Single-CpG-resolution methylation analysis identifies clinicopathologically aggressive CpG island methylator phenotype clear renal cell carcinomas

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To clarify the significance of DNA methylation alterations during renal carcinogenesis, methylation analysis using single-CpG-resolution Infinium array was performed on 29 normal renal cortex tissue (C) samples, 107 non-cancerous renal cortex tissue (N) samples and 109 tumors tissue (T) samples. DNA methylation levels at 4830 CpG sites were already altered in N samples compared with C samples. Unsupervised hierarchical clustering analysis based on DNA methylation levels at the 801 CpG sites, where DNA methylation alterations had occurred in N samples and were inherited by and strengthened in T samples, clustered clear cell RCCs into Cluster A (n = 90) and Cluster B (n = 14). Clinicopathologically aggressive tumors were accumulated in Cluster B, and the cancer-free and overall survival rates of patients in this cluster were significantly lower than those of patients in Cluster A. Clear cell RCCs in Cluster B were characterized by accumulation of DNA hypermethylation on CpG islands and considered to be CpG island methylator phenotype (CIMP)-positive cancers. DNA hypermethylation of the CpG sites on the FAM150A, GRM6, ZNF540, ZFP42, ZNF154, RIMS4, PCDHAC1, KHDRBS2, ASC2L, KCNQ1, PRAC, WNT3A, TRH, FAM78A, ZNF671, SLC12A5 and NKKX2-2 genes became hallmarks of CIMP in RCCs. On the other hand, Cluster A was characterized by genome-wide DNA hypomethylation. These data indicated that DNA methylation alterations at precancerous stages may determine tumor aggressiveness and patient outcome. Accumulation of DNA hypermethylation on CpG islands and genome-wide DNA hypomethylation may each underlie distinct pathways of renal carcinogenesis.

Introduction

Clear cell renal cell carcinoma (RCC) is the most common histological subtype of adult kidney cancer and frequently affects working-age adults in midlife. In general, RCCs at an early stage are curable by nephrectomy. However, some RCCs relapse and metastasize to distant organs, even if the resection has been considered complete (1). Such clinicopathological diversity may be attributable to distinct pathways of renal carcinogenesis (2). It is well known that clear cell RCCs are characterized by inactivation of the Von Hippel–Lindau tumor-suppressor gene (3). In addition, systematic resequencing and exome analysis of RCCs suggested that non-cancerous renal cortex tissue obtained from patients with RCCs is already at the precancerous stage associated with DNA methylation alterations, even though no remarkable histological changes are evident and there is no association with chronic inflammation or persistent infection with viruses or other pathogenic microorganisms. Genome-wide analysis using BAMCA revealed that DNA methylation status in non-cancerous renal cortex tissue at the precancerous stage was basically inherited by the corresponding clear cell RCC in individual patients (19). DNA methylation alterations at the precancerous stage may confer further susceptibility to genetic and epigenetic alterations and generate more malignant clear cell RCCs (2,13). However, in our previous studies using BAMCA, the resolution and the number of probes were limited. Therefore, further analysis is needed to clarify the significance of DNA methylation alterations in renal carcinogenesis.

Recently, methylation analysis using the Infinium array has made it possible to interrogate 27 000 highly informative CpG sites at single-CpG resolution (21). In order to clarify the significance of DNA methylation alterations during renal carcinogenesis, we used the Infinium BeadChip system to perform genome-wide DNA methylation analysis of 29 samples of normal renal cortex tissue (C) obtained from patients without any primary renal tumors, 107 samples of non-cancerous renal cortex tissue (N) from patients with clear cell RCCs and 109 samples of tissue from the tumors (T) themselves. Correlations between the genome-wide DNA methylation profiles and clinicopathological parameters were then examined.

