

QUANTIFY FUNCTIONAL DNA

Measure PCR-amplifiable DNA and discover PCR inhibition within the sample, guiding smarter sample input for NGS projects.

RESCUE LOW PERFORMING SAMPLES

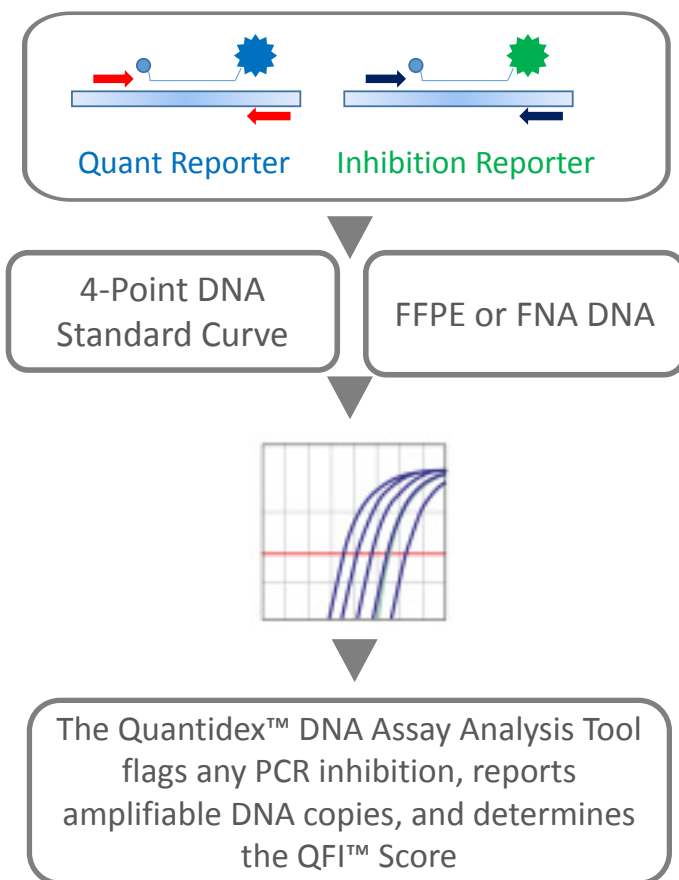
Calculate sample input based on functional DNA template rather than bulk DNA, ensuring enough sample complexity to avoid false calls and expensive run failures.

SIMPLE WORKFLOW

Rapid, single-reaction qPCR-based assay that can be set up in less than 10 minutes.

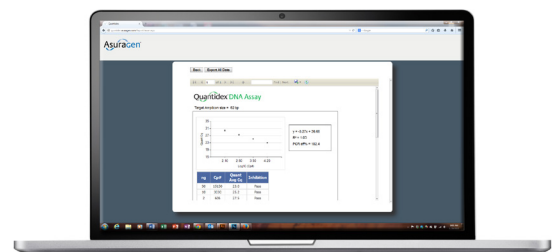
CONVENIENT REPORTING

Leverage the easy-to-use Quantidex™ DNA Assay reporting tool to automatically calculate functional DNA copy number and minimum sample input requirements.



The Quantidex™ DNA Assay is a multiplexed quantitative PCR assay that measures the absolute copy number of PCR-amplifiable DNA in a sample and reports PCR inhibition.

The Quantidex™ DNA Assay determines the functional quality of sample DNA using the Quantitative Functional Index (QFI™) Score, designating the fraction of total genomic DNA copies that can be successfully amplified. The QFI™ Score and amplifiable copy number provide actionable guidance that informs the input requirements for NGS target enrichment and helps assure a high level of analytical sensitivity and specificity. In addition, the Quantidex™ DNA Assay flags PCR inhibitors in the sample and provides an opportunity to salvage such samples through a subsequent clean-up step prior to further processing.



Convenient Reporting

Research Use Only. Not for use in diagnostic procedures.

KIT ORDERING INFORMATION

Quantidex™ DNA Assay*

[P/N 49539] 100 REACTIONS

*Research Use Only. Not for use in diagnostic procedures.

REFERENCES

1. Functional DNA quantification guides accurate next-generation sequencing mutation detection in formalin-fixed, paraffin-embedded tumor biopsies. Sah et al. Genome Med 2013;5(8):77
2. Evaluation of an integrated clinical workflow for targeted next-generation sequencing of low-quality tumor DNA using a 51-gene enrichment panel. Choudhary et al. BMC Med Genomics 2014;(7):62
3. Variation in pre-PCR processing of FFPE samples leads to discrepancies in BRAF and EGFR mutation detection: A diagnostic RING trial. Kapp et al. J Clin Pathol 2015;(68):111-118

WATCH THE TECH TALK

“Measure the DNA that Matters: The Quantidex™ DNA Assay as the Foundation for an Integrated NGS Workflow Solution”

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www.asuragen.com/tech-talks



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