ORIGINAL ARTICLE

Genome-wide DNA methylation profile of early-onset endometrial cancer: its correlation with genetic aberrations and comparison with late-onset endometrial cancer

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Abstract

The present study was performed to clarify the significance of DNA methylation alterations during endometrial carcinogenesis. Genome-wide DNA methylation analysis and targeted sequencing of tumor-related genes were performed using the Infinium MethylationEPIC BeadChip and the Ion AmpliSeq Cancer Hotspot Panel v2, respectively, for 31 samples of normal control endometrial tissue from patients without endometrial cancer and 81 samples of endometrial cancer tissue. Principal component analysis revealed that tumor samples had a DNA methylation profile distinct from that of control samples. Gene Ontology enrichment analysis revealed significant differences of DNA methylation at 1034 CpG sites between early-onset endometrioid endometrial cancer (EE) tissue (patients aged ≤40 years) and late-onset endometrioid endometrial cancer (LE) tissue, which were accumulated among ‘transcriptional factors’. Mutations of the CTNNB1 gene or DNA methylation alterations of genes participating in Wnt signaling were frequent in EEs, whereas genetic and epigenetic alterations of fibroblast growth factor signaling genes were observed in LEs. Unsupervised hierarchical clustering grouped EE samples in Cluster EA (n = 22) and samples in Cluster EB (n = 12). Clinicopathologically less aggressive tumors tended to be accumulated in Cluster EB, and DNA methylation levels of 18 genes including HOXA9, HOXD10 and SOX11 were associated with differences in such aggressiveness between the two clusters. We identified 11 marker CpG sites that discriminated EB samples from EA samples with 100% sensitivity and specificity. These data indicate that genetically and epigenetically different pathways may participate in the development of EEs and LEs, and that DNA methylation profiling may help predict tumors that are less aggressive and amenable to fertility preservation treatment.

Introduction

Endometrial cancer is one of the most common gynecological malignancies in the developed countries, with ~320 000 new cases diagnosed annually worldwide (1). Although it is typically a disease of postmenopausal women, ~2–14% of cases occur in women aged 40 years or less (2–4), and in fact recently endometrial cancer in younger patients has shown a marked increase (4). The standard treatment for endometrial cancer is hysterectomy and bilateral salpingo-oophorectomy with or without lymph node dissection, but for younger patients fertility-sparing treatment that avoids surgical menopause...
Patients and tissue samples
The 81 samples of cancerous tissue were obtained from 81 patients with primary endometrial cancer who underwent hysterectomy at Keio University Hospital between October 2005 and August 2016. Among the patients, 35 were aged 40 years or less (early-onset endometrial cancer) and 46 were older than 40 years (late-onset endometrial cancer). None of these patients had received any preoperative treatment. Supplementary Table 1A, available at Carcinogenesis Online, gives details of the ages and clinicopathological backgrounds of these patients. Histological diagnosis and grading were based on the 2003 World Health Organization classification (24) and the Tumor-Node-Metastasis classification (25). The clinical stage of the disease was based on the 2008 revised International Federation of Gynecology and Obstetrics classification (26). Recurrence was diagnosed by clinicians on the basis of physical examination and imaging modalities such as computed tomography and positron emission tomography.

For comparison, 31 samples of normal control endometrial tissue were obtained from materials that had been surgically resected from patients without endometrial cancer (Supplementary Table 1B, available at Carcinogenesis Online): 17, 6, 6 and 2 patients with uterine leiomyoma, ovarian cancer, cervical cancer and lobular endocervical glandular hyperplasia, respectively.

Tissue specimens were frozen immediately after surgery and preserved in the Keio Women’s Health Biobank in accordance with the Japanese Society of Pathology Guidelines for the handling of pathological tissue samples for genomic research (27). This study was approved by the ethics committee of Keio University Hospital and was performed in accordance with the Declaration of Helsinki. All patients included in this study provided written informed consent for use of their materials.

Infinium analysis
High-molecular-weight DNA was extracted from fresh frozen tissue samples using phenol–chloroform, followed by dialysis. DNA methylation status at 866 895 CpG loci was examined at single-CpG resolution using the Infinium MethylationEPIC BeadChip (Illumina, San Diego, CA) (28). Thereafter, 500 ng aliquots of DNA were subjected to bisulfite conversion using an EZ DNA Methylation-Gold Kit (Zymo Research, Irvine, CA). After hybridization, the specifically hybridized DNA was fluorescence-labeled by a single-base extension reaction and detected using an iScan Reader (Illumina) in accordance with the manufacturer’s protocol. The data were then assembled using GenomStudio Methylation software (Illumina). At each CpG site, the ratio of the fluorescence signal was measured using a methylated probe relative to the sum of the methylated and unmethylated probes, i.e. the so-called β-value, which ranges from 0.00 to 1.00 reflecting the methylation level of an individual CpG site.