Materials and methods

Patients and tissue samples

The 109 T samples and corresponding 107 N samples showing no remarkable histological changes were obtained from materials that had been surgically resected from 110 patients with primary clear cell RCCs. These patients did not receive preoperative treatment and underwent nephrectomy at the National Cancer Center Hospital, Tokyo, Japan. There were 79 men and 31 women with a mean (±SD) age of 62.8 ± 10.3 years (range 36–85 years). Histological diagnosis was made in accordance with the World Health Organization classification (22) (Supplementary Figure S1, available at Carcinogenesis Online). All the tumors were graded on the basis of criteria described previously (23) and classified according to the pathological Tumor-Node-Metastasis (TNM) classification (24). The criteria for macroscopic configuration of RCC (17–19) followed those established for hepatocellular carcinoma (HCC): type 3 (contiguous multinodular type). HCCs show poorer histological differentiation and a higher incidence of intrahepatic metastasis than type 1 (single nodular type) and type 2 (single nodular type with extranodal growth) HCCs (25). The presence or absence of vascular involvement was examined microscopically on slides stained with hematoxylin–eosin and elastica van Gieson. The presence or absence of tumor thrombi in the main trunk of the renal vein was examined macroscopically.
RCC is usually enclosed by a fibrous capsule and well demarcated, and hardly ever contains fibrous stroma between cancer cells. Therefore, we were able to obtain cancer cells from surgical specimens, avoiding contamination with both non-cancerous epithelial cells and stromal cells.

For comparison, 29 samples of normal renal cortex tissue (C1–C29) were obtained from materials that had been surgically resected from 29 patients without any primary renal tumor. These patients included 18 men and 11 women with a mean (±SD) age of 61.4 ± 10.8 years (range 31–81 years). Of these patients, 22 had undergone nephrectomy for urothelial carcinomas of the renal pelvis and ureter, 6 had undergone nephrectomy with resection of retroperitoneal sarcoma around the kidney, and the remaining 1 had undergone paracystic lymph node dissection for metastatic germ cell tumor, which resulted in simultaneous nephrectomy because it was not possible to preserve the renal artery. All patients included in this study provided written informed consent, and the study was approved by the Ethics Committee of the National Cancer Center, Tokyo, Japan.

**Infinium assay**

High-molecular-weight DNA from fresh frozen tissue samples was extracted using phenol-chloroform, followed by dialysis (26). Five-hundred-nanogram aliquots of DNA were subjected to bisulfite conversion using an EZ DNA Methylation Kit (Zymo Research, CA). Subsequently, the methylation status at 27 578 CpG loci was examined at single-CpG resolution using the Infinium HumanMethylation27 Bead Array (Illumina, San Diego, CA). This array contains CpG sites located within the proximal promoter regions of the transcription start sites of 14 475 consensus coding sequences in the National Center for Biotechnology Information Database. On average, two assays were selected per gene, and from 3 to 20 CpG sites for more than 200 cancer-related and imprinted genes. Forty control probes were employed for each array; these included staining, hybridization, extension, bisulfite conversion and negative controls. An Evogen rotor (Tecan, Switzerland) was used for automated sample processing. Whole-genome amplification was performed using the Infinium Assay Kit (Illumina (21)). After hybridization, the specifically hybridized DNA was fluorescence labeled by a single-base extension reaction and detected using a BeadScan reader (Illumina) in accordance with the manufacturer’s protocols. The data were then assembled using GenomeStudio methylation software (Illumina). At each CpG site, the ratio of the fluorescent signal was measured using a methylation probe relative to the sum of the methylated and unmethylated probes, i.e. the so-called \( B \)-value, which ranges from 0.00 to 1.00, reflecting the methylation level of an individual CpG site.

**Statistics**

In the Infinium assay, the call proportions (\( P \)-values for detection of signals above the background <0.01) for 32 probes (shown in Supplementary Table S1, available at Carcinogenesis Online) in all of the tissue samples examined were less than 90%. Since such a low proportion may be attributable to polymorphism at the probe CpG sites, these 32 probes were excluded from the present assay. In addition, all CpG sites on chromosomes X and Y were excluded, to avoid any gender-specific methylation bias, leaving a final total of 26 454 autosomal CpG sites.

