

**LABORATORY TEST REPORT**

Name	: Mrs. SUMITA DEKA		
Sample ID	: B3585185	Reg. No	: 0692508260444
Age/Gender	: 38 Years/Female	SPP Code	: SPL-AS-177
Referred by	: Dr. NIKUNJA DEKA	Collected On	: 26-Aug-2025 03:00 PM
Referring Customer	: LIFE CARE DIAGNOSTICS MALIGAON	Received On	: 27-Aug-2025 03:53 PM
Primary Sample	: Whole Blood	Reported On	: 28-Aug-2025 06:51 PM
Sample Tested In	: Serum	Report Status	: Final Report
Client Address	:		

**CLINICAL BIOCHEMISTRY**

Test Name	Results	Units	Biological Reference Interval
Free -Beta -HCG (Method: CLIA)	38.64	ng/mL	< 2 :Non-Pregnant 5.4 - 393.4 : Pregnant
PAPP-A (Method: CLIA)	0.82	mIU/mL	< 0.1 : Non-Pregnant 0.1-19.5 : Pregnant

[PDF Attached](#)

**Double Marker**

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(Method: CLIA)			
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Risk analysis for Trisomy 21 is 1:82 is positive. PAPP-A value low. Adv: NIPT, FISH and karyotyping

**Interpretation:**

DISORDER	SCREEN POSITIVE/HIGH RISK CUT OFF
Trisomy 21 (Down)	< 1:250
Trisomy 18/13	< 1:100
DISORDER	SCREEN NEGATIVE/LOW RISK CUT OFF
Trisomy 21 (Down)	> 1:250
Trisomy 18/13	> 1:100

**Note:** Statistical evaluation has been done using CE marked PRISCA 5 software. · Screening tests are based on statistical analysis of patient demographic and biochemical data. They simply indicate a high or low risk category. Confirmation of screen positives is recommended by Chorionic Villus Sampling (CVS). · The interpretive unit is MoM (Multiples of Median) which takes into account variables such as gestational age (ultrasound), maternal weight, race, insulin dependent Diabetes, multiple gestation, IVF (Date of Birth of Donor, if applicable), smoking & previous history of Down syndrome. Accurate availability of this data for Risk Calculation is critical. · Ideally all pregnant women should be screened for Prenatal disorders irrespective of maternal age. The test is valid between 9-13.6 weeks of gestation, but ideal sampling time is between 10-13 weeks gestation. · First trimester detection rate of Down syndrome is 60% with a false positive rate of 5%. A combination of Nuchal translucency, Nasal bone visualization and biochemical tests (Combined test) increases the detection rate of Down syndrome to 85% at the same false positive rate.

**Comments:** First trimester screening for Prenatal disorders (Trisomy 21, 18 & 13) is essential to identify those women at sufficient risk for a congenital anomaly in the fetus to warrant further evaluation and followup. For Open neural tube defects, second trimester screening before 20 weeks is recommended. These are screening procedures which cannot discriminate all affected pregnancies from all unaffected pregnancies. Screening cutoffs are established by using MoM values that maximize the detection rate and minimize false positives.

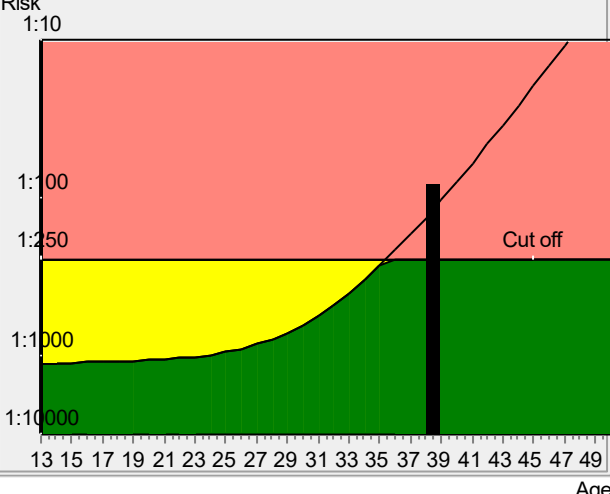
\*\*\* End Of Report \*\*\*



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DR. LAVANYA LAGISSETTY  
MD BIOCHEMISTRY

NA

Patient data			
Name	Mrs. SUMITA DEKA	Patient ID	0692508260444
Birthday	20-03-1987	Sample ID	B3585185
Age at sample date	38.4	Sample Date	26-08-2025
Gestational age	12 + 3		
Correction factors			
Fetuses	1	IVF	no
Weight	51	diabetes	no
Smoker	no	Origin	Asian
Biochemical data		Ultrasound data	
Parameter	Value	Corr. MoM	
PAPP-A	0.82 mIU/mL	0.16	Gestational age 12 + 1
fb-hCG	38.64 ng/ml	0.91	Method CRL Robinson
Risks at sampling date		Scan date 24-08-2025	
Age risk	1:119	Crown rump length in mm 57.5	
Biochemical T21 risk	>1:50	Nuchal translucency MoM 0.66	
Combined trisomy 21 risk	1:82	Nasal bone present	
Trisomy 13/18 + NT	1:77	Sonographer NA	
Risk		Qualifications in measuring NT MD	
		<b>Trisomy 21</b> <b>The calculated risk for Trisomy 21 (with nuchal translucency) is above the cut off, which indicates an increased risk.</b> After the result of the Trisomy 21 test (with NT) it is expected that among 82 women with the same data, there is one woman with a trisomy 21 pregnancy and 81 women with not affected pregnancies. The PAPP-A level is low. The calculated risk by PRISCA depends on the accuracy of the information provided by the referring physician. Please note that risk calculations are statistical approaches and have no diagnostic value! The patient combined risk presumes the NT measurement was done according to accepted guidelines (Prenat Diagn 18: 511-523 (1998)). The laboratory can not be held responsible for their impact on the risk assessment! Calculated risks have no diagnostic value!	
Trisomy 13/18 + NT			
<b>The calculated risk for Trisomy 13/18 test (with nuchal translucency) is 1:77, which represents an increased risk.</b>			

Sign of Physician

 below cut off

 Below Cut Off, but above Age Risk

 above cut off