

Name : Dr G V Hema Sudha (30Y/F)

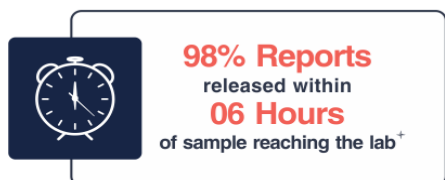
Date : 30 Aug 2025

Test Asked : Double Marker First Trimester 8-13 Week (Fmf Certified)

Report Status: Complete Report



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NABL From 2005[#]



ISO 9001: 2015 – From 2015



CAP From 2007⁻

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Hyderabad-500 044



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NAME : DR G V HEMA SUDHA (30Y/F)
REF. BY : DR HEMA SUDHA
TEST ASKED : DOUBLE MARKER FIRST TRIMESTER 8-13 WEEK (FMF CERTIFIED)

Sample Collected At
(5022206459),city centre,MULTI SPECIALITY HOSPITAL OPP BHARATH GAS OFFICE ROAD ZAHEERABAD DIST SANGAREDDY,502220

Report Availability Summary

Note: Please refer to the table below for status of your tests.

1 Ready

0 Ready with Cancellation

0 Processing

0 Cancelled in Lab

TEST DETAILS

REPORT STATUS

DOUBLE MARKER FIRST TRIMESTER 8-13 WEEK (FMF

Ready

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Adikmet Road, Near SBH,
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NAME : DR G V HEMA SUDHA (30Y/F)
REF. BY : DR HEMA SUDHA
TEST ASKED : DOUBLE MARKER FIRST TRIMESTER 8-13 WEEK
 (FMF CERTIFIED)

Sample Collected At
 (5022206459), city centre, MULTI SPECIALITY
 HOSPITAL OPP BHARATH GAS OFFICE ROAD
 ZAHEERABAD DIST SANGAREDDY, 502220

TEST NAME	TECHNOLOGY	VALUE	UNITS
FREE BETA HCG	E.C.L.I.A	26.3	IU/L

Bio. Ref. Interval. :-

Pregnancy:	Cut off risk for Downs syndrome or Trisomy 21		
Weeks Ranges	<35 years	>35 years	
10-11 : 45.60 - 54.53	Weeks	Ranges	Weeks
11-12 : 37.30 - 44.28	10-11 :	>210	10-11 :
12- 13 : 31.27 - 36.30	11-12 :	>162	11-12 :
13-14 : 27.52 - 30.60	12- 13 :	>130	12- 13 :
	13-14 :	>110	13-14 :
		>60	>60

Interpretation:

Free BHCG is a glycoprotein hormone normally found in blood and urine only during pregnancy. It is secreted by placental tissue, beginning with primitive trophoblast, almost from the time of implantation, and serves to support the corpus luteum during early weeks of pregnancy. HCG or HCG like material is also produced by a variety of trophoblastic and non-trophoblastic neoplasia. Intact HCG is 39,500 Da molecule composed of two non-identical subunits alpha and beta that are bound to each other in a non-covalent manner. These subunits can also occur in a free or unbound form. Only intact HCG has biological activity. The HCG-alpha subunit is structurally identical to the alpha subunit of homologous pituitary glycoprotein hormones, luteinizing hormones, follicle stimulating hormone and thyroid stimulating hormone. The beta subunit is specific for each of these hormones, and confers upon them their differing biological activities. Maternal serum free BHCG assessment is reported to have significant utility in first and second trimester prenatal screening for Down syndrome and other chromosomal anomalies.

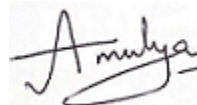
Specifications: Precision: Intra assay (%CV): 1.6, Inter assay (%CV): 3.8, Sensitivity: 0.3 IU/mL

Reference : Karl O. Kagan, Dave Wright, Catalina Valencia, Nerea Maiz, Kypros H. Nicolaides, Screening for trisomies 21, 18 and 13 by maternal age, fetal nuchal translucency, fetal heart rate, free β -hCG and pregnancy-associated plasma protein-A, Human Reproduction, Volume 23, Issue 9, 1 September 2008, Pages 1968-1975

Please correlate with clinical conditions.

Method:- SANDWICH ELECTROCHEMILUMINESCENCE IMMUNOASSAY

Sample Collected on (SCT) : 29 Aug 2025 19:19
Sample Received on (SRT) : 30 Aug 2025 14:08
Report Released on (RRT) : 30 Aug 2025 18:52
Sample Type : SERUM
Labcode : 3008082865/HYD93 Dr Amulya MD (Path)
Barcode :  EL364883



Page : 1 of 3

Scan QR code to verify authenticity of reported results; active for 30 days from release time.

