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#### Research paper

# Deciphering the complex interplay between microbiota, HPV, inflammation and cancer through cervicovaginal metabolic profiling



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#### ABSTRACT

*Background:* Dysbiotic vaginal microbiota have been implicated as contributors to persistent HPV-mediated cervical carcinogenesis and genital inflammation with mechanisms unknown. Given that cancer is a metabolic disease, metabolic profiling of the cervicovaginal microenvironment has the potential to reveal the functional interplay between the host and microbes in HPV persistence and progression to cancer.

Methods: Our study design included HPV-negative/positive controls, women with low-grade and high-grade cervical dysplasia, or cervical cancer (n=78). Metabolic fingerprints were profiled using liquid chromatographymass spectrometry. Vaginal microbiota and genital inflammation were analysed using 16S rRNA gene sequencing and immunoassays, respectively. We used an integrative bioinformatic pipeline to reveal host and microbe contributions to the metabolome and to comprehensively assess the link between HPV, microbiota, inflammation and cervical disease.

Findings: Metabolic analysis yielded 475 metabolites with known identities. Unique metabolic fingerprints discriminated patient groups from healthy controls. Three-hydroxybutyrate, eicosenoate, and oleate/vaccenate discriminated (with excellent capacity) between cancer patients versus the healthy participants. Sphingolipids, plasmalogens, and linoleate positively correlated with genital inflammation. Non-Lactobacillus dominant communities, particularly in high-grade dysplasia, perturbed amino acid and nucleotide metabolisms. Adenosine and cytosine correlated positively with Lactobacillus abundance and negatively with genital inflammation. Glycochenodeoxycholate and carnitine metabolisms connected non-Lactobacillus dominance to genital inflammation.

*Interpretation:* Cervicovaginal metabolic profiles were driven by cancer followed by genital inflammation, HPV infection, and vaginal microbiota. This study provides evidence for metabolite-driven complex host-microbe interactions as hallmarks of cervical cancer with future translational potential.

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#### 1. Introduction

Persistent human papillomavirus (HPV) infection, the most common sexually transmitted disease, is the etiologic agent for the development of precancerous cervical dysplasia and cervical cancer [1,2]. Even though screening and HPV vaccination have significantly reduced the number of cases in developed countries, cervical cancer is still the fourth

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most commonly diagnosed cancer globally among females [3]. After HPV infection, carcinogenesis does not progress uniformly; precancerous dysplasia may regress without treatment for reasons that are uncertain [4]. Regardless, persistent HPV infection initiates a chain of reactions that suppresses the cell-mediated immune response and hinders clearance of the infection and destruction of abnormal cells [5]. Additionally, in spite of the rapidly accumulating data on host immune responses and vaginal microbiome in patients across cervical carcinogenesis [6–11], our understanding of the molecular and metabolic changes in the cervicovaginal microenvironment relevant to microbiome has only recently begun to be elucidated [12,13].

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#### Research in context

#### Evidence before this study

Cervical cancer is a common gynaecological malignancy in women caused by persistent oncogenic HPV infection. Recent studies have demonstrated that dysbiotic microbiota is involved in HPV persistence, progression of cervical neoplasia, and genital inflammation. Metabolomic studies of the cervicovaginal microenvironment in bacterial vaginosis and pre-term birth suggest that vaginal microbiota drastically alter the metabolome. However, metabolic interactions between host and microbial communities in cervical carcinogenesis remain to be elucidated.

#### Added value of this study

In the present study, we identified metabolic signatures that effectively distinguish-HPV infected individuals, women with cervical dysplasia and cancer from HPV-negative women. Membrane lipids distinguished the cancer group from the HPV negative group and were highly associated with genital inflammation. In low-grade and principally high-grade dysplasia groups, we observed polymicrobial communities perturb amino acid and nucleotide metabolism in a similar manner to bacterial vaginosis. Our integrative and comprehensive bioinformatic approach pinpointed the microbial or host origin of specific amino acid metabolites and nucleotides. In addition, a novel aspect of our multi-omic analyses was to reveal the level to which specific metabolites were associated with Lactobacillus abundance as well as genital inflammation. An increase in the Lactobacillus abundance was positively associated with the levels of anti-inflammatory nucleotides. Collectively, we revealed that features of the cervicovaginal microenvironment (HPV, genital inflammation and vaginal microbiota) profoundly impact cervicovaginal metabolomes.

#### Implications of all the available evidence

This first report on cervicovaginal metabolomes in HPV-mediated cervical dysplasia and cancer demonstrate the predictive value of metabolic fingerprinting in abnormal cell metabolism. Observed perturbations in lipids, an emerging hallmark of cancer, link inflammation and cervical cancer. Additionally, we identified specific metabolites that were associated with *Lactobacillus* abundance and genital inflammation independently from the cancer. Amino acid and nucleotide metabolism connect vaginal dysbiosis to cervical dysplasia. The cervicovaginal metabolome can be a useful target for future diagnosis, prevention strategies and therapeutic intervention to positively impact women's health outcomes.

The advances in the metabolomics field during the last decade have proven that cancer is a metabolic disease [14] and led to the rediscovery of the metabolome as an emerging target for cancer detection and therapies. Early studies have shown that cervical cancer patients harbour a unique collection of molecules ranging from amino acids to nucleic acids in the serum [15], tumour tissues [16], and faeces [17]. Moreover, research on cancer sniffing dogs indicated that there are cancer-specific volatile organic compounds being released in cervical cancer [18]. In comparison to serum or faeces, metabolomes of the cervicovaginal microenvironment have the added benefit of reflecting the metabolic processes occurring in the mucosal environment instead of metabolic outcomes of the entire body or other systems. However, to the best of our knowledge, metabolomes of the local microenvironment in patients with HPV, cervical dysplasia, or invasive cervical carcinoma (ICC), have not been previously reported.

Cervicovaginal metabolomes throughout cervical carcinogenesis can be influenced by several factors that are host-, microbiome-, and environment-associated. For example, HPV suppresses cytokine production [19], which alters the microbial dynamics in the cervical microenvironment. Our group and others have shown a strong association between genital immune mediators and a decrease in Lactobacillus abundance in cervical carcinogenesis [6,8]. Shifts in cervicovaginal microbial communities due to dysbiosis or environmental factors can alter the cervicovaginal metabolome. For instance, a shift from healthassociated Lactobacillus-dominated microbiota towards complex anaerobic and microaerophilic communities in bacterial vaginosis (BV) is highly correlated with an increase in amino acid catabolites and polyamines [12,13]. A similar metabolic shift was also observed in smokers in comparison to non-smokers in connection to imbalances in the vaginal microbiota [20]. Additionally, alterations in cell physiology caused by HPV infection and neoplastic disease could transform the dynamics of the cervicovaginal microenvironment. For example, cancer cells with a drastically increased glucose demand ferment glucose into lactate instead of carbon dioxide, in contrast to healthy differentiated cells [21]. As such, secretions from cancer cells, as well as microbial fermentation products, could ultimately modulate the microenvironment and affect both host and microbial metabolism. In the context of HPV infection and cervical carcinogenesis, these metabolic shifts in the cervical microenvironment warrant further investigation. We hypothesized that cervicovaginal metabolic profiling would reveal HPV, inflammation, host and microbe driven metabolic signatures across cervical

Herein, we characterized differences in cervicovaginal metabolomes in HPV-negative and HPV-positive individuals, women with low-grade and high-grade cervical dysplasia, and newly diagnosed ICC using a global metabolomics approach. We report unique differences in metabolic profiles of participants with HPV infection, cervical dysplasia and ICC that discriminated these patient groups from healthy HPV-negative controls. The magnitude of the differences observed among the patient groups was on the metabolic super pathway level rather than on the individual metabolite level. However, we identified 3-hydroxybutyrate, eicosenoate, and oleate/vaccenate as key metabolites that robustly discriminated patients with ICC from healthy HPV-negative individuals, and many amino acid metabolism products such as glutamine, pyroglutamine, and *N*-acetyltaurine as discriminators of healthy HPV-negative individuals from participants that were HPV-positive or with cervical dysplasia.

We also interrogated the host- and microbiota-associated factors that shape the local microenvironment and found that genital inflammation and vaginal microbiota composition are strong drivers of the cervicovaginal metabolomes. Our comprehensive bioinformatic analysis of microbial contributions to the cervicovaginal metabolome pinpointed the metabolic effects of certain BV-associated microorganisms including Sneathia, Streptococcus, Prevotella, and Gardnerella on catabolism of amino acids, which differentiated high-grade cervical dysplasia from ICC. Genital inflammation, mainly in the ICC group was highly associated with various groups of lipids. Nucleotides with antiinflammatory properties positively correlated with *Lactobacillus* abundance and negatively correlated with genital inflammation. Overall, our results provide metabolic insights into complex virus-hostmicrobiome interactions in the context of a common gynaecologic malignancy, cervical cancer that are essential for the development of future metabolome targeted diagnostics and therapies.

#### 2. Materials and methods

#### 2.1. Study subjects

We recruited 78 premenopausal, non-pregnant women and grouped as follows: healthy HPV-negative [Ctrl HPV (-), n=18] and HPV-positive participants [Ctrl HPV (+), n=11] as controls,

participants with low-grade squamous intraepithelial lesions (LSIL, n =12), high-grade squamous intraepithelial lesions (HSIL, n = 27) and newly diagnosed invasive cervical carcinoma (ICC, n = 10) at three clinical sites in Phoenix, AZ: St. Joseph's Hospital and Medical Center, University of Arizona Cancer Center and Maricopa Integrated Health System. Our study was approved by the Institutional Review Boards of each participating site and written informed consent was attained from each study participant. The inclusion criteria and participant classification for each of the groups were described previously [6]. Briefly, women that were pregnant, post-menopausal, or currently menstruating, women that were on antibiotics and antifungals within the previous three months of the study and women with the following conditions: vaginal infection, bacterial vaginosis, vulvar infection, urinary tract infection or sexually transmitted infections including chlamydia, gonorrhoea, trichomoniasis, and genital herpes, women with type I or II diabetes, or that were HIV positive, women with unusual or foulsmelling vaginal discharge, and women who use douching substances, within the previous three weeks of the study, were also excluded from the study. Sexual intercourse or vaginal lubricant use within 48 h of the sample collection also excluded women from participation. Physician's pelvic exam, medical records and surveys were used to verify exclusion criteria.

#### 2.2. Sample collection and processing

Cervicovaginal lavages (CVL) and vaginal swabs were collected from study participants. Vaginal pH measurements, HPV genotyping, and microbiome analysis from vaginal swabs were described previously [6]. Briefly, CVLs were collected by the clinicians using 10 mL of sterile 0.9% saline solution (Teknova, Hollister, CA), immediately placed on ice and frozen at -80 °C. The CVLs were thawed on ice and clarified by centrifugation (700 ×g for 10 min at 4 °C) prior to aliquoting and aliquots were sent to Metabolon, Inc. (Durham, NC) for global metabolomics identification and quantification. Samples were randomized across the extraction and instrumentation platforms. Briefly, upon receipt, samples were inventoried, and each sample identifier was accessioned into the Metabolon LIMS system and assigned a unique identifier. Sample preparation was achieved using the MicroLab STAR® system from Hamilton company. Aliquots of 100 µL were mixed with ethanol and vigorously shaken for 2 min (Glen Mills GenoGrinder 200) to precipitate the proteins and dissociate the small molecules bound to proteins. The metabolome extract was divided into five aliquots and four of them were analysed. Samples were placed briefly on a TurboVap® (Zymark) to remove organic solvent. The extracts were stored overnight under nitrogen before preparation for analysis. The sample's extract was dried then reconstituted in solvents compatible to each of the four methods described below.

