## **PKR JAIN HEALTHCARE INSTITUTE** NASIRPUR, Hissar Road, AMBALA CITY- (Haryana) A PIONEER DIAGNOSTIC CENTRE

【 0171-2532620, 8222896961 🛛 🖾 pkrjainhealthcare@gmail.com

NAME	: Mrs. PREETI SHARMA			
AGE/ GENDER	: 24 YRS/FEMALE	PATIE	INT ID	: 1540086
COLLECTED BY:REFERRED BY:BARCODE NO.: 12503464CLIENT CODE.: P.K.R JAIN HEALTHCARE INSTITUTE		REG. N	IO./LAB NO.	: <b>122407060014</b> : 06/Jul/2024 10:36 AM
		REGIS	TRATION DATE	
		COLLECTION DATESTITUTEREPORTING DATE		: 06/Jul/2024 10:53AM : 08/Jul/2024 11:03AM
Test Name		Value	Unit	Biological Reference interval
		ENDOCRINO	LOGY	
		IAL MARKER MATERI	NAL SCREENING	
DUAL MARKER TEST				
PATEINT SPECIFICAT	IONS			
DATE OF BIRTH		2000-07-16		
MATERNAL AGE		24.48	YEARS	
WEIGHT		66	Kg	
		ASIAN		ASIAN
H/O IVF H/O SMOKING		ABSENT		
H/O INSULIN DEPEN	DANT DIARETES	ABSENT		
H/O TRISOMY 21 SCI		ABSENT		
ULTRA SOUND SCAN		ABOEIT		
DATE OF ULTRASOU		06/07/2024		
by ULTRASOUND SCA	N			
METHOD FOR GESTA by ULTRASOUND SCA	TION AGE ESTIMATION	ULTRASOUND SCAN DETAILS		
FOETUS (NOS)	rv -	1		
by ULTRASOUND SCA				
GA ON THE DAY OF SAMPLE COLLECTION		13.5	WEEKS	
by ULTRASOUND SCAN CROWN RUMP LENGTH (CRL)		78.2	mm	38 - 84
by ULTRASOUND SCAN				
GESTATIONAL AGE BY CRL		13.5		
by ULTRASOUND SCAN NUCHAL TRANSLUCENCY (NT)		2.2	mm	0.1 - 6.0
by ULTRASOUND SCAN				
NUCHAL TRANSLUCE by ULTRASOUND SCA		1.16		
	V CHEMICAL MARKERS			

**DR.VINAY CHOPRA** CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY & MICROBIOLOGY) MBBS , MD (PATHOLOGY)

DR.YUGAM CHOPRA CONSULTANT PATHOLOGIST

**NOT VALID FOR MEDICO LEGAL PURPOSE** 

440 Dated 17.5.2012 u/s 80 G OF INCOME TAX ACT. PAN NO. AAAAP1600. **REPORT ATTRACTS THE CONDITIONS PRINTED OVERLEAF (P.T.O.)** 



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<b>CLIENT ADDRESS</b> : NASIRPUR, HISSAR ROAD, AMBA		ALA CITY - HARYANA	A	
Test Name		Value	Unit	Biological Reference interval
PROTEIN A (PAPP-A)				
	IESCENCE IMMUNOASSAY)			
BETA HCG - FREE: SERUM		163.72	ng/mL	
by CLIA (CHEMILUMIN MULTIPLE OF MEDIA	IESCENCE IMMUNOASSAY)			
	AN (MON) VALUES			
PAPP-A MOM		1.64		
by CLIA (CHEMILUMINESCENCE IMMUNOASSAY) BETA HCG - FREE MOM		4.64		
	IESCENCE IMMUNOASSAY)	1.01		
TRISOMY 21 SCREE	NING (DOWNS SYNDROME) RISK	ASSESSMENT		
TRISOMY 21 SCREENING RISK RESULT		NEGATIVE (-ve)		NEGATIVE (-ve)
by CLIA (CHEMILUMINESCENCE IMMUNOASSAY)				
		1:1405 NEGATIV	'E (-ve)	
by CLIA (CHEMILUMINESCENCE IMMUNOASSAY) TRISOMY 21 BIOCHEMICAL RISK		1:641 NEGATIVE (-ve)		RISK CUT OFF 1:150
by CLIA (CHEMILUMINESCENCE IMMUNOASSAY)		1.041 NEG/TIVE	. ( VC)	
TRISOMY 21 COMBINED RISK (BIOCHEMICAL + NT)		< 1:10000 NEGATIVE (-ve)		RISK CUT OFF 1:150
2	IESCENCE IMMUNOASSAY)			
TRISOMY 18 SCREE	NING RISK ASSESSMENT			
TRISOMY 18 AGE RISK		NEGATIVE (-ve)		
by CLIA (CHEMILUMINESCENCE IMMUNOASSAY)				
TRISOMY 13/18 SCREENING RISK by CLIA (CHEMILUMINESCENCE IMMUNOASSAY)		< 1:10000 NEGA	IIVE (-Ve)	RISK CUT OFF 1:300
	ILSOLINGE IMMUNUNUASSAT)			

