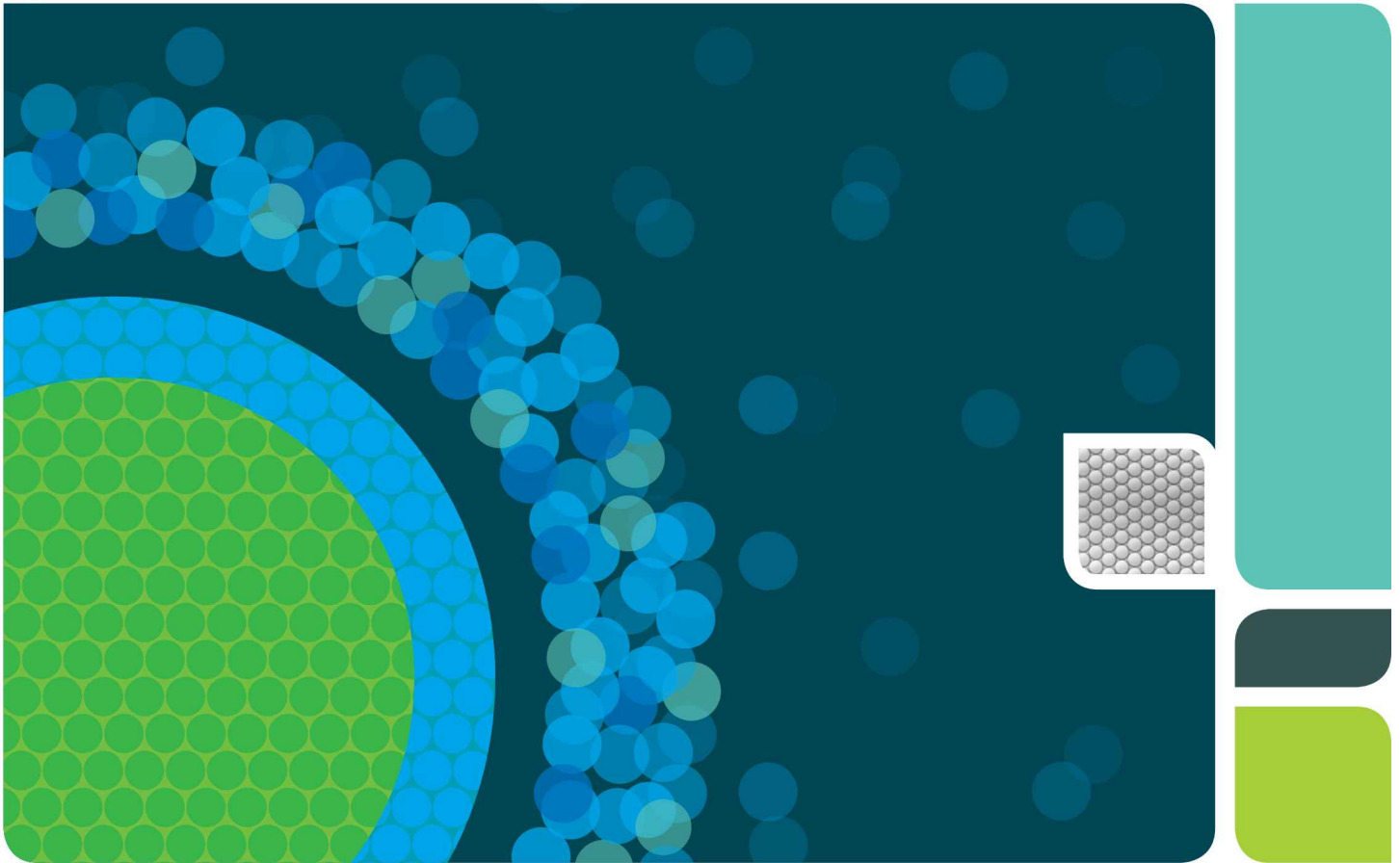




QX200™ Droplet Digital™ PCR System



IT STARTS WITH A
DROPLET
AND ENDS IN
DISCOVERY.



Droplet Digital PCR (ddPCR™) is Bio-Rad's unique digital PCR technology. With unrivaled precision, Droplet Digital PCR provides absolute quantification of target DNA or RNA molecules without the use of standard curves. It addresses the lack of scalable and practical technologies for digital PCR implementation. The new QX200 Droplet Digital PCR System puts this powerful technology in your hands, ready to unveil new worlds of research at previously unattainable levels.

PCR becomes “digital” through sample partitioning and the subsequent statistical analysis of target detection across the partitions. The QX200 System uses advanced microfluidics technology to achieve partitioning on a massive scale, generating 20,000 highly uniform nanoliter-sized droplets per sample. For even higher sensitivity, it is possible to combine wells and analyze millions of droplets for a single sample.

Simple and reliable, Droplet Digital PCR technology has already led to several research breakthroughs in fields such as cancer biomarker discovery, infectious diseases, genomic alterations, and gene expression. This research is only the beginning. What discoveries will you make using Droplet Digital PCR?

WHAT'S IN A DROPLET?

A VERSATILE AND SCALABLE WORKFLOW



Droplet Digital PCR has a simple, user-friendly experiment setup designed to process eight samples at a time. The process easily scales up to run a 96-sample experiment in 5 hours with minimal hands-on time. When higher throughput is required, multiple 96-sample experiments can be run in a day.

