Gene Promoter-Methylation Signature as Biomarker to Predict Cisplatin-Radiotherapy Sensitivity in Locally Advanced Cervical Cancer

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Despite efforts to promote health policies focused on screening and early detection, cervical cancer continues to be one of the leading causes of mortality in women; in 2020, estimated 30,000 deaths in Latin America were reported for this type of tumor. While the therapies used to treat cervical cancer have excellent results in tumors identified in early stages, those women who are diagnosed in locally advanced and advanced stages show survival rates at 5 years of <50%. Molecular patterns associated with clinical response have been studied in patients who present resistance to treatment; none of them have reached clinical practice. It is therefore necessary to continue analyzing molecular patterns that allow us to identify patients at risk of developing resistance to conventional therapy. In this study, we analyzed the global methylation profile of 22 patients diagnosed with locally advanced cervical cancer and validated the genomic results in an independent cohort of 70 patients. We showed that BRD9 promoter region methylation and CTU1 demethylation were associated with a higher overall survival (p = 0.06) and progression-free survival (p = 0.0001), whereas DOCK8 demethylation was associated with therapy-resistant patients and a lower overall survival and progression-free survival (p = 0.025 and p = 0.0001, respectively). Our results suggest that methylation of promoter regions in specific genes may provide molecular markers associated with response to treatment in cancer; further investigation is needed.

Keywords: gene promoter methylation, chemoradioresistance, cervical cancer, biomarkers, Cisplatin-Radiotherapy sensitivity
1 INTRODUCTION

Cervical cancer (CC) is the fourth most common type of cancer in women worldwide (1). In developing countries, mainly in Latin America, about 30,000 deaths per year are caused by this disease (2). The high mortality rates are due to the fact that 50% of patients are diagnosed in locally advanced cervical cancer stages (LACC); the overall survival (OS) rate to 5 years is approximately 60% (1) and a recurrence rate from 15% to 40% (3). Conventional treatment for LACC patients consists of concomitant chemoradiotherapy. Unfortunately, treatment resistance is observed in approximately 30% of patients (4).

Treatment resistance involves several molecular alterations such as genetic mutations, dysregulated microRNAs, dysregulated long noncoding RNAs expression profiles, and epigenetic modifications (5–9). Several reports described that the aberrant DNA methylation that involves hypo- or hypermethylation is also associated with tumor progression and therapy resistance (10, 11). For example, the hypermethylation of PTEN, MYOD1, RASSF1A, APC1A, PTGS2, and VIM genes, which are associated with OS of CC patients, covered all stages (12–16). Nevertheless, the expanding knowledge about methylation profiles in patients with LACC is pertinent and is focused on the treatment resistance in these particular patients.

The goal of this study was to obtain the global methylation pattern of tumor biopsies from 92 LACC patients treated with chemoradiotherapy to identify the methylation status of specific gene promoters with predictive potential to the cisplatin-radiotherapy response. For this purpose, we analyzed the methylation profile in 22 patients and found global changes in methylation patterns in 7,957 gene promoter regions that distinguish responsive and resistant LACC patients to chemoradiation. Next, by means of bioinformatics tools, we selected promoter sequences with a CpG density higher than 60%; these regions corresponded to the promoters of the bromodomain containing 9 (BRD9), dedicator of cytokinesis 8 (DOCK8), and cytosolic thiouridylase subunit 1 (CTU1) genes. Then, promoter regions were experimentally validated by methylation-specific PCR (MSP) in an independent cohort of 70 LACC patients. Strikingly, we found a correlation between BRD9 promoter region methylation and CTU1 demethylation with complete response to chemoradiotherapy in addition to higher overall survival (OS) (p = 0.06) and progression-free survival (PFS) (p = 0.0001). Moreover, demethylation of DOCK8 promoter region was associated with patients who developed treatment resistance and lower OS and PFS (p = 0.025 and p = 0.0001, respectively). These data point to the methylation status of BRD9 CTU1 and DOCK8 as potential biomarkers for predicting survival and response to chemoradiotherapy in LACC patients.

