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					Date of Report PRISCA	31/1/2023 5.1.0.17
Patient Data						
Name		N	ARS. KOMAL	Patient ID		012301290058
Birthday			31/1/1995	Sample ID		11571493
Age at term			28.6	Sample Date		29/1/2023
Gestational age			11+5			
Correction factors						
Fetuses	1	IVF		unknown	Previous trisomy 21	unknown
Weight in kg	55	Diabetes		NO	Pregnancies	unknown
Smoker	NO	Origin		Asian		
Biochemical Data				Ultrasound Data		
Parameter	Value		Corr Mom			11+4
PAPP-A	5.62	mIU/ml	1.3	Method		CRL (<>Robinson)
fb-hCG	164.2 ng/ml		3.49	Scan date		28/1/2023
Risks at sampling date				Nuchal translucency 0.7		
Age Risk	age Risk		1:772	Nuchal translucency MoM		0.54
Biochemical T21 risk		1:356	Nasal bone		Present	
Combined trisomy 21 risk	ζ.		1:1967			
Trisomy 13/18 + NT			<1:10000			
Risk				Down's Syndro	ome Risk (Trisomy 21	Screening)
Risk 1:10 1:100 1:250			Cut off	cut off, which After the result expected that a same data, the pregnancy and	1 risk for Trisomy 21(v represents a low risk. t of the Trisomy 21with mong 1967 women wi re is one woman with a . 1966 women with not	n NT test it is th the trisomy 21 affected pregnancies.
1:1000 1:10000 13 15 17 19 21 23 25 27 29 31 33 35 37 39 41 43 45 47 49 Age Trisomy 13/18+NT The calculated risk for Trisomy 13/18 (with NT) is <1:10000, which indicates a low risk				The calculated risk by PRISCA depends on the accuracy of the information provided by the referring physician. Please note that the risk calculations are statistical aapproaches and have no diagnostic value! The laboratory cannot be hold responsible for their impact on the risk assessment! Calculated risks have no diagnostic values		
	Above Cı			Risk above Ag	e Risk	Risk below Age risk