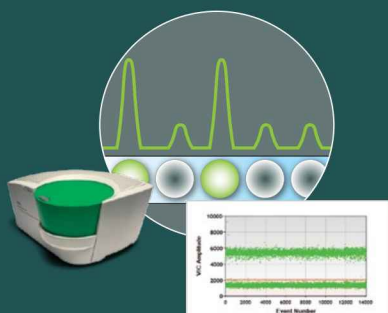
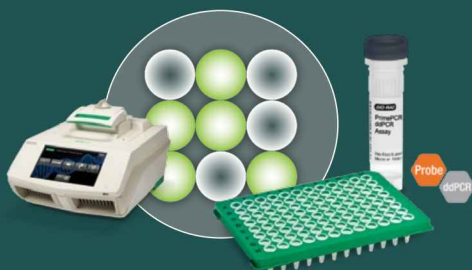
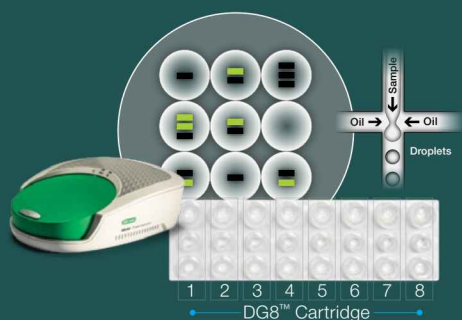
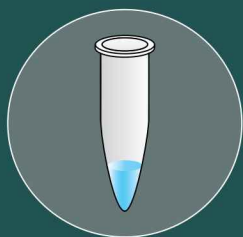
Bio-Rad's QX200 ddPCR System combines water-oil emulsion droplet technology with microfluidics. The QX200 System consists of two instruments — a droplet generator and a droplet reader — and associated consumables. The droplet generator partitions each sample into 20,000 uniform nanoliter-sized droplets in which nucleic acid molecules are distributed in a random fashion. Each droplet serves to partition the reactions. Droplets are transferred to a 96-well PCR plate and PCR is performed to end point in a thermal cycler. The droplets stream single file through the reader for fluorescence analysis. Positive droplets, which contain at least one copy of the target DNA or RNA molecule, exhibit increased fluorescence compared to negative droplets. The fraction of PCR-positive droplets enables the target to be quantified according to the Poisson distribution.

The QX200 System works with both hydrolysis probes and EvaGreen fluorescence detection chemistries, and its flexible design allows for high throughput and ultra-sensitive detection.



KEY BENEFITS

- Achieve absolute quantification without the use of a standard curve
- Design scalable assays for high sensitivity or high throughput
- Expand applications using flexible ddPCR chemistry — EvaGreen or probes



1

Prepare ddPCR reaction mix

- Combine DNA sample, primers, and/or probes with one of Bio-Rad's ddPCR Supermixes
- Fully validated PrimePCR ddPCR Assays can be used

2

Generate droplets

- Load the ddPCR reaction mix into the wells of a droplet generator cartridge
- 8 x 20,000 droplets are generated from each run in the QX200 Droplet Generator
- Target DNA (■) and background DNA (■) are randomly distributed in droplets

3

Perform PCR with EvaGreen or hydrolysis probes

- Transfer the droplets to a 96-well PCR plate and seal the plate
- Run the PCR protocol

4

Read and analyze results

- After PCR, load the 96-well PCR plate into the QX200 Droplet Reader
- Positive and negative droplets in each sample are read
- Analyze concentrations with QuantaSoft™ Software

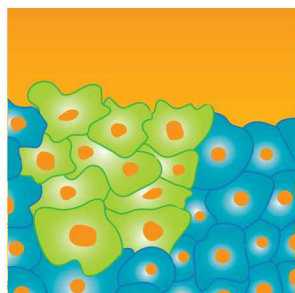
WHAT'S IN A DROPLET?

QUANTIFYING BIOMARKERS FOR CANCER



The level of sensitivity offered by the QX200 ddPCR System in quantifying cancer biomarkers overcomes the limitations posed by other methods. Using Droplet Digital PCR technology, researchers are now able to observe finer quantitative distinctions among mutations, better identifying their potential roles in cancer.

