





To:	Pallavi Maternity Home And General Hospital-Ulwe
	115 AND 116, 1st Floor,
	NMS Icon, Plot Ni. 194, Sector 19,
	Maharashtra
	Ulwe - 410206
	Contact:
	Report Of: Mrs. CHAITALI MANJREKAR
	Pt. Contact:

Referred by Sonography by	DR.PRAKASH KUDALE
Poforrod by	DR.PALLAVI A. JADHAV
Reported on	04/08/2022 11:27
Registered on	03/08/2022 19:35
Received on	02/08/2022 20:54
PatientID	1002255789
SampleID	2200095391

EVICOSCREEN - EVIDENCE BASED COMPREHENSIVE PRENATAL SCREENING REPORT

Patient Name: Mrs. Cl	HAITALI MANJREKAR	Patient DOB: <u>18/10/1982</u>	Patient DOB: <u>18/10/1982</u>		
Ethnicity: Asian	City: MUMBAI	Hospital ID:			

Sample Type:Serum

Risk assessment: Algorithm validated by SURUSS 2003, N.J Wald

Method:Electrochemiluminescence

EVIC Screen" is an evidence based prenatal screening program curated by Lilac Insights in accordance with the Fetal Medicine Foundation

(UK) guidelines for First Trimester Screening to determine the probability of most common chromosomal aneuploidies in a pregnancy. It utilizes:

• Hormonal values from the pregnancy measured on Fetal Medicine foundation (UK) accredited analyzers and reagents

• Robust indigenous medians from over 5 lac+ pregnancies for different gestation ages

Risk calculations from evidence based algorithms validated through large international studies

• External audit of the prenatal screening program by United Kingdom National External Quality Assessment Service (UKNEQAS) scheme and Randox International Quality Assessment Scheme (RIQAS)

RISK ASSESSMENT					
T21 (Down syndrome)	1:2100	Low Risk	LOW	INTERMEDIATE HIGH	MEDIAN (MoM)
T18 (Edwards' syndrome)	1: 100000	Low Risk	LOW	HIGH	Free ß-hCG 1.71
T13 (Patau syndrome)	1:12000	Low Risk	LOW	HIGH	PAPP-A 1.15
		INTERPRETAT	ION		

The First Trimester Screening for the given sample is found SCREEN NEGATIVE.

Verified by Dr. Suresh Bhanushali MD (Path), Consultant Pathologist Page **1** of 3



UK NEQAS Lab Reg. No. 90968

Verified by Mr. Pradip Kadam Incharge Biochemistry



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PREGNANCY DETAILS



Patient name: Mrs. CHAITALI MANJREKAR

Sample ID: 2200095391

				PREGNANC					
No. of fetuses : 1				EDD	: 14/02/2023	Age at Terr			
GA is Based		: CRL 52mm at 01/	/08/2022	LMP Date	: 12/05/2022			: Regular	
Smoking:None Parity :				Height	:	Weight	: 57.00	: 57.00 Kg	
FHR :									
F	revio	us pregnancy histo	ory	Pre-eclampsia history			Other findings		
Down syndrome Edwards' syndrome Patau syndrome NTD syndrome			PE in previous pregnancy Pat. mother had PE		Insu	Insulin dependent diabetes			
					Chronic hypertension				
EDD: Estimat	ted Due	Date GA: Gestation Age	LMP: Last M	enstrual Period FHF	R: Fetal Heart Rate NTL): Neural Tube Def	ect PE: Pre-ed	clampsia DOB: D	
				ofBir	th				
				SPECIMEN	DETAILS				
Sample ID		:2200095391	CRL	: 52 mm	Test Name	Conc.	Unit	Corr. Mom	
Collection [Date	:01/08/2022	CRL2	:	Free-ß-hCG	70.08	ng/mL	1.71	
Scan Date		:01/08/2022	BPD	:	NB	Present			
GA at Coll [Date	: 11 Weeks 6 Days	BPD2	:	NT	1.1	mm	0.92	
GA at Scan	Date	: 11 Weeks 6 Days	HC	:	PAPP-A	3216.00	mIU/L	1.15	
Received or	า	:02/08/2022	HC2	:					
GA: Gestatic	on Age C	CRL: Crown Rump Length NT: I			Head Circumference fre gnancy-associated Plasm		Human Chor	ionic Gonadotroj	
				RISK	(S				
	own Sv	ndrome			R	esult:	Low Risk	()	
Disorder: D	,								
Disorder: D Final risk:	1:21		Age risk:	1:90					
	-		Age risk: Risk type	1:90 Risk At Term					
Final risk: Cutoff	1:21 1:25	0	-		R	esult:	Low Risk		
Final risk: Cutoff Disorder: E d	1:21 1:25 dwards	0	-		R	esult:	Low Risk		
Final risk: Cutoff Disorder: E o Final risk:	1:21 1:25 dwards	0 ' Syndrome 0000	Risk type	Risk At Term	R	esult:	Low Risk		
Final risk:	1:21(1:25) dwards 1:10(1:10)	0 ' Syndrome 0000 0	Risk type Age risk:	Risk At Term 1:780		esult: esult:	Low Risk		
Final risk: Cutoff Disorder: E Final risk: Cutoff	1:21(1:25) dwards 1:10(1:10)	0 ' Syndrome 0000 0 ndrome	Risk type Age risk:	Risk At Term 1:780					



