RESEARCH ARTICLE

PREVALENCE OF HUMAN PAPILLOMA VIRUS IN CERVICAL CELLS AMONG PATIENTS WITH ABNORMAL PAP SMEARS.

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Abstract

Human papillomavirus (HPV) is the most common viral infection of the reproductive tract. Cervical cancer is by far the most common HPV-related disease. Nearly more cases of cervical cancer can be attributable to HPV infection. This study designed foe molecular detections and genotyping of human papilloma virus (HPV) by designing methods - Endpoint real time PCR for HPV detections and genotyping of HPV 16 and 18. The study includes 90 Pap smear samples of women suffering from abnormal Pap smears and 10 healthy women as a control, were collected from the Baghdad hospital (Medical City), Oncology hospital and Al-Amal hospital (in Baghdad), during the period from April, 2015 to April, 2016. The age of women was ranged from 16 to 60 years. The molecular detection method (Designed PCR) in our study was showed that the 35% of women have HPV infections.

Introduction:-

Cervical cancer is a cancer arising from the cervix. It is due to the abnormal growth of cells that have the ability to invade or spread to other parts of the body. At the beginning of the process the neoplastic lesions are confined to the epithelium where there no symptoms. Later symptoms may include abnormal vaginal bleeding, pelvic pain or pain during sexual intercourse [1].

Although the incidence of cervical cancer in Iraq is relatively low (Iraqi Cancer Board, 2012) as in most other Islamic countries, most of the cases usually present in late advanced stages. Studies from Iraq have reported significant rates of CIN lesions among Iraqi patients complaining of gynecological problems [2] [3]. Therefore, promoting the level of awareness among the Iraqi population through screening is mandatory to control the disease.

Human papilloma virus (HPV) infection appears to be involved in the development of more than 90% of cases; most people who have had HPV infections, however, do not develop cervical cancer. Other risk factors include smoking, a weak immune system, birth control pills, starting sex at a young age and having many sexual partners, but these are less important [4].

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The aims of this study is molecular detections of HPV-HR and genotyping by using Designed methods and.

Materials and Methods:

Subjects:
This study was carried out during the period between the 1st of April 2015 until the end of April 2016 within the University of Baghdad / Institute of Genetic Engineering & Biotechnology for post Graduate Studies and the National Cancer Research Center. One study group has been investigated:

Study group (With abnormal Pap smear):
This study has included 100 Iraqi women with different gynecologic complaints whose Pap smears revealed abnormal cytological findings. The patients were examined in the Outpatient Gynecology department of the Medical City Teaching Hospital. For each patient, a structured questionnaire containing different variables was completed.

A Healthy Control Group (a sub group from the study population) was categorized to consist of 10 healthy Iraqi women with different ages; all of whom were chosen depending on the following criteria [5]:
1. Having regular menstrual cycles in women with reproductive age (26 to 30 days)
2. Age range between 25 to 60 years
3. No history of endocrine disease
4. No use of medication or oral contraceptives.

Samples collection:
Examined specimens were collected cells from the cervix of each women in both Patients and Healthy Control Groups.

The Papanicolaou test (abbreviated as Pap test) is a method of cervical screening used to detect potentially precancerous and cancerous processes in the cervix.

The collected cells are examined under a microscope to look for abnormalities. The test aims to detect potentially pre-cancerous changes (called cervical intraepithelial neoplasia (CIN), cervical dysplasia or squamous intraepithelial lesion system (SIL) according to the Bethesda Terminology. These lesions are often caused by sexually transmitted human papilloma viruses [6].

Genomic DNA isolation:
Total DNA (genomic, mitochondrial, and viral) isolated from the Pap smear sample for molecular studies was applied using genomic DNA purification kits of Qiagen ( QIAamp DNA Mini Kit / Germany). This kit has been used by other researchers in similar study [7].

Agarose Gel Electrophoresis:
After genomic DNA extraction, agarose gel electrophoresis was adopted to confirm the presence and integrity of the extracted DNA [8].

Multiplex Polymerase Chain Reaction (PCR) for HPV detections:

Primer selections:
Multiplex polymerase Chain Reaction (PCR) for HPV type 16 and 18 - E6 gene detections were done by using a specific primer designed in our study.

The detections of L1 region for HPV as general (HPV- high risk) by using a specific primers [9].