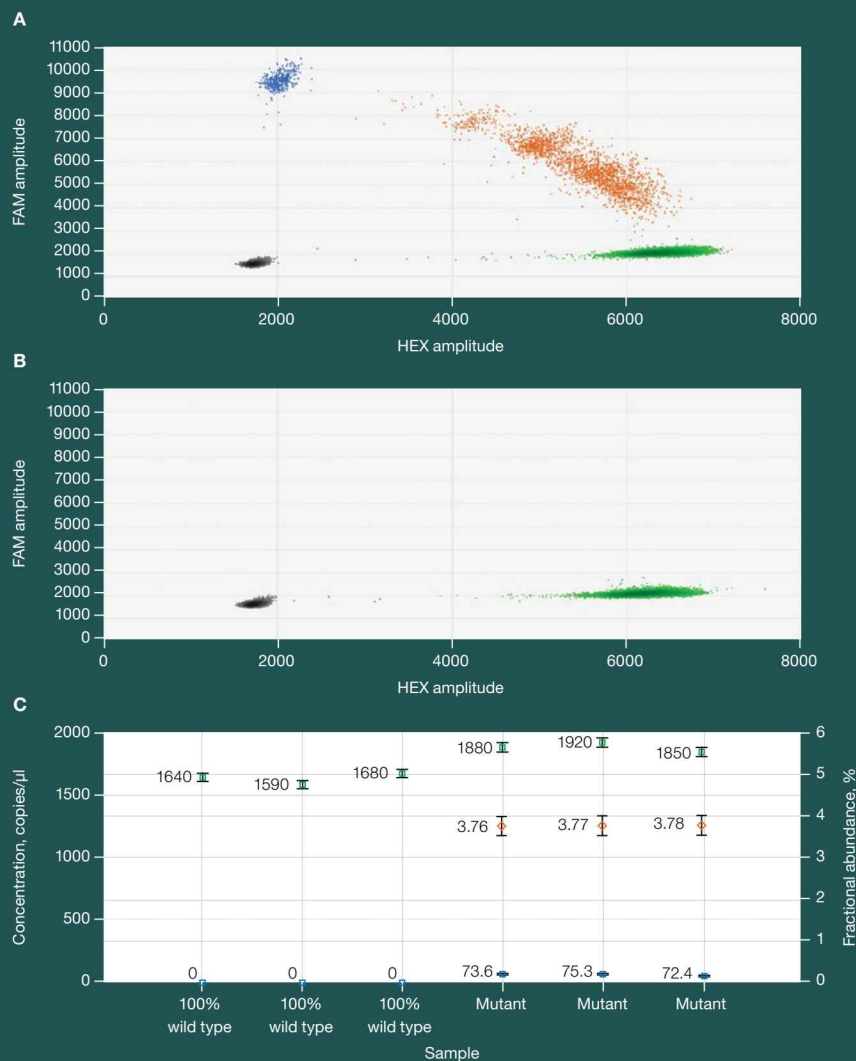
Cancer-associated mutations often evade detection due to their low concentrations relative to the background of wild-type DNA in a given sample. With its high sensitivity, the QX200 System can easily scale to quantify target concentrations as low as one out of 1,000,000 (0.0001%) total copies. What was previously undetectable with other methods can now be quantified with Droplet Digital PCR.



SUCCESS STORY

Detecting *T790M* mutation in epidermal growth factor receptor (*EGFR*), an important therapeutic target in some lung cancers, allows for a better understanding of resistance to tyrosine kinase inhibitor therapies. However, other techniques lacked the sensitivity to reliably identify the *T790M* allele in a high background of *EGFR* wild type. Using Droplet Digital PCR, a researcher in the biotechnology industry was able to develop a highly accurate assay to quantify *T790M* concentration.

DETECTION OF *BRAF* V600E MUTATION



High sensitivity of the QX200 System allows quantification of *BRAF* V600E mutation in the presence of wild-type DNA using PrimePCR ddPCR Assays. **A**, 2-D fluorescence amplitude plot generated by QuantaSoft Software shows triplicate wells of a mixed mutant and wild-type sample. The black cluster on the plot represents the negative droplets, the green cluster represents the droplets that are positive for wild-type DNA only, the blue cluster represents the droplets that are positive for mutant DNA only, and the orange cluster represents the droplets that are positive for both mutant and wild-type DNA. **B**, 2-D fluorescence amplitude plot shows three replicates of a wild-type-only sample. **C**, fractional abundance plot shows the percentage frequency (orange markers) of the mutant DNA in a wild-type DNA background. The blue markers indicate the concentration of mutant DNA (copies/ μ l) and the green markers indicate the concentration of wild-type DNA (copies/ μ l) in each of three replicate samples. All error bars are generated by QuantaSoft Software and represent a 95% confidence interval.

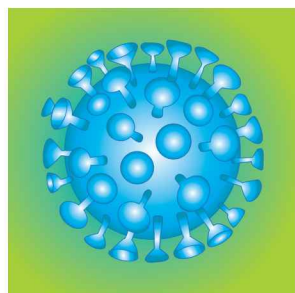
WHAT'S IN A DROPLET?

INSIGHT INTO VIRAL RESERVOIRS



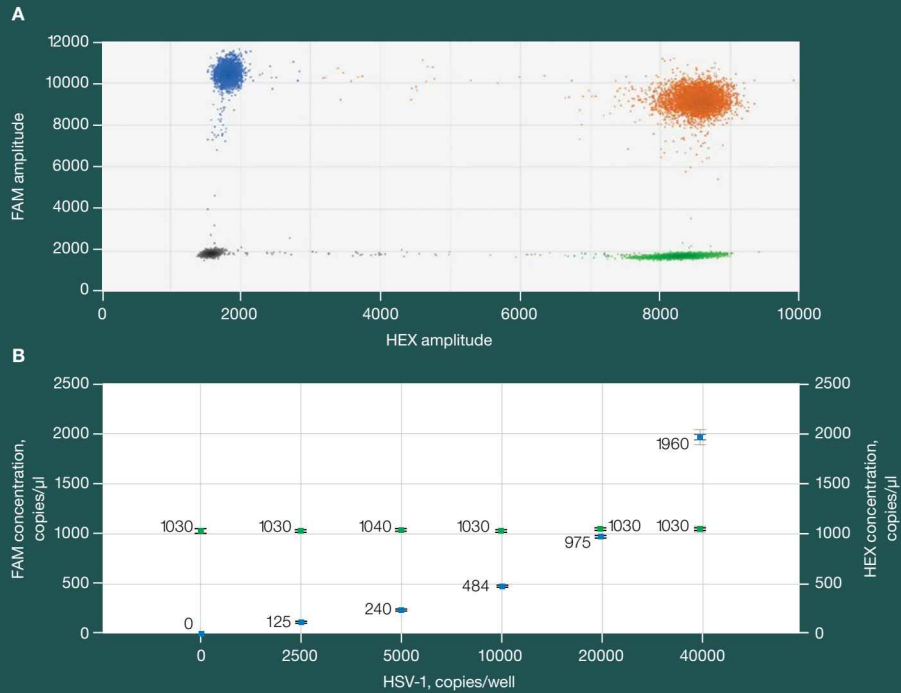
Precisely quantifying viral load is crucial for characterizing disease states and developing and validating therapies. Viral reservoirs fluctuate, sometimes dropping to very low levels that are nonetheless significant. Obtaining exact and reproducible measurements of viral DNA or RNA often represents the difference between success and failure in studying many different pathogens.

