

Patient Name : Mr MR.DUMMY	Sample Collected : Apr 26, 2024, 01:00 PM
DOB/Age/Gender : 23 Y/Male	Report Date : May 03, 2024, 04:57 PM
Patient ID / UHID : 8053272/RCL7249297	Barcode No : SA041269
Referred By : Dr. X	Report Status : Final Report
Sample Type : Serum	

Test Description	Value(s)	Unit(s)	Reference Range
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Combined Screening (Intermediate)- AutoDELFIA (Astraia)

Combined Screening- AutoDELFIA (Astraia)

Beta HCG Free <i>TRFI</i>	45.0	ng/mL	
Pregnancy Associated Plasma Protein(PAPP-A) <i>TRFI</i>	4526	mU/L	

Interpretation:

Risk factor calculated by : Astraia

Disclaimer:

1. This is a risk estimation test and not a diagnostic test. An increased risk result does not mean that the fetus is affected and a low risk does not mean that the fetus is unaffected, reported risk should be correlated and adjusted to the absence/presence of sonographic markers observed in the anomaly/malformation scan.
2. This interpretation assumes that patient and specimen details are accurate and correct.
3. The testing laboratory does not bear responsibility for the ultrasound measurements.

Interpretation guidelines

Disorder	Screen positive Cut off (ACOG 2007)	MOM Cut off (ACOG 2007)	Remarks
Trisomy-21	1:250	Free BHCG: > or = 1.98 PAPPa:< or = 0.43	Confirmatory tests needed under doctor's advise
Trisomy-18 / Trisomy-13	1:100	Free BHCG: < or = 0.5 PAPPa: < or = 0.4	Level-III ultrasound needed for confirmation

Note

All Lab results are subject to clinical interpretation by a qualified medical professional & This report is not subject to use for any medico-legal purpose.
 FMF Accredited Autodelfia Platform is used to measure the biochemical Marker
 Statistical evaluation has been done by Astraia
 Maternal Biochemical Marker's Screening is based on Statistical analysis & demographic & biochemical data of the Patient & only indicates a high or low-risk category, CUS is recommended for Confirmation & Screen Positives.
 Multiples & Median (MOM) are measured by accounting. Variables like Gestational age / Maternal weight / Multiple gestation / IVF or Not / Ultrasound / Smoking stand Previous history & T21, hence Accurate availability & this data is very important for risk Calculation.
 Ideal sampling time is between 10 weeks to 13 weeks plus 6 days of gestation and requires a crown-rump length between approximately 40mm to 85 mm.
 The detection rate for Down syndrome is 60% with a false positive rate of 5% if the only biochemical risk is estimated. A combination of Nuchal translucency and biochemical tests (Combined test) has a detection rate of Down syndrome 82 to 87% at a 5% false-positive rate. The addition of absent nasal bone status can improve the detection rate up to 93% at false-positive rate of 2.5%

1. Statistical risk factor calculation for Trisomy 21 (Down's syndrome), Trisomy 18 (Edward Syndrome) and Trisomy 13 (Patau Syndrome) has been done using Fetal Medicine Foundation (FMF) approved assays using Autodelfia Analyser
2. The statistical risk evaluation requires Maternal age to be decimalised for months, to be represented as Age at sampling & conversion of maternal hormonal values to mean of medians(MOMs). The MoMs are further calculated using Indian medians,



Dr. Dummy



Booking Centre :- DEMO PARTNER CHENNAI, DEMO PARTNER CHENNAI
 Processing Lab :-

📞 928-909-0609

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*** End Of Report ***

Disclaimer: Method given in report are only indicative and can be changed depending upon type of machine and kit available at time of testing.

Not all tests at all locations are under NABL scope. Availability of tests under NABL scope varies from lab to lab.



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2. It is to be presumed that the tests performed pertain to the specimen/sample attributed to the Customer's name or identification. It is presumed that the verification particulars have been cleared out by the customer or his/her representation at the point of generation of said specimen / sample. It is hereby clarified that the reports furnished are restricted solely to the given specimen only.
3. It is to be noted that variations in results may occur between different laboratories and over time, even for the same parameter for the same Customer. The assays are performed and conducted in accordance with standard procedures, and the reported outcomes are contingent on the specific individual assay methods and equipment(s) used, as well as the quality of the received specimen.
4. This report shall not be deemed valid or admissible for any medico-legal purposes.
5. The Customers assume full responsibility for apprising the Company of any factors that may impact the test finding. These factors, among others, includes dietary intake, alcohol, or medication / drug(s) consumption, or fasting. This list of factors is only representative and not exhaustive.

DISCLAIMER

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