Infinium probes showing significant differences in DNA methylation levels between the 29 C and 107 N samples were identified by a logistic model adjusted by sex, age and experimental batch. Ordered differences from 29 C to 107 N and then to 109 T samples themselves were examined by the cumulative logit model adjusted by sex, age and experimental batch. Differences of DNA methylation status between 104 paired samples of N and the corresponding T obtained from a single patient and assayed in the same experimental batch were examined by Wilcoxon matched pairs test. A false discovery rate (FDR) of \( q = 0.01 \) was considered significant. Unsupervised hierarchical clustering (Euclidan distance, Ward method) based on DNA methylation levels (\( \Delta \)KIl) was performed in patients with clear cell RCCs. Correlations between clusters of patients and clinicopathological parameters were examined using Wilcoxon rank sum test and Fisher’s exact test. Survival curves of patients belonging to each cluster were calculated by the Kaplan–Meier method, and the differences were compared by the log-rank test. The number of Infinium assay probes showing DNA hyper- or hypomethylation in each cluster and the average DNA methylation levels (\( \Delta P_{C,N} \)) of each cluster were examined using Wilcoxon rank sum test at a significance level of \( P < 0.05 \). The CpG sites discriminating the clusters were identified by Fisher’s exact test and random forest analysis (27).

**Results**

**DNA methylation alterations during renal carcinogenesis**

First, DNA methylation levels of representative CpG sites based on the Infinium assay were clearly verified using a highly quantitative pyrosequencing method (Supplementary Figure S2, available at Carcinogenesis Online). With regard to the well-known methylation-silencing Von Hippel–Lindau tumor-suppressor gene (probe Target ID: cg22782492), DNA hypermethylation (\( \Delta P_{C,N} > 0.1 \)) was detected in 12 (12%) of 104 patients, for whom both N and T samples were assayed in the same experimental batch. This incidence corresponded to that in previous reports (28,29). Taken together, the data confirmed the reliability of the present Infinium assay.

Although precancerous conditions in the kidney have been rarely described, our previous study suggested that N samples are already at precancerous stages, from the viewpoint of altered DNA methylation, despite the absence of any remarkable histological changes and the lack of association with chronic inflammation and persistent infection with viruses or other pathogenic microorganisms (17–20). (a) In fact, the logistic model adjusted by sex, age and experimental batch revealed that DNA methylation levels on 4830 probes were already altered in N samples compared with those in C samples (FDR, \( q = 0.01 \), Table I). (b) In order to reveal DNA methylation alterations inherited by clear cell RCCs themselves, ordered differences of DNA methylation level from C to N and then to T samples were examined by the cumulative logit model adjusted by sex, age and experimental batch. Ordered differences from C to N and then to T samples were observed on 11 089 probes (FDR, \( q = 0.01 \), Table I). (c) In order to reveal the cancer-prone DNA methylation alterations, differences in DNA methylation levels between 104 paired samples of N and T assayed in the same experimental batch were examined using the Wilcoxon matched pairs test. Significant differences between N and the corresponding clear cell RCCs themselves were observed on 10 870 probes (FDR, \( q = 0.01 \), Table I).

DNA hypermethylation frequently occurred at the very early stages of renal carcinogenesis (\( a \) in Table I), whereas DNA hypomethylation was also observed during progression to established cancers (\( b \) and \( c \) in Table I). Eight hundred and one probes satisfied all of the above criteria (\( a \)–\( c \)) (Table I): DNA methylation alterations on these 801 probes (Supplementary Table S2, available at Carcinogenesis Online) were already evident in N samples, and were inherited and strengthened in T samples.