In this assay, the call proportions (P < 0.01 for detection of signals above the background) for 801 probes on autosomes in all of the 112 tissue samples were >90%. As such low proportions may have been attributable to polymorphism at the probe CpG sites, these probes were excluded from the present assay. In addition, to avoid any gender-specific methylation bias, all 19 681 probes on chromosomes X and Y were excluded, leaving a final total of 846 413 autosomal CpG sites. Correlations between levels of DNA methylation and messenger RNA expression were examined using the dataset for endometrial cancer deposited in the TCGA database (https://tcga-data.nci.nih.gov/docs/publications/ucsc_2013/) (23).

Targeted sequencing analysis of 50 tumor-related genes
Targeted sequencing of 50 tumor-related genes was performed in 81 endometrial cancer tissue samples. Library preparation was performed using an Ion AmpliSeq Library Kit 2.0–96LV and Ion AmpliSeq Cancer Hotspot Panel v2 in accordance with the manufacturer’s protocol (Thermo Fisher Scientific, Waltham, MA). The panel is designed to amplify 207 amplicons covering 2849 mutations of 50 oncogenes and tumor suppressor genes deposited in the Catalogue of Somatic Mutations in Cancer (COSMIC) database (https://cancer.sanger.ac.uk) (Supplementary Table 2, available at Carcinogenesis Online). After library preparation, each amplicon library was quantified using an Ion Library Quantitation Kit on the 7500 Fast Real-Time PCR System (Thermo Fisher Scientific) using the relative standard curve method. Next, emulsion PCR and Ion Sphere™ particle enrichment were carried out using the Ion OneTouch™ 2 system and Ion OneTouch™ enrichment system (Thermo Fisher Scientific). Purified Ion Sphere™ particles were loaded on an Ion 316 or 318 Chip and sequenced using the Ion PGM™ sequencer (Thermo Fisher Scientific). Data were analyzed using Torrent Suite Software v5.2.2 and Ion Reporter.
Somatic mutations and copy number alterations in endometrial cancer

Five hundred and thirty-five somatic mutations were detected in 81 endometrial cancer tissue samples (6.6 ± 2.9 mutations per sample): 230 synonymous and 305 non-synonymous. The non-synonymous mutations consisted of 242 missense mutations (79%), 44 nonsense mutations (14%), 14 frameshift deletions (5%), 4 frameshift insertions (1%) and 1 frameshift block substitution (0.3%). Genes showing the highest incidences of somatic mutations included PTEN in 47 patients (58%), PIK3CA in 44 patients (54%), CTNBB1 in 25 patients (31%) and TP53 in 15 patients (19%). When compared with data for endometrial cancer deposited in the TCGA database (https://tcga-data.nci.nih.gov/docs/publications/ucec_2013/) (23), the incidences of somatic mutations of the MLH1 (P = 1.08 × 10−3), SMARCB1 (P = 3.01 × 10−3), AKT1 (P = 0.017) and STK11 (P = 3.47 × 10−3) genes were higher, and the incidences of those of the FBEW7 (P = 0.037), ATM (P = 0.049), EZH2 (P = 0.043), GNAS (P = 0.043) and JAK2 (P = 0.044) genes were lower in our cohort, whereas significant differences were not found in the remaining 41 genes (Figure 2A). The incidences of copy number alterations of all the 50 genes examined were lower than 10%, and this low incidence was again in accordance with the data deposited in the TCGA database (https://tcga-data.nci.nih.gov/docs/publications/ucec_2013/) (23) (Figure 2B).

Differences in DNA methylation profile between early- and late-onset endometrioid endometrial cancers

In order to avoid the bias caused by differences in histological subtypes, we then focused on the major subtype, i.e. endometrioid endometrial cancer. We again identified 63 033 probes that showed significant differences in DNA methylation levels between the 31 normal control endometrial tissue samples and 74 endometrioid endometrial cancer tissue samples (Welch’s t-test, adjusted Bonferroni correction [α = 1.18 × 10−4] and ΔR̂_{endometrial endometrial cancer tissue − normal control endometrial tissue }value of >0.25 or <−0.25).

Next, we focused on differences between early-onset endometrioid endometrial cancer (EE, patients aged <40 years) and late-onset endometrioid endometrial cancer (LE, patients aged >40 years). Clinicopathological parameters of early- and late-onset cases are summarized in Supplementary Table 3, available at Carcinogenesis Online. Among the 63 033 probes, 1034 showed significant differences in DNA methylation levels between the 34 samples of EE tissue and 40 samples of LE tissue (Welch’s t-test P < 0.01; ΔR̂_{EE − LE }Value >0.25 or <−0.25; Supplementary Table 4, available at Carcinogenesis Online). As several patients (n = 11) aged <50 years were included in the LE group, a relatively small number of CpG sites showed significant differences in DNA methylation levels between EE and LE samples. All 1034 probes again showed significant differences in DNA methylation levels between LE samples and normal control endometrial tissue samples (Welch’s t-test P < 0.01; ΔR̂_{LE − normal control endometrial tissue }value >0.25 or <−0.25), whereas only 102 of 1034 probes showed significant differences in DNA methylation levels between EE samples and normal control endometrial tissue samples (Welch’s t-test P < 0.01; ΔR̂_{EE − normal control endometrial tissue }value >0.25 or <−0.25). Among these 102 probes, the DNA methylation levels of 101 probes were higher in EE samples than in normal control endometrial tissue samples and even higher in LE samples.

