

*Free Home Sample Collection 9999 778 778



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Date of Report 20-03-19 PRISCA 5.0.2.37

					PRISCA	5.0.2.37
Patient Data						
Name	Mrs Poonam			Patient ID		011903180272
Birthday			06-01-92	Sample ID		10453784
Age at delivery			27.2	Sample Date		18/03/19
Gestational age by			11+3			
Correction factors	1					
Fetuses	1	IVF		unknown	Previous trisomy 21	unknown
Weight in kg	55	Diabetes		no	Pregnancies	
Smoker	no	Origin		Asian		
iochemical Data			Ultrasound Data			
Parameter	Value		Corr Mom	Gestational ag	ge	11+2
PAPP-A	1.31	mIU/ml	0.48	Method		CRL (<>Hadlock)
fb-hCG	29.71	ng/ml	0.57	Scan Date		18-03-19
				CRL measurm	nents	45.9
Risks at sampling date				Nuchal translucency MoM 0.71		
Age Risk			1:813	Nasal bone		Present
Biochemical T21 Risk			1:2796	Sonographer		DR.SUBUHI RIJVI
Combined Trisomy 21 Risk			<1:10000	Qualifications	in measuring NT	MBBS
Trisomy 13/18			<1:10000			
Risk					rome Risk (Trisomy 21	
1:10				The calculated risk for Trisomy 21 (with nuchal translucency) is below the cut off, which indicates a		
				low risk.	is below the cut off, w	men mulcates a
		100		After the resul	It of the Trisomy 21 test	(with NT) it is
100				expected that among more than 10000 women with the same data, there is one woman with a trisomy 21		
1:250			Cut off	same data, the pregnancy.	re is one woman with a	trisomy 21
					l risk by PRISCA depen	ds on the accuracy of
				the information provided by the referring physician. Please		
				note that risk calculations are statistical approach and have no		
				diagnostic values! The patient combined risk presumes the NT		
					vas done according to acc	
Trisomy 13/18 + NT						
The calculated risk for trisomy 13/18 (with nuchal translucency) is < 1:10000, which represents a low risk.						
transfucency) is > 1.10000, which represents a low risk.						