The QX200 System has the sensitivity to register extremely minute quantities of viral genetic material and distinguish it from complex mixtures. Investigators can obtain actual copy numbers of viral genome by generating complementary DNA through reverse transcription. Along with precision and reliability, the QX200 System also offers users the throughput needed to process samples with great efficiency.



SUCCESS STORY

A researcher compared quantitative PCR (qPCR) and Droplet Digital PCR approaches for detecting residual human immunodeficiency virus (HIV) infection in a clinical sample set, which included an infant who was functionally cured of HIV. The researcher found that Droplet Digital PCR was five times more sensitive than qPCR when measuring HIV DNA copies/million cells, and more than 20 times more accurate when measuring viral long terminal repeats (2-LTR circles).

DETECTION OF HERPES SIMPLEX VIRUS AND $\beta 2$ MICROGLOBULIN

Droplet Digital PCR enables precise and reproducible detection of herpes simplex virus 1 (HSV-1) and $\beta 2$ microglobulin (*B2M*) targets in a duplex assay. **A**, 2-D fluorescence amplitude plot shows three merged replicate samples of 10,000 copies/well of HSV-1 duplexed with 20,000 copies/well of *B2M*. The black cluster on the plot represents the negative droplets, the green cluster represents the droplets that are positive for HSV-1 only, the blue cluster represents the droplets that are positive for *B2M* only, and the orange cluster represents the droplets that are positive for both HSV-1 and *B2M* targets. **B**, concentration plot shows merged triplicate wells across an HSV-1 sample dilution series from 0 to 40,000 copies in a constant background of 20,000 copies (66 ng human genomic DNA) of *B2M*. The blue markers indicate HSV-1 copies/ μ l and the green markers indicate *B2M* copies/ μ l. All error bars are generated by QuantaSoft Software and represent a 95% confidence interval.

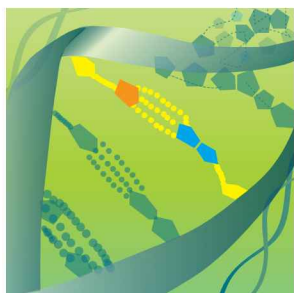
WHAT'S IN A DROPLET?

DISTINGUISHING GENOMIC VARIATIONS



Copy number variation (CNV) is a prominent source of interindividual variability in the human genome. CNV has also been associated with cancers, neurological and autoimmune diseases, and adverse drug responses. Reliably identifying such CNVs represents an important capability in cutting-edge research.

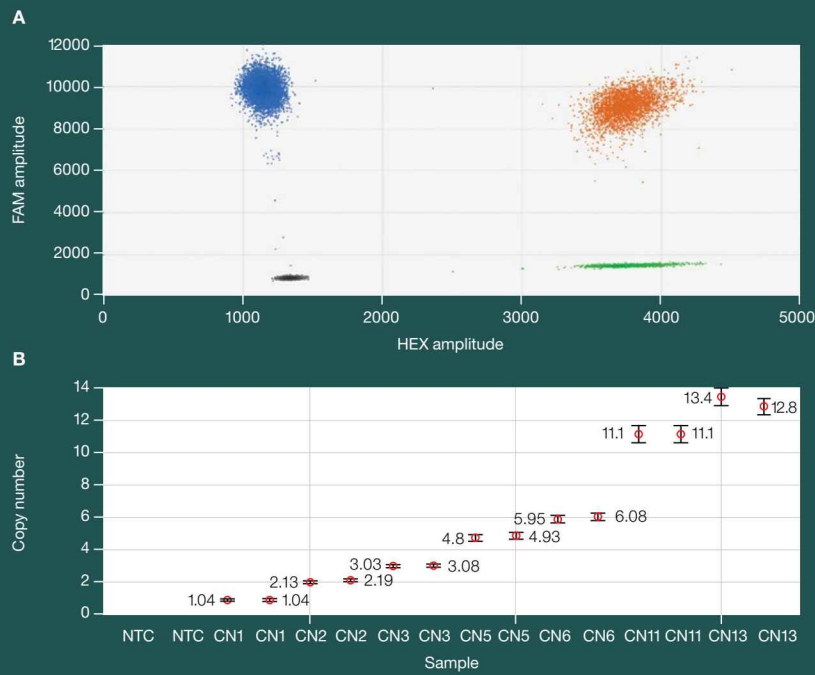
The major technical hurdle in copy number (CN) assessment is the ability to discriminate, with statistical confidence, between consecutive CN states. Fundamentally, as CN state increases, the percentage difference in target genomic material between states decreases, making it harder to measure CNV at higher orders. By partitioning the amplification reaction across thousands of droplets, the QX200 System can resolve high consecutive copy number states (such as five vs. six) with an excellent degree of statistical confidence.