The positive control for human beta-globin gene used in this study by using a specific primers (TAKARA company, Japan). Primers sequences are listed below, table (1).

Table (1): Primer sequences used in PCR amplification of E6-HPV16 and 18, L1 of HR-HPV and human beta-globin gene.

| Set  | Name            | Sequence (5' → 3')                  | Gene amplified | size  |
|------|-----------------|------------------------------------|----------------|-------|
| 1a   | MZNMI_F         | TTAGCTTTGCGGGATTTATGC              | E6-HPV16       | 204 bp|
|      | MZNMI_R         | CAGGACACAGTGGCTTTTGA               |                |       |
Primers have been designed in this study based on the Bioinformatics tools by using the international databases (NCBI, EMBL-EBI and DDBJ) and a number of tools that are available on website (online tools and software).

PCR Reaction setups:-
Multiplex PCR was performed in a 30 µl total volume, as described in Table (2).

Table 2:- PCR reaction setup.

| Component                                      | Reaction volume (µl) |
|------------------------------------------------|----------------------|
| PCR Mastermix (Biland):TaqDNA polymerase , dNTPs, MgCl2 and reaction buffer (pH 8.5) | 12 (µl)              |
| Primer forward                                 | 1 µl, total=3 µl     |
| Primer reverse                                 | 1 µl, total=3 µl     |
| Template DNA                                   | 4 (µl)               |
| RNase–free water                               | 8 (µl)               |
| Total reaction volume                          | 30 (µl)              |

Cycling Conditions:-
Cycling parameters for multiplex PCR are presented in Table (3).

Table (3):- Cycling conditions for multiplex PCR.

| Steps            | Temperature | Time   | No. of cycles |
|------------------|-------------|--------|---------------|
| Denaturation 1   | 95°C        | 3 min. | 1 cycle       |
| Denaturation 2   | 94°C        | 1 min  | 35 cycles     |
| Annealing        | 55°C        | 1 min  |               |
| Extension        | 72°C        | 1 min  |               |
| Final extension  | 72°C        | 5 min. | 1 cycle       |

End point RT-PCR for HPV detections:-
The last part of the designed PCR was end point RT-PCR for HPV detections, This part was conducted for the purpose of high precision and to make sure the true positive and true negative results.

The end point RT-PCR for HPV detections part includes the same targets of the multiplex PCR (E6 gene for each HPV type 16 and type 18, L1 gene for HPV-HR and human beta globin) but it was replaced PCR Mastermix (Biland) with SYBR green Mastermix (Biland) for real time PCR with same Cycling conditions for each gene (Table 4).

Table (4):- End point RT-PCR reaction setup.

| Component                   | Reaction volume (µl) |
|-----------------------------|----------------------|
| RT-PCR SYBR green Mastermix| 10 (µl)              |
| Primer forward              | 1 µl                 |
| Primer reverse              | 1 µl                 |
| Template DNA                | 4 (µl) , (0.7- 3.6µg)|
| RNase–free water            | 4 (µl)               |
| Total reaction volume       | 20 (µl)              |
Results and Discussion:

Subjects data:
Samples of cervical cells were collected by the examining doctor from 100 Iraqi individuals through Pap smear tests. The study population was divided into two subgroups:

Sub group I (Patient Group) comprised 90 patients who visited the Gynecology Oncology clinic with different gynecologic complaints.

Sub group II (Healthy Control Group) included 10 healthy individuals with normal Pap smear findings.

Age distribution of the samples:
The age of all women was categorized as those who were less than 30 years old versus those equal to and over 30. (Table 5) revealed that 28% of the study women in general were less than 30 years old while 72% were equal or over 30. This points out that may leads to conclude that group 2 constituted the greatest number groups in the present study. They were two subgroups in this study, the first group the women with abnormal pap smear revealed that 28.88% of the women were less than 30 (<30) years and followed by 71.12% of women whose age about more than 30 years old (>30) years. The second group the healthy women (with normal pap smear) revealed that 20% of the women were about (<30) years and followed by 80% of women whose age about (>30) years. The results agreed with those reported by Wright (2014) who displayed that the aging is a risk factor for persistent infection. The rate of persistent high-risk infection for women older than age 40 is 50%, compared with a persistence rate of 20% in women younger than age 25.