**Table I. DNA methylation alterations during renal carcinogenesis**

<table>
<thead>
<tr>
<th>(a) The probes on which DNA methylation levels were altered in samples of non-cancerous renal cortex tissue (N) obtained from patients with clear cell RCCs relative to those in samples of normal renal cortex tissue (C) obtained from patients without any primary renal tumor. (Logistic model adjusted by sex, age and experimental batch; FDR, ( q = 0.01 ))</th>
<th>DNA hypermethylation (( \beta_C \rangle \beta_N ))</th>
<th>4580</th>
<th>DNA hypomethylation (( \beta_C \rangle \beta_N ))</th>
<th>241</th>
<th>Total</th>
<th>4830</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Cumulative logit model adjusted by sex, age and experimental batch; FDR, ( q = 0.01 ))</td>
<td>DNA hypermethylation (( \beta_C \rangle \beta_N ))</td>
<td>6563</td>
<td>DNA hypomethylation (( \beta_C \rangle \beta_N ))</td>
<td>4436</td>
<td>Total</td>
<td>11 089</td>
</tr>
<tr>
<td>(b) The probes on which DNA methylation levels showed ordered differences from C to N and then to tumorous tissue (T) samples</td>
<td>DNA hypermethylation (( \Delta P_{C,N} &gt; 0 ))</td>
<td>5408</td>
<td>DNA hypomethylation (( \Delta P_{C,N} &lt; 0 ))</td>
<td>5462</td>
<td>Total</td>
<td>10 870</td>
</tr>
</tbody>
</table>

\( ^a \)Among the 4589 probes, 2675 showed DNA hypermethylation in T samples than in C samples (\( \beta_C \rangle \beta_N \); FDR, \( q = 0.01 \)).

\( ^b \)Among the 2414 probes, 126 showed DNA hypomethylation in T samples than in C samples (\( \beta_C \rangle \beta_N \); FDR, \( q = 0.01 \)).
Fig. 1. Unsupervised hierarchical clustering using DNA methylation levels ($\Delta \beta_{T-N}$) on the 801 probes in 104 patients with clear cell RCCs. The 801 probes satisfied all of the criteria (a), (b) and (c) in ‘DNA methylation alterations during renal carcinogenesis’ in Results and Table I. On the 801 probes, DNA methylation alterations occurred at the precancerous stages and were inherited by and strengthened in clear cell RCCs themselves. (A) 104 patients with clear cell RCCs were hierarchically clustered into Cluster A ($n = 90$) and Cluster B ($n = 14$). The DNA methylation levels ($\Delta \beta_{T-N}$) are shown in the color range maps. The cluster trees for patients and probes are shown at the top and left of the panel, respectively. (B) The cancer-free ($P = 3.59 \times 10^{-6}$) survival rates of Stage I–III patients in Cluster B were significantly lower (log-rank test) than those of patients in Cluster A. Overall ($P = 1.32 \times 10^{-2}$) survival rates of all patients in Cluster B were significantly lower (log-rank test) than those of patients in Cluster A.
Epigenetic clustering of clear cell RCCs

Unsupervised hierarchical clustering using DNA methylation levels ($\Delta \beta_{T-N}$) on the above 801 probes, on which DNA methylation alterations occurred at the precancerous stages and may continuously participate in renal carcinogenesis, subclustered 104 patients with clear cell RCCs, of whom both N and T samples were assayed in the same experimental batch, into Cluster A ($n = 90$) and Cluster B ($n = 14$). The clinicopathological parameters of clear cell RCCs belonging to Clusters A and B are summarized in Table II. The number of samples for each TNM stage is also described in Supplementary Table S3, available at Carcinogenesis Online. Epigenetic clustering of clear cell RCCs was dependent on neither age nor sex of the patients (Table II). Clear cell RCCs belonging to Cluster B had a larger diameter, more frequent macroscopically evident extranodular (type 2) or multinodular (type 3) growth, vascular involvement, renal vein tumor thrombi, infiltrative growth, tumor necrosis and renal pelvic invasion, and also had higher histological grades and pathological TNM stages than those in Cluster A (Table II). Figure 1B shows the Kaplan–Meier survival curves of patients belonging to Clusters A and B. The period covered ranged from 42 to 4024 days (mean, 1821 days). The cancer-free and overall survival rates of patients in Cluster B were significantly lower than those of patients in Cluster A ($P = 3.59 \times 10^{-6}$ and $P = 1.32 \times 10^{-7}$, respectively; Figure 1B).

