Interpretation

Test Performed: Chromosome FISH, Prenatal (CHR FISHP)
Specimen Type: Direct amniocytes (uncultured cells collected from pour-off at first feed of cultures)
Indication for Testing: Mono / di pregnancy affected with TTT syndrome; sample taken from twin A

RESULT
Normal FISH Result (Male)

INTERPRETATION
There was no evidence for aneuploidy of chromosomes 13, 18, 21, X and Y in 50 interphase cells scored.

This panel will not detect approximately one third of prenatal chromosome abnormalities. Aneuploidy of other chromosomes, structural abnormalities, and mosaicism have not been ruled out by this analysis. Additional testing is recommended for the final interpretation of this result; pending results will be reported separately.

This analysis was performed with chromosome enumeration probes for 13, 18, 21, X and Y using the FDA-approved AneuVysion probe kit (Abbott Molecular).

Cytogenomic Nomenclature (ISCN):

\text{nuc ish(DXZ1x1,DYZ3x1,D18Z1x2),(RB1,D21S259/D21S341/D21S342)x2}

This result has been reviewed and approved by

This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the U.S. Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes.