







Patient	Mrs. KRISTINABEN AMBALAL VANKAR	Lab No/ManualNo	5356053/
UHIDNo/IPNO	300441899	CollectionDate	12/02/2024 10:31AM
Age/Gender	35 Years/Female	Receiving Date	12/02/2024 2:47PM
Bed No/Ward	OPD	Report Date	12/02/2024 3:47PM
Referred By	Dr. Casualty Medical Officer	Report Status Sample Quality	Final Normal

Test Name	Result	Unit	Bio. Ref. Range	Method	Sample
	ArcoFemi H	Biochemistry			
					Serum
Gamma GT	22.90	U/L	6.00 - 42.00	Enzymatic metho	bd
					Serum
Creatinine	0.62	mg/dL	0.50 - 0.90	Jaffe Kinetic Cor	npensated
					Serum
Uric Acid	5.3	mg/dL	2.4 - 5.7	Uricase / Peroxid (Colorimetric)	dase
					Serum
Post prandial Glucose As per ADA Guideline For Fasting Plasma Glucose	119.6	mg/dL	< 140.0	Hexokinase	
Normal : Less than 100 mg/dL Prediabetes : 100 mg/dL to 125 mg/dL Diabetes : 126 mg/dL or Higher For 2 hrs Plasma Glucose after 75 Gms (Normal : Less than 140 mg/dL Prediabetes : 140 to 199 mg/dL Diabetes : 200 mg/dL or higher For Random Plasma Glucose	Glucose load				
Diabetes is diagnosed at blood glucose grea	ater than or equa	al to 200 mg/dL			

Serum

Lepon

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Random Glucose	88.6	mg/dL	< 140.0	Hexokinase
As per ADA Guideline				
For Fasting Plasma Glucose				
Normal : Less than 100 mg/dL				
Prediabetes : 100 mg/dL to 125 mg/dL				
Diabetes : 126 mg/dL or Higher				
For 2 hrs Plasma Glucose after 75 Gms	Glucose load			
Normal : Less than 140 mg/dL				
Prediabetes : 140 to 199 mg/dL				
Diabetes : 200 mg/dL or higher				
For Random Plasma Glucose				
Diabetes is diagnosed at blood glucose grea	ater than or equal to	200 mg/dL		

LIVER FUNCTION TEST (LFT) SERUM

SGPT(ALT)	н	39.00	U/L	0.00 - 33.00	IFCC without pyridoxal phosphate
SGOT(AST)	Н	32.80	U/L	0.00 - 32.00	IFCC without pyridoxal phosphate
Alkaline Phosphatase		106.3	U/L	35.0 - 140.0	PNP-Standardize
Bilirubin Total		0.32	mg/dL	0.00 - 1.00	Diazo Method
Bilirubin Direct		0.14	mg/dL	0.00 - 0.20	Diazo Method
Bilirubin Indirect		0.18	mg/dL	0.00 - 1.10	Calculate from Total and Direct Billirubin
Protein Total		7.43	g/dL	6.40 - 8.20	Biuret Method
Albumin		4.58	g/dL	3.97 - 4.95	BCG Endpoint
Globulin		2.85	g/dL	2.20 - 3.50	Calculated
A/G Ratio		1.61	Ratio	0.90 - 2.80	Ratio

EDTA Blood

Serum

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L-Low H-High CH -Critical High CL - Critical Low

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Patient UHIDNo/IPNO Age/Gender Bed No/Ward Referred By	Mrs. KRISTINABEN AI 300441899 35 Years/Female OPD Dr. Casualty Medical C			Lab No/ManualNo CollectionDate Receiving Date Report Date Report Status Sample Quality	5356053/ 12/02/2024 10:31AM 12/02/2024 11:04AM 12/02/2024 3:47PM Final Normal	
HbA1c (Glyco Hb)		6.09	%	4.8 % - 5.9 % Norr 5.9 % - 7.0 % Goo diabetic Control 7.0 % - 10.00 % Fa Diabetic Control >10.0 % Poor diab Control	d air	
Mean Plasma Gluco	ose	139.3	mg/dL	80.0 - 140.0		Serum
Blood Urea BUN*		25.5 11.9	mg/dL mg/dL	16.6 - 48.5 6.0 - 20.0	Urease,Kinetic,GLDH Ureas with UV	Serum
TOTAL T3* TOTAL T4* THYROID STIMUL4	ATING HORMONE	1.180 7.440 1.580	ng/mL ug/dL uIU/mL	0.850 - 2.020 5.130 - 14.060 0.270 - 4.200	ECLIA. ECLIA. ECLIA.	