Three hundred and seventy-one genes, for which the 1034 probes showing significant differences of DNA methylation

GO enrichment analysis and pathway analysis

In order to reveal the function of proteins encoded by genes showing DNA methylation alterations and gene mutations, and to reveal the biological processes in which such proteins participate, Gene Ontology (GO) enrichment analysis and pathway analysis were conducted using the GeneGO Metacore™ software package, version 6.7 (Thomson Reuters, New York, NY).

Statistics

Infinium probes showing significant differences in DNA methylation levels between sample groups were defined by Welch’s t-test and adjusted by Bonferroni correction. The DNA methylation profiles of the 112 tissue samples were analyzed using principal component analysis. Differences in the incidences of genetic aberrations between sample groups were examined using Fisher’s exact test. Unsupervised hierarchical clustering based on DNA methylation levels (Euclidean distance, Ward method and Canberra distance, complete linkage method) was performed. The associations between clinicopathological variables and DNA methylation alterations were evaluated using Fisher’s exact test, Mann–Whitney U test or Welch’s t-test. The receiver operating characteristic (ROC) curve was generated, and the Youden index (30) for each probe was used as a cutoff value for examining distinctions between the sample groups. Programming language R and IBM SPSS statistics 20.0 were used to analyze the data.

Results

DNA methylation alterations during endometrial carcinogenesis

First, to obtain a general overview of endometrial cancers, we identified 58 958 probes that were aberrantly methylated in 81 endometrial cancer tissue samples in comparison with the 31 normal control endometrial tissue samples (Welch’s t-test, adjusted Bonferroni correction [α = 1.18 × 10−4] and ΔR̂_{endometrial cancer tissue − normal control endometrial tissue }value of >0.25 or <−0.25), indicating that DNA methylation alterations had occurred during endometrial carcinogenesis. Principal component analysis using the DNA methylation levels of these 58 958 probes revealed that endometrial cancer tissue samples had a DNA methylation profile that differed distinctly from that of normal control endometrial tissue samples (Figure 1A). The leading 10 genes, i.e. TRIM15, HIST2H3P52, NBPF19, L1TD1, HIST2H2BA, NKPAL, DLX2-AS1, GRMI, ADAM12 and CFA4P6L, showing significant differences in DNA methylation levels between endometrial cancer tissue and normal control endometrial tissue are labeled in the Manhattan plot (Figure 1B). To the best of our knowledge, no previous English-language articles have yet reported correlations between these 10 genes and endometrial cancers; this study may have revealed novel genes potentially associated with endometrial carcinogenesis, although the expression levels and functions of these genes will need to be further examined in connection with endometrial cancers.
levels between EE samples and LE samples were designed, were evaluated for protein function by enrichment analysis using the MetaCore software and compared with the protein function distribution of genes within the GeneGo database. These genes were clearly overrepresented by ‘transcriptional factors’ ($P = 1.67 \times 10^{-32}$), being 6.004 times more abundant than expected. Indeed 66 of 371 genes were categorized as transcription factors by MetaCore software (shown by asterisks in Supplementary Table 4, available at Carcinogenesis Online). The top 20 statistically significant GO molecular functions in which the 371 genes participated are listed in Table 1. Fifteen of the top 20 functions were related to DNA binding or transcriptional regulation, and are shown by asterisks, and transcription factors included in the top 20 GO molecular functions are underlined in Table 1. All 189 statistically significant GO molecular functions ($P < 0.05$) are listed in Supplementary Table 5, available at Carcinogenesis Online.

**Differences in genetic aberrations between EE and LE**

Two hundred and one somatic mutations (90 synonymous and 111 non-synonymous) were detected in 34 samples of EE ($5.9 \pm 2.2$ mutations per sample), whereas 300 mutations (119 synonymous and 181 non-synonymous) were detected in 40 samples of LE ($7.5 \pm 3.3$ mutations per sample). The incidence of somatic mutation of the $CTNNB1$ gene was significantly higher in EE samples than in LE samples ($P = 4.26 \times 10^{-4}$), and the incidence of that of the $FGFR2$ gene was significantly higher in LE samples than in EE samples ($P = 8.77 \times 10^{-3}$), whereas significant differences were not found in the remaining 48 genes (Figure 3A). Average SIFT scores for the $CTNNB1$ and $FGFR2$ genes were both 0 in EE samples and LE samples, respectively, indicating that these somatic mutations impair protein function (a SIFT score of <0.05 means ‘damaging’ (29)).
Correlation between DNA methylation alterations and genetic aberrations in EE and LE

In order to clarify the correlation between DNA methylation alterations and genetic aberrations in EE samples, we identified 2908 probes that showed significant differences in DNA methylation levels between 19 EE samples with somatic mutation of the CTNNB1 gene (tissue samples of early-onset endometrioid endometrial cancer with somatic mutation of the CTNNB1 gene, CTNNB1-M) and 15 EE samples without it (tissue samples of early-onset endometrioid endometrial cancer...
Table 1. Top 20 statistically significant GO molecular functions revealed by MetaCore software analysis using the 371 genes, for which the 1034 probes showing differences in DNA methylation levels between samples of early-onset endometrioid endometrial cancer tissue (patients aged <40 years) and late-onset endometrioid endometrial cancer tissue were designed (listed in Supplementary Table 4, available at Carcinogenesis Online).

| Molecular functions | P value | Included genes showing differences in DNA methylation levels |
|---------------------|---------|------------------------------------------------------------|
| Sequence-specific DNA binding* | 2.05 × 10^{-19} | ASCL1, ASCL2, ASCL4, BARHL2, DMARTA2, DRGX, EN1, EVX1, EVX2, FEZF2, FLI1, FOXD3, FOXE1, FOXD3, DMARTA2, DRGX, EVX1, FOXD3, GATA4, GCM2, HAND2, HIC1, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, NKKX6-2, OTX1, OTX2, OTX3, PAX5, PAX6, PAX7, PHOX2A, PITX2, POU3F3, POU4F2, PRDM14, PROX1, RAX, SALL3, SATB2, SIX6, SOX1, SOX11, SOX14, SOX2, TBX5, TBX15, TBX18, TXL2, TP73, VSX1, ZFHX4, ZIC5, ZNF516 |
| Transcription factor activity, sequence-specific DNA binding* | 2.39 × 10^{-19} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, BHHE32, BN1, DMARTA2, DRGX, DXB1, EN1, EVX1, EVX2, FEZF2, FLI1, FOXD3, FOXE1, FOXI2, SKOR1, GATA4, GCM2, GFI1, HAND2, HIC1, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, NKKX6-2, OTX1, OTX2, OTX3, PAX5, PAX6, PSX7, PHOX2A, PITX2, POU3F3, POU4F2, PRDM14, PROX1, RAX, SALL3, SATB2, SIM1, SIX6, SOX1, SOX11, SOX14, SOX2, TBX7, TXB15, TXB18, TXL2, TP73, VSX1, ZFHX4, ZIC5, ZNF516 |
| Nucleic acid binding transcription factor activity* | 2.48 × 10^{-19} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, BHHE32, BN1, DMARTA2, DRGX, DXB1, EN1, EVX1, EVX2, FEZF2, FLI1, FOXD3, FOXE1, FOXI2, SKOR1, GATA4, GCM2, GFI1, HAND2, HIC1, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, NKKX6-2, OTX1, OTX2, OTX3, PAX5, PAX6, PSX7, PHOX2A, PITX2, POU3F3, POU4F2, PRDM14, PROX1, RAX, SALL3, SATB2, SIM1, SIX6, SOX1, SOX11, SOX14, SOX2, TBX7, TXB15, TXB18, TXL2, TP73, VSX1, ZFHX4, ZIC5, ZNF516, ZSCN1, ZSCN23 |
| RNA polymerase II transcription factor activity, sequence-specific DNA binding* | 3.14 × 10^{-16} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, BHHE32, BN1, DMARTA2, DRGX, DXB1, EN1, EVX1, EVX2, FEZF2, FLI1, FOXD3, FOXE1, FOXI2, SKOR1, GATA4, GCM2, GFI1, HAND2, HIC1, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, NKKX6-2, OTX1, OTX2, OTX3, PAX5, PAX6, PSX7, PHOX2A, PITX2, POU3F3, POU4F2, PRDM14, PROX1, RAX, SALL3, SATB2, SIM1, SIX6, SOX1, SOX11, SOX14, SOX2, TBX7, TXB15, TXB18, TXL2, TP73, VSX1, ZFHX4, ZIC5, ZNF516, ZSCN1, ZSCN23 |
| DNA binding* | 2.05 × 10^{-10} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, BHHE32, BN1, DMARTA2, DRGX, DXB1, EN1, EVX1, EVX2, FEZF2, FLI1, FOXD3, FOXE1, FOXI2, SKOR1, GATA4, GCM2, GFI1, HAND2, HIC1, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, NKKX6-2, OTX1, OTX2, OTX3, PAX5, PAX6, PSX7, PHOX2A, PITX2, POU3F3, POU4F2, PRDM14, PROX1, RAX, SALL3, SATB2, SIM1, SIX6, SOX1, SOX11, SOX14, SOX2, TBX7, TXB15, TXB18, TXL2, TP73, VSX1, ZFHX4, ZIC5, ZNF516, ZSCN1, ZSCN23 |
| Transcription regulatory region sequence-specific DNA binding* | 2.15 × 10^{-10} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, EN1, FEZF2, FLI1, FOXD3, GATA4, HAND2, IRF4, LHX1, LHX4, LHX5, MNX1, MSC, MKX1-1, MYOD1, NKKX6-2, OTSCU1, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Transcription factor activity, RNA polymerase II core promoter proximal region sequence-specific binding* | 2.81 × 10^{-10} | ASCL1, ASCL2, ASCL4, BARHL2, EN1, FLI1, GCM2, GFI1, HAND2, IRF4, MYOD1, NKKX6-2, ONECUT1, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Neurokinin receptor binding | 5.08 × 10^{-10} | TAC1 |
| Substance P receptor binding | 5.08 × 10^{-10} | TAC1 |
| Sequence-specific double-stranded DNA binding* | 7.65 × 10^{-10} | ASCL1, ASCL2, ASCL4, BARHL2, EN1, FEZF2, FLI1, FOXD3, GATA4, HAND2, IRF4, LHX1, LHX3, MCKX1-1, MYOD1, NKKX6-2, OTSCU1, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Transcription regulatory region DNA binding* | 9.47 × 10^{-10} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, EN1, FEZF2, FLI1, FOXD3, GATA4, GFI1, HAND2, IRF4, LHX1, MCKX1-1, MYOD1, NKKX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Regulatory region DNA binding* | 1.01 × 10^{-9} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, EN1, FEZF2, FLI1, FOXD3, GATA4, GFI1, HAND2, IRF4, LHX1, MCKX1-1, MYOD1, NKKX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Regulatory region nucleic acid binding* | 1.13 × 10^{-9} | ASCL1, ASCL2, ASCL4, BARHL2, BHHE22, EN1, FEZF2, FLI1, FOXD3, GATA4, GFI1, HAND2, IRF4, LHX1, MCKX1-1, MYOD1, NKKX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, POU4F2, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX12, TBX7, TXB15, TXB18, TXL2, TP73, ZISC, ZNF516 |
| Neuropeptide receptor activity | 3.30 × 10^{-9} | GALR1, QRFPR, GPR139, NPFFR2, NPY2R, NPY5R, PROK2, SSTR1, SSTR4, SORCS3 |
Table 1. Continued

| Molecular functions                        | P value     | Included genes showing differences in DNA methylation levels |
|-------------------------------------------|-------------|-------------------------------------------------------------|
| Gated channel activity                    | 6.48 × 10⁻⁹ | CACNG7, CNGA3, GABRA1, GABRA2, GABRB1, GLRA3, GRIAA, KCNAB1, KCNHB, KCNJ9, KCNQ5, KCNA6, KCNJ3, KCNA1, KCNA3, KCNA4, KCNV1, GRIN3A, CACNA1A, RYR2, KCNT2, TTYH1, CHRNA4 |
| Double-stranded DNA binding               | 7.49 × 10⁻⁹ | ASCL2, ASCL4, BARHL2, EN1, FEZF2, FLI1, FOXD3, FOXE1, GATA4, HAND2, IRF4, LHX1, MKX1-1, MYOD1, NX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, PO4U2F, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, SOX2, TRX7, TRX15, TRX18, TBX5, TP73, ZIC5, ZNF516 |
| RNA polymerase II regulated region DNA binding | 9.49 × 10⁻⁹ | ASCL2, ASCL4, BARHL2, EN1, FEZF2, FLI1, FOXD3, GATA4, HAND2, IRF4, LHX1, MKX1-1, MYOD1, NX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, PO4U2F, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, TRX7, TRX15, TRX18, TBX5, TP73, ZIC5 |
| Voltage-gated potassium channel activity   | 1.06 × 10⁻⁸ | CNGA3, KCNAB1, KCNB8, KCNK5, KCNQ5, KCNA6, KCNJ3, KCNA1, KCNA3, KCNA4, KCNV1, KCNT2 |
| RNA polymerase II regulated region DNA binding | 1.19 × 10⁻⁸ | ASCL2, ASCL4, BARHL2, EN1, FEZF2, FLI1, FOXD3, GATA4, HAND2, IRF4, LHX1, MKX1-1, MYOD1, NX6-2, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, PO4U2F, PRDM14, PROX1, RAX, SATB2, SOX1, SOX11, TRX7, TRX15, TRX18, TBX5, TP73, ZIC5 |
| Transcriptional activator activity         | 1.24 × 10⁻⁸ | BARHL2, FEZF2, FLI1, GATA4, GCM2, HAND2, IRF4, LHX4, MYOD1, ONECUT1, ONECUT2, OTX1, PAX5, PAX6, PHOX2A, PITX2, PO4U2F, RAX, SATB2, SIX6, SOX1, SOX11, SOX2, TRX5, TLX2, TP73 |

*GO molecular functions involved in DNA binding or transcriptional regulation. Genes for which the protein class is a transcription factor are indicated by underlining.