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Verified by **Mr. Pradip Kadam** Incharge Biochemistry



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Sample ID: 2200095391

Patient name : Mrs. CHAITALI MANJREKAR

PRENATAL SCREENING BACKGROUND

Every pregnant woman carries a certain degree of risk that her fetus/baby may have certain chromosomal defect/ abnormalities. Diagnosis of these fetal chromosomal abnormalities requires confirmatory testing through analysis of amniocytes or Chorionic Villous Samples (CVS). However, amniocentesis and CVS procedures carry some degree of risk for miscarriage or other pregnancy complications (Tabor and Alfirevic, 2010). Therefore in routine practice, prenatal screening tests are offered to a pregnant woman to provide her a personalised risk for the most common chromosomal abnormalities (T21-Down syndrome, T18- Edwards' syndrome, T13- Patau syndrome) using her peripheral blood sample. Based on this risk assessment, if the risk is high or intermediate, you can take informed decision of opting for invasive procedure such as amniocentesis or CVS followed by confirmatory diagnostic test(s), as per discussion with your clinician.

PRENATAL SCREENING TESTS ARE NOT CONFIRMATORY TESTS. THEY ARE LIKELIHOOD ASSESSMENT TESTS.

You may get your prenatal screening result as either of the following:-

High Risk

High Risk or Screen Positive Result: A High Risk Result does not mean that the pregnancy is affected with the condition. It means that the likelihood of the pregnancy having a condition is higher than the cut-off (Most commonly used cut-off is 1:250 and this represents the risk of pregnancy loss from confirmatory testing through CVS or amniocentesis).

Low Risk

Intermediat

Low Risk or Screen Negative Result: A Low Risk result does not mean that the pregnancy is not affected with a condition. It means that the likelihood of the pregnancy having a condition is lower than the cut-off.

Intermediate Risk result: An intermediate Risk result means that the pregnancy has an equivocal or a borderline risk of being affected with a condition. In this case, you may want to choose a second stage screening modality like an Integrated Screening Test that is done between 16 to 20 weeks of pregnancy or a Non-invasive Prenatal Screening Test between 12 to 20 weeks of pregnancy before taking a decision on an invasive confirmatory testing. This will help you improve the sensitivity of the screening test keeping an invasive test a last option were you to come as a high risk in the second stage screening test.

SIGNIFICANCE OF MULTIPLE OF MEDIANS (MoMs)

Prenatal Screening determines the likelihood of the pregnancy being affected with certain conditions by analysing levels of certain hormones. These hormones are Feto placental products (released by Fetus or placenta). Their levels not only indicate propensity of the fetus being affected with certain chromosomal conditions, they also provide indication of placental insufficiency that can potentially lead to pregnancy complications like Pre-Eclampsia or Intra-Uterine Growth Restriction. It is therefore important to take cognisance of the Reported MoMs alongside the Risk results.

For more information, visit our website at: www.lilacinsights.com/faq-pns

DISCLAIMERS

Limitations of the Test:

As prenatal screening tests are not confirmatory diagnostic tests, the possibility of false positive or false negative results can not be denied. The results issued for this test does not eliminate the possibility that this pregnancy may be associated with other chromosomal or sub- chromosomal abnormalities, birth defects and other complications.

Nuchal Translucency is the most prominent marker in screening for Trisomy 13, 18, 21 in the first trimester and should be measured in accordance with the Fetal Medicine Foundation (UK) guidelines. Nuchal Translucency or Crown Rump Length measurement, if not performed as per FMF (UK) imaging guidelines may lead to erroneous risk assessments and Lilac Insights bears no responsibility for errors arising due to sonography measurements not performed as per these criteria defined by international bodies such as FMF (UK), ISUOG.

It is assumed that the details provided along with the sample are correct. The manner in which this information is used to guide patient care is the responsibility of the healthcare provider, including advising for the need for genetic counselling or additional diagnostic testing like amniocentesis or Chorionic Villus Sampling. Any diagnostic test should be interpreted in the context of all available clinical findings. As with any medical test, there is always a chance of failure or error in sample analysis though extensive measures are taken to avoid these errors.

Note:

- Quality of the Down syndrome screening program (Biochemical values, MoMs and Risk assessments) is monitored by UKNEQAS on an ongoing basis.
- This interpretation assumes that patient and specimen details are accurate and correct.
- Lilac Insights does not bear responsibility for ultrasound measurements like CRL,NT,NB etc. We strongly recommend that ultrasound measurements are performed as per FMF (UK)/ISUOG practice guidelines.
- It must be clearly understood that the results represent risk and not diagnostic outcomes. Increased risk does not mean that the baby is affected and further tests must be performed before a firm diagnosis can be made. A Low Risk result does not exclude the possibility of Down's syndrome or other abnormalities, as the risk assessment does not detect all affected pregnancies.

END OF REPORT



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