#### 2.3. Metabolome analysis

All methods utilized a Waters ACQUITY ultra-performance liquid chromatography (UPLC) and a Thermo Scientific Q-Exactive high resolution/accurate mass spectrometer interfaced with a heated electrospray ionization (HESI-II) source and Orbitrap mass analyser operated at 35,000 mass resolution. Two of the aliquots were analysed with two separate reverse phase UPLC/MS/MS methods with positive ion mode electrospray ionization (ESI) optimized for hydrophilic compounds. In this method, the extract was gradually eluted from a C18 column (Waters UPLC BEH C18–2.1  $\times$  100 mm, 1.7  $\mu$ m) using water and methanol containing 0.05% perfluoropentanoic acid (PFPA) and 0.1% formic acid (FA). The second extracts were evaluated for more hydrophobic compounds by eluting the sample with methanol, acetonitrile, water, 0.05 PFPA, and 0.1% FA. The third aliquots were analysed using basic negative ion conditions by conditioning the sample with 6.5 mM ammonium bicarbonate at pH 8.0 and eluting with methanol and water through a dedicated separate C18 column. The last aliquots were analysed using a negative ionization following elution from HILIC column (Waters UPLC BEH Amide 2.1  $\times$  150 mm, 1.7  $\mu m)$  using a gradient consisting of water and acetonitrile with 10 mM ammonium formate, pH 10.8. Those four analytical methods were optimized according to the polarity or hydrophilicity of the compounds. The MS analysis alternated between MS and data-dependent MS^n scans using dynamic exclusion. The scan range varied slightly between methods but covered 70–100 m/z.

Variability and performance of extraction and instrumentation were validated by the utilization of recovery and internal standards. A pooled composite sample consisting of a small volume of each experimental sample served as a technical replicate and extracted water samples served as process blanks. Those control samples were spaced evenly among the injections to monitor instrument stability.

#### 2.4. Metabolome data collection

Data extraction was performed using Metabolon's Laboratory Information Management System (LIMS). Compounds were identified by comparison to library entries, which contain over 5000 commercially available purified standard or mass spectral entries of recurrent unknown entries. Mass-spectrometry based levels of metabolites were calculated based on area-under-the-curve of the peaks, that allows for quantitative analysis between the groups but not for the absolute quantification. Additionally, each biochemical's ionization coefficient depends on the environment. Hence, the threshold of detection is experiment, matrix, and compound dependent. However, average detection limits of the compounds were in the nanomolar range, and for some compounds, the detection limits were in the picomolar range. The threshold of detection is compound specific and determined by Metabolon's algorithm.

#### 2.5. Statistical analysis

Welch's two-sided two-sample t-test was used to perform a pairwise comparison of the means of the metabolites among the groups [Ctrl HPV (-), Ctrl HPV (+), LSIL, HSIL, and ICC]. The p values below 0.05 were corrected using false discovery rate (FDR) method and q values were reported. The fold changes observed among the groups were visualized using Cytoscape network analysis [22]. A trend analysis was performed to visualize the differences observed in number of metabolites between the groups. The metabolites that distinguished the patients in the ICC group from Ctrl HPV (-) group and Ctrl HPV (+) group from LSIL and HSIL groups were identified using Receiver Operating Characteristics (ROC) analysis and the strength of the discriminators were measured with the Area Under the Curve (AUC) values. AUC values above 0.8 were considered as good and above 0.9 were considered as excellent discriminators. To measure the strength and direction of the linear associations between the metabolites and other sample data including the inflammation scores, vaginal pH, and Lactobacillus dominance, Spearman's rank correlation coefficients were calculated, unadjusted and bootstrapped p values were reported. Based on the sample size (all samples (n = 78) versus without the ICC samples (n = 78)= 68)), Spearman's rho ( $\rho$ ) critical value of 0.223 and 0.239 were selected.

The dissimilarities in the metabolome dataset among the groups were analysed using principal component analysis (PCA). Multivariate analysis of variance (MANOVA) analysis was performed to determine the composite differences for the first two components among the clusters based on participant characteristics, *Lactobacillus* dominance, and genital inflammation. Additionally, the distances among the groups on PC1 and PC2 axes were calculated using Mann-Whitney's *U* test. Using R software and ggplot2 library [23], we overlaid shaded ellipses on clusters to reflect 95% confidence intervals from the centroids of the clusters. A hierarchical clustering analysis (HCA) was performed based on Euclidean distances observed among the metabolomes. The differences

in the patient clusters identified by HCA were tested using the chisquare test and p values were reported. The participant demographics, features of the cervicovaginal environment, genital inflammation and pH of the patient clusters were compared using Mann-Whitney's Utest and p values were reported. PCA and HCA were performed using metabolome composition of 475 metabolites that were identified by Metabolon Inc.

#### 2.6. Bioinformatic analysis to integrate metabolomes with microbiomes

Quantitative Insights into Microbial Ecology (QIIME) v1.9 pipeline [24] was used for preparing 16S rRNA gene sequences for metagenome predictions. Operational Taxonomic Units (OTUs) with 97% sequence similarity were picked using a closed OTU picking method with Uclust [25] against the Greengenes database [26]. OTUs that were normalized based on gene copy numbers using QIIME v1.9 [24] were used for prediction of the metagenomes using the Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) [27]. Relative contribution of the human and microbial genomes to the expression of metabolites detected were calculated using Annotation of Metabolite Origins via Networks (AMON) [28]. Briefly, using the cervicovaginal metabolome datasets, host genome and microbial genomes downloaded from the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (https://www.genome.jp/kegg/), and PICRUSt predicted metagenomes, we revealed microbial and host contribution to the metabolites. Homo sapiens (has) was used for the host genome and Sneathia amnii (sns), Prevotella denticola (dns), Gardnerella vaginalis ATCC 1409 (gdv), Atopobium parvulum (apv), Streptococcus sanguinis (sng), Lactobacillus crispatus (lcr) were used for the microbial genomes to represent the six most abundant bacteria observed in our dataset. Model-based Integration of Metabolite Observations and Species Abundances (MIMOSA) [29] was used to reveal the potential contribution of microbial taxa to the expression of metabolites that distinguish groups. For the MIMOSA analysis, PICRUSt-predicted metagenomes and the metabolome datasets were used. MIMOSA utilizes a predicted relative metabolic turnover (PRMT) model to estimate gene abundances from PICRUSt-predicted metagenome data to produce community-wide metabolic potential (CMP) scores for each metabolite and sample using metabolites observed, predicted metagenomes and datasets from KEGG database including genomes, reactions, and pathways. Comparing CMP scores to metabolite variations, the metabolites were classified as well predicted by certain taxa or anti-predicted. For the visualization of the dataset, only well-predicted metabolites with false discovery rate < 0.1 were presented.

#### 2.7. Data and software availability

16S rRNA gene sequences were deposited in the National Center for Biotechnology Information (NCBI) database on BioProject accession number: PRJNA518153 and under Sequence Read Archive (SRA) accession IDs SAMN10855179 to SAMN10855255. De-identified metabolomics data is available in Supplementary files.

#### 3. Results

3.1. Cervicovaginal metabolite diversity was reduced in Ctrl HPV (+) and cervical dysplasia groups whereas diversity was increased in cervical cancer patients compared to Ctrl HPV (-) controls

We characterized cervicovaginal metabolomes in premenopausal non-pregnant healthy HPV-negative individuals [Ctrl HPV (-)] (n=18), healthy HPV-positive individuals [Ctrl HPV (+)] (n=11), women with low-grade squamous intraepithelial lesions (LSIL) (n=12), high-grade squamous intraepithelial lesions (HSIL) (n=27), and newly diagnosed ICC (n=10) using a global metabolomics approach. The patient characteristics and demographic information are

summarized in Supplementary Table 1. Participants were assigned to the patient groups based on histology of colposcopy-directed biopsy samples that were evaluated by a pathologist. If histology was not available, cytology results were evaluated. As described in our previous publication [6], a two-tiered system of cervical intraepithelial neoplasia (CIN) was used: CIN1 and CIN2/3 were categorized as LSIL and HSIL, respectively. Our participant group consisted of 37 Hispanic and 41 non-Hispanic women and there was no significant difference in race/ethnicity among the patient groups (Fisher's exact test, p = .15). Sixty-eight percent of the participants had body mass index (BMI) >25, however BMI did not significantly differ among the groups (ANOVA, p = .47). The mean age across the groups was 38  $\pm$  8 years and age distribution among the groups was not significantly different (ANOVA, p = .46). Vaginal pH was significantly different among the groups, pH 4.9  $\pm$  0.6 for Ctrl HPV (-), pH 5.7  $\pm$  1.09 for Ctrl HPV (+), pH 5.2  $\pm$  0.6 for LSIL, pH 5.8  $\pm$  0.9 for HSIL and pH 6.6  $\pm$  0.9 for ICC (mean  $\pm$  standard deviation, 5.6  $\pm$  0.9) (ANOVA, p = .003). Predominant HPV genotypes in our study, determined with Linear Array HPV Genotyping Tests (Roche) on DNA extracts from vaginal swabs, were HPV 16 (n = 43), HPV 31 (n = 13), and HPV 45 (n = 13). Fifty seven out of 60 HPV (+) participants [Ctrl HPV (+), LSIL, HSIL, and ICC] were positive for multiple HPV genotypes.

Our metabolome dataset was composed of 475 compounds of known identity from eight different metabolic pathways; 127 lipid, 147 amino acid, 34 nucleotide, 82 xenobiotic, 17 cofactor and vitamin, 10 energy, 24 carbohydrate, 32 peptide, and 2 partially-characterized molecules. Principal component analysis (PCA) of cervicovaginal lavages (CVLs) illustrated that global metabolomic profiles of ICC patients compared to other patient groups were distinct on principal component 1 (PC1) and PC2, which explained 46.57% of variation in the dataset (Fig. 1A; MANOVA, Wilk's Lambda test, p = .006; Mann-Whitney U test, p = .0009 and p = .0225, for PC1 and PC2, respectively). Due to the complex nature of the cervicovaginal lavages, the unexplained variation (53.43%) can be accounted for by the heterogeneity in participant genetic background, diet, lifestyle, etc. However, other collected participant metadata, for example, age, ethnicity, and BMI did not have a statistically significant effect on clustering (MANOVA  $p \ge .05$ , Wilk's Lambda test, Supplementary Fig. 1). Samples from the HSIL group also formed a cluster that was significantly distinct from the Ctrl HPV (-)group samples, but only on PC1 (Mann-Whitney U test, p = .046). As shown in Fig. 1B, differences in the metabolic profiles were driven by the number of different metabolites observed in the groups (Mann-Whitney U test, p = .001). ICC patients had a significantly greater number of metabolites (378  $\pm$  36) than the Ctrl HPV (+) (326  $\pm$  33, p=.037), LSIL (324  $\pm$  45, p = .022), and HSIL (316  $\pm$  43, p = .0009) groups, but not the Ctrl HPV (-) group (348  $\pm$  36, p= .363). Overall, Ctrl HPV (+), LSIL, and HSIL groups had a lower number of metabolites in the CVLs compared to Ctrl HPV (—) group; however, this finding did not reach significance possibly due to the limited number of samples per group. However, when we performed a trend analysis, we found that due to a reduction in the number of metabolites present in samples from HPV positive participants with and without cervical dysplasia [Ctrl HPV (+), LSIL, and HSIL], and an increase in the number of metabolites in ICC, the trend was significantly non-linear (p = .0004). Based on curve analysis, we found that the metabolite diversity trend was polynomial (p = .01).

In order to evaluate similarities and dissimilarities in the datasets, we compared the number of shared and unique metabolites among the groups. Patient groups shared 86% of the identified metabolites (426/475), and the ICC group had the greatest number of unique metabolites (n=12) (Fig. 1C). The unique metabolites from ICC belonged to lipid and xenobiotic super-pathways, especially from xenobiotic/drug metabolism (See Supplementary Table 2). When we visualized significantly enriched or depleted metabolites on Cytoscape networks and performed pairwise comparisons, we observed a significant enrichment of metabolites in lipid, xenobiotics, and carbohydrate super-pathways

in the ICC group compared to Ctrl HPV (-), Ctrl HPV (+), LSIL, and HSIL groups. Additionally, the ICC group had an enrichment of amino acid metabolites in comparison to other groups that were HPV positive with and without cervical dysplasia [Ctrl HPV (+), LSIL, and HSIL] (Fig. 1D). Compared to the Ctrl HPV (-) group, Ctrl HPV (+), LSIL, and HSIL groups showed a depletion in metabolites mainly in the amino acid super-pathway. Interestingly, the sub-pathways under the amino acid super-pathway and the level of depletion was different for Ctrl HPV (+), LSIL, and HSIL groups when compared to the Ctrl HPV (-) group. For example, Ctrl HPV (+) participants had metabolites mainly depleted in lysine, polyamine, and phenylalanine metabolism, while LSIL participants had metabolites depleted in histidine and the urea cycle. The HSIL group had depletions in many amino acid metabolites; however, the depletions were at a lower magnitude. Additionally, all the dipeptide metabolites were depleted in the HSIL group, which might explain the differences observed on PCA (Fig. 1A). This comparative analysis revealed that HPV infection, cervical dysplasia, and ICC had a divergent impact on cervicovaginal metabolomes.