without somatic mutation of the CTNNB1 gene, CTNNB1-W) (Welch’s t-test P < 0.01; ΔP_{Carest,MCTNNB1-W} value >0.15 or < −0.15; Supplementary Table 6, available at Carcinogenesis Online). MetaCore pathway analysis revealed that 1419 genes for which these 2908 probes were designed were significantly accumulated in the Wnt signaling pathway (P = 3.45 × 10⁻⁴; Figure 3B). We identified 1133 probes that showed significant differences in DNA methylation levels between 10 LE samples with somatic mutation of the FGFR2 gene (tissue samples of late-onset endometrioid endometrial cancer with somatic mutation of the FGFR2 gene, FGFR2-M) and 30 LE samples without it (tissue samples of late-onset endometrioid endometrial cancer without somatic mutation of the FGFR2 gene, FGFR2-W) (Welch’s t-test P < 0.01; ΔP_{Carest,MCTNNB1-W} value >0.15 or < −0.15; Supplementary Table 7, available at Carcinogenesis Online). Three of the 567 genes for which these 1133 probes were designed were included in the fibroblast growth factor signaling pathway according to the MetaCore software (Supplementary Figure 1, available at Carcinogenesis Online).

Epigenetic clustering of endometrioid endometrial cancer based on DNA methylation profile

Unsupervised hierarchical clustering using 40 589 probes showing significant differences in DNA methylation levels between the 31 normal control endometrial tissue samples and 34 EE samples (Welch’s t-test, adjusted Bonferroni correction [α = 1.18 × 10⁻⁴]; ΔP_{Carest} - normal control endometrial tissue, value >0.25 or < −0.25) subclustered 34 patients with EE into samples in Cluster EA (n = 22) and samples in Cluster EB (n = 12) (Figure 4B). The clinicopathological parameters of the patients in these clusters are summarized in Supplementary Table 8B, available at Carcinogenesis Online. Patients in Cluster EA tended to show more frequent lymphovascular invasion and to be more frequently diagnosed as histological grade 3 than those in Cluster EB.

The cancer-free survival rate of patients belonging to Cluster EB tended to be lower than that of patients in Cluster EB, although the difference did not reach a statistically significant level (P = 0.24 and 0.53, respectively, log-rank test), probably due to the small number of deaths or recurrent cases (Supplementary Figure 2, available at Carcinogenesis Online). Moreover, all recurrence-positive (n = 4) and disease-specific death-positive (n = 1) cases were included in Cluster A (Supplementary Table 8A, available at Carcinogenesis Online).

Unsupervised hierarchical clustering using 40 589 probes showing significant differences in DNA methylation levels between the 31 normal control endometrial tissue samples and 34 EE samples (Welch’s t-test, adjusted Bonferroni correction [α = 1.18 × 10⁻⁴]; ΔP_{Carest} - normal control endometrial tissue, value >0.25 or < −0.25) subclustered 34 patients with EE into samples in Cluster EA (n = 22) and samples in Cluster EB (n = 12) (Figure 4B). The clinicopathological parameters of the patients in these clusters are summarized in Supplementary Table 8B, available at Carcinogenesis Online. Patients in Cluster EA tended to show more frequent lymphovascular invasion and to be more frequently diagnosed as histological grade 3 than those in Cluster EB.

The cancer-free survival rate of patients belonging to Cluster EB tended to be lower than that of patients in Cluster EB, although the difference did not reach a statistically significant level (P = 0.37, log-rank test), probably due to the small number of recurrent cases (Supplementary Figure 2, available at Carcinogenesis Online). Only one recurrence case was included in Cluster EA (Supplementary Table 8B, available at Carcinogenesis Online). As the overall survival rate was 100% in both Clusters EA and EB, the log-rank test was not performed. Moreover, all patients belonging to Cluster EA were included in Cluster A and all patients belonging to Cluster EB were included in Cluster B without exception (Figure 4C).

