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Host-cell DNA methylation patterns during high-risk HPV-induced carcinogenesis reveal a heterogeneous nature of cervical pre-cancer


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ABSTRACT
Cervical cancer development following a persistent infection with high-risk human papillomavirus (hrHPV) is driven by additional host-cell changes, such as altered DNA methylation. In previous studies, we have identified 12 methylated host genes associated with cervical cancer and precancer (CIN2/3). This study systematically analyzed the onset and DNA methylation pattern of these genes during hrHPV-induced carcinogenesis using a longitudinal in vitro model of hrHPV-transformed cell lines (n = 14) and hrHPV-positive cervical scrapings (n = 113) covering various stages of cervical carcinogenesis. DNA methylation analysis was performed by quantitative methylation-specific PCR (qMSP) and relative qMSP values were used to analyze the data. The majority of genes displayed a comparable DNA methylation pattern in both cell lines and clinical specimens. DNA methylation onset occurred at early or late immortal passage, and DNA methylation levels gradually increased towards tumorigenic cells. Subsequently, we defined a so-called cancer-like methylation-high pattern based on the DNA methylation levels observed in cervical scrapings from women with cervical cancer. This cancer-like methylation-high pattern was observed in 72% (38/53) of CIN3 and 55% (11/20) of CIN2, whereas it was virtually absent in hrHPV-positive controls (1/26). In conclusion, hrHPV-induced carcinogenesis is characterized by early onset of DNA methylation, typically occurring at the pre-tumorigenic stage and with highest DNA methylation levels at the cancer stage. Host-cell DNA methylation patterns in cervical scrapings from women with CIN2 and CIN3 are heterogeneous, with a subset displaying a cancer-like methylation-high pattern, suggestive for a higher cancer risk.

Introduction
Following a persistent infection with a high-risk (hr) type of human papillomavirus (HPV), additional genetic and epigenetic changes in the host cell genome are necessary for progression to cervical cancer [1]. Part of these host cell alterations are induced by expression of viral oncogenes E6 and E7 and include DNA methylation of tumor suppressor genes [2]. Methylation of cytosines at CpG-sites in promoter regions can lead to gene silencing. The DNA methyltransferases (DNMTs) responsible for CpG methylation can be activated by both hrHPV E6 and E7. E7 can directly bind to and activate DNMT1, whereas E6 can upregulate DNMT1 via p53 [3,4]. Conversely, silencing of E6 and E7 has been shown to reduce DNA methylation of tumor suppressor genes and to restore the transformed phenotype in cervical cancer cells [5].

Increased DNA methylation levels of several (candidate) tumor suppressor genes are associated with cervical cancer and a subset of its high-grade precursor lesions, i.e., cervical intraepithelial neoplasia grade 2 and 3 (CIN2 and CIN3) [1,6,7]. Using both targeted and genome-wide approaches,
we previously identified 12 genes, including ANKRD18CP, C13orf18, EPB41L3, and JAM3 [8,9]; SOX1 and ZSCAN1 [9,10]; GHSR, SST, and ZIC1 [11]; and FAM19A4, PHACTR3, and PRDM14 [12]. These genes were considered promising biomarkers to identify hrHPV-positive women and/or women with abnormal cytology at risk for cervical cancer [1,7–14].

So far, the majority of these genes has been evaluated individually or in small sets in separate studies. Herein, we performed a systematic analysis of 12 genes on the same cohort to compare their DNA methylation onset and patterns during cervical carcinogenesis. To this end, we analyzed a well-characterized longitudinal in vitro model of primary keratinocytes immortalized by HPV16 or HPV18 [15], complemented with a large series of clinically annotated hrHPV-positive cervical scrapings covering various stages of cervical carcinogenesis. The consecutive passages of hrHPV-immortalized cells, which are characterized by an initial acquisition of telomerase activity, followed by a more transformed phenotype characterized by anchorage independent growth, were shown to closely mimic the progressive stages of cervical precancerous disease with respect to both genetic and epigenetic changes [2,15–17].

