

Revised charges effective from 15th July 2017

Disease	Charges per sample	Service Tax @ (18%)	Total Charges	Reporting Time
<b>Beta-Thalassemia</b> Seven common mutations (IVS 1-5 G>C, IVS 1-1 G>T, 619 bp deletion, Codon 15 G>A, Codon 30 G>A, FS 8/9 +G, FS 41/42 - CTTT) Screening for specific mutation (as per the request of doctor) Detection of novel mutations (by sequencing)	Rs 1,525/- Rs 1,017/- Rs 2,034/-	Rs 275/- Rs 183/- Rs 366/-	Rs 1,800/- Rs 1,200/- Rs 2,400/-	2 weeks 10 days 4 weeks
<b>Sickle Cell Anemia</b> <b>Hemoglobin-E Disease</b>	Rs 1,017/- Rs 1,017/-	Rs 183/- Rs 183/-	Rs 1,200/- Rs 1,200/-	10 days 10 days
<b>Duchenne and Becker Muscular Dystrophy</b> Deletion analysis for 24 exons Deletion analysis for 79 exons by MLPA analysis Carrier analysis using MLPA	Rs 1,525/- Rs 2,542/- Rs 2,542/-	Rs 275/- Rs 458/- Rs 458/-	Rs 1,800/- Rs 3,000/- Rs 3,000/-	2 weeks 2 weeks 2 weeks
<b>Spinal Muscular Atrophy</b> Deletion analysis (exon 7 and 8 only) Carrier analysis using MLPA *SMN gene sequencing (exons 1 to 8)	Rs 1,525/- Rs 1,525/- Rs 4,576/-	Rs 275/- Rs 275/- Rs 824/-	Rs 1,800/- Rs 1,800/- Rs 5,400/-	2 weeks 2 weeks 4 weeks
<b>Fragile-X Syndrome</b> *CGG repeat tract analysis by Amplidex Kit <b>Myotonic Dystrophy</b> TP-PCR based analysis	Rs 3,559/- Rs 1,525/-	Rs 641/- Rs 275/-	Rs 4,200/- Rs 1,800/-	10 days 4 weeks
<b>Triplet Repeat Expansion Associated Diseases</b> Huntington's Disease, Spinocerebellar Ataxias 1, 2, 3, 6 and 12, DRPLA, Friedrich's Ataxia, SBMA	Rs 1,525/- for single disease Rs 3,559/- for panel (SCA 1, 2, 3)	Rs 275/- Rs 641/-	Rs 1,800/- Rs 4,200/-	2 weeks 4 weeks
<b>Hemophilia-A</b> Carrier analysis @ intragenic markers (BclI & XbaI RFLP) <b>Hemophilia-B</b> Carrier analysis @ intragenic markers (HhaI, DdeI, TaqI)	Rs 1,525/- Rs 1,525/-	Rs 275/- Rs 275/-	Rs 1,800/- Rs 1,800/-	3 weeks 3 weeks
<b>Chronic Pancreatitis (N34S SPINK1 mutation)</b>	Rs 1,017/-	Rs 183/-	Rs 1,200/-	10 days
<b>Cystic fibrosis</b> delta F508 mutation Four common mutations (dF508, G542X, G551D, R553X) 5T mutation <b>Glucose 6 phosphate dehydrogenase deficiency</b> Orissa, Mediterranean and Kerala-Kalyan mutations <b>Pre-coagulation profile</b> Factor V Leiden and Prothrombin (G20210A) mutation Single mutation analysis (any of the above two mutations) <b>MTHFR gene polymorphisms</b> 677C>T and 1298A>C mutations <b>Hereditary Haemochromatosis</b> H63D and C282Y mutations <b>Mitochondrial encephalopathy</b> <b>LHON</b> - 3 mutations (G3460A, G11778A, T14484C) <b>Leigh' disease</b> -3 mutations (T12706C, A13084T, G13513A) Specific mutation (as per the request of referring doctor)	Rs 1,017/- Rs 1,525/- Rs 1,017/- Rs 2,034/- Rs 1,525/- Rs 1,017/- Rs 1,525/- Rs 2,034/- Rs 2,034/- Rs 1,017/-	Rs 183/- Rs 275/- Rs 183/- Rs 366/- Rs 275/- Rs 183/- Rs 275/- Rs 366/- Rs 366/- Rs 183/-	Rs 1,200/- Rs 1,800/- Rs 1,200/- Rs 2,400/- Rs 1,200/- Rs 1,200/- Rs 1,800/- Rs 2,400/- Rs 2,400/- Rs 1,200/-	10 days 2 weeks 10 days 2 weeks 2 weeks 10 days 2 weeks 2 weeks 3 weeks 3 weeks 10 days
<b>Non-Syndromic Hearing Loss (NSHL)</b> Connexin 26 (GJB2) exon 2 screening (by sequencing)	Rs 1,525/-	Rs 275/-	Rs 1,800/-	2 weeks
<b>DNA Isolation and Storage up to two years</b>	Rs 595/-	Rs 105/-	Rs 700/-	1 week
<b>Prenatal Diagnosis</b>	Rs 3,559/-	Rs 641/-	Rs 4,200/-	3-7 days
<b>Maternal cell contamination</b>	Rs 1,525/-	Rs 275/-	Rs 1,800/-	3-7 days
<b>Prenatal Diagnosis &amp; Maternal cell contamination</b>	Rs 5,085/-	Rs 915/-	Rs 6,000/-	3-7 days