



		Chopra gy & Microbiology) Consultant Pathologist	Dr. Yugam C MD (Pa CEO & Consultant Pa	ithology)
NAME	: Mrs. PRIYA			
AGE/ GENDER	: 27 YRS/FEMALE	PATI	ENT ID	: 1715709
COLLECTED BY	:	REG.	NO./LAB NO.	: 012501040030
REFERRED BY	: LOOMBA HOSPITAL (AM	BALA CANTT) REGI	STRATION DATE	: 04/Jan/2025 12:54 PM
BARCODE NO.	:01523426	COLI	ECTION DATE	: 04/Jan/2025 03:24PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPO	DRTING DATE	: 04/Jan/2025 03:46PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROA	AD, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
HAEMOGLOBIN (H by CALORIMETRIC INTERPRETATION:-	~,	11.6 ^L	gm/dL	12.0 - 16.0
Hemoglobin is the pr tissues back to the lu	ngs.		m the lungs to the body	rs tissues and returns carbon dioxide from th
A low nemoglobin lev ANEMIA (DECRESED	vel is referred to as ANEMIA o HAEMOGLOBIN):			
1) 6 - 1 / 1	matic injury, surgery, bleedin	ng, colon cancer or stomac	h ulcer)	
1) LOSS OF DIOOD (trat 2) Nutritional deficie		/		
 2) Nutritional deficie 3) Bone marrow prot 	lems (replacement of bone m	arrow by cancer)		
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by re- 	lems (replacement of bone m blood cell synthesis by cher	arrow by cancer) notherapy drugs		
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by res 5) Kidney failure 6) Abnormal hemogl 	lems (replacement of bone m d blood cell synthesis by cher bbin structure (sickle cell ane	notherapy drugs		
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by ree 5) Kidney failure 6) Abnormal hemogi POLYCYTHEMIA (INCI 	lems (replacement of bone m d blood cell synthesis by cher bbin structure (sickle cell ane REASED HAEMOGLOBIN):	notherapy drugs		
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by ree 5) Kidney failure 6) Abnormal hemogi POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda) 	lems (replacement of bone m d blood cell synthesis by cher bbin structure (sickle cell ane REASED HAEMOGLOBIN): Ititudes (Physiological) ry Polycythemia)	notherapy drugs emia or thalassemia).		
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by ree 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda 3) Dehydration prod 	lems (replacement of bone m d blood cell synthesis by cher bbin structure (sickle cell ane REASED HAEMOGLOBIN): Ititudes (Physiological) ry Polycythemia) uces a falsely rise in hemoglo	notherapy drugs mia or thalassemia). bin due to increased haem	oconcentration	
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by res 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda 3) Dehydration prod 4) Advanced lung dis 5) Certain tumors 	lems (replacement of bone m d blood cell synthesis by cher obin structure (sickle cell ane REASED HAEMOGLOBIN): Ititudes (Physiological) ry Polycythemia) uces a falsely rise in hemoglo ease (for example, emphysem	notherapy drugs mia or thalassemia). bin due to increased haem a)	oconcentration	
 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by ref 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda 3) Dehydration prodid 4) Advanced lung dis 5) Certain tumors 6) A disorder of the k 7) Abuse of the drug 	lems (replacement of bone m d blood cell synthesis by cher bbin structure (sickle cell ane REASED HAEMOGLOBIN): lititudes (Physiological) ry Polycythemia) uces a falsely rise in hemoglo ease (for example, emphysem one marrow known as polycy	notherapy drugs mia or thalassemia). bin due to increased haem a) themia rubra vera, nletes for blood doping pur		nount of oxygen available to the body by

NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD





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DR.YUGAM CHOPRA CONSULTANT PATHOLOGIST MBBS , MD (PATHOLOGY)







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BARCODE NO.	:01523426	COLL	ECTION DATE	:04/Jan/202503:24PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPO	RTING DATE	:04/Jan/202504:19PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	CLINIC	CAL CHEMISTRY	/BIOCHEMISTI	RY
		GLUCOSE RAN		

A random plasma glucose level below 140 mg/dl is considered normal.
 A random glucose level between 140 - 200 mg/dl is considered as glucose intolerant or prediabetic. A fasting and post-prnadial blood test (after consumption of 75 gms of glucose) is recommended for all such patients.
 A random glucose level of above 200 mg/dl is highly suggestive of diabetic state. A repeat post-prandial is strongly recommended for all such patients. A fasting plasma glucose level in excess of 125 mg/dl on both occasions is confirmatory for diabetic state.



