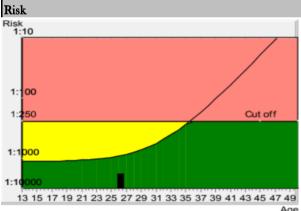




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Date of Report 23/3/2022 5.1.0.17 PRISCA Patient Data MRS. SHRIYA Patient ID 012203210131 Name 011119151 1/1/1996 Sample ID Birthday Age at term 26.08 Sample Date 22/3/2022 Gestational age Correction factors 1 IVF Fetuses unknown Previous trisomy 21 unknown 60 Diabetes NO Pregnancies Weight in kg unknown NO Origin Smoker Asian **Biochemical Data** Ultrasound Data 12+3 Value Corr Mom Gestational age **Parameter** 3.28 mIU/ml0.59 CRL (<>Robinson) Method 0.84 19/3/2022 43.4 ng/mlScan date Nuchal translucency 1.3

PAPP-A fb-hCG Risks at sampling date Age Risk 1:914 0.84 Nuchal translucency MoM Biochemical T21 risk 1:2392 Nasal Bone Presesnt Combined trisomy 21 risk <1:10000 Trisomy 13/18+NT <1:10000



Down's Syndrome Risk (Trisomy 21 Screening)

The calculated risk for Trisomy 21 with NT is below the cut off, which represents a low risk.

After the result of the Trisomy 21 test it is expected that among more than 10000 women with the same data, there is one woman with a trisomy 21 pregnancy 9999 women with not affected pregnancies. 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!

Trisomy 13/18+NT The calculated risk for Trisomy 13/18 with NT is <1:10000, which indicates a low risk

The laboratory cannot be hold responsible for their impact on the risk assessment! Calculated risks have no diagnostic values



Risk Above Cut Off

Risk above Age Risk



Risk below Age risk