SUCCESS STORY

A researcher investigating a highly duplicated gene involved in human brain size evolution needed a method to validate small CN variations in test samples. Droplet Digital PCR proved much more effective than other methods the researcher had previously used, showing greater precision in detecting small differences at high CN states, as well as excellent reproducibility. These improvements in CNV quantification will allow the researcher to ask more complex questions regarding neurogenetic disorders such as micro/macrocephaly, schizophrenia, and autism.

DETECTION OF COPY NUMBER FOR MYELOCYTOMATOSIS ONCOGENE AND β -GLOBIN



The QX200 System provides superior resolution with copy number (CN) calls from CN1 to CN13. **A**, 2-D fluorescence amplitude plot shows duplicate wells of a CN6 sample duplexed with myelocytomatosis oncogene (*MYC*) and β -globin (*HBB*) assays. The black cluster on the plot represents the negative droplets, the green cluster represents the droplets that are positive for *MYC* only, the blue cluster represents the droplets that are positive for *HBB* only, and the orange cluster represents the droplets that are positive for both *MYC* and *HBB* targets. **B**, copy number plot shows samples ranging from CN1 to CN13 with precise duplicate well values. All error bars are generated by QuantaSoft Software and represent a 95% confidence interval. NTC, no template control.

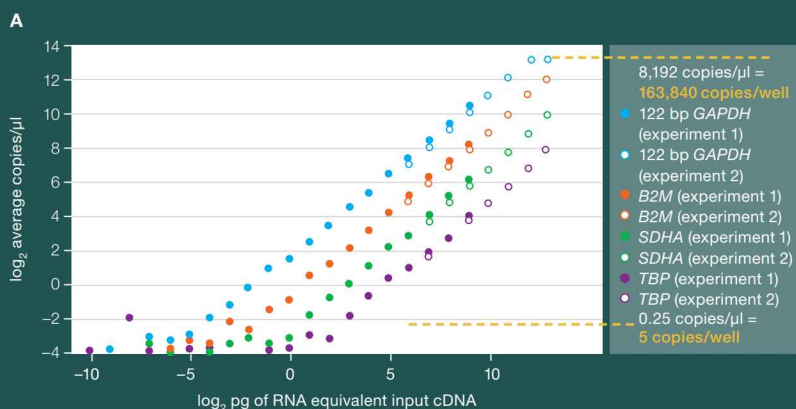
NGS VALIDATION AND LIBRARY QUANTIFICATION



GENE EXPRESSION STUDIES



VALIDATION FOR NEXT-GENERATION SEQUENCING



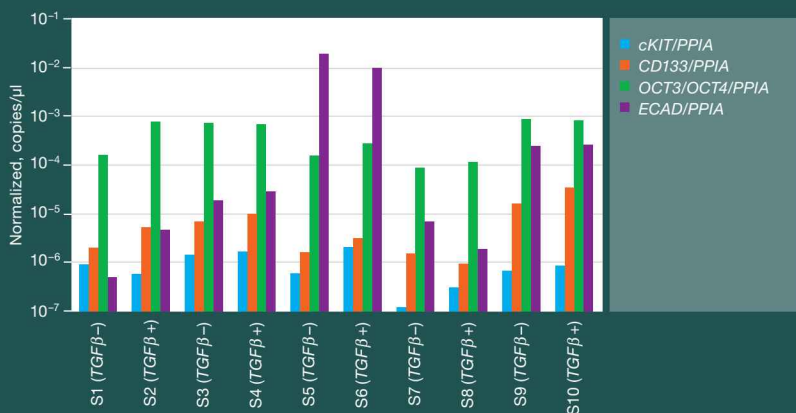
B

Gene	Droplet Digital PCR, copies/well		MiSeq, RPKM			
	100 ng RNA 4 replicates	1,000 ng 4 replicates	100 ng 2 replicates		1,000 ng 2 replicates	
<i>GAPDH</i>	1,671 ± 115*	16,275 ± 479*	974	953	1,077	1,061
<i>B2M</i>	504 ± 46*	3,450 ± 155*	233	251	229	237
<i>SDHA</i>	139 ± 14*	1,131 ± 81*	51	50	59	65
<i>HPRT1</i>	15,781 ± 2,310	140,705 ± 11,059	34	22	25	25
<i>TBP</i>	3,650 ± 178	31,625 ± 1,010	3	3	6	3
<i>GUSB</i>	1,794 ± 53	15,731 ± 1,134	9	12	13	12

* Obtained with cDNA diluted 200-fold because of its high abundance.