Table II. Correlation between the subclassification of patients with clear cell RCCs based on DNA methylation profiles and the clinicopathological parameters

<table>
<thead>
<tr>
<th>Clinicopathological parameters</th>
<th>Cluster A ($n = 90$)</th>
<th>Cluster B ($n = 14$)</th>
<th>$P^*$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>62.08± 6.736±</td>
<td>8.36×10^-2</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>Male</td>
<td>63± 11</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>27± 3</td>
<td></td>
</tr>
<tr>
<td>Tumor diameter (cm)</td>
<td>5.10± 8.75±</td>
<td>1.07×10^-4</td>
<td></td>
</tr>
<tr>
<td>Cancer configuration</td>
<td>Type 1</td>
<td>37± 1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Type 2</td>
<td>29± 2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Type 3</td>
<td>24± 11</td>
<td></td>
</tr>
<tr>
<td>Predominant histological grade</td>
<td>G1</td>
<td>47± 1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G2</td>
<td>35± 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>7± 7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>1± 2</td>
<td></td>
</tr>
<tr>
<td>Highest histological grade</td>
<td>G1</td>
<td>8± 0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G2</td>
<td>43± 1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>24± 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>15± 9</td>
<td></td>
</tr>
<tr>
<td>Vascular involvement</td>
<td>Negative</td>
<td>54± 1</td>
<td>2.45×10^-4</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>36± 13</td>
<td></td>
</tr>
<tr>
<td>Renal vein</td>
<td>Negative</td>
<td>69± 5</td>
<td>3.38×10^-3</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>21± 9</td>
<td></td>
</tr>
<tr>
<td>Tumor thrombi</td>
<td>Expansive</td>
<td>84± 7</td>
<td>1.86×10^-4</td>
</tr>
<tr>
<td>Most aggressive growth pattern</td>
<td>Expansive</td>
<td>57± 4</td>
<td>2.06×10^-3</td>
</tr>
<tr>
<td>Tumor necrosis</td>
<td>Negative</td>
<td>71± 2</td>
<td>4.86×10^-6</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>19± 12</td>
<td></td>
</tr>
<tr>
<td>Invasion to renal pelvis</td>
<td>Negative</td>
<td>83± 10</td>
<td>3.98×10^-2</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>7± 4</td>
<td></td>
</tr>
<tr>
<td>Pathological TNM stage</td>
<td>Stage I</td>
<td>50± 0</td>
<td>5.41×10^-3</td>
</tr>
<tr>
<td></td>
<td>Stage II</td>
<td>1± 1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Stage III</td>
<td>23± 9</td>
<td></td>
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<tr>
<td></td>
<td>Stage IV</td>
<td>16± 4</td>
<td></td>
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</table>

The number of samples in each TNM stage was described in Supplementary Table S3, available at Carcinogenesis Online. $^*$P-values of $<0.05$ are in italics.

DNA methylation profiles of clear cell RCCs belonging to each cluster

The distribution of DNA methylation levels ($\Delta \beta_{T-N}$) in all 26 454 probes for 104 clear cell RCCs belonging to Cluster A or B is summarized along chromosomes in Figure 2A. Clear cell RCCs belonging to Cluster B clearly showed accumulation of DNA hypermethylation ($\Delta \beta_{T-N} > 0.1$) relative to DNA hypomethylation ($\Delta \beta_{T-N} < 0.1$), whereas clear cell RCCs belonging to Cluster A showed greater DNA hypomethylation ($\Delta \beta_{T-N} < -0.1$) relative to DNA hypermethylation (Figure 2A).

Hallmark CpG sites of CIMP-positive clear cell RCCs

Scattergrams of DNA methylation levels ($\beta_{T}$-values) in T and the corresponding N samples from representative patients with clear cell RCCs belonging to Clusters A and B (Supplementary Figure S3, available at Carcinogenesis Online) indicated that probes for which DNA methylation levels were low in the N samples and for which the degree of DNA hypermethylation in T samples relative to the corresponding N samples was prominent (marked by red circles in panels E to H in Supplementary Figure S3, available at Carcinogenesis Online) were obvious only in Cluster B, and not in Cluster A. Therefore, in order to discriminate clear cell RCCs belonging to Cluster B from those belonging to Cluster A, we first focused on the probes for which the average \( \tilde{\beta} \) value in all N samples was less than 0.2 and the incidence of more than 0.4$\Delta \beta_{T-N}$ was markedly higher in Cluster B relative to Cluster A ($P < 1.98 \times 10^{-6}$; Fisher’s exact test). Among such probes, 16 (the FAM150A, GRM6, ZNF540, ZFP42, ZNF154, RIMS4, PCDHAC1, KHRD852, ASC12, KCNQ1, PRAC, ZNF154, WNT3A, TRH, FAM78A and ZNF671 genes) showed $>0.4\hat{\beta}_{T-N}$ in 6 (42.2%) or more RCCs among the 14 belonging to Cluster B, but only in 2 (2.2%) or fewer RCCs among the 90 belonging to Cluster A (Table IIIA). DNA methylation levels ($\Delta \beta_{T-N}$) on the 16 CpG sites differed completely between Clusters A and B (Supplementary Figure S4, available at Carcinogenesis Online).
random forest analysis (27) (Supplementary Figure S5, available at Carcinogenesis Online) using the 869 probes on which DNA methylation levels \((\Delta \beta_{T-N})\) differed significantly between Clusters A and B [FDR \((q = 0.01)\)] identified the top four probes that were able to discriminate Cluster B from Cluster A in Table IIIB. Two probes were shared by Tables IIIA and IIIB. Thus, CpG sites on the 18 probes can be considered as hallmarks of CIMP-positive clear cell RCCs, i.e., clear cell RCCs belonging to Cluster B.

Discussion

Here, we have reported the results of methylome analysis of 245 renal tissue samples at single-CpG resolution. To our knowledge, no study involving Infinium analysis of such a large number of renal tissue samples has been reported to date. We have been focusing on DNA methylation alterations at the precancerous stage: our previous studies using methylation-specific PCR, combined bisulfite restriction enzyme analysis and bacterial artificial chromosome arrays suggested that N samples are already at the precancerous stage associated with DNA methylation alterations (17–20). First, we identified the probes on which DNA methylation status in N samples were significantly altered relative to those in C samples. The single-CpG-resolution analysis revealed that the DNA methylation status of 4830 CpG sites was actually altered at the precancerous stage in comparison to normal renal cortex tissue samples. In addition, it was revealed that alterations at the precancerous stages tended to involve DNA hypermethylation (Table I(a)). Among the 801 probes we selected, DNA methylation alterations occurred at the precancerous stage and were inherited by, and strengthened in, clear cell RCCs themselves, indicating that DNA methylation alterations on the 801 probes may participate continuously in renal carcinogenesis from the precancerous stage until cancers have become established. The DNA methylation profiles of these 801 probes clustered clear cell RCCs into clinicopathologically valid subclusters: clear cell RCCs belonging to Cluster B showed clinicopathological parameters reflecting tumor aggressiveness, and patients with Cluster B tumors showed a poorer outcome. Quantitative reverse transcription–PCR analysis indicated that DNA hypermethylation may result in significantly reduced expression of representative genes listed in Tables II A and II B and Supplementary Table S4, available at Carcinogenesis Online (Supplementary Table S5, available at Carcinogenesis Online). These findings suggest that DNA

![Fig. 2.](https://academic.oup.com/carcin/article-abstract/33/8/1487/2463738)
The probes satisfied the following criteria: (i) the average \( \beta \) value for all samples of non-cancerous renal cortex tissue (N) was <0.2, (ii) >0.4Δ\( \beta \) was observed in six or more clear cell RCCs (\( \geq 42.9\% \)) in Cluster B, whereas >0.4Δ\( \beta \) in two or fewer clear cell RCCs (\( \leq 2.2\% \)) in Cluster A, and (iii) the incidence of >0.4Δ\( \beta \) was markedly higher in Cluster B than in Cluster A \( (P < 1.98 \times 10^{-6}, \text{Fisher’s exact test}) \).

The present single-CpG-resolution analysis identified such hallmark CpG sites for the first time. Using the 18 CpG sites in Tables IIIA and IIIB, CIMP-positive RCCs or RCCs equivalent to those in the present study were differed significantly between Clusters A and B (Wilcoxon rank sum test).