(*) Not in NABL Scope

End Of Report

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Bed No/Ward	OPD	Report Date	12/02/2024 3:47PM
Referred By	Dr. Casualty Medical Officer	Report Status Sample Quality	Final Normal

Test Name	Result	Unit	Bio. Ref. Range	Method	Sample
	ArcoFemi I	Biochemistr Healthcare Ltd B			
LIPID PROFILE (WITH DIRECT LDL					Serum
Sample Type	Random				
Cholesterol Total	161.80	mg/dL	Less than 160 mg/dL Excellent Less than 200 mg/dL Desirabale 200-239 mg/dL Borderline High 240 mg/dl & over high	Enzymatic (CHE/CHO/POD)	
Triglycerides	109.00	mg/dL	Less than 150 mg/dL Normal 150-199 mg/dL Borderline High 200-499 mg/dL High 500 mg/dL or greater very HIgh	GPO-PAP	
HDL Cholesterol	33.10	mg/dL	Less than 40 mg/dL Low 60 mg/dL or Above Excellent	Homogenous Enz	ymatic
LDL Cholesterol (Direct)	114.10	mg/dL	Less than 80 mg/dL Excellent Less than 100 mg/dL Optimal 100-129 mg/dL Near or above optimal 130-159 mg/dL Borderline High 160-189 mg/dL High 190 mg/dL & above Very High	Homogenous Enz	ymatic
VLDL Cholesterol	21.8	mg/dL	< 30	Calculated	
LDL/HDL RATIO	3.45	-	< 3.50	Calculated	
Cholesterol Total / HDL Ratio	H 4.89		< 4.50		

Lego

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L-Low H-High CH -Critical High CL - Critical Low









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Bed No/Ward	OPD	Report Date	12/02/2024 3:47PM
Referred By	Dr. Casualty Medical Officer	Report Status	Final
		Sample Quality	

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UHIDNo/IPNO	300441899	CollectionDate	12/02/2024 10:31AM
Age/Gender	35 Years/Female	Receiving Date	12/02/2024 12:29PM
Bed No/Ward	OPD	Report Date	12/02/2024 2:14PM
Referred By	Dr. Casualty Medical Officer	Report Status	Final
		Sample Quality	

Result	Unit	Bio. Ref. Range	Method	Sample
ArcoFemi He	ealthcare Ltd E	Below 40 Female		
				Urine
20 ml			Visual method	
Pale Yellow			Visual method	
Slightly Hazy			Visual method	
5			Reflectance ph	otometer
1.015		1.015 - 1.030	Reflectance photoe reaction	otometer/Enzymatic
			Reflectance photometer/Ma	nual
Nil				
Absent				
Absent				
Absent				
			Microscopy	
Occasional		/H.P.F.		
1-2		/H.P.F.		
1-2		/H.P.F.		
Absent				
	20 ml Pale Yellow Slightly Hazy 5 1.015 Nil Absent Absent Absent 1-2 1-2 Absent Absent Absent Absent	Clinical Patho ArcoFemi Healthcare Ltd E 20 ml Pale Yellow Slightly Hazy 5 1.015 Nil Absent Absent Absent Absent 1-2 1-2 Absent Absent Absent Absent Absent Absent	Clinical Pathology ArcoFemi Healthcare Ltd Below 40 Female	Clinical Pathology ArcoFemi Healthcare Ltd Below 40 Female 20 ml Visual method Pale Yellow Visual method Slightly Hazy Visual method 5 Reflectance ph 1.015 1.015 - 1.030 Reflectance ph reaction Reflectance photometer/Ma Nil Absent Absent Microscopy Occasional /H.P.F. 1-2 /H.P.F. 1-2 /H.P.F. Absent Absent Absent Absent Absent Absent Absent Absent Absent Absent Absent Absent

