COMPUTATIONAL META-ANALYSIS OF CERVICAL CANCER USING AVAILABLE 16S RRNA NGS DATA

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ABSTRACT

Cervical cancer is one of the most frequently occurring and deadliest gynaecological cancer which develops in cervical cells. Since it develops in tissues lining the internal organs it is a Carcinoma. Human papilloma virus infection is found to top the list of carcinogenic factors. Overexpression of certain proteins due to HPV integration in host body over a time can result in carcinoma. However vaginal microbiota plays a key role in development, persistence and progression of infections leading to diseases such as cervical cancer. Some recent studies have revealed potential roles of microbiome in cervicovaginal diseases. Thus a comparative metagenomic study among such samples can uncover microbial diversities present in these samples. Due to presence of highly conserved regions as well as hyper variable regions 16s rRNA gene sequence is selected for identification and classification of bacterial diversity. For the purpose of metagenomic analysis 16s rRNA gene sequences were analysed using QIIME pipeline.

Keywords: 16s rRNA, metagenomics, microbiota, cervical cancer.

INTRODUCTION

Cervical cancer is one of the most frequently occurring and deadliest gynaecological cancer which develops in the cells of cervix [5]. Since it develops in tissues lining the internal organs it is a Carcinoma. Normal cells undergo some changes due to some external factors or mutations. The abnormal changes occurring in cervical cells are called Squamous Intraepithelial Lesions (SIL). Cells become abnormal which can later result in cancer. These conditions are also known as cervical intraepithelial neoplasia (CIN). Depending on the extent of damage CIN can be CIN 1, CIN 2, or CIN 3 [6].

In both men and women, presence of Human papillomaviruses HPV is considered as a cause for most of the sexually transmitted diseases. Papillomaviruses are reported as a causative agent for many reproductive tract infections or STIs. Till now, on the basis of genomic differences more than 200 types of HPV have been recognised. Type 16, 18, 31 and 45 have been reported most commonly occurring in malignant cells of cervical cancer. Type 16 accounting for about 50% of the cases. 99.7% of the cervical squamous cell carcinoma cases worldwide have shown the implication of HPV.

Metagenomics is an unbiased and reasonable method of recognising and reviewing microorganisms within their environment and surroundings. Metagenomics has expanded the scope of targeting microbes inducing various diseases and infections including different types of cancers. The meta-genomic analyses of micro-organisms customarily aids in recognizing the microbial communities in samples and their evolutionary relationships. Genetics, environmental factors, lifestyle collectively cause different diseases. One such factor for causing different types of cancers is presence of different microbial communities in different proportions in an individual.

A healthy vaginal microbiota mainly includes Lactobacillus spp., which upholds vaginal wellbeing by dropping pH in vagina by secreting lactic acid. Bacterial Vaginosis (BV) can result in the loss of inherent Lactobacillus spp. Also promotes over-population of anaerobic bacteria which can cause inequity in vaginal microbiome allied with various health issues like vaginal
discharge syndrome, poor pregnancy outcomes, pelvic inflammatory disease, post-operative wound infections. The frequency of Bacterial vaginosis makes a person more susceptible to HPV infection. Persistent HPV infection is the cause for the development and progression of cervical cancer to a higher grade. Study conducted among women with and without HPV infection uncovers the presence of Lactobacillus in healthy women in abundance. Some other studies reveal low frequency of Lactobacillus spp. and notable presence of Fusobacteria spp., including Sneathia spp. in HPV positive cases [17,18].