Results

Early onset and gradual increase in DNA methylation during hrHPV-induced transformation in vitro

DNA methylation patterns of the 12 genes (ANKRD18CP, C13orf18, EPB41L3, FAM19A4, GHSR, JAM3, PHACTR3, PRDM14, SST, ZIC1, SOX1, and ZSCAN1) were analyzed in an in vitro model of hrHPV-induced transformation, using HFKs, consecutive passages of hrHPV-transformed cell lines reflecting the progressive stages of cervical precancerous disease [2,15], and cervical cancer cell lines. In general, the 12 genes showed no DNA methylation in HFKs, a progressive increase in DNA methylation levels from early to late passages of hrHPV-immortalized keratinocytes, and highest levels in the cervical cancer cell lines (Figure 1). The onset of DNA methylation varied between the 12 genes from early immortal passages (ANKRD18CP, FAM19A4, GHSR, JAM3, PRDM14, SST, and ZSCAN1) to late immortal passages (C13orf18, EPB41L3, PHACTR3, SOX1, and ZIC1) (Supplementary Figure 1).

A cancer-like methylation-high pattern in a subset of cervical scrapings from women with CIN2/3

Next, DNA methylation patterns of the 12 genes in hrHPV-positive cervical scrapes, covering different underlying histology (no disease, CIN2, CIN3, and cancer) were analyzed. Alike the in vitro model, a progressive and significant increase in DNA methylation levels towards cervical cancer was observed for all 12 genes (P < 0.05, Figure 2).

The DNA methylation signature of the 12 genes depicted by predicted probabilities is shown in Figure 3. All cancer cases (n = 14) had an average predicted probability of ≥0.19, which was further considered to be a cancer-like methylation-high pattern. Using this threshold, 72% (38 of 53) of CIN3 and 55% (11 of 20) of CIN2 showed a cancer-like methylation-high pattern. The remaining CIN2 and CIN3 lesions had predicted probabilities that were similar to those detected in the far majority (25 of 26) of hrHPV-positive controls, and were considered as methylation-low.

A few differences in DNA methylation patterns were observed between genes. ANKRD18CP and C13orf18 were the only 2 genes with low predicted probabilities in the single hrHPV-control with a cancer-like methylation-high pattern. Furthermore, C13orf18 showed either very low or very high predicted probabilities, whereas the other 11 methylated genes showed a gradual range of values. PHACTR3 had generally very low predicted probabilities in CIN2 and CIN3 lesions. This is also illustrated by unsupervised clustering, revealing a similar cluster for most genes, except for ANKRD18CP, C13orf18, and PHACTR3 (Supplementary Figure 2).

Discussion

This study systematically analyzed the onset and DNA methylation pattern of 12 genes during hrHPV-induced carcinogenesis using consecutive passages of hrHPV-immortalized cells, reflecting
Figure 1. DNA methylation levels of 12 genes in the in vitro model. HFK, human foreskin keratinocytes (n = 3); Early, early immortal passages of FK16A, FK16B, FK18A and FK18B p32–p52 (n = 4); Late, late immortal passages of FK16A, FK16B, FK18A and FK18B p129-p156 (n = 4); CCL, cancer cell lines (n = 3). Each of the individual cell lines is indicated by a different symbol. Only for the FK16A, FK16B, FK18A, and FK18B in the 2nd and 3rd column the symbols refer to the early and late passages of the same cell line.
Figure 2. DNA methylation level distributions in hrHPV-positive cervical scrapings across histological subgroups for 12 genes. Normal, hrHPV-positive controls (n = 26); CIN2 and CIN3, cervical intraepithelial neoplasia grade 2 (n = 20) and 3 (n = 53); (mi)Ca, micro-invasive cervical cancer (n = 14). *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; NS, not significant.
the progressive stages of cervical precancerous disease, complemented with clinically annotated specimens. A progressive increase in DNA methylation levels represented a general phenomenon during hrHPV-induced carcinogenesis. All genes showed increased DNA methylation levels during hrHPV-induced transformation in vitro and in clinical specimens proportional to disease severity. Highest DNA methylation levels were consistently seen in cervical cancer cell lines and cervical scrapings from women with cervical cancer. Using consecutive stages of hrHPV-transformed cells, we showed that the onset of DNA methylation occurred either at the early immortal stage or late immortal stage, when cells are still pre-tumorigenic. In a series of hrHPV-positive cervical scrapings, we showed that the majority of genes displayed a comparable DNA methylation pattern with an onset of a cancer-like methylation-high pattern being detected at the CIN2/3 stage.

