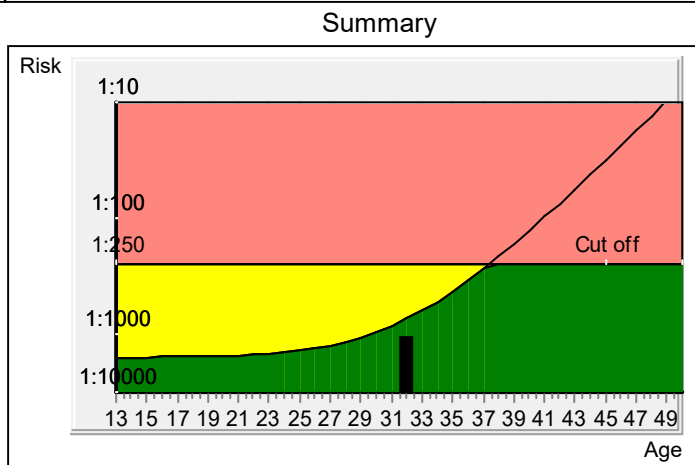


Results for:
Mrs. SANTHOSHASample no
A1699485Date of report:
06-01-2025

Referring Doctors



Patient data	
Age at delivery	32.0
WOP	15 + 1
Weight	45 kg
Patient ID	0012501010171
Ethnic origin	Asian

Risks at term	
Biochemical risk for Tr.21	1:1052
Age risk:	1:738
Neural tube defects risk	<1:10000

For Mrs. SANTHOSHA , born on 21-06-1993, a screening test was performed on the 01-01-2025. Prisca screens for Trisomy 21, Trisomy 18 and Neural tube defects (NTD).

MEASURED SERUM VALUES

	Value	Corr. MoMs
AFP	25.3 ng/mL	0.69
HCG	56222.6 mIU/mL	1.30
uE3	0.69 ng/mL	1.62
Gestation age	15+ 1	
Method	Scan	

The MoMs have been corrected according to:
maternal weight
ethnic origin

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 1052 women with the same data, there is one woman with a trisomy 21 pregnancy and 1051 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!

TRISOMY 18 SCREENING

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

NEURAL TUBE DEFECTS (NTD) SCREENING

The corrected MoM AFP (0.69) is located in the low risk area for neural tube defects.

Risk above
Cut off

Risk above
Age risk

Risk below
Age risk