed Fisher’s exact test}) \). When CIN was viewed as a gradient distribution of high- to low-degree aberrations (criterion 2), a significant correlation was also found between CIMP and low-degree aberrations \( (P = 0.013) \). As shown in Fig. 4A, the relationships between MSI, CIMP, and CIN are depicted in a Venn diagram, which illustrates that CIMP-positive samples generally comprise MSI and low-CIN tumors. We conclude that both quantitative methylation assays consistently identify CIMP-positive groups. The significance of the correlation between CIN and CIMP is
independent of the method used for quantitative methylation detection.

In the course of studying genome instability in colorectal cancer samples, we tested whether a different conclusion on the inverse relationship between CIMP and CIN could be made using selected LOH at loci of the 15 tumor suppressor genes mentioned above. LOH was examined at each gene locus using the Affymetrix 50K Human Mapping array (Fig. 4B). In general, LOH events were found at any of the analyzed tumor suppressor loci in nearly 80% of the tumors analyzed; however, tumors with a high level of methylated loci tended to have fewer LOH events. Interestingly, the majority of CIMP-positive samples (8 of 12) had low-degree or no LOH. This result is consistent with the above observation that colorectal cancers with high-level methylation have a different type of genetic abnormality.

Discussion

We have used a recently developed, accurate, and quantitative bisulfite/PCR/LDR/Universal Array approach to test the CIMP in colorectal cancers. Our data are consistent with previous studies that have indicated that the CIMP-positive tumors correlate with MSI and BRAF mutation and are more likely to occur in tumors of the proximal colon. We have also shown that the correlation between CIMP-positive and BRAF V600E mutation is independent of MSI status. A subset of colorectal cancers were assayed using MethyLight and markers of CACNA1G, IGF2, NEUROG1, RUNX3, and SOCS1, which showed the same correlation with clinicopathologic features. All these findings suggest that CIMP-positive tumors are a distinct subgroup of colorectal cancers. We also reported that a subset of colorectal cancers determined using MethyLight and the five designated markers rendered a bimodal CIMP pattern (Supplementary Fig. S4), which is contrary to the continuous phenotype measured using the p16INK4a, p14ARF, TIMP3, MGMT, and RARβ genes. Addressing the extent of methylation distribution is important not only for understanding the underlying biology but also for CIMP classification and data analysis. Some previous studies have implied that the number of MSI tumors and the methylation markers used in a study influenced CIMP distribution (18, 19). Because a consistent number of MSI tumors were present in both the cohort of 161 colorectal cancers and the subset of 60 tumors (20-24%), the diverse distribution patterns most likely resulted from the choice of methylation markers in the analysis. Our study did not attempt to rediscover CIMP markers, as several robust ones have already been published, and a unified list is essential for future studies. Rather, we sought to emphasize the importance of using quantitative assays for methylation profiling.

In our study, CIMP correlated with neither gender nor KRAS mutations, regardless of the methods used for CIMP scoring. Associations between CIMP and either gender or KRAS mutations are controversial. In a large population-based study, Samowitz et al. (22) showed that CIMP correlated with KRAS mutations but not with gender. However, recent studies reported that CIMP significantly associated with wild-type KRAS and female gender (18, 19). The discrepancies may result from the use of different methylation markers and/or sampling variation. We favor the latter explanation, as similar results of the absence of correlations were obtained whether the tumor cohort was analyzed by the LDR/Universal Array or the MethyLight assay (Supplementary Fig. S4).

Our findings provide additional nuances to a recent study suggesting that CIN and CIMP represent two independent mechanisms of genomic instability in sporadic colorectal cancers (14). First, in our study, promoter methylation status was determined using quantitative methylation assays. These methods have advantages over a nonquantitative and highly sensitive assay (e.g., methylation-specific PCR) that tends to detect genomic loci with biologically insignificant levels of methylation. Such an approach has often resulted in an overestimation of CIMP or even failure to identify it. Second, when the CIN status of a tumor was assessed based on the presence of one or more LOH events at eight microsatellite sequences located on chromosomes 2p, 3p, 5q, 17p, and 18q, we confirmed a previous report of an inverse correlation between CIN and CIMP. This observation is independent of the method of CIMP scoring. Nevertheless, because CIN leads to aneuploidy, in which chromosomal regions are either missing or duplicated, aberrations found in any chromosome may be viewed as CIN. Indeed, by using high-throughput assays, such as CGH and SNP array technology, we and others have shown common and broad allelic imbalance events in chromosomes 4p, 4q, 5q, 12q, 14q, 15q, 17p, 18q, and 20p (24, 25, 29) and frequent gains in chromosomes 7p, 7q, 8q, 13q, and 20q (32). These results allow a genome-wide analysis of CIN phenotype and provide an alternative for determination of CIN status in one third of colorectal cancers that had moderate- to low-level chromosomal aberrations. Third, the degree of aneuploidy in the analyzed colorectal cancer samples is spread over a wide spectrum. Thus, a limited LOH loci approach to determine CIN can become challenging and will likely identify tumors with the most chromosomal abnormalities, but not those that tend to have low-degree chromosomal aberrations, such as MSI and CIMP-positive tumors. Because MSI constitutes a good proportion of CIMP-positive samples (>50%), our findings suggest that the reported inverse correlation between CIN and CIMP may reflect the relationship between MSI and CIN (Fig. 4A). It is noteworthy that a subset of CIMP-positive tumors, albeit of small size, has a high degree of chromosomal aberrations. Whether this subset has a different clinical outcome needs to be further investigated. In summary, we conclude that CIMP-positive tumors are not free of chromosomal abnormalities; rather, they have significant low-degree chromosomal aberrations. This tendency of CIMP-positive tumors to have low-degree CIN most likely results from the contribution of their MSI feature. Our results indicate that the stratification of sporadic colorectal cancers based on epigenetic and genetic instabilities is best described as a quantitative assay and a genome-wide approach, respectively.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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References

CpG Island Methylator Phenotype Associates with Low-Degree Chromosomal Abnormalities in Colorectal Cancer
