



Garg Pathology


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National Accreditation Board For Testing & Calibration Laboratories
ISO 9001:2008
Garden House Colony, Near Nai Sarak, Garh Road, Meerut
Ph.: 0121-2600454, 8979608687, 9837772828

DR. MONIKA GARG

M.D. (Path) Gold Medalist

Former Pathologist :

St. Stephan's Hospital, Delhi

PUID : 220319/605 **C. NO:** 605 **Collection Time** : 19-Mar-2022 10:50AM
Patient Name : Mrs. HIMANI JAIN 35Y / Female **Receiving Time** : 19-Mar-2022 10:55AM
Referred By : Dr. BANK OF BARODA **Reporting Time** : 19-Mar-2022 12:25PM
Sample By : **Centre Name** : Garg Pathology Lab - TPA
Organization : 

Investigation	Results	Units	Biological Ref-Interval
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HAEMATOLOGY (EDTA WHOLE BLOOD)

COMPLETE BLOOD COUNT

HAEMOGLOBIN (Colorimetry)	10.1	gm/dl	12.0-15.0
TOTAL LEUCOCYTE COUNT (Electric Impedence)	7280	*10 ⁶ /L	4000 - 11000
DIFFERENTIAL LEUCOCYTE COUNT (Microscopy)			
Neutrophils	64	%.	40-80
Lymphocytes	30	%.	20-40
Eosinophils	04	%.	1-6
Monocytes	02	%.	2-10
Absolute neutrophil count	4.66	x 10 ⁹ /L	2.0-7.0(40-80%)
Absolute lymphocyte count	2.18	x 10 ⁹ /L	1.0-3.0(20-40%)
Absolute eosinophil count	0.29	x 10 ⁹ /L	0.02-0.5(1-6%)

Method:-((EDTA Whole blood,Automated /

RBC Indices

TOTAL R.B.C. COUNT (Electric Impedence)	3.88	Million/Cumm	4.5 - 6.5
Haematocrit Value (P.C.V.)	31.3	%	26-50
MCV (Calculated)	80.7	fL	80-94
MCH (Calculated)	26.0	pg	27-32
MCHC (Calculated)	32.3	g/dl	30-35
RDW-SD (Calculated)	43.5	fL	37-54



*THIS TEST IS NOT UNDER NABL SCOPE

Checked By Technician:

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MBBS, MD(Path)
(Consultant Pathologist)

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




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RDW-CV (Calculated)	13.0	%	11.5 - 14.5
Platelet Count (Electric Impedence)	2.27	/Cumm	1.50-4.50
MPV (Calculated)	9.4	%	7.5-11.5
GENERAL BLOOD PICTURE			
NLR 6-9 Mild stres 7-9 Pathological cause	2.13		1-3

-NLR is a reflection of physiologic stress,perhaps tied most directly to cortisol and catecholamine levels.
 -NLR can be a useful tool to sort out patients who are sicker, compared to those who are less sick (its not specific to infection).
 -NLR has proven more useful than white blood cell count (WBC) when the two are directly compared. Ultimately, NLR may be a logical replacement for the WBC. In some situations, NLR is competitive with more expensive biomarkers (e.g. procalcitonin,lactate).
 -With specific clinical contexts (e.g. pancreatitis, pulmonary embolism), NLR may have surprisingly good prognostic value.

BLOOD GROUP * "B" POSITIVE \$ \$



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GLYCATED HAEMOGLOBIN (HbA1c)*	4.8	%	4.3-6.3
ESTIMATED AVERAGE GLUCOSE	91.1	mg/dl	

EXPECTED RESULTS :

Non diabetic patients & Stabilized diabetics	: 4.3% to 6.30%
Good Control of diabetes	: 6.4% to 7.5%
Fair Control of diabetes	: 7.5% to 9.0%
Poor Control of diabetes	: 9.0 % and above

-Next due date for HBA1C test : After 3 months

-High HbF & Trig.level, iron def.anaemia result in high GHb

-Haemolytic anemia, presence of HbS, HbC and other Haemoglobinopathies may produce low values. three months.

