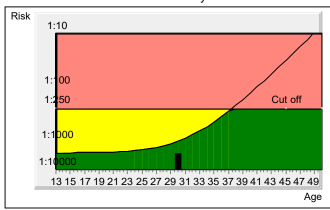
Results for:

MRS. MANOJ KUMARI

Referring Doctors

Sample no
2409051199/NOD
25/09/24

Summary



Patient data		
Age at delivery	30.0	
WOP	21 + 1	
Weight	45 kg	
Patient ID		
Ethnic origin	Asian	

For MRS. MANOJ KUMARI, born on 3/02/1995, a screening test was performed on the 24/09/2024. Prisca screens for Trisomy 21, Trisomy 18 and Neural tube defects (NTD).

MEASURED SERUM VALUES

	Valu	ie	Corr. MoMs	
AFP	94.1	ng/ml	1.10	
HCG	31174	mIU/ml	1.68	
uE3	2.8	ng/ml	1.07	
Gestation age 21+ 1				
Method BPD Hadlock				
The MoMs have been corrected according to:				
maternal we	ight			

TRISOMY 18 SCREENING

The calculated risk for trisomy 18 is < 1:10000, which indicates a low risk.

Risks at term		
Biochemical risk for Tr.21	1:1594	
Age risk:	1:832	
Neural tube defects risk	1:6973	

TRISOMY 21 SCREENING

The calculated risk for Trisomy 21 is below the cut off which represents a low risk.

After the result of the Trisomy 21 test it is expected that among 1594 women with the same data, there is one woman with a trisomy 21 pregnancy and 1593 women with not affected pregnancies.

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!

NEURAL TUBE DEFECTS (NTD) SCREENING

The corrected MoM AFP (1.10) is located in the low risk area for neural tube defects.



ethnic origin

