# 36180109715

### Ms. ANGELA WOOLLEY

PID NO: P36180009352

Age: 30 Year(s) Sex: Female

### Reference:

Sample Collected At: GILEAD MEDICAL & DENTAL CENTER HOUSE NO BALB NO C896/3,KANDA HIGHWAY NORTH RIDGE,ACCRA-14911. 014911 VID: 36180109715

Registered On: 23/07/2018 05:20 PM Collected On: 23/07/2018 Reported On:

Reported On 30/07/2018 04:47 PM

| <u>Investigation</u>                       | Observed Value | <u>Unit</u> | <b>Biological Reference Interval</b>  |
|--|----------------|-------------|---|
| FSH - Follicle Stimulating Hormone (Serum) | 6.80           | mIU/mL      | Normal Menstruating Women<br>Follicular Phase: 3.0 - 12.0<br>Mid Cycle Phase: 8.0 - 22.0<br>Luteal Phase: 2.0 - 12.0<br>Post Menopausal: 35.0 - 151.0 |
| LH- Luteinizing hormone (Serum,CMIA)       | 4.50           | mIU/mL      | Follicular phase: 2.4-12.6<br>Midcycle peak: 14.0-95.66<br>Luteal phase: 1.0-11.4<br>Post menopausal: 7.7-58.5<br>Post Menopausal: 7.7-58.2           |

# **Interpretation:**

Intact PTH has been demonstrated to be labile and is susceptible to fragmentation. This instability depends on both time and temperature . In room temperature EDTA sample stability is 8 hours and serum is for 4 hours. At 4degree C. EDTA sample stability is 72 hours and serum is for 48 hours.

# **E2 Estradiol Serum**

(Serum, CMIA)

**E2 - Estradiol level** 42.00 pg/mL Follicular phase: 12.5-166

Ovulating: 85.8-498 Luteal phase: 43.8-211 Post Menopausal: 5-54.7



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# **HPV DNA Detection And Typing**

Test Principle : PCR - Sequencing

Specimen : Material In LBC Container

Result :

| HPV DNA Detection | NOT DETECTED |  |
|-------------------|--------------|--|
| HPV GENOTYPE      |              |  |

# **Result Interpretation:**

- A "NOT DETECTED" result indicates the absence of HPV virus in the specimen.
- A "DETECTED" result indicates the presence of HPV virus in the specimen.
- All the result should always be correlated by clinical status and history of the patient.

# Clinical Background:

- · Molecular detection of HPV DNA is currently the gold standard for identification of HPV.
- The viral DNA is amplified in vitro by DNA polymerase to generate adequate amount of target, which is then directly visualized on the gel and sequenced to detect the specific genotype.
- The sensitivity of PCR based method is about 100 HPV viral genomes in a background of 100ng cellular DNA with a specificity of >98%. An internal control of 268bp is run for every sample to validate the assay.

# **Clinical Utility:**

- Over 120 HPV types have been identified and over 30 types are transmitted sexually making HPV the most common sexually transmitted disease (STD).
- High-risk HPV includes 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58,59, 68, 69 of which type 16 and 18 cause about 70 percent of cervical cancers.
- However, the low-risk types which rarely develop into cancer include 6, 11, 40, 42, 43, 44, 53, 54, 61, 72, 73 and 81 of which type 6 and 11 are linked to about 90 percent of genital warts.
- HPV status and the genotype involved in infection have an important clinical significance.
- Persistent infection of specific types of high risk HPV is essential for progression of cervical lesions that are likely to develop cancer.

**Limitation of the Assay:**Presence of PCR inhibitors in the sample prevents DNA amplification for HPV detection. Unknown risk genotypes are identified by this assay.

**Note:** This test has been developed and its performance validated at Molecular Biology Department, Metropolis Healthcare Ltd.

Reference: Husman et. al., 1995. Journal of General Virology, 76:1057-1062.;

Winder et. al., 2009. BMC Cancer, 9:440.

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Dr. Niranjan Patil MD( Micro)

HOD - Microbiology & Molecular Biology



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Reports to follow - Kindly await following pending reports :

<u>Investigation</u>: <u>Status</u>

PAP smear Liquid based cytology (LBC)

-- End of Report --

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