INTERPRETATION: HbA1c is an indicator of glycemic control.HbA1c represents average glycemia over the past six to eight weeks.Glycation of hemoglobin occurs over the entire 120 day life span of the red blood cell, but with in this 120 days. Recent glycemia has the largest influence on the HbA1c value. Clinical studies suggest that a patient in stable control will have 50% of their HbA1c formed in the month before sampling, 25% in the month before that, and the remaining 25% in months two to four. Mean Plasma Glucose mg/dl = (HbA1c x 35.6) - 77.3 Correlation between HbA1c and Mean Plasma Glucose (MPG) is not "perfect" but rather only this means that to predict or estimate average glucose from Hb-A1c or vice-versa is not "perfect" but gives a good working ballpark estimate. Afternoon and evening results correlate more closely to HbA1c than morning results, perhaps because morning fasting glucose levels vary much more than daytime glucose levels, which are easier to predict and control.

As per IFCC recommendations 2007, HbA1c being reported as above maintaining traceability to both IFCC (mmol/mol) & NGSP (%) units.

BIOCHEMISTRY (FLORIDE)

PLASMA SUGAR FASTING 92.0 mg/dl 70 - 110
(GOD/POD method)

PLASMASUGAR P.P. 141.0 mg/dl 80-140
(GOD/POD method)

BIOCHEMISTRY (SERUM)

SERUM CREATININE 0.7 mg/dl 0.6-1.4
(Enzymatic)

BLOOD UREA NITROGEN 15.40 mg/dL. 8-23



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




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LIVER FUNCTION TEST

SERUM BILIRUBIN

TOTAL 0.7 mg/dl 0.1-1.2
(Diazo)

DIRECT 0.3 mg/dl <0.3
(Diazo)

INDIRECT 0.4 mg/dl 0.1-1.0
(Calculated)

S.G.P.T. 27.0 U/L 8-40
(IFCC method)

S.G.O.T. 29.1 U/L 6-37
(IFCC method)

SERUM ALKALINE PHOSPHATASE 88.0 IU/L 37-103
(IFCC KINETIC)

SERUM PROTEINS

TOTAL PROTEINS 6.9 Gm/dL 6-8
(Biuret)

ALBUMIN 3.8 Gm/dL 3.5-5.0
(Bromocresol green Dye)

GLOBULIN 3.10 Gm/dL 2.5-3.5
(Calculated)

A : G RATIO 1.23 1.5-2.5
(Calculated)



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LIPID PROFILE

SERUM CHOLESTEROL (CHOD - PAP)	155.0	mg/dl	150-250
SERUM TRIGYCERIDE (GPO-PAP)	97.0	mg/dl	70-150
HDL CHOLESTEROL * (PRECIPITATION METHOD)	46.0	mg/dl	30-60
VLDL CHOLESTEROL * (Calculated)	19.4	mg/dl	10-30
LDL CHOLESTEROL * (Calculated)	89.6	mg/dL.	0-100
LDL/HDL RATIO * (Calculated)	01.9	ratio	<3.55
CHOL/HDL CHOLESTROL RATIO* (Calculated)	3.4	ratio	3.8-5.9

Interpretation :

Patient Should be Fast overnight For Minimum 12 hours and normal diet for one week

NOTE :

Lipid Profile Ranges As PER NCEP-ATP III :

SERUM CHOESTEROL : Desirable : < 200 Borderline : 200 - 239 Elevated : > 240 mg/dl
HDLCHOLESTEROL : Desirable : > 60 Borderline : 40- 60 Decreased : < 40 mg/dl
LDL CHOLESTEROL : Desirable : 100 mg/dl, Borderline : 100- 159 Elevated : >160 mg/dl
Triglycerides : Desirable : 150 Borderline : 150- 199 High : 200 - 499 Very High : >500

Friedwald Equation, VLDL & LDL values are not applicable for triglyceride > 400 mg/dl.