3.2. HPV infection depleted metabolites in specific amino acid and nucleotide super pathways and ICC enriched metabolites from multiple lipid pathways

In order to explore the potential effect of HPV infection on the cervicovaginal metabolome, we performed receiver operating characteristics (ROC) analysis using samples from Ctrl HPV (-) and Ctrl HPV (+) individuals. The analysis did not reveal metabolites that discriminate Ctrl HPV (+) samples from the Ctrl HPV (-) samples (AUC < 0.8). However, when Ctrl HPV (-) samples were compared to Ctrl HPV (+) samples, we observed three amino acids (*N*-acetyltaurine, N6-acetyllysine, C-glycosyltryptophan), two nucleotides (guanine and urate), phosphate and deoxycarnitine had discriminatory ability with AUC values >0.8 as shown in Fig. 2A, suggesting a significant depletion of these metabolites in the Ctrl HPV (+) group. Indeed, the concentrations of these metabolites were significantly lower in the Ctrl HPV (+)group compared to Ctrl HPV (-) and ICC groups (Supplementary Table 3). When we performed ROC analysis to identify metabolites that discriminate Ctrl HPV (-) from LSIL, we found three amino acid metabolism products (pentose acid, 1-methyl-4-imidazoleacetate, pyroglutamine), one nucleotide product (1-methylhypoxanthine), and a xenobiotic (tartrate) as discriminators, as shown in Fig. 2B. Interestingly, the discriminators of Ctrl HPV (-) from HSIL were not the same metabolites as identified for LSIL. Only a phospholipid (phosphoethanolamine) and amino acid (glutamine) had >0.8 AUC (Fig. 2C). In summary, HPV infection and cervical dysplasia led to depletion of metabolites from certain amino acid and nucleotide metabolisms. These amino acid and nucleotide metabolites distinguished Ctrl HPV (-) group from the Ctrl HPV (+) and cervical dysplasia (LSIL and HSIL) groups.

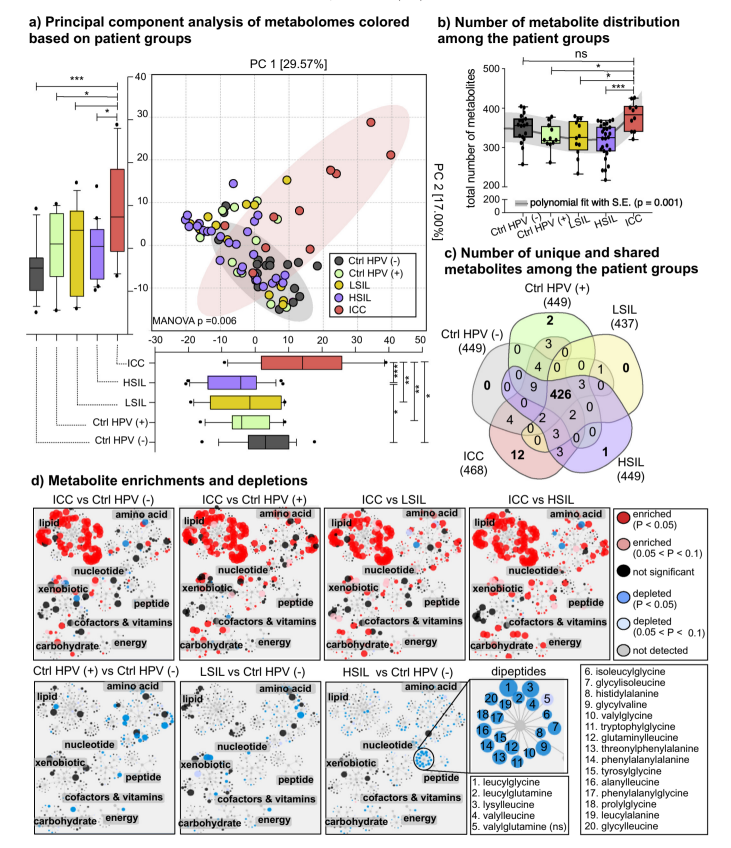
Since PCA analysis showed distinct metabolomes of ICC patients compared to the other groups, we next investigated the signatures of ICC using ROC analysis. As illustrated in Fig. 2D, 50 metabolites were either equal to or greater than AUC value of 0.80 indicating ICC significantly changes the expression of metabolomes in the cervicovaginal microenvironment and these metabolites can accurately distinguish participants with ICC from the Ctrl HPV (-). When ICC was compared to Ctrl HPV (+), most of the metabolites were similar to those that discriminated ICC from Ctrl HPV (-). Interestingly, many of the metabolites discriminated ICC group from the Ctrl HPV (-) group belonged to the lipid super-pathway (Fig. 2D). These metabolites were mainly long chain fatty acids, ketone bodies, steroids, ceramides, and plasmalogens. Among the detected molecules as discriminators of ICC, oleate/vaccinate, eicosenoate, and 3-hydroxybutyrate had AUC values >0.9, which provided more accurate discriminator and better predictive estimates than any other metabolite that was detected. As visualized in Fig. 2D, the abundance of these metabolites in the ICC group was not only significantly greater than the Ctrl HPV (-) and Ctrl HPV (+) groups, but also from the LSIL and HSIL groups (q < 0.05). When we calculated the AUC based on ICC compared to HSIL group, 71 metabolites had AUC >0.80 that distinguished ICC from HSIL. Similar to ICC and HPV comparisons, the majority of the metabolites were long chain saturated fatty acids, ketone bodies, plasmalogens, ceramides, and sphingomyelins (See Supplementary Table 3 and Supplementary Table 4). Overall, ROC analysis showed that ICC patients harbour elevated levels of lipid metabolites from different subclasses that can accurately distinguish them from all the other groups. Furthermore, those signature molecules did not show any gradation among the Ctrl HPV (+) and dysplasia groups when compared to Ctrl HPV (-) group (See Supplementary Table 3).

3.3. Microbially produced metabolites explained the dynamic shifts observed in the peptide and amino acid metabolisms

Microorganisms colonizing the cervicovaginal microenvironment produce a broad range of metabolites that are important for cervicovaginal health. Therefore, we investigated whether differences in the vaginal microbiota, in connection to the severity of cervical neoplasm, explain variations observed in the metabolome dataset. When we colour coded samples on PCA plots based on microbiota composition, metabolomes from patients with Lactobacillus-dominated (LD) vaginal microbiota formed a cluster significantly different than the metabolomes from patients with non-Lactobacillus dominance (NLD) (Fig. 3A). In accordance with the Lactobacillus dominance, vaginal pH revealed a similar clustering pattern on PCA (MANOVA, Wilk's Lambda test, *p* < .001, Supplementary Fig. 2). Grouping based on *Lactobacillus* dominance showed significance when PC1 and PC2 were analysed together (MANOVA, Wilk's Lambda test, p < .001). The difference was more apparent on PC2 (Mann-Whitney U test, p = .002), which explained 17% of the variation in the dataset. This observation provided further proof that vaginal microbiomes significantly modulate cervicovaginal metabolomes.

Considering that metabolomes of patients with LD microbiomes diverged from metabolomes of patients with NLD microbiomes, we examined their impact on metabolic super-pathways. Fig. 3B shows metabolic networks visualizing NLD vs LD metabolomes using Cytoscape. Many metabolites from lipid, amino acid, and nucleotide pathways were significantly enriched in the NLD metabolomes compared to LD. Additionally, all metabolites from dipeptide metabolism were significantly depleted, similar to our observation in the HSIL group compared to Ctrl HPV (—) group (Fig. 1D).

We also evaluated the origins of the cervicovaginal metabolites using Annotation of Metabolite Origins via Networks (AMON) software [28]. Based on PICRUSt metagenome predictions and quantified metabolites in our dataset that have Kyoto Encyclopedia of Genes and Genomes (KEGG) identifiers, 48% of the identified metabolites were predicted to be produced by the cervicovaginal microorganisms, which is slightly less than the metabolites that can be produced by the human metabolism (54.4%) (Fig. 3C). We found that 39.1% of the metabolites could not be explained by the host or the vaginal microbiota. We did not observe any differences in number of only microbiallyproduced metabolites among Ctrl HPV (-), Ctrl HPV (+), LSIL, HSIL, and ICC groups, since this analysis strictly depends on presence or absence of microbial phylotypes rather than their abundances. We also investigated the individual contribution of the most abundant microbial genera detected in our dataset: Lactobacillus, Prevotella, Sneathia, Atopobium, Streptococcus, and Gardnerella [6]. Microbial genomes downloaded from KEGG showed that Prevotella was the genus capable of producing most of the metabolites in our dataset (Fig. 3D) (n = 60; 23% of metabolites detected and exist in the KEGG database) and Gardnerella was the genus capable of producing the least number of metabolites (n = 38; 14% of metabolites detected and exist in the KEGG database) compared to the other genera tested. The metabolites predicted



to be produced by the most abundant bacteria were mainly amino acids, nucleic acids, or their metabolic end-products (Fig. 3D). Microbial contribution analysis did not explain the differences observed in lipid metabolism when LD and NLD groups were compared. Samples that belong to the NLD group are composed of mainly cervical dysplasia

and ICC patients, hence enrichments in the lipid metabolism in NLD could be coincidental.

In order to mechanistically associate cervicovaginal metabolites in our dataset to microbiome composition, we performed Model Based Integration of Metabolite Observations and Species Abundances

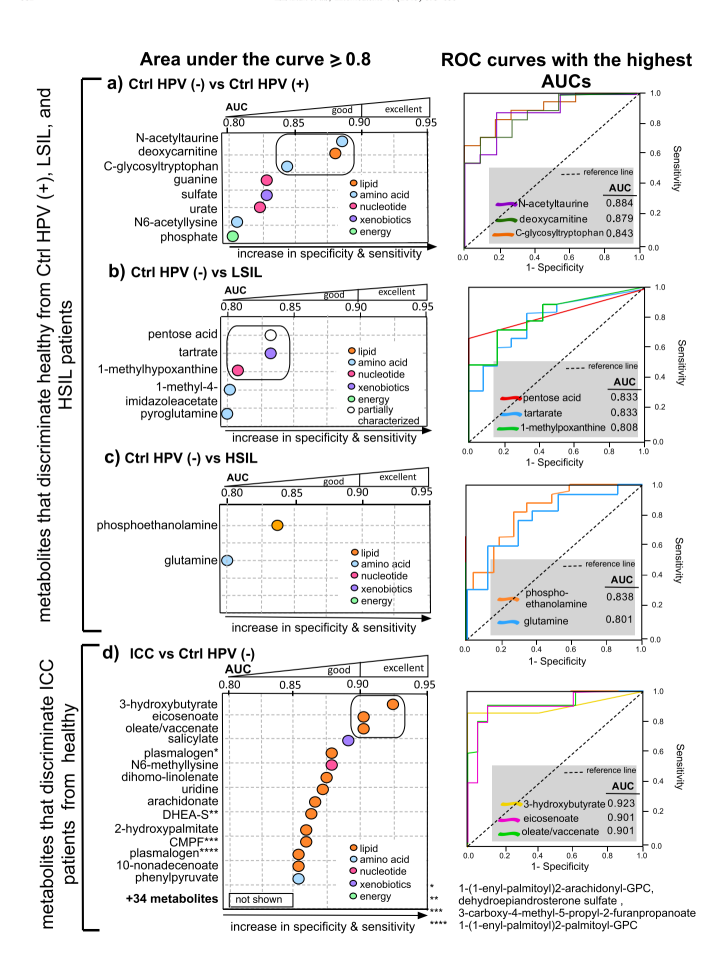
(MIMOSA) [29] analysis instead of correlation analysis which could lead to misinterpretation of the microbe-metabolite interactions. Correlation analysis between microbial taxa and host-associated metabolomes were found to lead to a poor explanation of the interactions and inaccurate interpretations [30]. Based on MIMOSA analyses, illustrated in Fig. 3E, of 127 metabolites that were predicted to be produced by the microbiota, 36 of them were significantly consistent with the metabolic potential (FDR < 0.1). As expected, mainly amino acids and amino acid degradation products, such as polyamines, were well-predicted to be modulated by the microorganisms. This model revealed that amino acids and their degradation products were well-predicted, and there were shifts in the products that can be produced by BV-associated microorganisms in comparison to Lactobacillus species. Lactobacillus species were the sole predictors of a secondary bile acid (glycochenodeoxycholate), amino acid (L-proline), and an amino acid product (glutathione disulphide), molecules that exert beneficial effects on the host based on metabolite fluxes. On the other hand, BVassociated bacteria explained the metabolism of specific metabolites. Atopobium contributed to the metabolism of L-threonine, adenosine, imidazole-4 acetate, glutathione disulphide, orthophosphate, L-lysine, 4-acetamidobutanoate, methylmalonate, and succinate. Gardnerella was the main contributor to the hippurate metabolism, NAD+ flux was mainly explained by the presence of Sneathia in our data set. Streptococcus contributed to 2,3-dihydroxy-3-methylbutanoate, a secondary metabolite in leucine, isoleucine, and valine metabolism, alphaisopropylmalate, and cytosine. Interestingly, non-abundant genera including Dialister, Roseomonas, Staphylococcus, and Shuttleworthia were predicted to contribute to metabolism of a diverse set of metabolites as orthophosphate, adenosine, L-lysine, methylmalonate, and imidazole-4 acetate (Fig. 3E). Overall, our comprehensive analysis of microbiome-metabolome relationships indicated that displacement of LD with NLD potentially induced various shifts in the amino acid metabolism that can be mostly explained by six most abundant genera detected in our dataset.

### 3.4. Lipid and xenobiotic metabolites were highly associated with genital inflammation

Since inflammation is a critical component of tumour progression, we investigated the link between cervicovaginal metabolomes and genital inflammation. Genital inflammation scoring strategy was previously reported [6,9,31,32]. Briefly, expression levels of seven cytokines (IL-1 $\alpha$ , IL-1 $\beta$ , IL-8, MIP-1 $\beta$ , CCL20, RANTES, and TNF $\alpha$ ) were evaluated in CVLs and the patients were assigned a cumulative score based on whether the level of each cytokine was in the upper quartile. Patients that scored five to seven were considered to have high genital inflammation, whereas one to five was assigned low genital inflammation and patients with a score of 0 were categorized as no inflammation. We overlaid genital inflammation scores on metabolomes and visualized on PCA as shown in Fig. 4A. Based on distances visualized on PCA, samples showed a gradation based on the genital inflammation category. Patients in the high genital inflammation category formed a cluster significantly distant from the none and low genital inflammation groups on both PC1 (Mann-Whitney *U* test, *p* values 0.0004 and 0.0143, respectively) and PC2 (p < .0001 and p = .0016, respectively). Based on MANOVA results, genital inflammation led to significant grouping on PCA (Wilk's Lambda p < .001). The difference between the low and no genital inflammation groups on PC2 was also significant (Mann-Whitney U test, p = .0228). Additionally, as indicated in Fig. 4A, the samples with high genital inflammation were mainly ICC patients (6/10). When we performed correlation analysis, 32% (187/575) of the metabolites significantly correlated with genital inflammation scores with Spearman's rho greater than the critical value (0.23 for n = 78, and bootstrapped p <.05). About 29% (168/575) and 3% (19/575) of the metabolites showed positive and negative correlation with genital inflammation, respectively (Fig. 4B). Metabolites that positively correlated with genital inflammation were mainly from lipids (n = 99) followed by amino acids (n = 23), nucleotides (n = 14) and xenobiotics (n = 11). Lipids that positively correlated ( $\rho > 0.68$ , p < .05) with genital inflammation were sphingolipids, plasmalogens, phosphatidylcholines, phospatidylethanolamines and long-chain-poly-unsaturated fatty acids (Fig. 4B, Supplementary Table 4), which are essential components of host and bacterial cellular membranes. The relative concentrations of the metabolites from these five lipid sub-families were significantly higher in the ICC group compared to other groups as shown in Fig. 4C. These data strengthen the link between genital inflammation, cancer, and lipids. The correlation between genital inflammation and lipids was stronger ( $\rho > 0$ . 74 for the first 20 highest ranked correlations) than the correlation between other metabolites and genital inflammation (Supplementary Table 4). Most metabolites that negatively correlated with genital inflammation were xenobiotics (n = 11), amino acids (n = 4), and nucleotides (n = 2).

Since genital inflammation is linked to vaginal microbial community composition, we investigated the relationship between the metabolites that correlated with genital inflammation and Lactobacillus abundance. Fig. 5A illustrates Spearman's rho coefficients between the metabolites and genital inflammation versus metabolites and Lactobacillus abundance. Many of the lipids that highly correlated with genital inflammation did not correlate with Lactobacillus abundance. However, one of the sphingomyelins, linoleate, glycochenodeoxycholate and deoxycarnitine were negatively correlated with Lactobacillus abundance. Glutathione synthesis intermediate, 2-hydroxybutyrate, branched chain amino acid metabolism product, alpha-hydroxy-isovalerate, and L-isoleucine metabolism product, 2-hydroxy-3-methyl-valerate moderately and positively correlated with genital inflammation and negatively correlated with Lactobacillus abundance. These amino acid metabolites were significantly higher in the ICC group compared to other groups (Welch's two-sided *t*-test, q < 0.05, Fig. 5B). Metabolites that discriminated ICC from the Ctrl HPV (-), eicosenoate, 3-hydroxybutyrate, and oleate/vaccinate highly correlated with genital inflammation (Fig. 5A). Metabolites that positively correlated with Lactobacillus abundance and genital inflammation included amino acids/products such as arginine and betaine, a polyamine, N(1)-acetylspermine, and lipids mainly involved in carnitine metabolism such as decanoylcarnitine, hexanoylcarnitine, and acetylcarnitine. The levels of these metabolites were depleted in the Ctrl HPV (+), LSIL, and HSIL groups compared to Ctrl HPV (—) and ICC groups as shown in Fig. 5C and E. There were only a few metabolites that negatively correlated with genital

**Fig. 1.** HPV infection, dysplasia, and invasive cervical carcinoma (ICC) divergently impacted the cervicovaginal metabolic profile. A) Principal component analysis (PCA) showed a distinct cervicovaginal metabolome profile in ICC patients compared to Ctrl HPV (-), Ctrl HPV (+), LSIL, and HSIL patient groups. The difference was significant on both PC1 and PC2 axes. HSIL group was significantly different than Ctrl HPV (-) group on PC2 axis. Boxplots represent median, first and third quartite, minimum and maximum values in the dataset. Shaded ellipses represent 95% confidence intervals of the cluster centroids. Mann-Whitney U test p values represented as \*p < .05, \*\*p < .01, \*\*\*p < .001. B) Number of metabolites detected in each group. ICC patients had significantly greater number of metabolites in comparison to patients in the Ctrl HPV <math>(+), LSIL, and HSIL groups. However, there was no significant difference between the ICC and Ctrl HPV (-) groups. n = n or significant. The trend analysis showed that the trend was polynomial (p = .01). C) Number of unique and shared metabolites among the groups visualized on a Venn diagram. Majority of the metabolites were detected in the all groups. ICC group had the greatest number of metabolites that were not detected in any of the participants in other groups. Supplementary Table 2 contains the list of metabolites that are different in each group. D) Enrichment and depletions of metabolites among patient groups visualized by Cytoscape metabolic network analysis. The node size is proportional to the magnitude of differences observed among the groups. Red and blue nodes represent enriched and depleted metabolites, respectively. In comparison to all other patient groups, ICC patients had an enrichment of metabolites that belong to lipid, amino acid, carbohydrate, and xenobiotic metabolities. Amino acids and their metabolites were depleted in Ctrl HPV (+), LSIL, and HSIL groups. Dipeptides were significantly depleted in the HSIL group compared to Ctrl HPV (-



inflammation but positively correlated with *Lactobacillus* abundance. These metabolites included nucleotides adenosine and cytosine and xenobiotics such as 2-keto-3-deoxy-gluconate and 1,2,3-benzenetriol. The relative levels of these nucleotides or xenobiotics were lower in the ICC group compared to other groups and the difference in cytosine levels between Ctrl HPV (-) and ICC group and 2-keto-3-deoxy-gluconate between HSIL and ICC were statistically significant (Welch's two-sided ttest, q < 0.05) (Fig. 5D).

We also observed many biogenic amines (cadaverine, putrescine, tyramine, tryptamine, and agmatine) negatively correlated with Lactobacillus abundance as predicted but did not correlate with genital inflammation. In general, levels of the biogenic amines did not significantly differ among the groups (See Supplementary Fig. 3). Additionally, lactate levels strongly and positively correlated with Lactobacillus abundance. Interestingly, non-Lactobacillus dominated HSIL group had significantly lower levels of lactate compared to Ctrl HPV (-) and ICC groups. A major Gardnerella associated metabolite, hippurate, did not correlate with either genital inflammation or Lactobacillus abundance. A metabolite that was mainly predicted to be modulated by Sneathia was NAD+ based on MIMOSA analysis. Additionally, NAD+ levels positively correlated with *Lactobacillus* abundance. Interestingly, NAD+ levels were significantly lower in HSIL compared to Ctrl HPV (-). To better understand the associations independent from cancer, we performed the correlation analysis by excluding all ICC samples. As shown in Supplementary Fig. 4, we observed the same trend between metabolites that were associated with inflammation and Lactobacillus when the ICC samples were excluded. However, we observed minor differences in correlation coefficients of key metabolites, for example,  $\rho$  for lactate and *Lactobacillus* abundance was greater ( $\rho = 0.65$  versus  $\rho =$ 0.69) whereas  $\rho$  for alpha-hydroxy-isovalerate and *Lactobacillus* abundance was slightly smaller ( $\rho = -0.59$  versus  $\rho = -0.58$ ) when the ICC samples were excluded.

## 3.5. Metabolomes formed three distinct clusters mainly driven by cervical cancer, vaginal pH and microbial communities

Metabolites detected in the CVLs can be impacted by many biological variables including viruses, microbial communities, host-derived factors, and outcomes of host-microbe interactions including genital inflammation and vaginal pH. We performed unsupervised hierarchical clustering using all identified metabolites to reveal whether any of these covariates drive the similarities between the metabolomes. Based on Euclidean distances among the metabolomes, samples were assembled into three clusters as shown in Fig. 6A. Cluster 1 (n = 4) composed of only ICC patients, whereas Clusters 2 (n = 29) and Cluster 3 (n = 45) were not dominated by any particular participant group. The participant group composition of Cluster 1 was significantly different than Cluster 2 (Chi-Square test, p = .002) or Cluster 3 (chi-square test, p < .0001) (Fig. 6B). As ICC explained only Cluster 1, we evaluated other factors that might be defining Cluster 2 and 3. The genital inflammation score was greater in Cluster 1 compared to Clusters 2 and 3 as shown in Fig. 6C, which did not distinguish Cluster 2 from Cluster 3. Next, we evaluated *Lactobacillus* dominance. As in depicted in Fig. 6D, clusters had significantly different vaginal microbiota composition. Cluster 1 was solely dominated by NLD microbiota, whereas Cluster 2 was mainly consisted of participants with LD microbiota. Similar to Cluster 1, Cluster 3 mainly consisted of patients with NLD microbiota as well. Cluster 2 was dominated by Lactobacillus crispatus and Lactobacillus iners. On the other hand, Cluster 3 was dominated by polymicrobial communities followed by *L. iners.* Since vaginal pH and microbial composition are linked [6], as expected, vaginal pH of the patients in each cluster differed significantly as well. Individuals in Cluster 1, that was dominated by multiple non-*Lactobacillus* genera, had significantly higher vaginal pH compared to the individuals in Clusters 2 and 3. Cluster 2 was composed of participants with significantly lower vaginal pH (Fig. 6E) in accordance with *Lactobacillus* dominance, when compared to Cluster 1 (Mann-Whitney U test, p=.002) or Cluster 3 (Mann-Whitney U test, p=.046). Consistent with our previous observations, *Lactobacillus* dominance and vaginal pH better explained the clustering than the other variables. In summary, these results indicate that cancer, vaginal microbiota, and genital inflammation are robust drivers of cervicovaginal metabolomes.