2 MATERIAL AND METHODS

2.1 Tissue Samples

This study was approved by the Central Ethics and Scientific Committee at the National Cancer Institute in Mexico City (INCan) (015/012/ICI, CEI/961/15) and has been conducted in agreement with the ethical standards as laid down in the 1964 Declaration of Helsinki and its later amendments. A total of 92 biopsies from patients with LACC cancer were obtained. Tumor samples were collected from 2014 to 2018 by the Pathology Department, INCan, Mexico City. After confirmed diagnosis, all patients received concurrent chemoradiotherapy using cisplatin [weekly cis-diaminedichloroplatinum (II) at a dose of 40 mg/m²] for a total of five or six cycles and radiation (external irradiation and intracavitary brachytherapy, for a total dose of 64–66 Gy over 67 days) (17). The patients’ therapy response was assessed according to RECIST criteria defined as follows: the disappearance of all target lesions was assigned as complete response (CR); meanwhile, patients with partial response, progressive disease, or stable disease were considered as therapy resistant (TR). The biopsies were divided into two cohorts: the first with 22 patients (12 CR and 10 TR) used as a discovery cohort to generate a microarray specific for CpG islands Array-Based Profiling of Reference-Independent Methylation Status (aPRIMES) (18); the second cohort, with 70 biopsies (40 CR and 30 TR), used for molecular data validation. The patient eligibility criteria consisted of (a) confirmed pathological diagnosis of CC stages from II-B to IV-B (LACC), (b) biopsies with a pathology report confirming more than 80% tumorous cells, (c) age range of 29–65 years, (d) high-quality DNA and RNA samples, (e) no other comorbidity, (f) no previous oncological treatment, and (g) patients able to receive the standard therapy based on concurrent chemoradiotherapy and radiotherapy.

2.2 Nucleic Acid Extraction

The DNA extraction from the 92 biopsies was performed as follows: 20 mg of fresh tissue was placed in a Fisherbrand Bead Mill homogenizer, and 2 ml soft tissue homogenizing Mix Tube was preloaded with lysis buffer [10 mM Tris–HCl, 2 mM ethylenediaminetetraacetic acid (EDTA), 1% sodium dodecyl sulfate (SDS)]. The tissue was homogenized using the MagNA Lyser instrument at 6,000 rpm for 1 min. To purify the genomic DNA, the QIAamp DNA Blood Kit (Qiagen, CA, USA) was used according to the manufacturer’s protocol. Finally, the purified DNA was stored at −20°C.

2.3 Microarray Differential Methylation Analysis (aPRIMES)

We employed the 3x720K CpG Island Plus RefSeq Promoter Arrays (Roche, Penzberg, Germany). These arrays cover the annotated CpG islands and the promoters of the RefSeq genes derived from the UCSC RefFlat files (Hg 38). Then, the hybridization probes were synthesized by aPRIMES assay. Briefly, genomic DNAs were digest by MseI, and the fragments obtained were subjected to linker-mediated PCR as described by Klein and coworkers (19); later, through enzymatic digestion by methylated-sensitive and methylated-specific enzymes, we obtained a methylated and unmethylated fraction of DNA, which were labeled with Cy5 and Cy3 fluorophores, respectively, and competitive hybridizing in a Human DNA Methylation 3x720K CpG Island Plus (Roche) as shown in Supplementary Figure 1. Then, arrays were scanned in
an MS200 scanner (Roche). Finally, the alignment of the images and the extraction of data were carried out using the software DEVA Project Manager—1.2.1 (Roche). Next, for each region in the array, we obtained a continuous numerical ratio that represents if a region is hyper- or hypomethylated; we termed this ratio as bi-weight (BW) and is represented with the following formula:

\[ BW = \frac{\log(2(Cy3))}{\log(2(Cy5))} \]

Finally, to determine the significative methylated regions between responsive and resistant tumors, we calculated the Student’s t-test for each region between both groups and considered as statistical significance those methylated regions with a p < 0.01. Then, selected regions were ranked in ascending and descending orders accordingly to the difference of the means for both groups. All statistical analysis were executed in R environment.

