

*Free Home Sample Collection 9999 778 778



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Date of Report	22-03-19
PRISCA	5.0.2.37

				P	PRISCA	5.0.2.37
Patient Data						
Name	Mrs Monika		Patient ID		011903200112	
Birthday	01-01-95		Sample ID		10410012	
Age at delivery			24.2	Sample Date		20/03/19
Gestational age by			12+4			
Correction factors						
Fetuses	1	IVF		unknown P	Previous trisomy 21	unknown
Weight in kg	53.5	Diabetes		no P	Pregnancies	
Smoker	no	Origin		Asian		
Biochemical Data				Ultrasound Dat	ta	
Parameter	Value		Corr Mom	Gestational age		12+2
PAPP-A	1.89	mIU/ml	0.42	Method		CRL (<>Hadlock)
fb-hCG	20.561	ng/ml	0.44	Scan Date		18-03-19
				CRL measurmer	nts	57.1
Risks at sampling date			Nuchal translucency MoM 0.73			
Age Risk			1:994	Nasal bone		Present
Biochemical T21 Risk			1:3877	Sonographer		DR.RAHUL
Combined Trisomy 21 R	isk		<1:10000	Qualifications in	n measuring NT	DNE
Trisomy 13/18			<1:10000			
Risk				Down's Syndro	ome Risk (Trisomy 21	1 Screening)
1:10			The calculated risk for Trisomy 21 (with nuchal translucency) is below the cut off, which indicates a			
				translucency) is low risk.	s below the cut off, w	inich indicates a
					of the Trisomy 21 test	(with NT) it is
1:100			expected that among more than 10000 women with the			
s		*	is one woman with a	trisomy 21		
1.250			Cut on	pregnancy.	: 1.1 DDIGGA 1	1 1 0
		The calculated risk by PRISCA depends on the accuracy of the information provided by the referring physician. Please				
		note that risk calculations are statistical approach and have no				
1:10000		diagnostic values!				
		The patient combined risk presumes the NT measurement was done according to accepted guidelines				
			Age	measurement wa	s done according to acc	sepied guideilnes
Trisomy 13/18 + NT The calculated risk for tri	somy 13/19	R (with no	rhal			
translucency) is < 1:10000, which represents a low risk.						