Original Research Article
doi: http://dx.doi.org/10.20546/ijcmas.2016.502.031

Microbial Vaginitis in Reproductive Women of Silk city of South India

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ABSTRACT

Vaginitis is caused by several microbial agents and non-infective irritants. It is usually associated with urethritis, dysperunia, dysmenorrhea and lower abdominal pain. Microbial agents are transmitted through sexual contact and unhygienic habits. Hence, this study is designed to rule out the incidence of Microbial agents in reproductive women by microscopy, staining &culture. 100 patients after sensitization of the project were taken for this study. Samples processed by microscopy, special staining and culture. The culture was performed with KUPFERBERG media. Out of 100 samples, 54 grew bacterial colonies, 30 were found to be positive for *Trichomonas vaginalis*, 2 for *Candida albicans* and the remaining 14 were having normal vaginal flora. The parasitic agents were found in 30 swabs by Giemsa stain. Preliminary wet mount microscopic examination revealed 15 cases to be positive. The analysis showed the correlation between bacterial and parasitic isolation, which was more common among sexually active female patients, which requires screening of patients in appropriate manner and treat accordingly to envisage a healthy quality life for the patients.

Keywords: *Trichomonas vaginalis*, Culture, Microscopy, Vaginitis, Silk city of South India

Article Info

Accepted: 15 January 2016
Available Online: 10, February 2016

Introduction

Vaginitis is a known infectious disease caused by several microbial agents and non-infective irritants. The disease often involves vulva with mixed microbial agents. Therefore it is necessary to know the causative agents for patient care and management.

Vagina has mixed anaerobic and aerobic bacterial flora including *Gardnerella vaginalis* causing bacterial vaginosis. Infections with *Clostridium perfringens* typically after illegal abortions with non-sterile instruments is also a well established entity (1).
Genital tract Infections of women are well known in reproductive age group and they are the commonest problem that makes a lady to visit her gynecologist with vaginal discharge. This can happen due to bacterial, fungal and parasitic microbial agents. It is usually associated with urethritis, dysperunia, dysmenorrhea and lower abdominal pain.

*Trichomonas vaginalis* is an important cause for sexually transmitted infection in reproductive age group. It is the most prevalent non-viral, non–bacterial cause for genital tract infection. It can also co-exist with HIV, Herpes simplex viruses, and can enhance the risk of Human Papilloma virus infection in women (2).

This disease is caused by a pear shaped protozoan *Trichomonas vaginalis*, giving rise to frothy, foul-smelling discharge with vulvo-vaginal irritation and lower abdominal pain (3).

It can be an important cause for Pelvic inflammatory Disease (PID) leading to complications like infertility (4).

In this era of infertility and sterility with long married life, we decided to take up this study among reproductive age group, among our patients attending OG clinic in our hospital.

The main aim of this study includes to know the incidence of Microbial agents in reproductive age group women attending our hospital with Leucorrhoea. Demonstration of the agent by microscopy, staining techniques. Demonstration of the agent by culture techniques. To demonstrate the sensitivity of culture isolation from clinical samples. And also to correlate the clinical findings.

**Materials and Methods**

100 willing patients after sensitization of the project, who were attending our OG-OPD were taken for this study.

**Inclusion Criteria**

Willing, Reproductive age group women with vaginal discharge.

**Exclusion Criteria**

Elderly (>50) and Young (<15), other vaginitis conditions.

**Methods**

A total of 100 samples were taken from 100 willing, sexually active female patients with an age group ranging from 20-50 years (Table: 1).

The vaginal swab with appropriate labelling and duly filled up request form was received in microbiology laboratory and subjected for microscopic examination, specific special staining and culture inoculation and incubation.

**For Bacterial Etiology**

The swab was inoculated onto the Mac Conkey agar, Blood agar and Chocolate agar and incubated at 37°C for 48 hours. The culture plates were examined for colony growth and processed up to species level identification and antibiotic sensitivity performed as per NCCLS guidelines (5).

**For Parasitic STD**

The direct wet mount of vaginal smear was examined by direct microscopic examination. The culture was performed
with KUPFERBERG media, though there are other media like, Feinberg, Whittington media. The swab was inoculated into the media incubated at 37ºC for 7 days. The media was examined on 3rd, 5th, 7th day and a strict aseptic precautions. Both wet mount microscopic and Giemsa staining were done (Table: 2)

The culture media and Giemsa stains were performed as per NCCLS guidelines (6).

For fungal Causes

Samples were inoculated onto Sabourauds Dextrose Agar and fungal colonies grown were identified up to species level.

Results and Discussion

Out of 100 samples, 54 samples grew bacterial colonies, 30 samples were found to be positive for *Trichomonas vaginalis*, 2 samples were positive for *Candida albicans* and the remaining 14 were having normal vaginal flora.

Among the bacterial colonies 52 samples had Gram negative bacterial growth and 2 samples had Gram positive cocci.

The parasitic agents were found to be positive in 30 swabs and identified by culture and staining techniques by Giemsa stain. Preliminary wet mount microscopic examination revealed 15 cases to be positive (Table: 3).

2 samples yielded *Candida albicans* were found commonly in the 4th decade of age group.

Both bacterial, parasitic agents were found commonly in the 3rd decade of age group (Table: 4).

Gram positive bacteria, isolates were MSSE (Methicillin Sensitive *Staphylococcus epidermidis*). The gram negative isolates had an array of bacteria: *Escherichia coli*, *Klebsiella oxytoca* and *Klebsiella pneumonia*, *Pseudomonas aeruginosa* and *Acinetobacter baumanii*.

The KUPFERBERG culture media was more fruitful as it yielded positive results on 3rd day, 5th day, and 7th day proving to be more specific and sensitive for clinching the parasitic STD agent.

Totally they were 30 % culture positive for *Trichomonas vaginalis*, which was further confirmed by demonstration of the pear shaped protozoan by Giemsa staining technique (Pic:1)

The analysis of the 100 samples showed the following microbial profile (Table.5)

We got 54 bacterial vaginitis with heavy growth of *Escherichia coli*, *Klebsiella oxytoca*, *Klebsiella pneumonia*, *Pseudomonas aeruginosa*, *Staphylococcus epidermidis* and *Acinetobacter baumanii*. Their distribution is shown in (Table: 6)

The vaginal flora has complex microbiota with varying quantities and relative proportions (7).

There is an intricate balance among the microbes that maintains the normal vaginal flora. It is mainly dominated by the genus Lactobacillus that gives acidic pH to the vagina which guards against vaginitis.

When this pH gets altered, or colonization due to other bacteria and unhealthy sexual contact gives rise to vaginitis, which leads to vaginal discharge, urethritis, dysmenorrhea, dysperunia, lower abdominal pain and infertility.
**Table 1** Age Distribution of the Patients (100)

| AGE GROUP (YEARS) | PERCENTAGE % |
|-------------------|--------------|
| 20-25             | 20           |
| 26-30             | 21           |
| 31-35             | 30           |
| 36-40             | 15           |
| 41-45             | 07           |
| 46-50             | 04           |
| 51 & Above        | 04           |

**Table 2** Culture Wet Mount Preparation and Giemsa Staining

| S: NO | INCUBATION DAYS | MICROSCOPY CULTURE | STAIN |
|-------|-----------------|--------------------|-------|
| 1     | 3rd day         | 15                 | 15    |
| 2     | 5th day         | 10                 | 10    |
| 3     | 7th day         | 5                  | 5     |

**Table 3** Direct Wet Mount Preparation:

| Diagnosis                     | NO. positive for *T. Vaginalis* | NO. negative for *T. Vaginalis* |
|-------------------------------|---------------------------------|----------------------------------|
| Wet mount preparation         | 15                              | 85                               |

**Table 4** Age Group Correlation with Parasitic STD

| S.No | Age     | Positive for T. vaginalis | Positive for Bacterial vaginitis |
|------|---------|---------------------------|---------------------------------|
| 1    | 20-30   | 21                        | 23                              |
| 2    | 31-40   | 6                         | 16                              |
| 3    | 41-50   | 3                         | 8                               |
| 4    | 50 and above | -                  | 7                               |
The analysis of the vaginal discharge with patients attending our OG department, gave rise to a wide range of microbial profile including bacterial, parasitic and fungal agents. About 54% of the samples had bacterial culture positivity, in which *Escherichia coli* was 39% positive (20/54) samples, *Klebsiella oxytoca* contributed for 24% with 13 samples, and *Klebsiella pneumoniae* had a share of 19% with (10/54) samples, the other isolates were *Pseudomonas aeruginosa* which was isolated from 5 samples contributing for 10% positivity and *Acinetobacter baumanii* and *Staphylococcus epidermidis* 4% with (2/54) samples in each. These bacterial isolates were sensitive to antibiotics as shown in Table 6.
isolates were found in the vaginal swabs due to the lack of personal hygiene, as majority of our patients come from low socio economic stata , with paddy field work pattern. which would have enhanced the colonization of this organism from the large intestine.

The next important isolate was parasitic agent *Trichomonas vaginalis* which contributed for 30%, was responsible for green coloured foul-smelling discharge, itching in vagina & vulva. It is the major sexually transmitted non-viral, non-bacterial disease, present all over the world. It inhabits the genetalia, urinary tract in female, urethra and prostate in male. It takes shelter and nutrition from another living organism causing vulvo-vaginitis, PID. It can also precipitate infertility and sterility.

Kupferburg Media was selected as it contains selective agent to inhibit the growth of GNB & GPC. In addition it contains nitrogenous compounds for ensuring the growth of *Trichomonas vaginalis*. The inclusion of 0.1% agar reduces the oxygen tension for the prolific growth of *Trichomonas vaginalis*. Our study shows 30% of the samples were positive for *Trichomonas vaginalis* (Table: 2).

The sensitivity pattern of the gram negative bacilli showed 100% sensitivity to Imipenem, Amikacin and Gentamycin.100% resistant for Cefazolin, 80% for Ceftazidime and 74% for Cefotaxime.

The analysis of Gram positive cocci (*Staphylococcus epidermidis*), had the following pattern, 100% oxacillin, linezolid and vancomycin sensitive. It had 50% resistant to Gentamycin and Ampicillin.

Other than the bacterial, parasitic agents, the study also showed the presence of fungal agent in the sample- *Candida albicans* (2/100) with 2%. This also may be associated with sexual transmission similar to *Trichomonas vaginalis*.

In conclusion, the present study reveals that the vagina of the reproductive age group female gets inhabited with a variety of microorganisms due to faulty hygienic habits, which enhances migration of bacteria from intestine to genital tract.

Coitus is another precipitating factor for vaginitis which aids in the parasitic, fungal vaginitis. As an health professional it is mandatory to give health education to the patients with regards to personal hygienic habits and safe coitus in order to prevent vaginitis from happening in young reproductive ladies as shown in our studies.

This analysis shows the correlation between bacterial and parasitic isolation, which was more common among sexually active female patients as shown in our study, which requires screening of patients in appropriate manner and treat accordingly to envisage a healthy quality life for the patients.

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How to cite this article:
Karthika Jayakumar, Wills shiela, S. Jayalakshmi, Sathya Pandurangan and M. Mohanambal. 2016. Microbial Vaginitis in Reproductive Women of Silk city of South India. Int.J.Curr.Microbiol.App.Sci.5(2): 272-278. doi: http://dx.doi.org/10.20546/ijcas.2016.502.031