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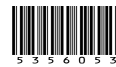
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Age/Gender	35 Years/Female	Receiving Date	12/02/2024 11:04AM
Bed No/Ward	OPD	Report Date	12/02/2024 12:57PM
Referred By	Dr. Casualty Medical Officer	Report Status	Final
		Sample Quality	

Test Name		Result	Unit	Bio. Ref. Range	Method	Sample
		Amore II	Haematology	our 40 Eomol-		
		ArcoFemi He	ealthcare Ltd Bel	ow 40 Female		
<u>CBC WITH ESR</u>						EDTA Blood
Haemoglobin		13.2	g/dL	12.5 - 16.0	SLS Method	
Hematocrit/PCV		42.3	%	37.0 - 47.0	H.focusing Meth	nod
RBC COUNT		5.12	mill/Cmm	4.20 - 5.40	H.focusing impe	edance
MCV	L	82.6	fl	83.0 - 101.0	Calculated	
МСН	L	25.8	pg	27.0 - 31.0	Calculated	
МСНС	L	31.2	g/dL	32.0 - 36.0	Calculated	
RDW-CV		13.5	%	11.5 - 14.0	Calculated	
Platelet count		369000	/cumm	150000 - 410000	H.focusing impe	edance
Mean Platelet Volume(MPV)*		10.7	fl	8 - 12	Calculated	
Total Leucocyte Count (TLC)		7800.00	/cumm	4000.00 - 10500.00	Flow Cytometry	
Differential Leucocyte Count					Flowcytometry/I	Microscopic
Neutrophils		65	%	40.0 - 70.0		
Lymphocytes		28	%	22 - 45		
Eosinophils		02	%	1.0 - 4.0		
Monocytes		05	%	1.0 - 6.0		
Basophils		00	%	0.0 - 1.0		
Immature Granulocytes		00	%	0 - 2		
Absolute Leucocyte Count						
Absolute Neutrophil Count*		5070	/cumm	1800 - 7700		
Absolute Lymphocyte count*		2184	/cumm	1000 - 4800		
Absolute Eosinophil Count (AEC)		156.0	/cumm	0.0 - 450.0		
Absolute Monocyte Count*		390	/cumm	0 - 800		
Peripheral Smear Study		RBCs are Normochromic & Normocytic.Platelets are adequate in number. Malarial Parasites are not seen.No Premature cells are seen.				
Erythrocyte Sedimentation Rate (ESR)	н	69	mm/hr	0 - 35	Photometric cap	

stopped flow kinetic 10-

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Age/Gender	35 Years/Female	Receiving Date	12/02/2024 2:10PM
Bed No/Ward	OPD	Report Date	12/02/2024 3:44PM
Referred By	Dr. Casualty Medical Officer	Report Status	Final
		Sample Quality	

CytoPathology

ArcoFemi Healthcare Ltd Below 40 Female

Cytopathology Pathology Report

Specimen Cervical PAP smear.

Clinical Diagnosis LMP: 26/01/2024, P2G2.

Gross Description Two fixed unstained slide received, PAP stain done.

Microscopic Description

Smears are satisfactory for evaluation. Many superficial, few intermediate cells and few parabasal cells seen. Severe inflammation with predominance of neutrophils seen. Mild lactobacilli are seen. No parasites/ fungi. No evidence of intraepithelial lesion or malignancy.

Diagnosis

Cervical smear - Severe inflammation and no evidence of intraepithelial lesion or malignancy.

Note- The pap test is a screening procedure to aid in the detection of cervical cancer and its precursors. Because false negative results may occur, regular PAP tests are recommended.

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Test Name	Result	Unit	Bio. Ref. Range	Method	Sample
		muno-Haemat lealthcare Ltd E	t ology Below 40 Female		
BLOOD GROUPING					EDTA Bloo
ABO Group	"A"			Tube Agglutina	tion Method

ABO Group Rh Type "A" Positive

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