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BARCODE NO.	: 01523426		COLLECTION D		: 04/Jan/2025 03:24PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		REPORTING D	ATE	: 05/Jan/2025 11:11AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	AMBALA CANT'I			
Test Name		Value		Unit	Biological Reference interva
		ENDOC	RINOLOGY		
	QUADRU	PLE MARKE	R MATERNA	L SCREE	NING
QUADRUPLE MAR	<u>KER</u>				
PATEINT SPECIFI					
DATE OF BIRTH		8/08/19	97		
MATERNAL AGE		27.9		YEARS	
WEIGHT		60		Kg	
ETHNIC ORIGIN		ASIAN		U	ASIAN
H/O IVF		ABSENT			
H/O INSULIN DEPI	ENDANT DIABETES	ABSENT			
H/O SMOKING		ABSENT			
H/O TRISOMY 21 S	SCREENING	ABSENT			
ULTRA SOUND SCA	AN DETAILS				
DATE OF ULTRASO	NN .	04-01-20			
METHOD FOR GES by ultrasound sca	TATION AGE ESTIMATION	ULTRAS	OUND SCAN DE	TAILS	
FOETUS (NOS) by ULTRASOUND SCA		1			
GA ON THE DAY OF	F SAMPLE COLLECTION	15.3		WEEKS	
BIPARIETAL DIAM	ETER (BPD)	29.6		mm	26 - 52
QUADRUPLE TEST	- BIOCHEMICAL MARKERS				
ALPHA FETO PROT PRENATAL SCREE	NING: SERUM	32.4		ng/mL	
ESTRIOL (uE3) UN	escence immunoassay) CONJUGATED escence immunoassay)	0.93		ng/mL	
BETA HCG	ESCENCE IMMUNOASSAY)	51895		mIU/mL	
INHIBIN A by CLIA (CHEMILUMIN	ESCENCE IMMUNOASSAY)	184		pg/mL	
	an		hopra		

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					_
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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMI	BALA CANTT			
Test Name		Value	Unit	Biological Reference interval	
MULTIPLE OF MED	IAN (MOM) VALUES				
AFP MOM		0.99			
	SCENCE IMMUNOASSAY)	0.00			
. ,		0.99			
BETA HCG MOM		1.38			
	SCENCE IMMUNOASSAY)	1 1			
	SCENCE IMMUNOASSAY)	1.1			
TRISOMY 21 SCREE	NING (DOWNS SYNDROME) RI	ISK ASSESSME	NT		
TRISOMY 21 SCREEN		NEGATIVE	(-ve)	NEGATIVE (-ve)	
		1.1176 NEC	ATIVE (NO)		
	SK SCENCE IMMUNOASSAY)	1.1170 NEG	ATIVE (-ve)		
TRISOMY 21 BIOCHI		1:4525 NEGATIVE (-ve)		RISK CUT OFF 1:270	
		NECATIVE	(-VA)		
	SCENCE IMMUNOASSAY)	NEGATIVE	(-vc)		
TRISOMY 18 SCREET	NING RISK scence immunoassay)	< 1:10000 N	NEGATIVE (-ve)	RISK CUT OFF 1:100	
	ECTS SCREENING RISK ASSESS	MENT			
	ECT SCREENING RISK	NEGATIVE	(-ve)	RISK CUT OFF 1:50	
		< 1:10000 N	JEGATIVE (-ve)	RISK CUT OFF 1.50	
by CLIA (CHEMILUMINES	SCENCE IMMUNOASSAY)	. 1.100001			
BARCODE NO. CLIENT CODE. CLIENT ADDRESS Test Name MULTIPLE OF MEDI AFP MOM by CLIA (CHEMILUMINES) ESTRIOL (uE3) MOM by CLIA (CHEMILUMINES) BETA HCG MOM by CLIA (CHEMILUMINES) INHIBIN A MOM by CLIA (CHEMILUMINES) TRISOMY 21 SCREEN by CLIA (CHEMILUMINES) TRISOMY 21 AGE RIS by CLIA (CHEMILUMINES) TRISOMY 18 SCREEN by CLIA (CHEMILUMINES) SPINA BIFIDA/ANEN	: 01523426 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AMI IAN (MOM) VALUES SCENCE IMMUNOASSAY) I SCENCE IMMUNOASSAY) SCENCE IMMUNOASSAY) SCENCE IMMUNOASSAY) SCENCE IMMUNOASSAY) SCENCE IMMUNOASSAY) SK SCENCE IMMUNOASSAY) EMICAL RISK SCENCE IMMUNOASSAY) EMICAL RISK SCENCE IMMUNOASSAY) SK SCENCE IMMUNOASSAY) EMICAL RISK SCENCE IMMUNOASSAY) EMICAL RISK SCENCE IMMUNOASSAY) EMICAL RISK SCENCE IMMUNOASSAY) ECTS SCREENING RISK ASSESSI ECT SCREENING RISK SCENCE IMMUNOASSAY) NCEPHALY SCREENING RISK	CO BALA CANTT Value 0.99 0.99 1.38 1.1 SK ASSESSME NEGATIVE 1:1176 NEG 1:4525 NEG NEGATIVE < 1:10000 N	DILECTION DATE EPORTING DATE Unit NT (-ve) GATIVE (-ve) GATIVE (-ve) (-ve)	: 04/Jan/2025 03:24PM : 05/Jan/2025 11:11AM Biological Reference interval NEGATIVE (-ve) RISK CUT OFF 1:270 RISK CUT OFF 1:100	