Materials and methods

2.1 Data retrieval
Based on literature study raw sequence files were downloaded from NCBI – SRA by using keyword, Cervical” (https://www.ncbi.nlm.nih.gov/sra/?term=cervical).

| Sr. No. | Source                           | AccessionNo. | HPV    | Region   | Other Conditions          |
|---------|----------------------------------|--------------|--------|----------|---------------------------|
| 1       | Homo sapiens                     | SRR6505152   | Positive | V4       |                           |
| 2       | Homo sapiens                     | SRR6505153   | Negative | V4       |                           |
| 3       | Homo sapiens                     | SRR3109486   | Positive | V3 & V4  | No cervicallesion         |
| 4       | Homo sapiens                     | SRR3109483   | Negative | V3 & V4  | Healthy                  |
| 5       | Homo sapiens                     | SRR3109479   | Positive | V3 & V4  | Cervical cancer          |
| 6       | Homo sapiens                     | SRR3109471   | Positive | V3 & V4  | SIL                      |
| 7       | Human metagenome                 | SRR7225960   | Positive | V3 & V4  | No bacterial vaginosis    |
| 8       | Human metagenome                 | SRR7225958   | Positive | V3 & V4  | Bacterial vaginosis       |
| 9       | Human metagenome                 | SRR7225944   | Negative | V3 & V4  | No bacterial vaginosis    |
| 10      | Human metagenome                 | SRR7225943   | Negative | V3 & V4  | Bacterial vaginosis       |
| 11      | Human reproductive system metagenome | ERR1684337 | -   | V4 & V5  | Healthy                  |
| 12      | Human reproductive system metagenome | SRR3506081 | Positive | V3 & V4  | Normal                   |
| 13      | Human reproductive system metagenome | SRR3504686 | Positive | V3 & V4  | CIN 1                    |

https://iabcd.org.in/
Table 1 Data selected from various sources

|   | metagenome | SRR3504755 | Positive | CIN 2 |
|---|------------|-------------|----------|-------|
| 14| Human reproductive system metagenome | SRR3504954 | Positive | V3 & V4 | CIN 3 |

2.2 Quality Filtering
Raw files downloaded in Sra format were converted to Fastq format using Sratoolkit version 2.9.2-ubuntu 64. Paired end fastq files were split into two files by using command fastq-dump --split-files. Quality check of these files was done by using FastQC v_11.8 tool [27].

2.3 Merge paired end reads
By using FLASH_1.2.11 the paired end files were merged.

2.4 High quality reads
The files obtained after using FLASH were again processed for Quality check so as to obtain only high quality data. The poor quality reads were trimmed by using trimmomatic software [28]. In some samples few reads were removed by using a PERL Script.

2.5 QIIME installation
Qiime includes python cod and some additional packages. For Mac OS environment MacQIIME can be used. QIIME virtual box can be installed for windows and linux environments. QIIME can be installed by using Miniconda. Command based installation is done.

2.6 Steps to run QIIME pipeline are as follows:
2.6.1 Quality control
It involves generating a mapping file, validate mapping file and demultiplexing of sequence. Quality is checked by FastQC. Demultiplexing of data is done by using _split_libraries and for FastQ files _fastq.py is used.

2.6.2 Chimera Removal
It involves identification of chimeric sequences and filtration of chimeric and non chimeric sequences. First a Usearch61 file is generated. USEARCH61 performs reference based chimera detection as well as de novo chimera detection on the basis of abundances of input sequences [24,25].

2.6.3 OTU clustering
Sequences are clustered into operational taxonomic units (OTU). QIIME provides different strategies for OTU clustering [24,25].

2.6.4 Selecting a representative sequence & Assign Taxonomy
Assignment of taxonomy to OTU representative sequences is included as a default step in the open-reference OTU-picking workflow. QIIME uses curated database to assign taxonomy when it uses closed-reference OUT-picking approach. Behaviour of taxonomy assignment can be changed by using parameters with the script assign_taxonomy.py.

