Droplet digital PCR is an accurate method to assess methylation status on FFPE samples

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ABSTRACT
Most tissue samples available for cancer research are archived as formalin-fixed paraffin-embedded (FFPE) samples. However, the fixation process and the long storage duration lead to DNA fragmentation and hinder epigenome analysis. The use of droplet digital PCR (ddPCR) to detect DNA methylation has recently emerged. In this study, we compare an optimized ddPCR assay with a conventional qPCR assay by targeting a dilution series of control DNA. In addition, we compare the ddPCR technology with results from Infinium arrays targeting two separate CpG sites on a set of colon adenoma FFPE samples. Our data demonstrate that qPCR and ddPCR assess methylation status equally well on dilution controls with a high DNA input. However, the methylation detection on low-input samples is more accurate using ddPCR. The proposed primer design (methylation-independent primers with amplification of solely the converted DNA target) will allow for methylation detection, independent of bisulfite conversion efficiency. Those data show that ddPCR can be used for methylation analysis on FFPE samples with a wide range of DNA input and that the precision of the assay depends largely on the total amount of amplifiable DNA fragments. Due to accessibility of the ddPCR technology and its accuracy on high- as well as low-DNA input samples, we propose the use of this approach for studies involving degraded FFPE samples.

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BRIEF REPORT

Introduction
Epigenetic control mechanisms play a central role in a wide variety of natural and disease processes [1]. The best known epigenetic mechanisms include DNA methylation, histone alterations, and noncoding RNA-associated gene silencing. Ablative DNA methylation is described in many diseases, including cancer, vascular diseases, and neurodegenerative disorders [2–4]. Epigenetic factors may also contribute to the interperson variation in drug response [5]. The ability to detect these altered methylation patterns is key in understanding the complex biology of diseases. To date, several methods for methylation analysis are available, each with their (dis)advantages [6,7], and the selection of the appropriate technology should be based on the nature of the clinical samples [blood sample, fresh-frozen tissue, formalin-fixed paraffin-embedded (FFPE) samples], the amount and quality of the DNA, the required sensitivity and precision, and the availability of the equipment. Whole-genome methylation profiling methods include (but are not limited to) array hybridization, bisulfite (next-gen) sequencing, immunoprecipitation-based methods, and chromatography methods [6,8,9]. Recently, bisulfite-free sequencing methods (Nanopore sequencing and Single-Molecule, Real-Time (SMRT) sequencing with the PacBio platform) have emerged to assess the methylation status [10–13]. PCR-based methods (including Methylight qPCR, pyrosequencing, methylation-sensitive restriction endonuclease methods) are widely used for the confirmation of the findings from initial screenings performed with one of the whole-genome profiling methods [7,14].

In recent years, the droplet digital PCR (ddPCR) system has gained interest for the detection of rare mutations, copy number variations, gene rearrangements, and pathogen detection [15]. ddPCR is based on sample partitioning, by randomly distributing a single sample into small droplets, such that some droplets have no template molecules and others have one or more. Each droplet undergoes PCR amplification and analysis separately. The droplets are then individually assessed and scored as positive or negative for fluorescence. The Poisson limit theorem allows for the estimation of the per-droplet distribution of the number of target copies. This then allows for the absolute quantitation of the target sequence [16]. The use of ddPCR for DNA methylation detection has recently emerged, mainly for use with liquid biopsies [17,18]; a limited number of studies have examined tissue samples or other body fluids [18–21]. Most tissue samples available for cancer research are archived as Formalin-Fixed Paraffin-Embedded (FFPE) samples for many years. Although those retrospective sample repositories are valuable for biomarker research, the fixation process and the long storage duration have a major impact on the DNA quality [22–24]. Bisulfite treatment, which involves the deamination of unmodified cytosine residues to uracil, will cause further DNA degradation. In this study, we compare an optimized ddPCR assay with a conventional qPCR assay by targeting a dilution series of control DNA. In addition, we compare the ddPCR technology with results from Infinium
arrays targeting two separate CpG sites on a set of colon adenoma FFPE samples.

**Results**

**Accurate methylation detection with droplet digital PCR**

We have evaluated the performance of conventional qPCR and ddPCR for methylation detection of the CpG sites cg07164631 and cg25249613 on two series of mixtures of methylated and unmethylated DNA controls. Primer pairs targeting only converted cytosine residues were designed. This allowed for the amplification of only the bisulfite-converted DNA sequence, but not of the non-converted DNA template. To enforce that the primers efficiencies are methylation-independent, no CpG sites were included in the primer pairs. Two probes per primer pair were designed with one probe, labeled with FAM, targeting the methylated sequence and the second probe, labeled with HEX, targeting the unmethylated sequence (Table 1). Identical primer/probe combinations were used for both technologies but not of the non-converted DNA template. To enforce that accurate methylation detection on low-input samples is more accurate using ddPCR.

