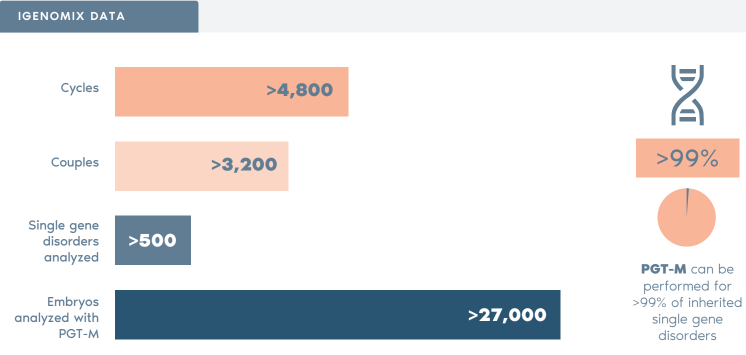
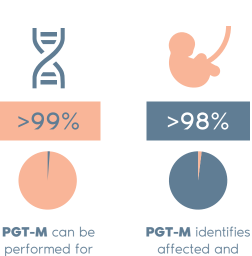


## PGT-M Update - Igenomix clinical results

PGT-M prevents the transmission of single gene disorders to offspring. This test is for couples with a family history or known carrier status of monogenic diseases such as cystic fibrosis, fragile X syndrome or Huntington's, among others.





## PGT-A and PGT-M can be performed on the same sample

## **Indications**

Monogenic disease

&

Advanced maternal age Recurrent miscarriage Severe male factor

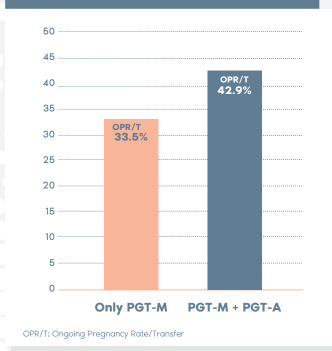
Previous pregnancy with trisomy

Previous pregnancy with trisomy

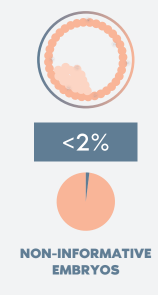
(XO, XXX, XXY, XYY)

Translocations and inversions analyzed only by aCGH (comparative genomic hybridization)

## ADVANTAGES OF PERFORMING PGT-M WITH PGT-A



50% of normal embryos for single gene disorders are affected by chromosomal abnormalities<sup>1</sup>





Igenomix internal data