2.6.5 Generate taxa Summary tables
OTU table file will be generated by _otu_table. By using this file downstream analysis is done. In this step out table is created, OTUs are filtered, taxonomy summary is generated and taxa plot are also generated [25,26].
RESULTS
Source 1 HPV negative

![Figure 1 bacterial diversity in HPV negative sample](https://iabcd.org.in/image1.jpg)

Source 1 HPV positive

![Figure 2 Bacterial diversity in HPV positive sample](https://iabcd.org.in/image2.jpg)

Source 2 Cervical cancer

![Figure 3 Bacterial diversity in cervical cancer sample](https://iabcd.org.in/image3.jpg)
Source 2 negative healthy

Figure 4 Bacterial diversity in HPV negative and healthy sample

Source 2 no lesion positive

Source 2 SIL

Figure 5 Bacterial diversity in HPV positive sample without lesion

Figure 6 Bacterial diversity in SIL sample
Figure 7 Bacterial diversity in HPV negative and BV negative sample

Figure 8 Bacterial diversity in HPV negative and BV positive sample

Figure 9 Bacterial diversity in HPV positive and BV negative sample

Source 3 HPV negative BV negative

Source 3 HPV negative BV positive

Source 3 HPV positive BV negative
Source 3 HPV positive BV positive

Figure 10 Bacterial diversity in HPV positive and BV positive sample

Source 4 healthy normal

Figure 11 Bacterial diversity in Healthy normal sample

Source 5 CIN 1

Figure 12 Bacterial diversity in CIN 1 sample
Source 5 CIN 2

Figure 13 Bacterial diversity in CIN 2 sample

Source 5 CIN 3

Figure 14 Bacterial diversity in CIN 3 sample

Source 5 Control
Discussion

Figure 15 Bacterial diversity in Control sample

Cervicovaginal microbiota plays a role in maintaining vaginal health. For identification of cervicalmicrobial diversities metagenomics approach was used. This meta-analysis includes comparative studies of healthy samples, HPV positive samples, HPV negative samples, samples with different degrees of CIN and bacterial vaginosis samples which were downloaded from SRA. Pre-processing and quality filtering provided high quality data (QV > / = 20) which was then analysed through QIIME pipeline. In this study 16s rRNA gene sequences were used for identification and taxonomical classification of microbial world occurring in healthy as well as infected samples so as to identify potential pathogens present in cervical cancer. Lactobacillus species which is known to maintain vaginal health was found in healthy samples with relative abundance > 99%. Bacterial vaginosis samples were also having greater abundance i.e. 80% of Lactobacillus species.

| Genus          | HPV negative | HPV positive |
|----------------|--------------|--------------|
| Bacteroides    | 48.87%       | --           |
| Anaerococcus   | 23.79%       | 1.04%        |
| Fusobacterium  | 15.76%       | 4.22%        |
| Porphyromonas  | 3.13%        | 12.00%       |
| Prevotella     | 1.91%        | 21.81%       |
| Campylobacter  | 1.72%        | 31.87%       |
| Lachnospiraceae| --           | 12.91%       |
| Peptoniphilus  | 1.71%        | --           |
| Moryella       | 1.07%        | 1.23%        |

Table 2 Comparative results of HPV negative and HPV positive

| Genus          | Cervical cancer | HPV negative | HPV positive |
|----------------|-----------------|--------------|--------------|
| Fusobacterium  | 76.16%          | --           | --           | 17.45%       |
| Dialister      | 9.10%           | --           | --           | --           |
| Mycoplasma     | 8.46%           | --           | --           | 42.52%       |
| Prevotella     | 2.16%           | --           | --           | --           |
| Granulicatella | 1.33%           | --           | --           | --           |
| Campylobacter  | 1.20%           | --           | --           | --           |
| Lactobacillus  | --              | 99.89%       | 99.93%       | 19.19%       |
| Bifidobacteriaceae | -- | -- | -- | 1.14% |