**Table 2. Summary of the median differences between expected and measured methylation status. Numbers between brackets represent the 2.5th and 97.5th percentiles.**

| ASSAY          | PCR       | Low-input control series | High-input control series |
|---------------|-----------|--------------------------|---------------------------|
| cg07164631    | ddPCR     | 0.00 (–0.461 –7.03)      | 1.04 (–1.49 – 4.87)       |
| cg07164631    | qPCR      | 9.28 (4.05 – 16.74)      | 5.52 (–4.92 – 8.20)       |
| cg25249613    | ddPCR     | 1.03 (–6.60 – 5.35)      | 1.16 (–0.84 – 3.92)       |
| cg25249613    | qPCR      | 4.18 (–2.08 – 12.14)     | 4.71 (–4.23 – 8.90)       |

**Table 1. Primer and probe sequences.**

| Assay                        | cg07164631 | cg25249613 | Cytosine-free priming assay |
|------------------------------|------------|------------|-----------------------------|
| Forward primer sequence (5'-3') | TTTATGGTAATTTTTGAAATGGTTT | TTTATATAGTTAGGGGTAGGT | TGG GTT AAA GTG ATT GAG TAA |
| Reverse primer sequence (5'-3') | TTATATGCATAAATCTGGTACCACTACT | ACTAAAAAAACCTAAATAATTTTACAAAC | TAT TCA TCC TTC AAT TTA CCC T |
| Probe sequence (5'-3')        | TTAGGATAAATCAGTTTTT | ACTAAAAAAACCTAAATAATTTTACAAAC | TGTTAATGTTAGGGGTAGGT |
|                              | FAM*: AATGATTGTTGATGAAA | FAM*: TAAAGGATAAATCAGTTTTT | FAM*: TGGTTAATGTTAGGGGTAGGT |
|                              | HEX*: AATGATTGTTGATGAAA | HEX*: TAAAGGATAAATCAGTTTTT | HEX*: TGTTAATGTTAGGGGTAGGT |

* probe targeting methylated (FAM) and unmethylated (HEX) sequence.

**No impact of low bisulfite conversion efficiency on the methylation detection**

The first step in the methylation detection of clinical samples is an optimal DNA extraction and bisulfite conversion method. To avoid any effect of an incomplete bisulfite conversion on the accurate methylation detection, the primers we have designed target only converted cytosine residues. Consequently, only the bisulfite converted DNA will be amplified and not the non-converted DNA. We have experimentally evaluated the performance of the two ddPCR assays (cg07164631 and cg25249613) on varying degrees of bisulfite converted control DNA (0%, 25%, 50%, 75%, and 100%) in a background of non-converted control DNA. As shown in Figure 2, an increase in the absolute amount of detected copies is observed by an increasing percentage of converted DNA in the samples. However, the methylation status is stable [mean methylation status ± 95% CI of 87.6 ± 3.8% (cg07164631) and 92.7 ± 2.5% (cg25249613); Figure 2]. These data confirmed that an optimal design of the primers/probe can avoid any potential impact of an incomplete bisulfite conversion on the accurate methylation detection.

**Droplet digital PCR can assess methylation statuses on FFPE samples**

The methylation statuses of the two CpGs (cg07164631 and cg25249613) were tested on FFPE tissue samples (n = 109) using ddPCR. To assess the success of the bisulfite conversion, an additional assay was designed using primers targeting cytosine-free sequences (adapted from [25]). This allowed for the amplification of both the converted and non-converted sequences. Two probes, one labeled with FAM and one with HEX, were designed to bind converted and non-converted DNA sequence, respectively. A mean bisulfite conversion efficiency of 99.57 ± 2.03% is detected on those clinical samples. One sample was excluded from further analysis due to low amount (<8000) of generated droplets in the ddPCR. As shown in Figure 3, a wide range of both the total copies per reaction and
the percentage methylation in those 108 clinical samples was detected. The mean 95% CI were small [3.09% (cg07164631) and 4.35% (cg25249613)], and the magnitude was comparable to the experiments with the control DNA mixtures. Only 13 (6%) out of the 216 generated data points have an associated 95% CI greater than 10% and all these have a low amount (<750) of total copies per reaction. The five data points with a 95% CI >15% have an even smaller amount (<250) of positive droplets in the reaction. Those data have demonstrated that droplet digital PCR can be used for methylation analysis on FFPE samples with a wide range of DNA input and that precision of the assay depends largely on total copies per reaction, i.e., total amount of amplifiable DNA fragments.

In addition, we have compared the methylation data for those 2 CpG sites from the Infinium arrays and the ddPCR assays. The correlation between those two technologies is demonstrated in Figure 4 (slope = 1.2 ± 0.03; r² = 0.78). The ddPCR technology could therefore be valuable for confirmation of genome-wide methylation screenings.