Droplet Digital PCR provides nearly 1,000-fold greater sensitivity and an amplification bias-free approach compared to RNA-Seq for validation. **A**, concentration plot shows a twofold dilution of cDNA generated using Bio-Rad's iScript™ Advanced cDNA Synthesis Kit for RT-qPCR. Two independent measurement sets were made: one at a high concentration range and one at a lower range, with four points overlapping. **B**, table shows comparative results between Droplet Digital PCR and MiSeq. Droplet Digital PCR detected thousands of copies per well with low-abundance transcripts (*TBP* and *GUSB* genes) using an input of 100 ng total RNA, which demonstrates enhanced sensitivity. The MiSeq Sequencer detected only single-digit reads per kilobase per million reads (RPKM) for these transcripts.

GENE EXPRESSION ANALYSIS OF STEM CELL MARKERS



High sensitivity and precision of the QX200 System allow reliable detection of small fold changes with rare transcripts. The *TGFβ* signaling pathway plays a role in ovarian stem cell regulation, and the addition of *TGFβ* will increase the number of stem cells in the population, increase the expression of putative stem cell markers, and decrease terminal differentiation markers like E-cadherin (*ECAD*). Data show expression levels of *cKIT*, *CD133*, *OCT3/OCT4*, and *ECAD* genes across ten samples with and without *TGFβ* treatment, normalized to *PPIA* in this study. Reproducible detection of cDNA copies as low as 0.5–1 target/μl in a 20 μl reaction was achieved. S, sample.

WHAT'S IN A DROPLET?

ABSOLUTE QUANTIFICATION WITH EvaGreen



Through droplet partitioning and statistical analysis, Droplet Digital PCR provides absolute quantification without the need for a standard curve, bringing substantial improvements to many applications that require detection of low-abundance target copies or distinction of small fold changes between samples. Combining Droplet Digital PCR with the power of EvaGreen detection chemistry enables genotyping and gene expression analysis for low-level target genes. The QX200 ddPCR System enables absolute quantification using EvaGreen chemistry, making such assays ideal for target DNA measurement, viral load analysis, and microbial quantification.

VALIDATED ddPCR ASSAYS



Bio-Rad's PrimePCR ddPCR* Assays are fully wet-lab validated assays, expertly designed to provide single-copy PCR resolution without the use of a standard curve. These mutation detection and copy number assays have been validated specifically for digital PCR.

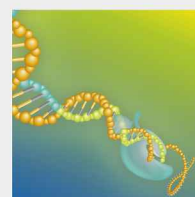
Bio-Rad's PrimePCR Real-Time PCR Assays* can also be used on the QX200 Droplet Digital PCR System with QX200 ddPCR EvaGreen Supermix. Choose from more than 40,000 existing human and mouse gene expression assays.

* Each assay for the human, mouse, and rat genomes was experimentally tested for optimal efficiency, specificity, sensitivity, and linear dynamic range.



SUCCESS STORY

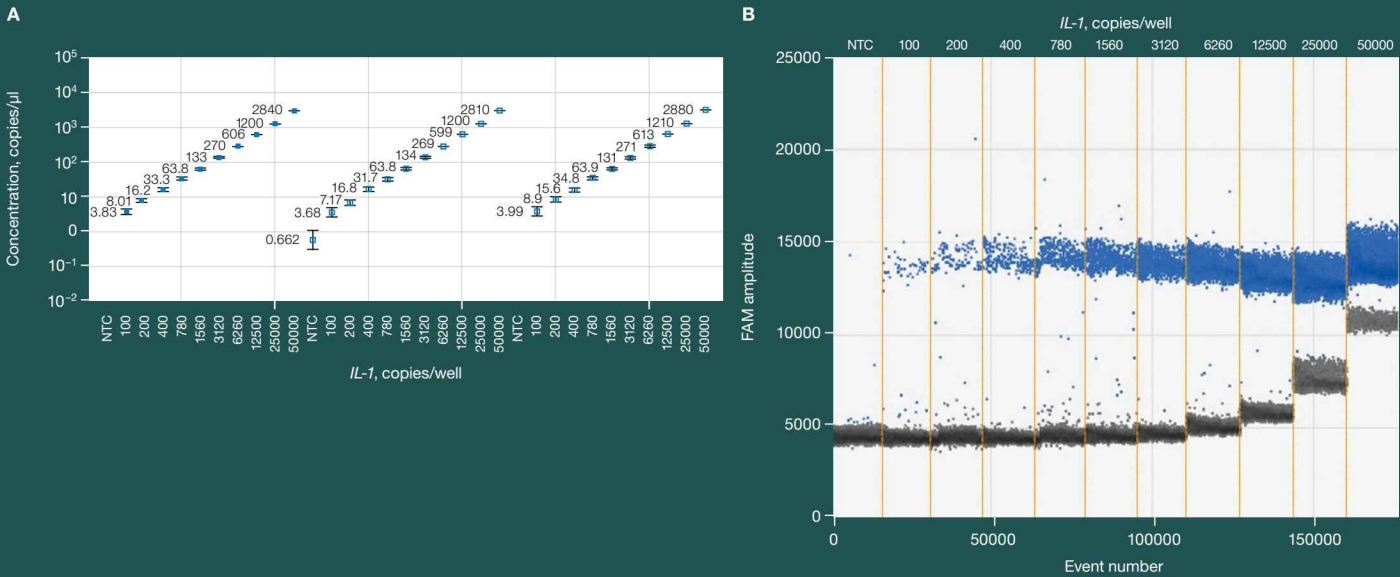
Precision cancer medicine involves rapid assessment of cancer mutations, with certain genetic errors serving as indicators for a specific therapy. A researcher was challenged with the composition and degradation of clinical samples that were used for assessment of *FLT3* mutations for gastric tumors (sporadic diffuse). By partitioning the samples' DNA templates, Droplet Digital PCR removed the effects of PCR inhibitors and bias in the data, resulting in a more reliable quantification of the mutations present in the samples.