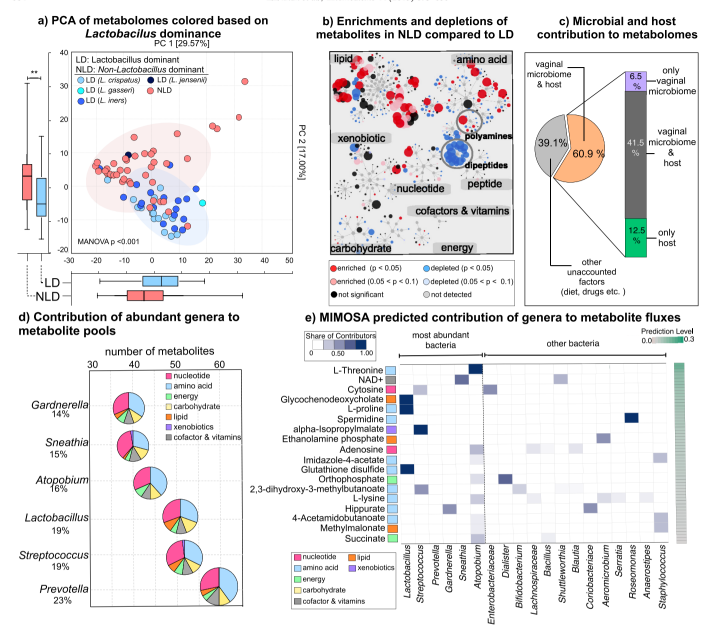
#### 4. Discussion

The metabolomic profiling offers a unique understanding of the complex interactions between the host and microbes in carcinogenesis. In addition, dysregulation of metabolism is an emerging hallmark of cancer [33]. A number of studies conducted on humans have shown a distinct metabolic fingerprint of cervical cancer in plasma [15], tumour tissues [16], faeces [17], and urine [34]. However, the impacts of HPV infection, cervical dysplasia and cancer on the metabolic profiles of the cervicovaginal microenvironment, a unique ecosystem where host epithelial and immune cells directly interact with viruses, bacteria and other environmental factors, have not been previously addressed. Unquestionably, metabolic fingerprinting of cervicovaginal lavages more accurately reflects the interactions between host and the microorganisms within the local microenvironment. Additionally, metabolic fingerprinting can provide insights into the interplay between persistent HPV infection, cervical neoplasm, and other features of the cervicovaginal microenvironment (i.e., genital inflammation, vaginal microbiome and vaginal pH) in the development of cervical dysplasia and progression to ICC. Here, we evaluated these metabolic interactions and characteristics of the cervicovaginal microenvironment (vaginal pH, vaginal microbiome, genital inflammation) in Ctrl HPV (-), Ctrl HPV (+), LSIL, HSIL, and newly diagnosed ICC patient groups. We show that unique metabolic profiles, impacted by genital inflammation, vaginal pH, and the vaginal microbiome, predict patient groups and reflect profound differences in the host and microbe co-metabolism during HPV infection, cervical dysplasia, and ICC.

Cervical cancer, dysplasia, and HPV infection exhibited divergent effects on the cervicovaginal metabolomes. A depletion of amino acid metabolites in HPV-positive and cervical dysplasia groups (LSIL and HSIL), and depletion of dipeptides in the HSIL group was replaced in ICC by enrichment of xenobiotics and lipids from plasmalogens, sphingomyelins, phosphatidylcholines, and long chain polyunsaturated fatty acids. Observation of xenobiotics, mainly anti-inflammatory and analgesic drugs such as ibuprofen, acetaminophen, nafloxin, and their metabolites in the CVLs of ICC patients indicated a possible translocation of these drugs from blood to the cervix and vagina. A higher abundance of lipid metabolites in ICC patients compared to healthy individuals can be explained by an increase in cell proliferation and cell membrane synthesis through activation of oncogenic pathways in the tumour microenvironment [35]. Hence, enrichment of lipids in the cervicovaginal microenvironment relevant to nutrient acquisition validated one of the emerging hallmarks of cancer metabolism [33].

Using a well-characterized and robust biomarker discovery analysis (ROC) [36], we identified three lipid molecules, 3-hydroxybutyrate,

**Fig. 2.** Multiple metabolites from diverse pathways robustly discriminated healthy, dysplasia, and ICC patient groups. The cut-off value for the Receiver Operating Characteristics (ROC) analysis was 0.8 to include only good  $(0.8 \le AUC < 0.9)$  and excellent  $(AUC \ge 0.9)$  biosignatures. A) ROC analysis comparing Ctrl HPV (-) to Ctrl HPV (+). ROC showed that amino acid, lipid, nucleotide, and xenobiotic metabolism products were indicators of absence of HPV infection. B) ROC comparing Ctrl HPV (-) to LSIL. Depletions in the amino acid metabolism products in LSIL discriminated Ctrl HPV (-) from LSIL. C) ROC comparing Ctrl HPV (-) to HSIL. Phosphoethanolamine and glutamine were discriminators of Ctrl HPV (-) from HSIL. D) ROC analysis performed between ICC and Ctrl HPV (-) groups revealed metabolites from different lipid classes with >0.9 area under the curve (AUC) values serve as strong discriminators of ICC. Those lipids were associated with cancer related inflammation. Supplementary Table 3 contains the median levels of the metabolites and Supplementary Table 4 contains the q values of Welch's t-test for group comparisons listed in this figure.



**Fig. 3.** Vaginal microbiota composition profoundly impacted amino acid and nucleotide metabolisms. A) Principal component analysis of metabolomes visualized based on *Lactobacillus* dominant (LD) and non-*Lactobacillus* dominant (NLD) metabolomes were significantly different on PC2. Boxplots represent median, first and third quartile, minimum and maximum values in the dataset. Shaded ellipses represent 95% confidence intervals of the cluster centroids. Mann-Whitney U test p values represented as \*p < .05, \*\*p < .01, \*\*\*p < .001. B) Metabolic enrichments and depletions in NLD vs LD visualized on Cytoscape networks. Metabolomes of patients that had NLD vaginal microbial communities had enrichments in lipid and amino acid metabolism. Dipeptides were depleted in patients with NLD communities. C) Prediction of vaginal microbiota or host origins of the metabolites from polyamines were enriched whereas some were 60% of the detected metabolites in cervicovaginal lavages in all groups. D) The six most abundant genera within the data set were predicted to contribute 14–23% of the metabolites that were detected. The microbially produced metabolites mainly belonged to amino acid and nucleotide metabolism. E) Well-predicted (FDR < 0.1) metabolites based on MIMOSA analysis and relative contribution of microbial phylotypes to the production or depletion of metabolites. The six most dominant genera within our dataset explained the observed concentrations of many amino acids and their metabolic end-products.

eicosenoate, and oleate/vaccenate with excellent discrimination capacity (AUC > 0.9) of ICC patients from healthy participants. A ketone body, 3-hydroxybutyrate, was elevated in ovarian cancer and was associated with invasion and migration of cancer cells [37,38]. Additionally, 3-hydroxybutyrate augments tumour growth in-vitro epithelial cell models [39]. A systematic review of 23 metabolome studies in ovarian cancer highlighted abnormally high levels of lipids including, oleate/vaccenate and eicosenoate in the serum [40], an observation consistent with the metabolic profiles we observed in the CVLs of ICC patients. In addition, elevated levels of oleic acid have been previously observed in cervical cancer biopsies [41] and oleic acid has been shown to promote cervical cancer cell growth in-vitro [42], thus validating our findings.

Interestingly, these markers are not only specific to cancer, but also linked to inflammation and obesity, two comorbidities related to cancer [43]. Genital inflammation and high BMI are also characteristics of our patients although BMI did not significantly differ among the patient groups.

ROC analysis also revealed that metabolites from multiple metabolic pathways were significantly depleted in HPV positive and cervical dysplasia groups. Interestingly, we observed unique discriminators that distinguished the Ctrl HPV (-) group from the Ctrl HPV (+), LSIL or HSIL groups. Depletion of metabolites from taurine, glutamine and lysine metabolism, key amino acids that are involved in cellular growth [44], in HPV-positive women and cervical dysplasia patients indicates

#### a) PCA of metabolomes colored based on genital b) Number of metabolites correlated with genital inflammation inflammation Negative correlation Positive correlation I CPUFA Sphingomyelins $(\rho = 0.70)$ $(\rho = 0.68)$ (3% of the metabolites) (29% of the metabolites) PC 1 [29.57%] genital inflammation peptide none (0) energy , PE cofactor and vitamins (p = 0.69) Olow (1-4) 30 carbohydrate high (5-7) xenobiotics $(\rho = 0.69)$ nucleotide 20 amino acid lipid Plasmalogens = 0.72) 20 10 0 10 20 30 40 50 60 70 10 c) Relative levels of lipids that were associed with genital inflammation 0 Sphingomyelins Plasmalogens PC LCPUFA PΕ -10 indicates samples from MANOVA p < 0.001 ICC patients -20 log10 transform relative peak a -10 -30 -20 10 20 30 40 50 low T\* hiah

**Fig. 4.** Levels of genital inflammation highly correlated with metabolite profiles and patient groups. A) Principal component analysis (PCA) demonstrated a gradual separation of metabolomes based on presence or intensity of genital inflammation. Metabolomes of patient with high genital inflammation score of 5–7) formed a separate cluster from patients without (inflammation score = 0) or low genital inflammation score of 1–4) on PC1 and PC2. Majority of the high genital inflammation samples belonged to ICC patients. Boxplots represent median, first and third quartile, minimum and maximum values in the dataset. Shaded ellipses represent 95% confidence intervals of the cluster centroids. Mann-Whitney U test p values represented as \*p < .05, \*p < .01, \*p < .01. B) Number of metabolites positively and negatively correlated with genital inflammation. Spearman's rho correlation coefficient greater than the critical value (0.23 for n = 78) with a bootstrapped p < .05 were considered significant. Metabolites from lipid metabolism positively correlated with genital inflammation whereas xenobiotics negatively correlated with the genital inflammation. Lipids that correlated with genital inflammation belonged to sphingomyelins, plasmalogens, phosphatidylcholines (PC), long-chain-polyunsaturated fatty acids, and phosp hatidylethanolamines (PE). Supplementary Table 5 includes the list of Spearman's correlation coefficients reflecting the strength of correlation between genital inflammation and metabolites. C) Relative levels of plasmalogens, sphingomyelins, phosphatidylcholines, phosphatidylcholamines, and long-chain polyunsaturated fatty acid metabolites were greater in the ICC group compared to the others. Boxplots represent median, first and third quartile, minimum and maximum values in the dataset. Welch's t-test p values represented as \*p < .05, \*\*p < .01, \*\*\*p < .001.

Ctrl HPV (-)

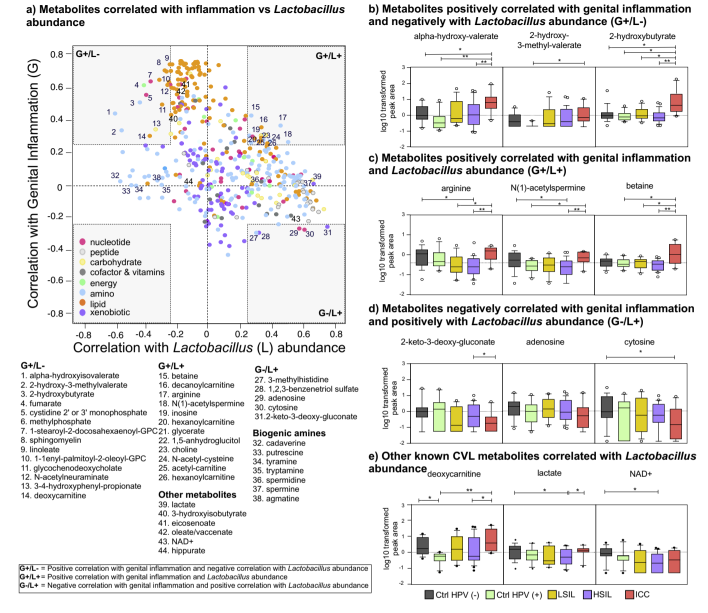
Ctrl HPV (+) LSIL

disruption of healthy cell metabolism as a functional consequence of potential HPV-mediated dysbiosis. Glutamine, a critical amino acid for tumour cell growth [33], and phosphoethanolamine, a well-established anti-tumour agent [45], were found to be strong discriminators of healthy patients vs. the HSIL patients. The impact of HPV on metabolite depletions serves as an example of changes in cervicovaginal microenvironment due to virus and host interactions.

Vaginal dysbiosis has emerged as a key risk factor in reproductive health [46], inflammation [47] and HPV acquisition, persistence, and cervical carcinogenesis [48]. However, the biological mechanisms and host-microbe interactions that drive persistence and carcinogenesis have not been elucidated. Thus, we integrated microbiome and metabolome data sets using the state-of-the-art bioinformatic tools (MIMOSA and AMON) to reveal a functional impact of microbiomes on metabolic fingerprints. Using a predicted relative metabolic turnover model on our measured community metabolome, we provide unique insights into the microbial contribution to observed metabolomes beyond correlations between microbial genera and metabolites. Our analysis based on microbial genomes indicated that differences in non-Lactobacillus dominated (NLD) communities compared to Lactobacillus-dominated (LD) communities specifically affect the flux of dipeptides, amino acid and nucleotide metabolites, which was consistent with a previous report that employed the same modelling technique on metabolomes in bacterial vaginosis (BV) [29]. Our NLD communities were enriched in multiple microbial genera including Gardnerella, Prevotella, Streptococcus and Atopobium, that were typically found in BV patients [13,49-51]. A decrease in the abundance of vaginal Lactobacillus species in BV patients was previously associated with an increase in the biogenic amine levels, amino acid degradation products and depletion of dipeptides [12,13,51,52]. Hence our findings agree with the observation that microbial diversity in the vaginal milieu alters amino acid metabolism.

Our study design excluded symptomatic BV patients, however we observed several metabolic signatures of BV in our dataset. For example, deoxycarnitine [12] and alpha-hydroxyisovalerate [52], previously identified as signatures of BV, were negatively correlated with Lactobacillus abundance. Interestingly, other acylcarnitines that are known proinflammatory cytokine inducers [53] were significantly associated with genital inflammation and Lactobacillus abundance. We also observed BV-associated Gardnerella was the main contributor to hippurate metabolism. Hippurate degradation is not only used as a test for Gardnerella identification [54], but also used in pathogen detection. Another important metabolic signature of BV is presence of biogenic amines. Three out of ten polyamines were significantly enriched in the NLD metabolomes and five BV-associated biogenic amines (cadaverine, putrescine, tyramine, tryptamine, and agmatine) [13,55] negatively correlated with Lactobacillus abundance, however their levels were not significantly different between LD and NLD communities. Notably, none of these biogenic amines were positively or negatively associated with genital inflammation. Arginine and its polyamine product N(1)-acetylspermine are indicative of rapidly proliferating cells [56] and positively correlated with Lactobacillus abundance and genital inflammation. Previously, arginine metabolism was previously highlighted as an emerging hallmark of cancer [33]. Interestingly, N(1)-acetylspermine levels are increased in severely inflamed sites in colorectal cancer patients [57], indicating a potential role of this polyamine in severe inflammation in accordance with our genital inflammation findings. However, other BV-associated biogenic amines did not correlate with genital inflammation. Metabolites that are precursors of biogenic amines including choline and betaine have anti-inflammatory properties [58,59]. Betaine enhances survival and provides protection to Lactobacillus species facing osmotic stress [60]. Future comparative studies that incorporate microbiomes to metabolomes in symptomatic and asymptomatic BV patients will

HSIL



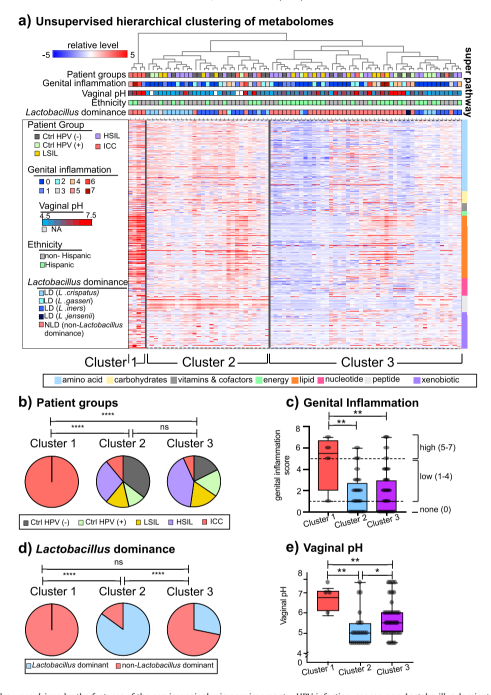
**Fig. 5.** A) Metabolites that connected genital inflammation to *Lactobacillus* abundance. Scatter plot of metabolites that correlated with genital inflammation and *Lactobacillus* abundance. Lipids (plasmalogens, sphingomyelin, phosphatidylcholine, and long chain polyunsaturated fatty acids (LCPUFA)) strongly and positively correlated with genital inflammation but not with *Lactobacillus* abundance. A few amino acids negatively correlated with *Lactobacillus* abundance positively correlated with genital inflammation. Carnitine metabolites positively correlated with both *Lactobacillus* abundance and genital inflammation. Biogenic amines negatively correlated with *Lactobacillus* abundance but did not correlate with genital inflammation. B) Relative levels of metabolites that positively correlated with genital inflammation among the patient groups. The concentrations were relatively higher in the ICC group. C) Relative levels of metabolites that positively correlated with genital inflammation and *Lactobacillus* abundance among the patient groups. The concentrations were relatively higher in the ICC group. D) Relative levels of metabolites that negatively correlated with genital inflammation and positively with *Lactobacillus* abundance among the patient groups. The concentrations were relatively lower in the ICC group. Metabolites with the three highest ρ values were visualized on graphs B, C, and D. E) Relative levels of metabolites that had significant associations with *Lactobacillus*-dominance or dysbiotic microbiota. Welch's t-test *p* values represented as \*p < .05, \*\*p < .01, \*\*\*p < .001.

further enhance the understanding of the role of microbes and microbehost interactions in the context of vaginal dysbiosis and BV.

Our analysis using a well-characterized prediction model (MIMOSA) also revealed that *Lactobacillus* species are the key contributors to the metabolism of glycochenodeoxycholate (GCDC) in our dataset. GCDC, a key metabolic product of host-microbe co-metabolism [61], can inhibit several pathogens that are also observed in vaginal dysbiosis and BV [62]. Similar to other bile acids, GCDC induces inflammatory and toxic responses and exhibits carcinogenic effects on host epithelial cells [63]. Our observation of negative correlation with *Lactobacillus* abundance and positive correlation with genital inflammation further strengthened the link between GCDC, inflammation, and *Lactobacillus* 

abundance. As such, future studies are needed to investigate the role of GCDC in maintaining *Lactobacillus*-mediated homeostasis, as well as pathogen exclusion in the lower reproductive tract.

It is important to note that the microbiome-based data integration was limited only to the metabolites and genomes available in the KEGG database. Overall, differences in the vaginal microbial communities in cervical dysplasia and ICC were reflected upon metabolomes. Microbe-microbe and host-microbe interactions through metabolic networks can shape the local tumour environment [64], hence dysbiotic vaginal microbiome-mediated metabolome has the potential to become a hallmark of cervical cancer. Additionally, the metabolites produced by the cancer microenvironment itself, including lipids, can enhance the



**Fig. 6.** Metabolome profiles were driven by the features of the cervicovaginal microenvironment – HPV infection, cancer, non-Lactobacillus dominated microbiota, and genital inflammation. A) Hierarchical clustering (HCA) dendrogram constructed using top-down approach presented with patient characteristics including HPV status, genital inflammation score, vaginal pH, and microbial community based on Lactobacillus abundance. HCA analysis revealed three main clusters. B) Distribution of patient groups in the clusters. Cluster 1 only contained ICC patients whereas Cluster 2 and 3 contained patients from all the groups. The patient group composition of Cluster 1 was significantly different than Cluster 2 based on chi-square test. C) Genital inflammation scores among the clusters. Genital inflammation score of Cluster 1 was significantly higher than the scores of Clusters 2 and 3. Boxplot plots demonstrate median, 25th and 75th quartiles. D) Percentage of samples belonging to patients with Lactobacillus dominant and non-Lactobacillus dominant microbiota. Cluster 1 and 3 were dominated by non-Lactobacillus dominated communities whereas Cluster 2 had greater number of samples that were Lactobacillus dominated. Based on chi-square test, Cluster 2 was significantly different than Cluster 3. E) Vaginal pH measurements among the clusters. Cluster 1 and 3 had samples from patients with higher vaginal pH compared to Cluster 2 patients. \*p < .5, \*\*p < .01, \*\*\*\*p < .0001.

crosstalk between microbiota and the host, and therefore the procarcinogenic activities of the microbiota.

Genital inflammation, another key feature of the cervicovaginal microenvironment, co-shaped the metabolic profiles along with the microbial composition in our dataset. Notably, the lipids that discriminated the ICC group from the healthy group showed a strong positive correlation to genital inflammation. However, our additional analysis (Supplementary Fig. 4) excluding ICC patients showed that lipids remained associated with genital inflammation independent from cancer. An

increase in lipid metabolites from plasmalogens and long chain polyunsaturated fatty acids in ICC, not only indicate abnormality in cellular metabolism, but also highlight the significance of lipids in cancer development, due to their roles as precursors of pro-inflammatory cytokines, inducers of abnormal gene expression and ability to dysregulate cytokine production [65]. Interestingly, phosphatidylcholines, another group of lipids that positively correlated with genital inflammation, are typically known for their anti-inflammatory properties [66], but also contribute to cellular proliferation and programmed cell death [67]. Adenosine and cytosine, nucleotides, that were well-predicted with MIMOSA analysis, negatively correlated with genital inflammation and positively correlated with *Lactobacillus* abundance. These well-recognized anti-inflammatory molecules [68] were also well-predicted to be modulated by the vaginal microbiome in our dataset. For instance, adenosine, an anti-inflammatory agent, has therapeutic potential because it inhibits phagocytosis, adhesion, and production of reactive oxygen species [69]. Additionally, animal and *in vitro* cell models demonstrated anti-inflammatory effects of adenosine through its impact on macrophages [70] and mast cells [71]. Taken together these studies indicate that the potent anti-inflammatory function of microbially modulated adenosine could be a molecule associated with genital tract health.

A key metabolite that strongly correlated with Lactobacillus abundance was lactate. When the correlation analysis was performed without the ICC patients included, the correlation between lactate and Lactobacillus was stronger and there was a trend towards negative correlation with genital inflammation (Supplementary Fig. 4). Lactic acid is the main end-product of Lactobacillus-mediated fermentation [72], and is a critical metabolite to maintain low vaginal pH (<4.5) and homeostasis in the vaginal microenvironment [73]. Higher levels of lactate and lower vaginal pH observed in Lactobacillus dominated communities compared to non-Lactobacillus-dominated communities are in agreement with other studies relating this molecule to vaginal health and homeostasis [12,73,74]. In a murine model, lactate was shown to inhibit the pro-inflammatory response of macrophages [75], hence our data highlight a potential anti-inflammatory role of lactate in the lower reproductive tract. Additionally, lactic acid can induce secretion of antiinflammatory IL-10, reduce the production of proinflammatory IL-12 in dendritic cells, and diminish the cytotoxicity of natural killer cells [76]. On the other hand, lactate is also one of the main cellular respiration products of cancer cells, previously established as a biomarker of cervical cancer tumours [77] and is associated with angiogenesis and immunotolerance [76]. Therefore, the dual role of lactate in health and disease, its origin, and relationship to genital inflammation in the context of gynaecologic cancer warrants further investigation.