Interestingly, the hrHPV-positive scrapings from women with CIN2/3 displayed a heterogeneous DNA methylation pattern, in which three-quarters of the CIN3 samples and half of the CIN2 samples had a cancer-like methylation-high pattern. These results correspond to the finding that only a subset of CIN2/3 will progress to cancer over a long time period [18,19]. Previous studies on copy number changes and DNA methylation levels of only a few genes also showed a cancer-like profile in only a few CIN2/3 lesions [11,14,20,21]. These lesions were characterized by a preceding hrHPV infection of >5 years and considered as more advanced lesions, which have a high short-term progression risk to cancer. Conversely, the methylation-low CIN2/3 may be more likely to regress, knowing that approximately ~30% of CIN3 and ~40–54% of CIN2 regress [19,22,23].

Most of the genes evaluated in this study were reported to have a (putative) tumor suppressive
function (i.e., C13orf18 [24], EPB41L3 [25–29], PRDM14 [30], SOX1 [31,32], SST [33,34], ZIC1 [35,36], and ZSCAN1 [37,38]). The remaining genes with a currently unknown function await further study.

A few genes (ANKRD18CP, C13orf18, and PHACTR3) showed a different DNA methylation pattern compared with the majority of genes. This may in part relate to a different onset of DNA methylation during cervical carcinogenesis. DNMTs can be activated by both hrHPV E6 and E7 [3,4], and specific promoter sequences might be more prone to DNA methylation compared to others [39,40], which may relate to the binding of either the polycomb repressor complex (PRC) 1 or 2 [41]. We purposely retained these 3 genes in our analysis on DNA methylation patterns in order to obtain an unbiased representative result. In cervical scrapings, only one hrHPV-positive control showed a cancer-like methylation-high pattern with very high individual predicted probabilities for 10 of 12 methylated genes. However, no gynecological diseases were identified in the follow-up data. Other unidentified underlying abnormalities may have resulted in increased DNA methylation levels. Considering a potential effect of age, i.e., 55 year for this control, DNA methylation levels of the markers tested were not or minimally affected by age in our previous studies using large cohorts [8–11,42,43]. Therefore, solely age is very unlikely to explain the cancer-like methylation-high pattern in this hrHPV-positive control.

A limitation is that the amount of dysplastic cells in the specimens is unknown, and a potential effect of varying sample constitution cannot be fully excluded.

The genes evaluated in this study may well serve as objective molecular tools to improve cervical cancer screening, especially as triage test after primary hrHPV testing [1,7,44]. In several countries, including The Netherlands, hrHPV testing is replacing cytology as primary screening method. Compared to cytology, hrHPV testing has a higher sensitivity for CIN2+ detection [45,46]. However, its 3–5% lower specificity makes triage testing of hrHPV-positive women necessary to prevent over-referral and overtreatment. In the Dutch population-based hrHPV screening program, cytology is included as triage method. However, cytology comes with some limitations, including its subjective nature and the required repeated cytology to ensure sufficient safety in the screening program. Moreover, prior knowledge of hrHPV-positivity influences cytology reading, which may result in an increase of false-positive referrals with simultaneously higher costs for the healthcare system [47–49]. Recent clinical validation studies in screening populations have shown that DNA methylation markers provide a good alternative for cytology [14,50–54].

In conclusion, this study showed that hrHPV-induced carcinogenesis is characterized by increased DNA methylation, with onset typically occurring at the pre-tumorigenic stage and highest DNA methylation levels at the cancer stage. Host-cell DNA methylation patterns of the 12 genes are comparable and reveal the heterogeneous nature of cervical pre-cancer, with a subset of CIN2 and CIN3 lesions displaying a cancer-like methylation-high pattern, suggestive for a higher risk of progression to cervical cancer.

**Materials and methods**

**Cell lines**

Cells representing the various stages of hrHPV-induced transformation, consisted of (i) primary human foreskin keratinocytes (HFK) from three donors, (ii) HPV16- (FK16A and FK16B) and HPV18- (FK18A and FK18B) immortalized keratinocytes, including early (passages 32 to 52) and late (passage 129 to 156) passages [15], and (iii) hrHPV-positive cervical cancer cell lines SiHa, HeLa and CaSki. HFKs, FK16A, FK16B, FK18A and FK18B were obtained and cultured as described before [15]. Cervical cancer cell lines SiHa, HeLa, and CaSki (American Type Culture Collection) were cultured as described previously [55]. Cell lines were authenticated using the PowerPlex 16 System (Promega) and were negative for mycoplasma.