INTERPRETATION:

1.Multiple marker serum has become standard tool used in obstetrica care to identify pregnancies that may have increased risk for certain birth defects such as NEURALTUBE DEFECTS (NTD'S), DOWN'S SYNDROME (TRISOMY 21) AND TRISOMY 18. The screen is performed by measuring analytes in maternal serum that are produced by the fetus and the placenta. The analytes values along with maternal demographic information such as age, weight, gestational age, diabetic status, and race are used together in mathematical model to derive risk estimate. 2. The laboratory establishes a specific cut off for each condition, which classifies each screen as either screen-positive or screen-negative. 3.A screen-positive result indicates that the value obtained exceeds the established cut off.

4. The estimated risk calculation and screen results are dependent on accurate information for gestation, maternal age, race, IDD, and weight. Inaccurate information can lead to significant alterations in the estimated risk. In particular, erroneous assessment of gestational age can result in false-positive or false-negative screen results. Because of its increased accuracy, we therefore recommend determination of gestational age by ultrasound, rather than by last menstural period (LMP), When possible.



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Test Name	Value	Unit	Biological Reference interva

4.A negative screen indicates a lower probability of having a baby with TRISOMY 21 , TRISOMY 18 and NEURAL TUBE DEFECTS, but does not completely exclude the possibility.

5.A positive screen on the contrary only indicates a higher probability of having a baby with TRISOMY 21, TRISOMY 18 and NEURAL TUBE DEFECTS, and needs confirmation by cytogenetic studies and/or level II scan.

NOTE:

1. Triplet and higher multiple pregnancies cannot be interpreted

2. The reportable range for Trisomy 21, Trisomy 18 and NTD : >1:50 to < 1:10000

3.TRISOMY 21: HIGH RISK: >1:50 - 1:250

4.TRISOMY 18: HIGH RISK: >1:50 - 1:100

5.NEURAL TUBE DEFECT (NTD'S): HIGH RISK: >1:50

6.Biological markers evaluated in this test have marked as H(HIGH) or L(LOW) since there is wide variation in Alpha Fetoprotein, HCG and Unconjugated Estriol ranges depending upon gestational age. "In Range" and "Out of Range" columns are not applicable for the parameters appearing in Multiple of Median (MoM) and Risk calcultion.

