

Prisca 5.1.0.17
Date of report: 18/01/25

KRANTHI DIAGNOSTICS

Patient data				
Name	Mrs. CHINNA JAMMAKKA		Patient ID	0352501170027
Birthday	01/01/05		Sample ID	A1175525
Age at sample date	20.0		Sample Date	17/01/25
Gestational age	12 + 4			
Correction factors				
Fetuses	1	IVF	no	Previous trisomy 21 pregnancies
Weight	43	diabetes	no	
Smoker	no	Origin	Asian	
Biochemical data			Ultrasound data	
Parameter	Value	Corr. MoM	Gestational age	12 + 3
PAPP-A	3.28 mIU/mL	0.55	Method	CRL Robinson
fb-hCG	32.35 ng/mL	0.69	Scan date	16/01/25
Risks at sampling date			Crown rump length in mm	
Age risk	1:1090		61	
Biochemical T21 risk	1:3677		Nuchal translucency MoM	
Combined trisomy 21 risk	<1:10000		1.01	
Trisomy 13/18 + NT	<1:10000		Nasal bone	
			present	
			Sonographer	
			N A	
			Qualifications in measuring NT	
			MD	
Risk			Trisomy 21	
			The calculated risk for Trisomy 21 (with nuchal translucency) is below the cut off, which indicates a low risk. After the result of the Trisomy 21 test (with NT) it is expected that among more than 10000 women with the same data, there is one woman with a trisomy 21 pregnancy. 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! The patient combined risk presumes the NT measurement was done according to accepted guidelines (Prenat Diagn 18: 511-523 (1998)). The laboratory can not be hold responsible for their impact on the risk assessment ! Calculated risks have no diagnostic value!	
Trisomy 13/18 + NT				
The calculated risk for trisomy 13/18 (with nuchal translucency) is < 1:10000, which represents a low risk.				

Sign of Physician