

\*Free Home Sample Collection 9999 778 778

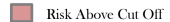


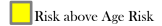
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Date of Report 21/02/2023 PRISCA 5.1.0.17

					PRISCA	5.1.0.17	
Patient Data	Value						
Name		MRS. PINK	Y YADA'	V	Patient ID	012302200012	
Birthday		08/0	1/198736.	5	Sample ID	11637908	
Age at delivery			3	0	Sample Date	20/02/2023	
Correction factors	S						
Fetuses	1	IVF		unknown	Previous trisomy 21	unknown	
Weight in kg	67	Diabetes		NO	Pregnancies	unknown	
Smoker	NO	Origin		Asian			
Biochemical Data				Risks at sampl	Risks at sampling date		
Parameter	Value	Cor	т МоМ	Age Risk		1:307	
AFP	66.2	ng/ml	1.02	Biochemical T	risomy 21 Risk	1:1958	
uE3	1.96	ng/ml	1.13	Neural Tube I	Defect Risk	Low risk area	
hCG	8551.8	mIU/ml	0.52	Trisomy 18		<1:10000	
Inhibin	227.3	IU/ml	1.37				
Ultrasound Data				Down's Syndro	ome Risk (Trisomy 2	21 Screening)	
Gestational age	20+4			The calculated risk for Trisomy 21 is below the cut off which represents a low risk.			
Method	BPD (<>Hadlock)			After the result of the Trisomy 21 test it is expected that			
				_	among 1958 women with the same data, there is one woman		
Risk				with a trisomy 21 pregnancy and 1957 women with not affected pregnancies.			
Risk 1:10				The calculated risk by PRISCA depends on the accuracy of			
1.10					the information provided by the referring physician. Please note that the risk calculations are statistical aapproaches and		
						itistical aapproaches and	
				have no diagno	osuc varue:		
1:100							
		/		TD: 10			
1:250		Out o	off	Trisomy 18			
				The calculated risk for Trisomy 18 is <1:10000, which			
1:1000				indicates a low risk Neural Tube Defect (NTD) Screening			
1:10000				Neural Tube I	Defect (N I D) Screen	ning	
13 15 17 19 21 23 25 27 29 31 33 35 37 39 41 43 45 47 49 Age				The corrected	MoM for AFP (1.02	2) is located in the low	

The laboratory can not be held responsible for their impact on the risk assessment! Calculated value has no diagnostic value!





risk area for neural tube defects.