2.4 Pathway Analysis
Differentially methylated regions were analyzed by Pathway enrichment analysis by using Webgestalt (20) and ReactomePA (21); we only considered pathways with a p < 0.05 as subject of regulation.

2.5 CpG Island Density Determination
We obtained from the Genome Browser database (22) a sequence of 2,000 bp (1,000 bp downstream; transcription start site, 1,000 bp upstream) that included the promoter region from each analyzed gene. These sequences were analyzed using MethPrimer web tool from Urogene to determine the CpG density (23).

2.6 Methylation-Specific PCR Assay
To determine the methylation status of selected genes (BRD9, CTU1, and DOCK8), genomic DNA from each sample was modified using Methylation-Direct EZ DNA Kit (ZYMO, CA, USA). DNA bisulfite treatment changed unmethylated cytosines to uracil, but the methylated bases remained as cytosines. Then, two PCR reactions were performed per sample using specific primers to determine the methylated (M) or unmethylated (U) DNA status. The list of primers and its characteristics are shown in Supplementary Table S1. The product of each reaction was analyzed in agarose gels and resolved in Minigel OWLTM Easy CastTM B2 system (Thermo Scientific, MA, USA). Later, the gel was stained with ethidium bromide and photo-documented using a Gel Doc EZ Imager transilluminator (Bio Rad, CA, USA).

2.7 Statistical Analysis
Chi-squared tests were employed to determine the differences in the distribution of the methylation status of the genes and the clinicopathological characteristics, considering p < 0.05 as statistically significant.

2.8 Survival Analysis
Kaplan–Meier plotter was calculated using the survival package in R, where the significance testing was assessed using the log-rank test. Significance was considered as p < 0.05.

3 RESULTS
3.1 Clinicopathological Characteristics of Patients
This study was approved by the Central Ethics and Scientific Committee at the National Cancer Institute in Mexico City (approval number 015/01271B/CEI/961/15). The 92 patients who were enrolled accepted and signed the informed consent. All patients received treatment based on cisplatin and radiotherapy as mentioned in Material and Methods. The median age was 48 years. Patients were classified following the last version of International Federation of Gynecology and Obstetrics (FIGO) staging criteria as II (51.8%), III (37%), and IV (11.2%) stages. According to the FIGO’s guidelines, 52 patients (56.52%) showed complete response (CR) to therapy; meanwhile, 40 (43.48%) exhibited therapy resistance (TR). The HPV-genotype of all patients was determined by nested PCR (24). Table 1 shows the clinicopathological characteristics; a supplementary table that compiles all clinical data is available as Supplementary Table 2.

3.2 Global Analysis of DNA Methylation in Locally Advanced Cervical Cancer Tumors
The determination of DNA methylation patterns has been proposed as a prognosis predictor in several types of cancer (25, 26). In this study, we employed aPRIMES arrays to obtain the genome-wide DNA methylation patterns on both groups, namely, responsive (CR) and therapy-resistant (TR) LACC patients. Next, to establish the differentially methylated regions (DMRs) on CR and TR groups, we compared the bi-weight ratio values from each analyzed region, using a Student’s t-test. The results showed a methylated DNA profile between the two groups composed of 16,538 DMRs that corresponded to 7,957 unique regions, where 2,833 of them were hypermethylated, 5,881 were hemimethylated, and 757 regions had both hyper and hypomethylated DMRs regions (p < 0.05) (Figure 1). As expected, a global hypomethylation pattern across the genome was observed, where the distribution of these DMRs varied to a considerable extent depending on the chromosome (Supplementary Figure S2). We noticed that chromosomes 1 and 19 had the higher number of gene promoters with DMRs, 859 and 753, respectively (Supplementary Figures S2, S3). Next, we observed that clustering of these DMRs using the Euclidean distance algorithm could distinguish the TR (black bar) from CR LACC patients (green bar) (Figure 1). Since the bi-weight ratios are continuous variables, we transformed them to a Z-score to visualize the DMRs in a heatmap, which clearly shows a DNA methylation profile that includes 3,533 DMRs hypermethylated and 13,005 DRMs hypomethylated (Figure 1).