Metabolic profiling of the cervicovaginal microenvironment offers three important benefits to cervical cancer research. First, it provides a snapshot of metabolic communications between the host, microbiome, and HPV. Correspondingly, metabolic profiling is the first step to understand host-microbiota interplay in modulating the hallmarks of cancer in the context of HPV. Second, CVL collection is non-invasive method that may provide more biological insights into the mechanisms of viral persistence and disease progression. Finally, the limitations and challenges in murine and non-human primate models are avoided since these models do not faithfully recapitulate the composition of vaginal microbiota in humans [78]. A limitation of our study is the relatively small sample size and future studies with larger sample sizes are needed to validate and extend our findings. However, the results presented herein generate additional questions on virus-host epithelia and immune system interactions that can be addressed by metabolic profiling in future clinical studies. In addition, studies employing physiologically relevant 3-dimensional human FRT models [78–81] could reveal biological mechanisms into how these microbes are dysregulating host metabolism and contributing to the hallmarks of cancer.

Multiple features of the cervicovaginal microenvironment, genital inflammation, vaginal pH, and vaginal microbiota composition explained differences in the metabolic profiles of the participant groups across cervical carcinogenesis. Our observations on genital inflammation- and microbiota-associated metabolic signatures connect the microbiome to inflammation, dysplasia, and cancer. Collectively, our analysis revealed that cancer had the greatest impact on metabolic profiles and genital inflammation, and vaginal microbiota had a secondary impact that explained metabolic profiles of healthy HPV-negative, HPV-positive individuals, and cervical dysplasia patients. Our analysis clearly demonstrated that metabolic fingerprinting of the cervicovaginal

microenvironment is an excellent approach to discriminate between individuals with HPV infection, cervical dysplasia and ICC, which may facilitate future discoveries of new therapeutic and prevention avenues for patients. This integrative analysis of the cervicovaginal metabolome, microbiome and immunoproteome revealed multiple emerging hallmarks of cancer in cervical carcinogenesis, including abnormalities in nitrogen and energy metabolism, increased cellular proliferation due to changes in amino acid and carbohydrate fluxes, and dysbiotic microbiota and host-microbe co-metabolism [33,82]. In conclusion, the complex virus-host-microbe interplay within the cervicovaginal microenvironment lead to unique metabolic fingerprints that may be exploited for future development of diagnostics, preventatives or therapeutics to positively impact women's health outcomes.

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#### **Declaration of interests**

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#### **Author contributions**

Conceptualization, M.M.H.K. and D.M.C.; Methodology, Z.E.I.; Formal Analysis, P.L., Z.E.I., N.T., Investigation, P.L., Z.E.I., Resources, D.M.C.; Data curation, Z.E.I., P.L., N.T.; Writing Original Draft, Z.E.I., P.L., and M.M.H.K.; Writing – Review & Editing, P.L., M.M.H.K., D.M.C., D.R., N.T.; Visualization, Z.E.I., P.L., M.M.H.K.; Supervision, M.M.H.K.; Project Administration, M.M.H.K.; Funding acquisition, D.M.C. and M.M.H.K.

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#### References

- Schiffman M, Doorbar J, Wentzensen N, De Sanjosé S, Fakhry C, Monk BJ, et al. Carcinogenic human papillomavirus infection. Nat Rev Dis Prim 2016;2.
- [2] Koshiol J, Lindsay L, Pimenta JM, Poole C, Jenkins D, Smith JS. Persistent human papillomavirus infection and cervical neoplasia: a systematic review and meta-analysis. Am J Epidemiol 2008;168(2):123–37.
- [3] Bray F, Jacques F, Soerjomataram I, Siegel R, Torre L, Jemal A. Global Cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin 2018;68:394–424.
- [4] Carter JR, Ding Z, Rose BR. HPV infection and cervical disease: a review. Aust New Zeal J Obstet Gynaecol 2011;51(2):103–8.
- [5] Moerman-Herzog A, Nakagawa M. Early defensive mechanisms against human papillomavirus infection. Clin Vaccine Immunol 2015;22(8):850–7.

- [6] Łaniewski P, Barnes D, Goulder A, Cui H, Roe DJ. Linking cervicovaginal immune signatures, HPV and microbiota composition in cervical carcinogenesis in non-Hispanic and Hispanic women. Sci Rep 2018;8(1):7593.
- [7] Brotman RM, Shardell MD, Gajer P, Tracy JK, Zenilman JM, Ravel J, et al. Interplay between the temporal dynamics of the vaginal microbiota and human papillomavirus detection. J Infect Dis 2014;210(11):1723–33.
- [8] Audirac-Chalifour A, Torres-Poveda K, Bahena-Román M, Téllez-Sosa J, Martínez-Barnetche J, Cortina-Ceballos B, et al. Cervical microbiome and cytokine profile at various stages of cervical cancer: a pilot study. PLoS One 2016;11(4).
- [9] Shannon B, Gajer P, Yi TJ, Ma B, Humphrys MS, Thomas-Pavanel J, et al. Distinct effects of the cervicovaginal microbiota and herpes simplex type 2 infection on female genital tract immunology. J Infect Dis 2017;215(9):1366–75.
- [10] Mitra A, MacIntyre DA, Lee YS, Smith A, Marchesi JR, Lehne B, et al. Cervical intraepithelial neoplasia disease progression is associated with increased vaginal microbiome diversity. Sci Rep 2015;5:1–11.
- [11] Passmore J-AS, Williamson A-L. Host immune responses associated with clearance or persistence of human papillomavirus infections. Curr Obstet Gynecol Rep 2016; 5(3):177–88 Current Obstetrics and Gynecology Reports. Available from: http:// link.springer.com/10.1007/s13669-016-0163-1.
- [12] Srinivasan S, Morgan MT, Fiedler TL, Djukovic D, Hoffman NG, Raftery D, et al. Metabolic signatures of bacterial vaginosis. MBio 2015;6(2):1–16.
- [13] Nelson TM, Borgogna JLC, Brotman RM, Ravel J, Walk ST, Yeoman CJ. Vaginal biogenic amines: biomarkers of bacterial vaginosis or precursors to vaginal dysbiosis? Front Physiol 2015:6:1–15.
- [14] Wishart DS, Mandal R, Stanislaus A, Ramirez-Gaona M. Cancer metabolomics and the human metabolome database. Metabolites 2016;6(1).
- [15] Yang K, Xia B, Wang W, Cheng J, Yin M, Xie H, et al. A comprehensive analysis of metabolomics and transcriptomics in cervical cancer. Sci Rep 2017;7:1–11 Nature Publishing Group. Available from: https://doi.org/10.1038/srep43353.
- [16] Sitter B, Bathen T, Hagen B, Arentz C, Skjeldestad FE, Gribbestad IS. Cervical cancer tissue characterized by high-resolution magic angle spinning MR spectroscopy. Magn Reson Mater Physics, Biol Med 2004;16(4):174–81.
- [17] Chai Y, Wang J, Wang T, Yang Y, Su J, Shi F, et al. Application of 1H NMR spectroscopy-based metabonomics to feces of cervical cancer patients with radiation-induced acute intestinal symptoms. Radiother Oncol 2015;117(2):294–301 Elsevier Ireland Ltd. Available from: https://doi.org/10.1016/j.radonc.2015.07.037.
- [18] Guerrero-Flores H, Apresa-García T, Garay-Villar Ó, Sánchez-Pérez A, Flores-Villegas D, Bandera-Calderón A, et al. A non-invasive tool for detecting cervical cancer odor by trained scent dogs. BMC Cancer 2017;17(1):1–8 BMC Cancer. Available from: https://doi.org/10.1186/s12885-016-2996-4.
- [19] Karim R, Tummers B, Meyers C, Biryukov JL, Alam S, Backendorf C, et al. Human papillomavirus (HPV) Upregulates the cellular Deubiquitinase UCHL1 to suppress the Keratinocyte's innate immune response. PLoS Pathog 2013;9(5).
- [20] Nelson TM, Borgogna JC, Michalek RD, Roberts DW, Rath JM, Glover ED, et al. Cigarette smoking is associated with an altered vaginal tract metabolomic profile. Sci Rep 2018;8(1):1–13.
- [21] Vender Heiden MG, Cantley LC, Thompson CB. Understanding the Warburg effect: the metabolic requirements of cell proliferation. Science (80-) 2009;324(May): 1029–33 Available from https://doi.org/10.1126/science.1160809.
- [22] Shannon P, Markiel A, Ozier O, Baliga NS, Wang JT, Ramage D, et al. Cytoscape: a soft-ware environment for integrated models of biomolecular interaction networks. Genome Res 2003;13(11).
- [23] Wickman H. ggplot2: Elegant graphics for data analysis. Springer; 2016.
- [24] Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 2010;7(5):335-6.
- [25] Edgar RC. Search and clustering orders of magnitude faster than BLAST. Bioinformatics 2010;26(19):2460–1.
- [26] DeSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, Keller K, et al. Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Appl Environ Microbiol 2006;72(7):5069–72.
- [27] Langille MGI, Zaneveld J, Caporaso JG, McDonald D, Knights D, Reyes JA, et al. Predictive functional profiling of microbial communities using 16S rRNA marker gene sequences. Nat Biotechnol 2013;31(9):814.
- [28] Shaffer M, Kevin Q, Doenges K, Zhang N, Bokatzian S, Reisdorph N, et al. AMON: Annotation of metabolite origins via networks. bioRvix 2018. https://doi.org/10.1101/439240.
- [29] Noecker C, Eng A, Srinivasan S, Theriot CM, Young VB, Jansson JK, et al. Metabolic model-based integration of microbiome taxonomic and metabolomic profiles elucidates mechanistic links between ecological and metabolic variation. mSystems 2016;1(1):e00013–5.
- [30] Noecker C, Chiu H-C, McNally CP, Borenstein E. Defining and evaluating microbial contributions to metabolite variation in microbiome-metabolome association studies bioRxiv; 2018; 402040 Available from: https://www.biorxiv.org/content/early/ 2018/08/28/402040.
- [31] Arnold KB, Burgener A, Birse K, Romas L, Dunphy LJ, Shahabi K, et al. Increased levels of inflammatory cytokines in the female reproductive tract are associated with altered expression of proteases, mucosal barrier proteins, and an influx of HIVsusceptible target cells. Mucosal Immunol 2016;9(1):194–205 Nature Publishing Group. Available from: https://doi.org/10.1038/mi.2015.51.
- [32] Shannon B, Yi TJ, Perusini S, Gajer P, Ma B, Humphrys MS, et al. Association of HPV infection and clearance with cervicovaginal immunology and the vaginal microbiota. Mucosal Immunol 2017;10(5):1310–9.
- [33] Pavlova NN, Thompson CB. The emerging hallmarks of Cancer metabolism. Cell Metab 2016;23(1):27–47 Elsevier Inc. Available from: https://doi.org/10.1016/j. cmet.2015.12.006.