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REF. BY : DR HEMA SUDHA
TEST ASKED : DOUBLE MARKER FIRST TRIMESTER 8-13 WEEK (FMF CERTIFIED)

Sample Collected At
 (5022206459),city centre,MULTI SPECIALITY HOSPITAL OPP BHARATH GAS OFFICE ROAD ZAHEERABAD DIST SANGAREDDY,502220

TEST NAME	TECHNOLOGY	VALUE	UNITS
PREGNANCY ASSOCIATED PLASMA PROTEIN A	E.C.L.I.A	1678	mIU/L

Bio. Ref. Interval. :-

Pregnancy:	Cut off risk for Downs syndrome or Trisomy 21			
Weeks Ranges	<35 years	>35 years		
10-11 : 1180 - 1534	Weeks	Ranges	Weeks	Ranges
11-12 : 1617 - 2301	10-11 :	<1000	10-11 :	<1200
12- 13 : 2445 - 3490	11-12 :	<1040	11-12 :	<1750
13-14 : 3694 - 5101	12-13 :	<1450	12-13 :	<2650
	13-14 :	<1800	13-14 :	<3390

Interpretation

Pregnancy-associated plasma protein-A (PAPP-A) was first identified as a high-molecular weight constituent in human pregnancy serum. PAPP-A is a metalloprotease which belongs to the metzincin super family of zinc peptidases. Insulin-like growth factor binding protein 4 (IGFBP-4) as well as IGFBP-5 have been found to be specific substrates for PAPP-A in vitro.

PAPP-A is widely recognized biochemical marker of Down syndrome (DS) used in the first trimester of pregnancy. PAPP-A level in maternal serum increases with gestational age until term. In case of DS pregnancy, PAPP-A concentration in the first trimester is markedly decreased.

Specifications: Precision: Intra assay (%CV): 5, Inter assay (%CV): 6.9, Sensitivity: <= 8 mIU/mL

References:

Patil M, Panchanadikar TM, Wagh G. Variation of papp-a level in the first trimester of pregnancy and its clinical outcome. J Obstet Gynaecol India. 2014 Apr;64(2):116-9.

Karl O. Kagan, Dave Wright, Catalina Valencia, Nerea Maiz, Kypros H. Nicolaides, Screening for trisomies 21, 18 and 13 by maternal age, fetal nuchal translucency, fetal heart rate, free β -hCG and pregnancy-associated plasma protein-A, Human Reproduction, Volume 23, Issue 9, 1 September 2008, Pages 1968-1975.

Please correlate with clinical conditions.

Method:- SANDWICH ELECTROCHEMILUMINESCENCE IMMUNOASSAY

~~ End of report ~~

Sample Collected on (SCT) : 29 Aug 2025 19:19
Sample Received on (SRT) : 30 Aug 2025 14:08
Report Released on (RRT) : 30 Aug 2025 18:52
Sample Type : SERUM
Labcode : 3008082865/HYD93 Dr Amulya MD (Path)
Barcode :  EL364883



Page : 2 of 3

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First Trimester Screening results

Patient data

Name and surname:	DR G V HEMA SUDHA	Weight:	96 Kg.
Lab ID:	EL364883	Height:	N/I
Race/Ethnicity:	INDIAN	Diabetes:	No
Date of birth:	24-11-1994 (31 years in the DoB)	Smoker:	No
Type of Pregnancy:	Spontaneous	Ovulation Ind.:	No
Prev. Obstetric History:	None	Referral Center:	HYD93
Prenatal Software:	SSDWLAB6	Referral Doctor:	DR HEMA SUDHA

Biochemical data

Sample date:	29-08-2025	Gestational age:	12 weeks and 3 days
Sample ID:	EL364883		
Free beta hCG 1T:	26.3 IU/L	1.15 MoM	
PAPP-A:	1678 mIU/L	0.85 MoM	

Ultrasound data

Ultrasound date:	28-08-2025	Gestational age:	12 weeks and 2 days
CRL:	57 mm		
Nuchal Translucency:	1.3 mm	0.87 MoM	

Dichotomous markers

Absent nasal bones=No.

Risk report (At term)

Risk type	Probability	Result	Graphic representation
Trisomy 21:	< 1/10000	Low Risk	< 1/10000 250
Trisomy 18/13:	< 1/100000	Low Risk	< 1/100000 100
Trisomy 21 age risk:	1/821		

Interpretation

Screen Negative.

- The risk calculations are statistical approaches and have limited diagnostic value.
- The calculated risk by the software depends on the accuracy of USG details and patient details provided.
- FMF-Certified Free beta hCG & PAPP-A kits are being used for analysis.
- Participants in UKAS-proficiency testing (EQAS) for maternal serum markers.
- The laboratory can not be held responsible for their impact on the risk assessment. Calculated risks have no diagnostic value.
- The screening risk estimates final risk using biochemical parameters results, maternal demographic characteristics, and maternal medical and obstetric history. The risk calculation is optimal when accurate critical information is provided and incorrect information ((TRF/ U.S scan report) may significantly alter the risk assessment.
- Risks cannot be calculated for triplets or higher order gestations. In twin pregnancies with fetal demise (vanishing twins) risk estimation can be calculated but may be unreliable.

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- Comparison with other screening software's and assay methodologies may give varying risk assessments. Risk assessment at term and sampling are both valid ways of estimating risk, but the risk score between the two varies because a correction factor of intrauterine mortality is applied for risk at sampling which is not taken into consideration while computing risk at term.
- Sophisticated software program SsdwLab6 works on statistical database to calculate this risk and hence any risk indicated should not be considered to be confirmatory evidence of fetal risk.
- This is a screening report and will need further confirmatory tests for diagnosis. Kindly consult your doctor for further.

First Trimester Screening	
Risk Analysis in Interpretation	Description according to the Risk Analysis in Interpretation means
Screen Negative	It means the result is statistically low risk for Trisomy 21 & Trisomy 18/13
Screen Positive	It means the result is statistically high risk for either both Trisomy 18/13 & Trisomy 21 or for any one of Trisomy

Refer the below table for statistical cut off ratio for interpretation of Trisomy 18/13 & Trisomy 21 risk:

Anomaly	Screen positive risk cut-off(ACOG 2007)	Remarks
Trisomy 21	1:250	Confirmatory tests needed
Trisomy 18/13	1:100	Confirmatory tests needed

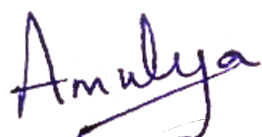
Cut-off for Trisomy 18/13 is in accordance to ACOG - Breathnach FM, Malone FD, Lambert-Messerlian G et al. First- and second-trimester screening: detection of aneuploidies other than Down syndrome. Obstet Gynecol. 2007 Sep;110(3):651-7. doi: 10.1097/01.AOG.0000278570.76392.a6. PMID: 17766613

Authorized by:

Adinath Suryavanshi

Printing date:

30-08-2025



Dr. Amulya Ravi
MBBS, MD (Pathology)

CONDITIONS OF REPORTING

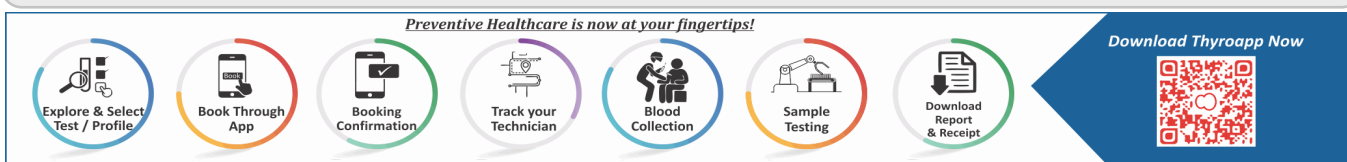
- v The reported results are for information and interpretation of the referring doctor only.
- v It is presumed that the tests performed on the specimen belong to the patient; named or identified.
- v Results of tests may vary from laboratory to laboratory and also in some parameters from time to time for the same patient.
- v Should the results indicate an unexpected abnormality, the same should be reconfirmed.
- v Only such medical professionals who understand reporting units, reference ranges and limitations of technologies should interpret results.
- v This report is not valid for medico-legal purpose.
- v Neither Thyrocare, nor its employees/representatives assume: (a) any liability, responsibility for any loss or damage that may be incurred by any person as a result of presuming the meaning or contents of the report, (b) any claims of any nature whatsoever arising from or relating to the performance of the requested tests as well as any claim for indirect, incidental or consequential damages. The total liability, in any case, of Thyrocare shall not exceed the total amount of invoice for the services provided and paid for.
- v Thyrocare Discovery video link :- <https://youtu.be/nbdYeRqYyQc>

EXPLANATIONS

- v Majority of the specimen processed in the laboratory are collected by Pathologists and Hospitals we call them as "Clients".
- v **Name** - The name is as declared by the client and recorded by the personnel who collected the specimen.
- v **Ref.Dr** - The name of the doctor who has recommended testing as declared by the client.
- v **Labcode** - This is the accession number in our laboratory and it helps us in archiving and retrieving the data.
- v **Barcode** - This is the specimen identity number and it states that the results are for the specimen bearing the barcode (irrespective of the name).
- v **SCP** - Specimen Collection Point - This is the location where the blood or specimen was collected as declared by the client.
- v **SCT** - Specimen Collection Time - The time when specimen was collected as declared by the client.
- v **SRT** - Specimen Receiving Time - This time when the specimen reached our laboratory.
- v **RRT** - Report Releasing Time - The time when our pathologist has released the values for Reporting.
- v **Reference Range** - Means the range of values in which 95% of the normal population would fall.

SUGGESTIONS

- v Values out of reference range requires reconfirmation before starting any medical treatment.
- v Retesting is needed if you suspect any quality shortcomings.
- v Testing or retesting should be done in accredited laboratories.
- v For suggestions, complaints, clinical support or feedback, write to us at customersupport@thyrocare.com or call us on **022-3090 0000**



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* T&C Apply, #As on 5th December 2024 (Applicable for all company owned labs except Bhagalpur & Vijayawada),
* As per survey on doctors' perception of laboratory diagnostics (IJARIIT, 2023), -Mumbai Reference Lab is CAP Accredited