**INTERPRETATION:** 

1.Double marker test (maternal serum screen – first trimester) is a prenatal test to screen for Trisomy 21 (down's syndrome) and Trisomy 13/18 during gestational period 8 – 13 weeks.

2.Besides the biochemical markers tested – maternal pregnancy associated plasma protein a (papp-a) & maternal free beta hcg, the risk is calculated combining usg measurement of nuchat translucency (nt), gestational age at the time of sample with other maternal factors as age, weight, h/o diabetes, smoking, race, twin pregnancies, use of assissted reproductive technologies (IVF).





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Test Name Value Unit **Biological Reference interval** 

## NOTE:

1. This is only screening test based purely on statistical analysis which is further based on the data submitted; hence the correctness of data is vital for risk analysis

2.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.

3.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.* 4.The detection rate by this test is about 60%, with 5% false positive rate when assessment is done for only biochemical parameters and increase to 85 % with 5% false positive rate when both biochemical parameters and nt are combined for analysis. 5.Correlation with patient history, family history and detailed USG scan is required to decide further course of action in cases who have high risk statistically called a state.

statistically calculated by this test.

End Of Report



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Basic Information		
Name: PREETI SHARMA	Contact:	Gender: Female
Weight: 66.00 Kg	Birthdate: 2000-07-16	Age of EDC: 24.48 Year
Race: Asian	Twins: No	GA calc method: CRL Robinson
LMP Day:	Sender:	
Sample information Send time: 2024-07-08	a 1. No. 10502464	G D ( 2024.07.0C
	Sample NO.: 12503464	Scan Date: 2024-07-06
Lab: Sample Date: 2024-07-		GA: 13+5
BPD: mm	CRL length: 78.20 mm	NT length: 2.20 mm
Assay		
NO. Item abbr Result	t Unit M	IOM Reference range
1 free-β-HCG <b>163.72</b>	2 ng/ml 4	.64
2 PAPP-A <b>10000.0</b>	00 mIU/L 1	.64
3 NT <b>2.20</b>	mm 1.	.16
Risk calculate		
Age risk: 1:1405		21-3 syndrome risk
Demonster Trisomv21		50
Parameter: Trisomy21		100 – Risk above cut off
Risk: 1:641	Risk	You risk 1:641
Cut Off: ( < 1:150 )		>5000
Screaning Result: Negative		50
5 5		Age
		18-3 syndrome risk
Parameter: Trisomy18/13		100
Risk: 1:107948978	×	Risk above cut off
Cut Off: ( < 1:300 )	Risk	200 - You risk 1: >10000
Screening Result: Negative		>5000
		50 Age

Advice: Diagnostic results with less risk

Note: \*The basic information on the basis of Down's risk assessment in this report is provided at the time of your onsite. When you get this report, please first check whether your relevant information is correct. If there is any discrepancy, please contact your doctor in time, so as to feedback us the correct information and documents, then obtain the correct report. \*The high risk and borderline risk of trisomy 21 or trisomy 18 requires further interventional prenatal diagnosis (from fetuses such as villus, amniotic fluid, cord blood, etc.); high risk of neural tube defect (NTD), please go to ultrasound prenatal diagnosis qualified hospitals use ultrasound to exclude.

\*The risk of NTD is only calculated at 14-22 weeks.

\*The screening result with low risk only shows that the chance of this kind of congenital abnormality in your fetus is less, and the possibility of this kind of abnormality or other abnormalities cannot be completely ruled out. Please consult a doctor in time after you get the report, and the doctor will follow your Risks and other conditions (whether you are older than 35 years old, whether you have had more than one child with other deformities, or have other diseases such as tumors) are comprehensively considered to suggest whether you need to take further examination to confirm the diagnosis.

\*\*This report only can be reference and assistant for doctor , cannot directly give conclusion by this \*\*

Doctor: