JITM Diagnostics

Patient data						
Name	MRS. PRATIKSHA			Patient ID		
Birthday			16/07/92	Sample ID		2305042712/NOD
Age at sample date	30.8		Sample Date		22/05/23	
Gestational age			12 + 2			
Correction factors						
Fetuses	1	IVF		no	Previous trisomy 21	no
Weight	65	diabetes		no	pregnancies	
Smoker	no	Origin		Asian		
Biochemical data				Ultrasound da	ata	
Parameter	Value		Corr. MoM	Gestational age 12 +		12 + 0
					9-	
PAPP-A	2.21 mIU/m	I	0.68	Method CRL		CRL Robinson
fb-hCG	89.4 ng/ml		1.99	Scan date		20/05/23
Risks at sampling date	C C			Crown rump length in mm		56
Age risk	e risk 1:572			Nuchal translucency MoM 0.61		
Biochemical T21 risk			1:284	Nasal bone		present
Combined trisomy 21 risk 1:1721			Sonographer DR. ANAMIKA			
Trisomy 13/18 + NT	risomy 13/18 + NT <1:10000			Qualifications in measuring NT M.D		
Risk				Trisomy 21		
1:10				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 1721 women with the same data,		
				there is one woman with a trisomy 21 pregnancy and 1720		
				women with not affected pregnancies.		
Cut off				The calculated risk by PRISCA depends on the accuracy of the information provided by the referring physician.		
				Please note	that risk calculations a	are statistical
				approaches	and have no diagnost	tic value!
1:1000				was done a	ccording to accepted g	es the NT measurement guidelines (Prenat Diagn
				18: 511-523		ponsible for their impact
1:10000				on the risk	assessment ! Calculate	sponsible for their impact ed risks have no
13 15 17 19 21 23 25	27 29 31 33 3	5 37 39 41 4	3 45 47 49	diagnostic v		_
			Age			

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