7.Individually, Alpha Fetoprotein or HCG or unconjugated Estriol levels do not correlate with risk assessment of Trisomy 18, Trisomy 21 or Neural Tube Defects





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Test Name		Value	Unit	Biological Reference interval
		CLINICAL PA	THOLOCY	
		UTINE & MICRO	DSCOPIC EXAMINA	ATION
PHYSICAL EXAMIN				
QUANTITY RECIEV	ED TANCE SPECTROPHOTOMETRY	10	ml	
COLOUR		AMBER YEL	LOW	PALE YELLOW
by DIP STICK/REFLEC TRANSPARANCY	CTANCE SPECTROPHOTOMETRY	CLEAR		CLEAR
	TANCE SPECTROPHOTOMETRY	OLEAR		
SPECIFIC GRAVITY	TANCE SPECTROPHOTOMETRY	1.01		1.002 - 1.030
CHEMICAL EXAMI				
REACTION		ACIDIC		
	TANCE SPECTROPHOTOMETRY	North		
PROTEIN by DIP STICK/REFLEC	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
SUGAR		Negative		NEGATIVE (-ve)
pH	TANCE SPECTROPHOTOMETRY	<=5.0		5.0 - 7.5
by DIP STICK/REFLEC	TANCE SPECTROPHOTOMETRY			
BILIRUBIN	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
NITRITE		Negative		NEGATIVE (-ve)
by DIP STICK/REFLEC UROBILINOGEN	TANCE SPECTROPHOTOMETRY.	Normal	EU/dL	0.2 - 1.0
	TANCE SPECTROPHOTOMETRY	Normai	EU/UL	0.2 - 1.0
KETONE BODIES	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
BLOOD	TANCE SPECTROPHOTOMETRT	Negative		NEGATIVE (-ve)
	TANCE SPECTROPHOTOMETRY	-		
ASCORBIC ACID by DIP STICK/REFLEC	TANCE SPECTROPHOTOMETRY	NEGATIVE (-	-ve)	NEGATIVE (-ve)
MICROSCOPIC EXA				
RED BLOOD CELLS		NEGATIVE (-	-ve) /HPF	0 - 3





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EXCELLENCE IN HEALTHCARE & DIAGNOSTICS

Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist

Dr. Yugam Chopra MD (Pathology) CEO & Consultant Pathologist

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CLIENT CODE.	: KOS DIAGNOSTIC LAB]	REPORTING DATE	: 04/Jan/2025 04:18PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AN	MBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
by MICROSCOPY ON (CENTRIFUGED URINARY SEDIMENT			
PUS CELLS by MICROSCOPY ON (CENTRIFUGED URINARY SEDIMENT	1-3	/HPF	0 - 5
EPITHELIAL CELLS		2-4	/HPF	ABSENT

by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	~ 1	/ 111 1	MODELLI
CRYSTALS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
CASTS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
BACTERIA by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
OTHERS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
TRICHOMONAS VAGINALIS (PROTOZOA) by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	ABSENT		ABSENT

** End Of Report ***



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KOS DIAGNOSTIC LAB 6349/1, NICHOLSON ROAD, AMBALA CANTT

			I	Result Down's sy	ndrome screening	g	
Name				Sample ID	2501220080/AMB	diabetes	n
		MR	S. PRIYA	D.O.B.	8/08/1997	Fetuses	
Patient ID				Age at delivery		Smoker	r
	um taking		/01/2025	Weight [kg]	60 kg		r
Date of re	-	5	5/01/2025			Ethnic origin	Asia
Previous to pregnancie			no				
orognarion							
			C	corrected MoM's a	Ind calculated risl	ks	
AFP	32.4	ng/ml	0.99	Corr. MoM	Gestational age at	sample date	15 + 3
uE3	0.93	ng/ml	0.99	Corr. MoM	determination meth		BPD Hadlock
HCG	51895	mIU/mI	1.38	Corr. MoM	Physician		
Inh-A	184	pg/ml	1.10	Corr. MoM			
Risk 1:10							
				/			Tr.21 risk
							at term
							1:4525
1:100							
1:250				Cut off			
							Age risk
1:1000							at term
							1:1176
1:10 <mark>000</mark>							
13 15 17	7 19 21 23 2	25 27 29 31	33 35 37	39 41 43 45 47 49 Ag	e		
Down's	Syndror	ne Risk					
	-			halaw the aut off	which represents a		
After the	result of the	ne Trisomy	/ 21 test it	t is expected that am	ong 4525 women wit		a, there is one woman
				women with not affe	ected pregnancies.	ided by the ref	
					and have no diagno		ening physician.
					· ·		

Neural tube defects risk	Risk for trisomy 18
The corrected MoM AFP (0.99) is located in the low risk area for neural tube defects.	The calculated risk for trisomy 18 is < 1:10000, which indicates a low risk.

	below cut off	Below Cut Off, but above Age Risk	above cut off	Prisca 5.2.0.13
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