Table (5): Distribution of the Study Group according to Age.

| Group NO. | Age group (years) | Study group | The sub groups |
|-----------|-------------------|-------------|---------------|
|           |                   | No. | %  | Women with abnormal pap smear | Healthy women (Normal) |
|           |                   | No. | %  | No. | %  | No. | %  |
| 1         | Less than 30      | 28  | 28 | 26  | 28.88 | 2   | 20  |
| 2         | More than 30      | 72  | 72 | 64  | 71.12 | 8   | 80  |
| Total     |                   | 50  | 100 | 90  | 100   | 10  | 100 |
Pap smear and sample collections:-
The distribution of sample study according to the histology examination is show in table (6) while the distribution
according to the cytology examination is illustrated in table (7).

Table 6:- Distribution of the Study Population according to Histological examination.

| Histology | Number | Percentage (%) |
|-----------|--------|----------------|
| WNL       | 40     | 40.00          |
| CIN I     | 26     | 26.00          |
| CIN II    | 12     | 12.00          |
| CIN III   | 9      | 9.00           |
| Carcinoma | 3      | 3.00           |
| Normal    | 10     | 10.00          |
| Total     | 100    | 100%           |

Chi-square value --- 11.926 **
P-value --- 0.00496

** (P<0.01).

WNL: With normal limited and CIN: Cervical intraepithelial neoplasia.

Table 7:- Distribution of sample study according to cytology.

| Cytology  | Number | Percentage (%) |
|-----------|--------|----------------|
| ASC-US    | 21     | 21.00          |
| HSIL      | 18     | 18.00          |
| LSIL      | 20     | 20.00          |
| WNL       | 38     | 38.00          |
| Carcinoma | 3      | 3.00           |
| Total     | 100    | 100%           |

Chi-square value --- 9.337 **
P-value --- 0.00783

** (P<0.01).

WNL: With normal limited, ASC-US: Atypical squamous cells of undetermined significance, LSIL: Low-grade squamous intraepithelial lesion and HSIL: High-grade squamous intraepithelial lesion.

Fifty years ago, for squamous histology, the cervical cellular abnormalities viewed as the precursor of cervical cancer were termed mild, moderate or severe dysplasia; severe dysplasia was distinguished from the more severe diagnosis of carcinoma in situ. In the late 1960s, Richart proposed the concept of intraepithelial neoplasia [9]. CIN3 encompassed severe dysplasia and carcinoma in situ, CIN2 replaced moderate dysplasia, and CIN1 later came to include both the cytologic evidence of HPV infection (koilocytotic atypia) and mild dysplasia. The severity of the diagnosis was based on the degree of replacement of the normal stratified epithelium with mitotically active basal-like epithelium (≤1/3 = CIN1, ≤2/3 = CIN2, >2/3 = CIN3). CIN was viewed as a stepwise progression, with a high probability of transition from the more minor to more serious cancer precursors [10].

Designed PCR - Multiplex Polymerase Chain Reaction and End point RT-PCR for HPV detections:-
The present study used Multiplex Polymerase Chain Reaction and end point RT-PCR technique for HPV detections. The multiplex PCR results revealed that identical bands related to the (E6) gene for HPV type 16 and type 18. L1 region for HPV high risk and human beta globin were present. (E6-HPV16) PCR amplified regions showed a molecular weight of 204 bp, (E6-HPV16) showed a molecular weight of 151 bp, (L1-HR-HPV) showed 450 bp and human beta globin about 345 bp (Figure 8).

Emanuella et al.,(2014) showed that the L1 HPV conventional PCR products were 450 bp and their findings were in line with the results of this study [11].

PCR is being increasingly used in clinical laboratories to diagnose HPV. In another study that used conventional PCR for HPV detections a wide range. Studies in the literature indicate variables rates of HPV detection. This differences in detecting DNA of HPV suggest a potential difference in the ability to amplify fragments of different sizes and specific types of HPV, depending on the methods used for DNA detection, and may also be attributable to
the difference in types of studied material (smears, frozen material, paraffin material), anatomical localization, population issue, and design of oligonucleotides [12].