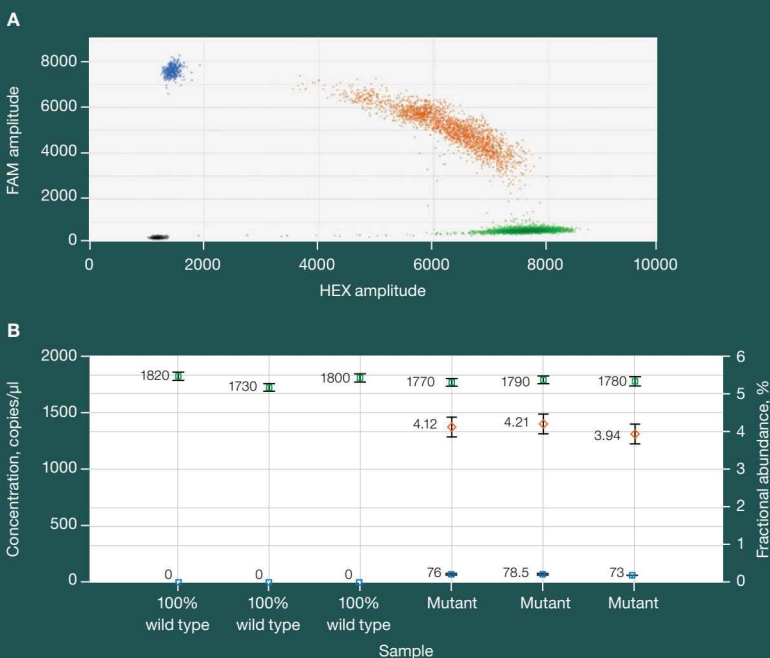
SUCCESS STORY

Designing an optimal probe assay may pose challenges and delay generation of reliable data. A researcher spent a great deal of time trying to design probes and primers for *EGFR L858R*, a receptor for epidermal growth factor, which is associated with lung tumors. Using the fully validated PrimePCR ddPCR Mutation Detection Probe Assay for *EGFR L858R* eliminated these tedious optimization steps, allowing the researcher to focus on the biology of the investigation, rather than assay design.

ABSOLUTE QUANTIFICATION USING EvaGreen



Droplet Digital PCR enables precise and reproducible detection calls across a range of sample concentrations. **A**, concentration plot shows data for both merged wells (solid blue markers) and individual replicate wells (open blue markers) across a dilution series. Interleukin 1 (*IL-1*) samples ranging from 0 to 50,000 copies were detected with an EvaGreen Assay and plotted on a log scale. There is excellent linearity and reproducibility between replicates. **B**, 1-D fluorescence amplitude plot shows a sample dilution curve detected with an EvaGreen *IL-1* Assay in the FAM/EvaGreen channel of the QX200 Droplet Reader. The data demonstrate good separation between positive (●) and negative (■) data points. All error bars are generated by QuantaSoft Software and represent a 95% confidence interval. NTC, no template control.

DETECTION OF *EGFR* L858R MUTATION

PrimePCR ddPCR Assays are fully validated for the detection of *EGFR* L858R mutation. **A**, 2-D fluorescence amplitude plot shows triplicate wells of a mixed mutant and wild-type sample. The black cluster on the plot represents the negative droplets, the green cluster represents the droplets that are positive for wild-type DNA only, the blue cluster represents the droplets that are positive for mutant DNA only, and the orange cluster represents the droplets that are positive for both mutant and wild-type DNA. **B**, fractional abundance plot shows the percentage frequency (orange markers) of the mutant DNA in a wild-type DNA background. The blue markers indicate the concentration of mutant DNA (copies/μl) and the green markers indicate the concentration of wild-type DNA (copies/μl) in each of three replicate samples. All error bars are generated by QuantaSoft Software and represent a 95% confidence interval.



More Droplet Digital PCR Resources

Visit bio-rad.com/web/QX200 for a wide range of videos, technical reports, published findings, and other resources describing Bio-Rad's unique Droplet Digital PCR technology and the QX200 ddPCR System.

Specifications

QX200 Droplet Generator	
Starting sample size	20 µl
Capacity	1–8 samples/cartridge
Droplets per sample	20,000
Dimensions (W x D x H)	28 x 36 x 13 cm (11 x 14 x 5 in.)
QX200 Droplet Reader	
Precision	±10%
Linear dynamic range	5 orders of magnitude
Capacity	1–96 samples
Droplets per 96-well plate	Approximately 1,500,000
Sample illumination	Light-emitting diodes
Sample detection	Multipixel photon counter
Detection channels	FAM (EvaGreen), HEX (VIC)
Dimensions (W x D x H)	66 x 52 x 29 cm (26 x 20 x 11 in.)