- [34] Ye N, Liu C, Shi P. Metabolomics analysis of cervical cancer, cervical intraepithelial neoplasia and chronic cervicitis by <sup>1</sup>H NMR spectroscopy. Eur J Gynaecol Oncol 2015;36(2):174–80.
- 35] Santos CR, Schulze A, Lipid metabolism in cancer, FEBS J 2012;279(15):2610-23.
- [36] Daskivich T, Luu M, Noah B, Fuller G, Anger J, Spiegel B. The meaning and use of the area under a receiver operating characteristic (ROC) curve. Radiology 1982;143: 29–36
- [37] Vettukattil R, Hetland TE, Flørenes VA, Kærn J, Davidson B, Bathen TF. Proton magnetic resonance metabolomic characterization of ovarian serous carcinoma effusions: chemotherapy-related effects and comparison with malignant mesothelioma and breast carcinoma. Hum Pathol 2013;44(9):1859–66 Elsevier Inc. Available from: https://doi.org/10.1016/i.humpath.2013.02.009.
- [38] Schmitt WD, Hilvo M, Gopalacharyulu P, Budczies J, Denkert C, Darb-Esfahani S, et al. Accumulated metabolites of hydroxybutyric acid serve as diagnostic and prognostic biomarkers of ovarian high-grade serous carcinomas. Cancer Res 2016;76(4): 796–804.
- [39] Bonuccelli G, Tsirigos A, Whitaker-Menezes D, Pavlides S, Pestell RG, Chiavarina B, et al. Ketones and lactate "fuel" tumor growth and metastasis: evidence that epithelial cancer cells use oxidative mitochondrial metabolism. Cell Cycle 2010;9(17): 3506–14.
- [40] Turkoglu O, Zeb A, Graham S, Szyperski T, Szender JB, Odunsi K, et al. Metabolomics of biomarker discovery in ovarian cancer: a systematic review of the current literature. Metabolomics 2016;12(4):1–16 Springer US.
- [41] Zietkowski D, Davidson RL, Eykyn TR, De Silva SS, DeSouza NM, Payne GS. Detection of cancer in cervical tissue biopsies using mobile lipid resonances measured with diffusion-weighted1H magnetic resonance spectroscopy. NMR Biomed 2010;23 (4):382–90.
- [42] Zhao L, Ruan XZ, Zeng H, Wei L, Varghese Z, Chen Y, et al. Dietary oleic acid-induced CD36 promotes cervical cancer cell growth and metastasis via up-regulation Src/ERK pathway. Cancer Lett 2018;438:76–85 Elsevier. Available from: https://doi.org/10. 1016/j.canlet.2018.09.006.
- [43] Meikle PJ, Summers SA. Sphingolipids and phospholipids in insulin resistance and related metabolic disorders. Nat Rev Endocrinol 2017;13(2):79–91 Nature Publishing Group. Available from: https://doi.org/10.1038/nrendo.2016.169.
- [44] Wang WW, Qiao SY, Li DF. Amino acids and gut function. Amino Acids 2009;37(1): 105–10.
- [45] Ferreira AK, Meneguelo R, Pereira A, Filho OMR, Chierice GO, Maria DA. Anticancer effects of synthetic phosphoethanolamine on Ehrlich ascites tumor: an experimental study. Anticancer Res 2012;32(1):95–104.
- [46] Anahtar MN, Gootenberg DB, Mitchell CM, Kwon DS. Cervicovaginal microbiota and reproductive health: the virtue of simplicity. Cell Host Microbe 2018;23(2):159–68 Elsevier Inc. Available from https://doi.org/10.1016/j.chom.2018.01.013.
- [47] Ness RB, Kip KE, Hillier SL, Soper DE, Stamm CA, Sweet RL, et al. A cluster analysis of bacterial vaginosis-associated microflora and pelvic inflammatory disease. Am J Epidemiol 2005;162(6):585–90.
- [48] Brusselaers N, Shrestha S, Van De Wijgert J, Verstraelen H. Vaginal dysbiosis, and the risk of human papillomavirus and cervical cancer: systematic review and metaanalysis. Am J Obstet Gynecol Dec 12 2018. https://doi.org/10.1016/j.ajog.2018.12. 011 Elsevier Inc. Available from https://linkinghub.elsevier.com/retrieve/pii/ S000293781832221X. (pii: S0002-9378(18)32221-X, [Epub ahead of print] Review. PMID: 30550767).
- [49] Onderdonk AB, Delaney ML, Fichorova N. The human microbiome during bacterial vaginosis. Clin Microbiol Rev 2016;29(2):223–38.
- [50] Fredricks DN, Fiedler TL, Thomas KK, Oakley BB, Marrazzo JM. Targeted PCR for detection of vaginal bacteria associated with bacterial vaginosis. J Clin Microbiol 2007; 45(10):3270-6.
- [51] Vitali B, Cruciani F, Picone G, Parolin C, Donders G, Laghi L. Vaginal microbiome and metabolome highlight specific signatures of bacterial vaginosis. Eur J Clin Microbiol Infect Dis 2015;34(12):2367–76.
- [52] McMillan A, Rulisa S, Sumarah M, Macklaim JM, Renaud J, Bisanz JE, et al. A multiplatform metabolomics approach identifies highly specific biomarkers of bacterial diversity in the vagina of pregnant and non-pregnant women. Sci Rep 2015;5: 1–14 Nature Publishing Group. Available from: https://doi.org/10.1038/srep14174.
- [53] Rutkowsky JM, Schneider D, McCoin CS, Huang S, Adams SH, Ono-Moore KD, et al. Acylcarnitines activate proinflammatory signaling pathways. Am J Physiol Metab 2014;306(12):E1378–87.
- [54] Briselden AM, Hillier SL. Longitudinal study of the biotypes of Gardnerella vaginalis. J Clin Microbiol 1990;28(12):2761–4.
- [55] Sujatha K, Mahalakshmi A, Solaiman DKY, Shenbagarathai R. Sequence analysis, structure prediction, and functional validation of phaC1/phaC2 genes of Pseudomonas sp LDC-25 and its importance in Polyhydroxyalkanoate accumulation. J Biomol Struct Dyn 2009;26(6):771–9.
- [56] Nakajima T, Katsumata K, Kuwabara H, Soya R, Enomoto M, Ishizaki T, et al. Urinary polyamine biomarker panels with machine-learning differentiated colorectal cancers, benign disease, and healthy controls. Int J Mol Sci 2018;19(3):1–14.
- [57] Weiss TS, Herfarth H, Obermeier F, Ouart J, Vogl D, Schölmerich J, et al. Intracellular polyamine levels of intestinal epithelial cells in inflammatory bowel disease. Inflamm Bowel Dis 2004;10(5):529–35.
- [58] Zhao G, He F, Wu C, Li P, Li N, Deng J, et al. Betaine in inflammation: mechanistic aspects and applications. Front Immunol 2018;9:1–13.
- [59] Rowley TJ, McKinstry A, Greenidge E, Smith W, Flood P. Antinociceptive and antiinflammatory effects of choline in a mouse model of postoperative pain. Br J Anaesth 2010:105(2):201–7.
- 60] Kets E, Bont J. Protective effect of betaine on survival of *Lactobacillus plantarum* subjected to drying. FEMS Microbiol Lett 1994;116(3):251–6.

- [61] Ridlon JM, Harris SC, Bhowmik S, Kang DJ, Hylemon PB. Consequences of bile salt biotransformations by intestinal bacteria. Gut Microbes 2016;7(1):22–39 Taylor & Francis. Available from: https://doi.org/10.1080/19490976.2015.1127483.
- [62] Fiorucci S, Distrutti E. Bile acid-activated receptors, intestinal microbiota, and the treatment of metabolic disorders. Trends Mol Med 2015;21(11):702–14.
- [63] Tatsugami M, Ito M, Tanaka S, Yoshihara M, Matsui H, Haruma K, et al. Bile acid promotes intestinal metaplasia and gastric carcinogenesis. Cancer Epidemiol Biomarkers Prev 2012;21(11):2101–7.
- [64] Fulbright LE, Ellermann M, Arthur JC. The microbiome and the hallmarks of cancer. PLoS Pathog 2017;13(9):e1006480 Available from https://doi.org/10.1371/journal. ppat.1006480.
- [65] Wallner S, Orso E, Grandl M, Konovalova T, Liebisch G, Schmitz G. Phosphatidylcholine and phosphatidylethanolamine plasmalogens in lipid loaded human macrophages. PLoS One 2018:13(10):1–21.
- [66] Treede I, Braun A, Sparla R, Kühnel M, Giese T, Jerrold R, et al. Anti-inflammatory effects of phosphatidylcholine. J Biol Chem 2007;282(37):27155–64.
- [67] Ridgway ND. The role of phosphatidylcholine and choline metabolites to cell proliferation and survival. Crit Rev Biochem Mol Biol 2013;48(1):20–38.
- [68] Hall IH, Burnham BS, Elkins A, Sood A, Powell W, Tomasz J, et al. Boronated Pyrimidines and purines as cytotoxic, Hypolipidemic and anti-inflammatory agents. Met Based Drugs 1996;3(3):155–60 Available from http://www.hindawi.com/archive/ 1996/694271/abs/.
- [69] Cronstein BN Adenosine, an endogenous anti-inflammatory agent, J Appl Physiol 1994;76(1):5–13 Available from http://www.ncbi.nlm.nih.gov/pubmed/8175547.
- [70] Fukuda T, Majumder K, Zhang H, Matsui T, Mine Y. Adenine has an antiinflammatory effect through the activation of adenine receptor signaling in mouse macrophage. J Funct Foods 2017;28:235–9 Elsevier Ltd. Available from https://doi. org/10.1016/j.jff.2016.11.013.
- [71] Feng C, Mery aG, Beller EM, Favot C, J a Boyce. Adenine nucleotides inhibit cytokine generation by human mast cells through a Gs-coupled receptor. J Immunol 2004; 173(12):7539–47 Available from http://www.jimmunol.org/cgi/doi/10.4049/ jimmunol.173.12.7539.
- [72] France MT, Mendes-soares H, Forney LJ. Genomic comparisons of LActobacillus crispatus and Lactobacillus iners reveal potential ecological drivers of community

- composition in the vagina. 2016;82(24):7063-73 Available from https://doi.org/10.4049/jimmunol.173.12.7539.
- [73] O'Hanlon DE, Moench TR, Cone RA. Vaginal pH and Microbicidal lactic acid when lactobacilli dominate the microbiota. PLoS One 2013;8(11).
- [74] Boskey ER, Cone RA, Whaley KJ, Moench TR. Origins of vaginal acidity: high D/L lctate ratio is consistent with bacteria being the primary source. Hum Reprod 2001;16(9):1809–13 Available from http://www.crime-expertise.org/17eme-congres-danthropologie-medico-legale-2017/.
- [75] Errea A, Cayet D, Marchetti P, Tang C, Kluza J, Offermanns S, et al. Lactate inhibits the pro-inflammatory response and metabolic reprogramming in Murine macrophages in a GPR81-independent manner. PLoS One 2016;11(11):1–11.
- [76] Sun S, Li H, Chen J, Qian Q. Lactic acid: no longer an inert and end-product of glycolysis. Physiology 2017;32(6):453–63 Available from http://physiologyonline. physiology.org/lookup/doi/10.1152/physiol.00016.2017.
- [77] Walenta S, Wetterling M, Lehrke M, Schwickert G, Sundfør K, Rofstad EK, et al. High lactate levels predict likelihood of metastases, tumor recurrence, and restricted patient survival in human cervical cancers. Cancer Res 2000;60(4):916–21.
- [78] Herbst-Kralovetz MM, Pyles RB, Ratner AJ, Sycuro LK, Mitchell C. New systems for studying intercellular interactions in bacterial vaginosis. J Infect Dis 2016;214: S6–13.
- [79] Radtke AL, Herbst-Kralovetz MM. Culturing and applications of rotating wall vessel bioreactor derived 3D epithelial cell models, Jove-J Vis Exp 2012(62).
- [80] Doerflinger SY, Throop AL, Herbst-Kralovetz MM. Bacteria in the vaginal microbiome alter the innate immune response and barrier properties of the human vaginal epithelia in a species-specific manner. J Infect Dis 2014;209(12):1989–99.
- [81] Radtke AL, Quayle AJ, Herbst-Kralovetz MM. Microbial products alter the expression of membrane-associated mucin and antimicrobial peptides in a three-dimensional human endocervical epithelial cell model1. Biol Reprod 2012;87(6):1–10 Available from: https://academic.oup.com/biolreprod/article-lookup/doi/10.1095/biolreprod. 112.103366.
- [82] Goodman B, Gardner H. The microbiome and cancer. J Pathol Nature Publishing Group 2018;244(5):667–76 Available from: https://doi.org/10.1038/nrc3610.