Unsupervised hierarchical clustering based on our previous study using bacterial artificial chromosome arrays also clustered clear cell RCCs into clinicopathologically valid subclusters: 14% of examined RCCs belonged to a subcluster showing clinicopathological parameters reflecting tumor aggressiveness and poorer patient outcome (19). DNA methylation profiles in N samples based on BAMCA data were also inherited by the corresponding clear cell RCC developing in the same patient. In this study, 14% of the clear cell RCCs subjected to Infinium analysis belonged to Cluster B. BAMCA is suitable for detecting DNA methylation alterations occurring in a coordinated manner on individual large regions of chromosomes (34–37), whereas 27 000 Infinium array is suitable for detecting DNA methylation alterations on promoter regions of specific genes. Different methodologies identified similar clinicopathologically valid subclusters of RCCs, indicating that such clustering based on DNA methylation profiles is not accidental but reproducible, and may reflect the distinct epigenetic pathway of renal carcinogenesis.

In contrast to Cluster A, which appeared to be characterized by accumulation of DNA hypomethylation (Figure 2A), Cluster B was clearly characterized by accumulations of DNA hypermethylation on CpG islands. Since Cluster B was significantly associated with both frequent DNA hypermethylation on CpG islands and distinct clinicopathological phenotypes of clear cell RCCs, RCCs belonging to Cluster B can be recognized as CIMP-positive clear cell RCCs on the basis of the definition of well-studied CIMP-positive cancers (30,31) such as colorectal cancer (32) and stomach cancer (33), although Morris et al. (28) previously considered that the relevance of the CIMP-positive phenotype to RCCs had not yet been clearly defined. Although McDonald et al. (29) suggested that a subset of RCCs might display CIMP based on findings indicating that the distribution of the number of methylated CpGs in individual tumors differed from the expected Poisson distribution, they did not identify distinct CpG sites that could become hallmark CIMP in the kidney. It has been suggested that, in order to identify CIMP-positive cancers in specific organs, marker CpG sites or genes that are specific to each organ or histological type of tumor should be used (38), rather than classical CIMP marker genes (30,31) that were originally identified in colorectal cancers. The present single-CpG-resolution analysis identified such hallmark CpG sites for the first time. Using the 18 CpG sites in Tables IIIA and IIIB, CIMP-positive RCCs or RCCs equivalent to those in the present study were reproducibly identified. These 18 CpG sites may be useful for further clarifying the molecular basis of the epigenetic pathway of renal carcinogenesis.
DNA methylation alterations are known to result in chromosomal instability through chromatin configuration changes (39). In fact, germline mutations of the de novo DNA methyltransferase DNMT3B gene have been reported in patients with immunodeficiency, centronuclear instability and facial anomalies (ICF) syndrome, a rare recessive autosomal disorder characterized by DNA hypomethylation of pericentromeric satellite regions (40). In HCCs and urothelial carcinomas, DNA hypomethylation of these regions is correlated with copy number alterations on chromosomes 1 (41) and 9 (42), respectively, where satellite regions are plentiful. Correlations between the clustering based on Infinium assay and copy number alterations should be further examined.

Taken together, the data suggest that in CIMP-positive clear cell RCCs belonging to Cluster B, DNA hypermethylation of distinct CpG islands participates even in the very early and precancerous stages. Such DNA methylation alterations occurring in the precancerous stages may induce more aggressive tumor phenotypes and poorer patient outcome in Cluster B. On the other hand, in the other pathway of renal carcinogenesis leading to clear cell RCCs in Cluster A, DNA hypomethylation may be a later event (Table I) than DNA hypermethylation on CpG islands. We are now performing exome, transcriptome and proteome analyses of RCCs belonging to both clusters. Such multiplatform omics analyses may identify the upstream genetic events inducing DNA methylation profiles and key signal pathways that characterize Clusters A and B.

Supplementary material
Supplementary Figures S1–S5 and Tables S1–S5 can be found at http://carcin.oxfordjournals.org/.

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