Ordering Information

Catalog # Description

QX200 Droplet Digital PCR System

1864001	QX200 Droplet Digital PCR System , includes droplet generator, droplet reader, laptop computer, software, associated component consumables
1864002	QX200 Droplet Generator , includes droplet generator, 1 pkg of 24 DG8 Cartridges, 1 pkg of 24 DG8 Gaskets, 2 cartridge holders, 1 power cord
1864003	QX200 Droplet Reader , includes droplet reader, ddPCR manual, 2 plate holders, USB cable, power cord
1864007	Droplet Generator Cartridges and Gaskets , includes 5 pkg of 24 DG8 Cartridges, 5 pkg of 24 DG8 Gaskets
1864008	DG8 Cartridges for QX200/QX100™ Droplet Generator , 1 pkg of 24 cartridges
1863009	DG8 Gaskets for QX200/QX100 Droplet Generator , 1 pkg of 24 gaskets
1863051	DG8 Cartridge Holder
1863005	Droplet Generation Oil for Probes , 10 x 7 ml
1864005	QX200 Droplet Generation Oil for EvaGreen , 2 x 7 ml
1864006	QX200 Droplet Generation Oil for EvaGreen , 10 x 7 ml
1863004	ddPCR Droplet Reader Oil , 2 x 1 L

ddPCR Reagents

1863026	ddPCR Supermix for Probes , 2 ml (2 x 1 ml), 200 x 20 µl reactions, 2x supermix
1863010	ddPCR Supermix for Probes , 5 ml (5 x 1 ml), 500 x 20 µl reactions, 2x supermix
1863027	ddPCR Supermix for Probes , 25 ml (5 x 5 ml), 2,500 x 20 µl reactions, 2x supermix
1863028	ddPCR Supermix for Probes , 50 ml (10 x 5 ml), 5,000 x 20 µl reactions, 2x supermix
1863023	ddPCR Supermix for Probes (No dUTP) , 2 ml (2 x 1 ml), 200 x 20 µl reactions, 2x supermix
1863024	ddPCR Supermix for Probes (No dUTP) , 5 ml (5 x 1 ml), 500 x 20 µl reactions, 2x supermix
1863025	ddPCR Supermix for Probes (No dUTP) , 25 ml (5 x 5 ml), 2,500 x 20 µl reactions, 2x supermix

Catalog #	Description
1863021	One-Step RT-ddPCR Kit for Probes , 2 ml (2 x 1 ml), 200 x 20 µl reactions, 2x RT-ddPCR mix, includes 1 manganese acetate tube
1863022	One-Step RT-ddPCR Kit for Probes , 5 ml (5 x 1 ml), 500 x 20 µl reactions, 2x RT-ddPCR mix, includes 2 manganese acetate tubes
1864033	QX200 ddPCR EvaGreen Supermix , 2 ml (2 x 1 ml), 200 x 20 µl reactions, 2x supermix
1864034	QX200 ddPCR EvaGreen Supermix , 5 ml (5 x 1 ml), 500 x 20 µl reactions, 2x supermix
1864035	QX200 ddPCR EvaGreen Supermix , 25 ml (5 x 5 ml), 2,500 x 20 µl reactions, 2x supermix
1864036	QX200 ddPCR EvaGreen Supermix , 50 ml (10 x 5 ml), 5,000 x 20 µl reactions, 2x supermix
1863052	ddPCR Buffer Control Kit , 9 ml (2 x 4.5 ml), 2x buffer
1864052	QX200 Buffer Control for EvaGreen , 9 ml (2 x 4.5 ml), 2x buffer
1863040	ddPCR Library Quantification Kit for Illumina TruSeq , 200 x 20 µl reactions, includes ddPCR Supermix for Probes (No dUTP) (2 x 1 ml vials), ddPCR Library Quantification Assay (1 x 200 µl vial), for quantification of Illumina TruSeq libraries using the QX200 or QX100 System
1863041	ddPCR Library Quantification Kit for Ion Torrent , 200 x 20 µl reactions, includes ddPCR Supermix for Probes (No dUTP) (2 x 1 ml vials), ddPCR Library Quantification Assay (1 x 200 µl vial), for quantification of Ion Torrent libraries using the QX200 or QX100 System
Thermal Cycler and Plate Sealer	
1851197	C1000 Touch™ Thermal Cycler with 96-Deep Well Reaction Module , includes C1000 Touch Thermal Cycler Chassis, 96-deep well reaction module, USB flash drive
1814000	PX1™ PCR Plate Sealer , includes heat sealing instrument, plate support block that holds 96-well and 384-well plates, sealing frame, power cord

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The QX100 or QX200 Droplet Digital PCR System and/or its use is covered by claims of U.S. patents, and/or pending U.S. and non-U.S. patent applications owned by or under license to Bio-Rad Laboratories, Inc. Purchase of the product includes a limited, non-transferable right under such intellectual property for use of the product for internal research purposes only. No rights are granted for diagnostic uses. No rights are granted for use of the product for commercial applications of any kind, including but not limited to manufacturing, quality control, or commercial services, such as contract services or fee for services. Information concerning a license for such uses can be obtained from Bio-Rad Laboratories. It is the responsibility of the purchaser/end user to acquire any additional intellectual property rights that may be required.

Bio-Rad's thermal cyclers and real-time thermal cyclers are covered by one or more of the following U.S. patents or their foreign counterparts owned by Eppendorf AG: U.S. Patent Numbers 6,767,512 and 7,074,367.

The QX200 Droplet Digital PCR System and Automated Droplet Generator are for research use only and are not intended for use in diagnostic procedures.



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