Table 3 Comparative results of Cervical Cancer, HPV negative, HPV positive-no lesion and SIL

| Genus             | HPV negativeBV negative | HPV negativeBV positive | HPV positiveBV negative | HPV positiveBV positive |
|-------------------|--------------------------|-------------------------|-------------------------|-------------------------|
| Lactobacillus     | 80.73%                   | --                      | 30.13%                  | --                      |
| Flavobacteriaceae | 5.86%                    | 1.83%                   | --                      | --                      |
| Pseudomonas       | 2.68%                    | --                      | --                      | --                      |
| Pedobacter        | 2.24%                    | --                      | --                      | --                      |
| Clostridium       | 1.14%                    | 6.57%                   | --                      | 2.65%                   |
| Prevotella        | --                       | 37.54%                  | 5.39%                   | 29.72%                  |
| Bifidobacteriaceae| --                       | 28.40%                  | 19.47%                  | --                      |
| Megasphaera       | --                       | 8.47%                   | 6.59%                   | 11.59%                  |
| Dialister         | --                       | 4.91%                   | --                      | 6.02%                   |
| Atopobium         | --                       | 4.55%                   | --                      | --                      |
| Sneathia          | --                       | 4.09%                   | --                      | 12.59%                  |
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Lactobacillus species result in cervical carcinoma.

In this study, 16s rRNA gene sequences were used from SRA for identification and taxonomic classification of cervical microbiota through the metagenomics approach. For meta-analysis of this data, QIIME pipeline was used. Relative abundance of cervical microbes was identified in HPV positive and negative samples. Exclusive presence and greater abundance of some species such as Prevotella and Campylobacter was found abundantly in HPV positive samples. Fusobacterium was present in maximum abundance i.e. 76% in samples with cervical cancer. BV positive samples were having a greater abundance of Prevotella which was rarely present in BV negative samples.

The abundance of lactobacillus was found to decrease with increase in CIN stage i.e lactobacillus were higher in CIN1 (70%) and their abundance decreased in CIN 3 (0.64%).

CONCLUSION

In this study, 16s rRNA gene sequences were used from SRA for identification and taxonomic classification of cervical microbiota through the metagenomics approach. For meta-analysis of this data, QIIME pipeline was used. Relative abundance of cervical microbes was identified in HPV positive and negative samples. Exclusive presence and greater abundance of some species such as Prevotella and Campylobacter was found abundantly in HPV positive samples which signifies that these could be potentially pathogenic and their prevalence for longer time could result in cervical carcinoma. Presence of Fusobacterium in greater abundance was observed in cervical cancer samples. Whereas healthy and HPV negative samples were rich in Lactobacillus species.

Table 4 Comparative results of HPV negative- BV positive & negative, HPV positive – BV positive & negative samples

| Genus       | Healthy Control |
|-------------|-----------------|
| Lactobacillus | 99.46%          |
| Sneathia    | 0.09%           |
| Prevotella  | 0.08%           |

Table 5 Microbial diversity in Healthy control sample

| Genus           | CIN 1 | CIN 2 | CIN 3 | Control |
|-----------------|-------|-------|-------|---------|
| Lactobacillus   | 79.20%| 38.31%| 0.64% | 27.41%  |
| Halomonas       | 2.30% | 4.66% | 4.44% | 8.34%   |
| Shewanella      | 1.24% | 1.91% | 1.81% | 3.87%   |
| Comamonadaceae  | 0.81% | --    | --    | --      |
| Prevotella      | --    | 24.63%| 1.37% | 2.95%   |
| Bifidobacteriaceae | -- | 2.00% | 79.03%| 24.33%  |
| Streptococcus   | --    | 2.36% | 1.53% | --      |

Table 6 Comparative results of CIN 1, CIN 2, CIN 3 and Control samples

Comparative analysis of HPV positive and negative samples reveals the presence of Bacteroides and Anaerococcus exclusively in HPV negative samples in a greater abundance. Whereas Prevotella, Campylobacter and Lachnospiraceae were found abundantly in HPV positive samples. Fusobacterium was present in maximum abundance i.e. 76% in samples with cervical cancer. BV positive samples were having a greater abundance of Prevotella which was rarely present in BV negative samples.

The abundance of lactobacillus was found to decrease with increase in CIN stage i.e lactobacillus were higher in CIN1 (70%) and their abundance decreased in CIN 3 (0.64%).
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