3.3 Gene Pathway Analysis
Furthermore, we were interested in evaluating the impact of the methylation profile in biological pathways. The gene set enrichment analysis using the Kyoto Encyclopedia of Genes and Genomes (KEGG) database revealed that multiple key carcinogenic pathways such as the PI3K-AKT signaling
pathway, nuclear factor (NF)-kappa B pathway, RNA polymerase, and pathways associated with breast cancer were dysregulated as a consequence of differentially methylation profile (Figure 2). As the PI3K-Akt pathway was the most enriched, we focused on analyzing it in more detail. As shown in Figure 2, multiple key genes in this pathway were hypomethylated, including the insulin receptor substrate-1 (IRS-1) and oncogene JAK3. In contrast, genes such as RELA that inhibit the tumor growing were hypermethylated.

3.4 The Methylation Status of BRD9, CTU1, and DOCK8 Gene Promoter Regions Is Associated With Clinical Outcomes of LACC Patients

To select methylated genes as potential biomarkers of response to chemoradiation, we further narrowed the methylation profile by considering only those DMRs that showed hyper- or hypomethylation status for further analysis. The results showed 4,463 DMRs with these methylation patterns that correspond to 1,439 unique genes. Then, median bi-weight values of methylation from each gene in TR and CR tumors were compared to calculate the median difference (MD). Promoter regions with an MD upper than 1.4 times, corresponding to the 13 genes enlisted in Table 2, were chosen for further analysis. The promoter sequence of the selected genes was analyzed as mentioned in Section Cpg Island Density Determination, and the three genes with highest Cpg density were selected for validation. A Cpg Island is defined as a DNA region higher than 500 bp that contains 50% or more of CG islands.
The promoter region of BRD9, CTU1, and DOCK8 genes showed 88%, 63%, and 90% of CpG density, respectively (Figure 3). Additionally, the MD for these promoter regions was 1.61, 2.25, and 1.73 for BRD9, CTU1, and DOCK8, respectively (Figure 3, boxplots). Interestingly, H3K4me3 mark (chromatin compaction mark) was found near to these promoter regions (Figure 3).

To validate the methylation levels of BRD9, DOCK8, and CTU1 promoter regions as therapy response biomarkers, an MSP assay was performed. Bisulfite-treated DNA from 30 TR and 40 CR tumors samples were used to analyze methylated status. The results showed that the BRD9 promoter region was methylated in all CR tumor samples (40 CR tumor samples, 100% of cases), whereas it was hemimethylated in 25 TR tumor samples and unmethylated in 5 TR tumor samples (83% and 17% of cases, respectively) (Figure 4A). Instead, the promoter region of CTU1 gene was detected to be unmethylated in all CR tumor samples (40 CR tumor samples, 100% of cases), while in 27 TR tumors samples, it was hemimethylated; in 2 TR tumor samples, it was unmethylated and only in 1 TR tumor sample that it was methylated (90%, 6.6%, and 3.4% of cases, respectively) (Figure 4B). On the other hand, the DOCK8 promoter region was unmethylated in 29 TR tumors samples (97% of cases), whereas in 31 CR tumors samples, it was hemimethylated, and in 9 CR tumors samples, it was unmethylated (77.5% and 22.5% of cases, respectively) (Figure 4C).

Additionally, a chi-square analysis was performed to compare the methylation status of BRD9, CTU1, and DOCK8 genes with demographic characteristics of LACC patients (Table 3). The methylation of the BRD9 gene was associated with tumor stages II and tumor size <5 cm. In contrast, unmethylation of the CTU1 promoter region gene was associate with stages II and with tumor size <5 cm. The unmethylation status of the DOCK8 promoter region showed an association with stages III–IV; however, no significant relationship was found between the methylation status of this promoter with tumor size.

3.5 A Gene Methylation Signature as Biomarker for Overall Survival and Progression-Free Survival in CC

Finally, we determined if the methylation status of BRD9, CTU1, and DOCK8 genes could be an OS and the PFS biomarker of LACC patients. The results showed a better OS (p < 0.0041) and PFS (2.28 months in the hemimethylated group, p < 0.0001) in patients with the methylation of BRD9 promoter (Figures 5A, D). In contrast, worse OS (p < 0.025) and PFS (3.12 months in the unmethylated group p < 0.0001) was observed in patients with the methylation of DOCK8 promoter (Figures 5B, E). Moreover, patients with a unmethylated CTU1 promoter showed a better OS and PFS (1.76 months in the hemimethylated group p < 0.0001) (Figures 5C, F). These data highlight that the methylation status of BRD9, CTU1, and DOCK8 have the potential as biomarkers of OS and PFS in LACC patients.

4 DISCUSSION

Despite global screening programs, CC remains a health problem in Latin American countries, with an estimated...
56,000 new cases and 28,000 cervical cancer deaths (2). Unfortunately, more than 50% of CC patients are diagnosed at locally advanced stages with a 5-year survival rate of 60% (28). Epigenetic processes are crucial in cellular homeostasis, and their dysregulation leads to cancer and progression (29). DNA methylation is a tag for chromatin remodeling factors that have a crucial role in transcription regulation; DNA methylation in promoter regions is considered as a transcriptional repression mark of gene expression (30). The aberrant methylation of genes is a relevant event during carcinogenesis, which could be a diagnostic biomarker of the disease (31, 32). However, few studies are focused on associating the methylation status with the response to treatments in CC patients. Therefore, expanding knowledge about methylation profiles in patients is decisive to build knowledge focused on treatment resistance. In this regard, we aimed to find gene methylation as a biomarker of response to chemoradiotherapy in LACC. Consequently, we performed a global analysis of DNA methylation from chemoradiotherapy-responsive tumor biopsies to establish DNA methylation patterns. Hence, we identified a gene methylation profile that distinguished between responsive patients and resistance to chemoradiotherapy. As mentioned previously, prognostic biomarkers based on chemoradiotherapy-related aberrant DNA methylation are limited. However, a study in head and neck squamous cell carcinoma described a characteristic promoter methylation pattern of ZNF10, TMPRSS12, ERGIC2, and RNF215 genes, which was proposed as a biomarker of response to radiotherapy treatment (33).

Another study performed in low-grade gliomas reported a consistent signature in the methylation of MGMT, MLH3, RAD21, and SMC4 promoter region predictive value for response to temozolomide (34). In breast cancer, the hypermethylation of IL15RA gene promoter induced the upregulation of genes involved in adhesion and ECM-interaction pathways correlating with the OS of patients (35). In CC, methylation patterns are used as biomarkers to distinguish between healthy and cancerous tissue (36–39). Besides, methylation of SOCS2 and hTERT promoter region was associated with early-stage tumors (40), while the methylation of the APC1A promoter was related to advanced stages (15). Therefore, methylation profiles could predict cancer stages. Elsewhere, reports indicated the role of gene methylation associated with survival, such as MYOD1 and VIM methylation status associated with more favorable disease-free survival and OS (12, 39, 41). Likewise, our results showed a correlation between the methylation status of BRD9, CTU1, and DOCK8 promoter regions with PFS, OS, and clinicopathological characteristics of LACC patients.