**Clinical samples and hrHPV testing**

A set of hrHPV-positive scrapings (n = 113) obtained from screening or gynecologic outpatient
populations was used, comprising scrapings from control women with normal cytology and/or without evidence of CIN2+ (n = 26) or from women who were histologically diagnosed with CIN2 (n = 20), CIN3 (n = 53) or (micro-invasive) cervical cancer [(mi)Ca; n = 14]. HrHPV was detected by GP5+/6+ PCR enzyme immunoassay [56] or HPV-Risk assay (Self-screen B.V.) [57]. This study followed the ethical guidelines of the Institutional Review Board of the VU University Medical Center and University Medical Center Groningen.

**DNA isolation, modification, and qMSP**

DNA extraction, quality assessment, sodium bisulfite-treatment, (multiplex) qMSP conditions, and DNA methylation ratio calculations were described previously for ANKRD18CP, C13orf18, EPB41L3, and JAM3 [8,9]; SOX1 and ZSCAN1 [9,10]; GHSR, SST and ZIC1 [11]; and FAM19A4, PHACTR3, and PRDM14 [12]. Quality control was routinely performed using a methylation-independent assay for the housekeeping gene ACTB [13].

**Data analyses**

The DNA methylation ratios of the 12 genes were relatively analyzed to obtain equivalent values from each of the individual genes in the datasets of both the in vitro model and the cervical scrapings.

To analyze the onset of DNA methylation using the in vitro model representing the various stages of hrHPV-induced transformation, we set the highest DNA methylation ratio of each methylated gene at 100% and subsequently defined the time point of DNA methylation onset at the stage where DNA methylation was detected in at least 2 out of 4 cell lines above 5%.

In cervical scrapings, the Kruskal-Wallis test was first performed on square-root transformed DNA methylation ratios of each methylated gene to assess the differences in DNA methylation levels among disease categories. Following a significant result from the test, post-hoc testing was performed using Wilcoxon rank sum test. The Bonferroni correction was subsequently applied for multiple testing, with a significance level of 0.05 (two-sided).

To obtain equivalent DNA methylation values in cervical scrapings, univariable logistic regression analysis was performed on the square-root transformed DNA methylation ratios of the 12 genes. For this analysis, scrapings from women with cervical cancer were categorized as cases and scrapings from hrHPV-positive control women (i.e., with normal cytology and/or without evidence of CIN2+) as controls. Subsequently, the logistic regression models were used to calculate the predicted probability (value range 0 to 1), representing the risk for an underlying cervical cancer. By this, equivalent values for the levels of DNA methylation were obtained for each of the individual genes allowing direct comparison of the genes. The average predicted probability of the 12 methylated genes was calculated for each sample. The lowest average predicted probability in cervical cancer was used as a threshold to define a cancer-like methylation-high pattern (i.e., threshold of ≥0.19). Hierarchical clustering was performed for the 12 methylated genes using the predicted probabilities.

All statistical analyses have been performed using R version 3.1.2.

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**Disclosure statement**

AGJvdZ, ES, and GBAW are inventors of patents related to the contents of the manuscript. RDMS, CJLMM, PJFS, DAMH have minority stake in Self-screen B.V., a spin-off company of VU University Medical Center. Self-screen B.V. holds patents related to this work. ES is a member of the scientific advisory board of Roche, Hologic and QCMD, and received travel reimbursements from Roche, Abbott, Hologic Inc. and QCMD. GBAW is a member of the scientific advisory board of CC Diagnostics. DAMH has been on the speaker’s bureau of Qiagen, and serves occasionally on the scientific advisory boards of Pfizer and Bristol-Meyer Squibb. CJLMM has received speakers’ fee from SPMSD/Merck, served occasionally on the scientific advisory board (expert meeting) of Qiagen, SPMSD/Merck. He has been co-investigator on a Sanofi Pasteur MSD sponsored trial, of which his institute received research funding. He is part-time director of Self-screen B.V. and has a very small number of Qiagen shares. Until April 2016 he had minority stock of Diassay B. V. PJFS has been on the speaker’s bureau of Roche, Abbott,
Gen-Probe, Qiagen and Seegene. He is consultant for Crucell Holland B.V.

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