Figure (1):- Multiplex PCR products of E6 gene for each of (HPV type 16 and type 18), L1 region – HPV-HR and human beta globin on 2% agarose gel at 75 voltages for (45-50) min. 
M= Marker (100 bp ladder), -C = Negative control , +C = Positive control and IC = Internal control

The last part of the designed PCR was (End point RT-PCR for HPV detections), This part was conducted for the purpose of high precision and to evaluate and confirm the true positive and true negative results.

The end point RT-PCR for HPV detections part includes the same targets of the multiplex PCR (E6 gene for each HPV type 16 and type 18, L1 gene for HPV-HR and human beta globin) (Figure 2) (Table 8).

Table 8: The keys of End point RT-PCR Run Results .

| No. | Colour | Name                  | Type        | Genotype | Cycling A.Green |
|-----|--------|-----------------------|-------------|----------|----------------|
| 1   | red    | +C                    | Positive Control | Reaction |
| 2   | green  | - C                   | Negative Control | No Reaction |
| 3   | blue   | P1-HPV 18 – E6        | Unknown     | Reaction |
| 4   | purple | P1-HPV 16 – E6        | Unknown     | No Reaction |
| 5   | pink   | P1-HPV HR – L1        | Unknown     | Reaction |
| 6   | blue   | P1-Human B globin     | Unknown     | Reaction |
| 7   | green  | P2-HPV 18 – E6        | Unknown     | No Reaction |
| 8   | red    | P2-HPV 16 – E6        | Unknown     | Reaction |
| 9   | green  | P2-HPV HR – L1        | Unknown     | Reaction |
| 10  | red    | P2-Human B globin     | Unknown     | Reaction |
| 11  | red    | P3-HPV 18 – E6        | Unknown     | No Reaction |
| 12  | red    | P3-HPV 16 – E6        | Unknown     | No Reaction |
| 13  | red    | P3-HPV HR – L1        | Unknown     | Reaction |
| 14  | red    | P3-Human B globin     | Unknown     | Reaction |
| 15  | red    | P4-HPV 18 – E6        | Unknown     | No Reaction |
It was illustrated that four negative results (in multiplex PCR) turned out to be positive (in end point RT-PCR), and therefore the total number of positive results by using multiplex PCR technique was 31/100. On the other hand, the total number of positive results by using end point RT-PCR technique was 35/100. The number and percentage of positive and negative results are shown in table (9).

The E6 gene for HPV 16 and HPV 18 was screened by sequencing from women samples. The determination of the sequences were for the purpose of proving the identity of the virus (HPV 16 and HPV 18) with international references.

All results were directly compared with human reference – E6 sequences (http:NCBI Reference Sequence) directly. The E6 gene for HPV 16 and HPV 18 sequences have been published in the international databases:
1. NCBI (National Center for Biotechnology Information)
2. ENA-EBI (European Nucleotide Archive)
3. DDBJ (DNA Data Bank of Japan).

**Table (9):** The number and percentage of results by using multiplex PCR and End point RT-PCR for HPV detections.

| Results | multiplex PCR | End point RT-PCR |
|---------|---------------|------------------|
|         | No. | %   | No. | %   |
| Positive| 31   | 31.00 | 35  | 35.00 |
| Negative| 69   | 69.00 | 65  | 65.00 |
| Total   | 100  | 100   | 100 | 100   |

Real-time PCR is carried out in a thermal cycler with the capacity to illuminate each sample with a beam of light of at least one specified wavelength and detect the fluorescence emitted by the excited fluorophore. The thermal cycler is also able to rapidly heat and chill samples, thereby taking advantage of the physicochemical properties of the nucleic acids and DNA polymerase. Then the reaction is run in a real-time PCR instrument, and after each cycle, the intensity of fluorescence is measured with a detector; the dye only fluoresces (such as dsDNA dyes - SYBR Green) will bind to all dsDNA PCR products. This method has the advantage of only needing a pair of primers to carry out the amplification, which keeps costs down and more sensitivity and high accuracy for real time PCR than conventional PCR [12].

**Conclusions:**
1. In this study a new method has been developed of high precision and more sensitivity for early detection of human papilloma virus (HPV) the main cause of cervical cancer compared with the global diagnosis kit (Sacace, Italy).
2. The study found variation in the results of PCR (RT-PCR/Sacace, Italy and designed PCR) for HPV detection depending on the histological and Cytological examinations.
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