In the present work, we ascertained a signature to predict chemoradiotherapy response in LACC patients. This signature consisted of the methylation of the BRD9 promoter region, the unmethylation of the CTU1 gene, and the unmethylation of the promoter region of DOCK8. Fascinatingly, the methylation of the BRD9 promoter region and unmethylation of CTU1 were related to CR, and the unmethylation status of DOCK8 was
related to TR. Furthermore, the methylation signature was validated in an independent cohort, allowing us to propose it as a potential biomarker to predict the response capacity of LACC patients to chemoradiotherapy. In this regard, CpG island methylation from DNA promoter regions leads to the inactivation of genes, some of which are tumor suppressors, whereas the demethylation of those repeat elements induces gene expression of oncogenes (5).

In our work, we detected the BRD9 gene promoter methylation pattern in CR tumors, suggesting low levels of expression of this gene, which could explain the response rates to chemoradiotherapy. The BRD9 gene encodes a protein that functions as a protein interaction module that recognizes lysine acetylation domains, a key event in the reading of epigenetic marks (40). The overexpression of this gene in lung cancer cells was associated with poor prognosis, and its oncogene role was demonstrated in synovial sarcoma (42).

In this work, we detected CTU1 unmethylated in LACC samples of patients that showed response to chemoradiotherapy and better OS. CTU1 plays a crucial role in the processing of transfer RNA by modifying nucleosides for the precise binding of the anticodon, thus guaranteeing the fidelity of the translation by the ribosome (43). However, its role in CC has not been analyzed yet, but in breast cancer, CTU1 overexpression promotes cell invasion (44). On the other hand, we found that the unmethylation of the promoter region of DOCK8 was detected in TR patients. The role of this gene is unknown in CC. Nevertheless, in a recent work, Biswas and colleagues (45) reported that DOCK8 is a gene that codifies to a nucleotide exchange factor (GEF) that activates the GTPase Cdc42, participating in cell migration and invasion. In addition, it was shown acute in myeloid leukemia that its pharmacological inhibition attenuates cell survival (45).

Then, we performed a multi-pathway analysis using the differential methylation pattern established from the comparison between CR and TR tumors. The results showed dysregulated pathways such as PI3K-Akt-mTOR. This pathway regulates multiple cellular and molecular functions like cell cycle progression, cellular growth, and protein synthesis and is altered in various cancer types including CC, which are crucial for tumor initiation, invasion, and metastasis (46). These data suggested that this pathway could be hyperactivated in chemoradiotherapy-resistant LACC patients (Figure 2). This is
the case of ovarian and breast cancers, where it was shown that hyperactivation of this pathway is related to chemoresistance and drug resistance, respectively (47, 48). There are no studies that corroborate the causality of hyperactivation of the pathway and chemoradiotherapy resistance in CC. Thus, the elucidation of molecular pathways altered by the differential methylation pattern between responsive and resistant cervical tumors remains a perspective to future studies.

In summary, this is the first study to report a molecular signature of promoter methylation of the BDR9, CTU1, and DOCK8 genes, which could distinguish LACC response patients resistant to chemoradiotherapy. In this regard, we propose them as potential biomarkers of response to chemoradiotherapy in LACC patients. Extending this study to other cohorts and deepening the biological role of these genes are of great interest.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Ethics committees of the National Cancer Institute
of Mexico (015/012/ICI, CEI/961/15). The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

Conceptualization, CC-R, CP-P; Experimentation CC-R, AM-G, GC-A; Analysis of results, CC-R, AM-G, E-AP-Y, AC-P, AZ-D, CP-P; Writing-Original Draft Preparation, CC-R, E-AP-Y, AM-G; Writing-Review & Editing, E-AP-Y; CP-P; AZ-D Project Administration, CP-P; Funding Acquisition, CP-P. Clinical follow up: JMC. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.773438/full#supplementary-material
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