SERUM SODIUM (Na) * 139.0 mEq/litre 135 - 155
(ISE method)
(ISE)



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




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THYROID PROFILE*

Triiodothyronine (T3) * (ECLIA)	1.101	ng/dl	0.79-1.58
Thyroxine (T4) * (ECLIA)	9.422	ug/dl	4.9-11.0
THYROID STIMULATING HORMONE (TSH) * (ECLIA)	1.982	uIU/ml	0.38-5.30

Hyperthyroid patient have suppressed TSH values, with the exception of those few individuals whos have hyperthyroidism caused by TSH producing pituitary tumor or other rare disorders such as pituitary resistance to thyroid hormones. Subclinical hyperthyroidism is defined as low TSH with levels of T4 and T3 within the reference interval. In most patients with hypothyroidism,serum TSH results are markedly elevated, but results are low in individuals with hypothyroidism caused by pituitary or hypothalamic disorders. An important cause of both incresed and decreased TSH results is NTI. Patients with NTI tend to have low TSH results during their acute illness ,then TSH rises to within or above the reference range with resolution of the underlying illness,and finally returns to within the reference range. The situation is complicated because drugs,including glucagon and dopamine,suppress TSH . Sensitive TSH assays are helpful in evaluation of treatment with thyroid hormone both for replacement therapy and suppressive doses for malignant thyroid disease.

SERUM POTASSIUM (K) * (ISE method)	3.8	mEq/litre.	3.5 - 5.5
SERUM CALCIUM (Arsenazo)	10.1	mg/dl	9.2-11.0

BIOCHEMICAL EXAMINATION

URIC ACID	3.2	mg/dL.	2.5-6.8
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




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CYTOLOGY EXAMINATION

SPECIMEN

Microscopic:

MG-134/22

SITE OF SMEAR: ECTOCERVIX AND POSTERIOR FORNIX OF VAGINA

METHOD OF EVALUATION: BETHSEDA SYSTEM

EVALUATION OF SMEAR : SATISFACTORY

REPORT: CELLULAR SPREAD SHOWS DESQUAMATED EPITHELIAL CELLS PREDOMINANTLY SUPERFICIAL AND INTERMEDIATE CELLS.

FEW ENDOCERVICAL CELLS SHOWING REACTIVE CHANGES ARE SEEN.

BACKGROUND SHOWS MILD INFLAMMATORY REACTION. LACTOBACILLI ARE SEEN. ANY DYSKARYOTIC CELL IS NOT SEEN.

ANY BUDDING SPORES OR TROPHOZOITE IS NOT SEEN.

INFERENCE: NEGATIVE FOR INTRAEPITHELIAL LESION OR MALIGNANCY

NOTE: This test has its own limitations. Please interpret the findings in light of clinical picture. not for medicolegal use



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




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CLINICAL PATHOLOGY

STOOL EXAMINATION

§PHYSICAL EXAMINATION

Colour	Brownish
Mucous	Absent
Consistency	Semi solid
PH	Acidic

Method:- Descriptive

§MICROSCOPIC EXAMINATION

R.B.C.s Nil

Method:-Microscopic

Pus Cells 0-2

Method:-Microscopic

Ova Nil

Method:-Microscopic

Cysts Nil

Method:-Microscopic

§SPECIAL EXAMINATION

Occult Blood Negative

Method:-Homospot Card Test



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URINE

PHYSICAL EXAMINATION

Volume	20	ml	
Colour	Yellow		
Appearance	Clear		Clear
Specific Gravity	1.020		1.000-1.030
PH (Reaction)	Acidic		

BIOCHEMICAL EXAMINATION

Protein	Nil		Nil
Sugar	Nil		Nil

MICROSCOPIC EXAMINATION

Red Blood Cells	Nil	/HPF	Nil
Pus cells	1-2	/HPF	0-2
Epithelial Cells	2-3	/HPF	1-3
Crystals	Nil		
Casts	Nil		
@ Special Examination			
Bile Pigments	Absent		
Blood	Nil		
Bile Salts	Absent		

-----{END OF REPORT }-----



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