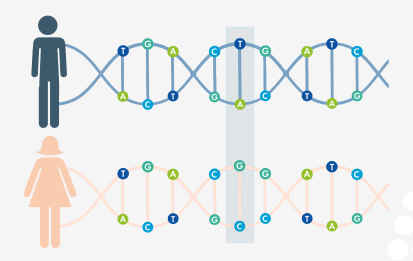
Smart



To enhance the genetic testing of IVF embryos and expand the diagnostic capabilities and clinical utility of our preimplantation genetic testing for aneuploidy (Smart PGT-A), we have developed and validated a parallel targeted Next-Generation Sequencing (NGS) strategy using the power of SNP technology without the need for parental samples.

Single Nucleotide Polymorphisms (SNPs) are changes in single nucleotides distributed throughout the genome and frequently vary at the same genomic position between individuals. Most SNPs have only two different alleles.

SNPs can be used for 'DNA fingerprinting' to detect ploidy differences, DNA contamination and the genetic relatedness of embryos.





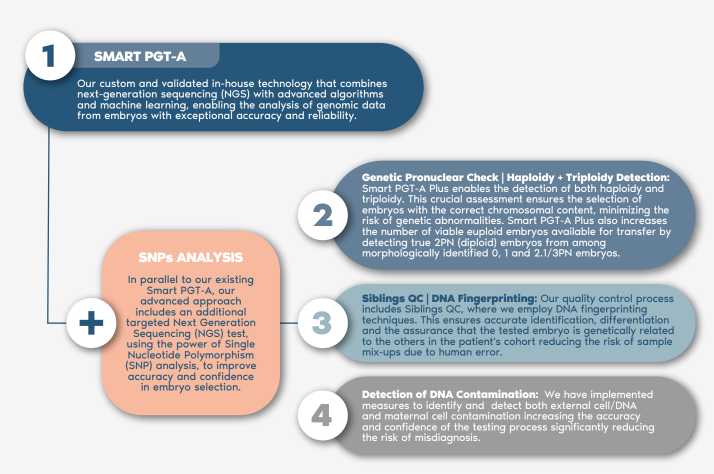
We call this dual assessment **Smart PGT-A Plus**. Building on our extensive expertise in the field of embryo genetic testing, this advanced solution goes beyond Smart PGT-A by incorporating additional features including ploidy analysis, sibling embryo genetic relatedness and DNA contamination detection into a standard PGT-A workflow.

Smart PGT-A & Smart PGT-A Plus **Compared**

		Smart PGT-A	Smart PGT-A PLUS
Technology	NGS	+	+
	SNPs	-	+
Results	Aneuploidies	+	+
	Mosaicism	+	+
	Segmentals*	≥ 10 MB	≥ 10 MB
	Accuracy	98%	98%
	Mitoscore	+	+
New features	Ploidy	-	+
	Contamination (maternal and external)	-	+
	Sibling QC	-	+
	Sibling QC	-	+



Experience the power of our new **Smart PGT-A Plus**, a 4-in-1 genetic test that empowers informed decision-making for embryo transfer







Smart PGT-A Plus uses two independent analyses on every sample to deliver a comprehensive 4-in-1 genetic test for aneuploidy in embryos.







Who should consider Smart PGT-A Plus?

While any couple can have an embryo with aneuploidy, the chances can increase with the following factors:

- Age of oocyte provider
- History of recurrent pregnancy loss
- Previous IVF failure
- Prior child or pregnancy with a chromosome abnormality
- Patients with severe male factor or high rate of diploidy in sperm
- Previous or recurrent triploid pregnancy
- Previous molar pregnancy
- Rescue of high quality embryos derived from abnormally fertilized oocytes (OPN, 1PN, 2.1PN/3PN)

What makes our Smart PGT-A Plus stand out from the rest?



Offers enhanced confidence with robust and accurate results, utilizing two independent analyses to detect abnormalities.



Strengthened by the power of big data and artificial intelligence, effectively overcoming the limitations of human subjectivity and greatly reducing the risk of human error.



Maximizes the likelihood of successful pregnancy by carefully identifying optimal embryos for transfer.



Reduces the risk of miscarriage due to previously undetected abnormalities (e.g. triploidy).



Increases the number of viable embryos available for transfer.



Enhances accuracy and reduces the risk of misdiagnosis by detecting external and maternal cell DNA contamination.



Provides confirmation that all embryos from the same patient are genetically related to each other without the need for additional parental samples.



Ensures enhanced quality control in the laboratory procedures conducted within your IVF lab, providing greater assurance.