As seen in Figure 4B, Probe Clusters I and III showed obvious differences in DNA methylation levels between Clusters EA and EB. Furthermore, the ratio of probes located in CpG islands, CpG island shores and CpG island shelves that were important for transcription regulation to all probes belonging to Probe Cluster I was significantly higher than that of probes in Probe Cluster A than in Cluster B, such differences did not reach statistically significant levels (P = 0.24 and 0.53, respectively, log-rank test), probably due to the small number of deaths or recurrent cases (Supplementary Figure 2, available at Carcinogenesis Online). Moreover, all recurrence-positive (n = 4) and disease-specific death-positive (n = 1) cases were included in Cluster A (Supplementary Table 8A, available at Carcinogenesis Online).
We then focused on Probe Cluster I, in which 927 of the 6415 probes were designed from the transcription start site to 1500 bp upstream of the transcription start site of the 470 genes. One hundred and one (Supplementary Table 9A, available at Carcinogenesis Online) of the 470 genes showed a significant ($P < 0.05$) inverse correlation ($r < -0.2$) between DNA methylation levels and the messenger RNA expression levels in the TCGA database (https://tcga-data.nci.nih.gov/docs/publications/ucec_2013/) (23). The 101 genes were also evaluated for protein function by enrichment analysis using the MetaCore software and compared with the protein function distribution of genes within the GeneGo database (Supplementary Table 9B, available...
Figure 4. Epigenetic clustering of endometrioid endometrial cancer. (A) Unsupervised hierarchical clustering of endometrioid endometrial cancer tissue samples using DNA methylation levels on 63,033 probes showing significant differences in DNA methylation levels between 31 samples of normal control endometrial tissue and 74 samples of endometrioid endometrial cancer tissue (Welch’s t-test, adjusted Bonferroni correction [\(\alpha = 1.18 \times 10^{-8}\) and \(\Delta\beta_{\text{endometrioid endometrial cancer tissue - normal control endometrial tissue}}\) value of >0.25 or <−0.25). On the basis of DNA methylation status, 74 patients were subclustered into Cluster A (\(n = 58\)) and Cluster B (\(n = 16\)). Correlations between this epigenetic clustering and clinicopathological parameters are summarized in Supplementary Table 8A, available at Carcinogenesis Online. (B) Unsupervised hierarchical clustering of EE samples using DNA methylation levels on the 40,589 probes showing significant differences in DNA methylation between 31 normal control endometrial tissue samples and 34 EE samples (Welch’s t-test, adjusted Bonferroni correction [\(\alpha = 1.18 \times 10^{-8}\) and \(\Delta\beta_{\text{EE - normal control endometrial tissue}}\) value of >0.25 or <−0.25). On the basis of DNA methylation status, the 34 EE patients were subclustered into Cluster EA (\(n = 22\)) and Cluster EB (\(n = 12\)). Correlations between this epigenetic clustering and clinicopathological parameters of the patients are summarized in Supplementary Table 8B, available at Carcinogenesis Online. Probe Clusters I–V are shown on the left side of the heatmap. (C) Venn diagram showing the relationship between Clusters A and B of endometrioid endometrial cancer tissue samples and Clusters EA and EB of EE samples. All samples belonging to Cluster EA are included in Cluster A, whereas all samples belonging to Cluster EB are included in Cluster B without exception.
at Carcinogenesis Online). Fifty-nine statistically significant GO molecular functions in which the 101 genes participated are listed in Supplementary Table 9C, available at Carcinogenesis Online. DNA methylation levels of 18 of the 101 genes, including HOXA9, HOXD10 and SOX11, were significantly correlated with a higher incidence of lymphovascular invasion of EE (Supplementary Table 10, available at Carcinogenesis Online).

Identification of marker CpG sites for distinguishing Cluster EB from Cluster EA

In order to identify diagnostic markers capable of distinguishing the less aggressive Cluster EB from Cluster EA, ROC analysis was performed using the top 100 probes showing the largest differences in DNA methylation levels between EA and EB samples, and the corresponding area under curve (AUC) values were calculated. Among the 100 probes, 11 showed AUC values of 1 (Supplementary Table 11, available at Carcinogenesis Online). The Youden index was used as a cutoff value for each of the 11 probes. As shown by the scattergrams in Figure 5, the use of such cutoff values was able to discriminate EB samples from EA samples with 100% sensitivity and specificity. The DNA methylation levels of these 11 candidate marker CpG sites were successfully verified using another quantification method, MassARRAY: significant correlations between DNA methylation levels demonstrated by Infinium assay and those by MassARRAY were confirmed (Supplementary Figure 3, available at Carcinogenesis Online).

Discussion

We focused on the differences between early- and late-onset cancers of the major histological subtype, i.e. endometrioid endometrial cancer. The incidence of somatic mutations of the CTNNB1 gene was significantly higher in EE samples than in LE samples, whereas somatic mutations of the FGFR2 gene were more frequent in LE samples than in EE samples. Moreover, DNA methylation alterations and genetic aberrations may cooperatively activate the Wnt signaling pathway (31) during the development of EE. Although an inverse association between immunohistochemically detected nuclear accumulation of β-catenin and the age of patients with endometrial cancer has been reported previously (32), and mutations of the CTNNB1 gene have been shown to be accumulated in a subset of endometrioid endometrial cancers arising in young and obese patients (33), this study has comprehensively revealed epigenetic and genetic alterations of genes participating in Wnt signaling in EE for the first time. Similarly, DNA methylation alterations and gene aberrations may cooperatively participate in the fibroblast growth factor signaling pathway in late-onset endometrioid endometrial carcinoma. Although activating mutation of the FGFR2 gene has been considered a therapeutic target for endometrioid endometrial cancer (34,35), its correlation with late-onset carcinogenesis has never been reported previously. Thus, the pathways contributing to the development of early- and late-onset endometrial cancer appear to partially differ.

With regard to the genome-wide DNA methylation profile, all 1034 probes showing significant differences in DNA methylation levels between EE and LE samples again showed significant differences between LE and normal control endometrial tissue samples. On the other hand, only 102 probes of the 1034 probes showed significant differences in DNA methylation levels between EE and normal control endometrial tissue samples, indicating that DNA methylation alterations for the majority of the 1034 probes occurred specifically in LE samples. GO enrichment analysis indicated that 371 genes, for which the 1034 probes were designed, were clearly overrepresented by ‘transcriptional factors’ and were accumulated in signaling pathways participating in transcriptional regulation. The late-onset-specific DNA methylation profile may modify the clinicopathological features of LE samples as a result of the differences in the gene expression profiles. In fact, myometrial invasion, one of the most important prognostic factors of endometrioid endometrial cancer (2), was more frequent in LE than in EE samples.

Epigenetic clustering based on DNA methylation profiles showed that endometrioid endometrial cancers belonging to Cluster A were clinicopathologically more aggressive than those belonging to Cluster B. This tendency was also evident in patients belonging to Cluster EA, and all patients in Cluster EA were included in Cluster A without exception. The 101 hallmark genes for Cluster EA may drive the development and progression of different tumor subtypes. For example, the DNA methylation levels of 18 genes, including HOXA9, HOXD10 and SOX11, were significantly correlated with more frequent lymphovascular tumor invasion. DNA hypermethylation of the HOXA9 homeobox gene is reportedly associated with a higher grade of serous ovarian cancer (36), recurrence of bladder cancer (37) and nonsmall cell lung cancer (38) and mortality in non-infant patients with neuroblastoma (39). Decreased expression of the HOXD10 homeobox gene has been reported in prostate (40), breast (41), thyroid (42), colorectal (43), ovarian (44) and endometrial (45) cancer. Promoter methylation of the SOX11 gene has reportedly been associated with cell growth, invasion or poor prognosis of gastric cancer (46), hematopoietic malignancies (47) and nasopharyngeal carcinoma (48). It is feasible that DNA methylation alterations of these genes participate in determining the clinicopathological characteristics of Cluster EA. These data indicate that clinicopathological features are strongly defined on the basis of DNA methylation profiles, even in EE.

ROC analysis identified markers at CpG sites that were able to distinguish epigenetic Cluster EB from Cluster EA: 11 CpG sites showed AUC values of 1 for such discrimination, and Cluster EB of this cohort was diagnosed using the 11 marker CpG sites with both sensitivity and specificity of 100%. Although some of the 11 marker CpGs were located in CpG islands around the transcription start sites, an inverse correlation between levels of DNA methylation and messenger RNA expression was not confirmed in the data for endometrial cancer deposited in the TCGA database (https://tcga-data.nci.nih.gov/docs/publications/ucec_2013/) (23). Therefore, DNA methylation alterations of such genes may not result in alterations of expression, and such genes may not be potential therapeutic targets. Instead, as the present data have indicated that the clinicopathological features of endometrial cancers may be determined by their DNA methylation profiles, these 11 markers may be able to reproducibly identify less aggressive cancers, such as those belonging to Cluster EB. Although a validation study will, of course, be needed, DNA methylation diagnostics of biopsy specimens using appropriate marker CpG sites, such as these 11 CpG sites, may help to indicate the feasibility of fertility preservation therapy for patients with EE.

As all samples belonging to Clusters EA and EB were included in Clusters A and B, respectively, it is not surprising that the 11 CpG sites showed excellent AUCs even in ROC analysis for discrimination of the less aggressive Cluster A from the less aggressive Cluster B (Supplementary Table 12, available at Carcinogenesis Online): The 11 CpG sites may also become good markers for prognostication of all age groups of patients.
with endometrioid endometrial cancers. Moreover, the DNA methylation levels of the 11 CpG sites were not significantly correlated with clinicopathological parameters reflecting tumor aggressiveness, i.e. histological grade and stage, or patient outcome, in cancers of the lung, stomach, breast, colon and ovary deposited in the TCGA database (data not shown), suggesting that the prognostic potential of the 11 CpG sites may be specific to endometrioid endometrial cancers.

Figure 5. Scattergrams of DNA methylation levels for all 11 probes showing an area under the curve value of 1 in receiver operating characteristic curve analysis for discrimination of EB samples (n = 12) from EA samples (n = 22). P values by Welch’s t-test for each probe are shown in each panel. Cutoff values (CVs) are shown by a dotted line in each panel. Using each probe and its CV, EB samples were discriminated from EA samples with 100% sensitivity and specificity.
In summary, genome-wide DNA methylation analysis using EPIC array and targeted sequencing of tumor-related genes for 112 endometrial tissue samples has indicated that genetically and epigenetically different pathways may participate in the development of early- and late-onset endometrial cancers. As DNA methylation profiles may determine the clinicopathological features of endometrial cancers, such profiling may predict tumors that are less aggressive and amenable to fertility preservation treatment.

**Supplementary material**

Supplementary data are available at Carcinogenesis online.

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