**Discussion**

A limited amount of reports regarding methylation detection using ddPCR are available in the literature, most of them with a focus on the sensitivity of this technology [19]. In most cases, methylation-specific PCR primers, which do contain CpG sites and specifically amplify the methylated DNA sequence, and cytosine-less primers (to control for DNA input) were used [19]. In this study, we have used a different approach for the primer/probe design. Since no CpG sites were included in the primer pairs, the primers were methylation-independent, which would allow for a proportional amplification of methylated and unmethylated bisulfite converted templates and minimize any PCR bias. Optimal bisulfite conversion should allow

![Figure 1. Bland-Altman plots for assessment of accuracy of the two CpG sites cg07164631 (A) and cg25249613 (B) using ddPCR and qPCR. The difference in measured and expected percentage methylation (y-axis) is plotted against the expected percentage methylation of the dilution series (x-axis). Two series of mixtures with methylated and unmethylated DNA controls were analyzed. The upper figures (control series 1) have a 5-fold higher DNA input. Dashed line = median; dotted lines = 2.5th and 97.5th percentile; curved blue solid line = Loess; dashed blue delimit 95% confidence band around Loess.](image-url)
for the sufficient conversion of cytosine residues while leading to only limited inappropriate conversion of methylated cytosine residues and limited DNA degradation. With the experiments in this study, we have demonstrated that the proposed primer design [i.e., methylation-independent primers with amplification of solely the converted DNA target, in combination with (un)methylation-specific probes] will allow for an accurate and reliable methylation detection, independent of bisulfite conversion efficiency.

Droplet digital PCR technology is based on sample partitioning by randomly diluting a single sample into many droplets, such that some droplets have no template molecules and others have one or more. This endpoint PCR method does not require standard curves for absolute quantification (unlike conventional qPCR), can be easily optimized and is therefore rapidly emerging in many applications [15]. Since each droplet in the ddPCR assays independently undergoes PCR amplification, there is less impact of PCR inhibition. By using series of mixtures of methylated and unmethylated DNA controls, we have demonstrated the similar performances of qPCR and ddPCR (for the two primer pairs used) for the methylation detection on high-input samples. Both methods have demonstrated a
linear and accurate detection over the complete methylation range. Moreover, the high accuracy of the methylation detection obtained with ddPCR was also obtained with low-DNA input samples, which was opposite to qPCR. Although it should be noted that the excellent amplification achieved may not be generalizable to all primer pairs, the primers used allow us to explore the characteristics of both methods. Due to the high accuracy on both high- and low-input samples, the convenient optimization and readily access to the ddPCR technology, ddPCR has many advantages for studies using highly-degraded FFPE samples. In this study, a precise methylation detection was obtained for most clinical samples (which contain a wide range of amplifiable DNA fragments). Since the fixation process and the long storage duration of such samples have a major impact on the DNA quality, care should be given to those parameters and for certain sample sets the DNA could be too degraded to obtain reliable methylation data. This precision could be further enhanced by running multiple replicates in separate wells and merging those data for the analysis (a well-known feature of the QuantaSoft software).

In conclusion, our data have demonstrated that droplet digital PCR can be a valuable tool for the precise and accurate methylation detection on clinical samples and can be used for confirmation of genome-wide screenings.

Materials and methods

Samples

Colon adenoma samples of patients undergoing surveillance colonoscopy were acquired from Avaden Biosciences. The median storage time of the samples was 12 years.

DNA extraction and bisulfite conversion

Paraffin was removed from one FFPE slice (10 μm) with deparaffinization solution (Qiagen) and DNA was extracted using the QiaAmp DNA FFPE Tissue. This was followed by bisulfite conversion using the EpiTect Fast Bisulfite Conversion Kit (Qiagen). Three extractions/bisulfite conversion reactions per sample were pooled into a total elution volume of 60 μl.

Controls

Synthetic DNA fragments (gBlocks Gene Fragments; stock concentration 5 ng/μl; IDT), corresponding to either the fully methylated or unmethylated bisulfite converted DNA sequence surrounding the CpG site of interest, were synthesized. A series of mixtures of methylated and unmethylated DNA controls (gBlock fragments: 10% methylation differences) were prepared fresh and stored in TE (10 mM Tris-Cl, 1 mM EDTA) buffer with 100 ng/μl carrier RNA. Two series were made; dilution series 2 has a 5-fold lower DNA input than dilution series 1. For the first series of methylation mixtures, a mean of 1157 and 2621 total copies per reaction (sum of the positive droplets for methylated and unmethylated target) are observed for the cg07164631 and cg2524961 ddPCR assays, respectively. For the second series of methylation mixtures, a mean of 222 and 537 total copies per reaction are observed for the cg07164631 and cg2524961 ddPCR assays. In addition, dilution series with varying degrees of converted and non-converted DNA (EpiTect PCR Control DNA; Qiagen) were prepared. EpiTect Control DNAs (completely methylated or completely unmethylated bisulfite converted DNAs and untreated, unmethylated genomic DNA) are added to each PCR reaction.

Primer/probe design

Three assays were designed: two for evaluation of the methylation status and the third assay for evaluation of the bisulfite
conversion efficiency. Sequences of the primers and probes of the above assays are provided in Table 1.

To evaluate the methylation status of two CpG sites (cg07164631 and cg25249613), primer pairs targeting only converted cytosine residues (optimal number ≥4) were designed. This allowed for the amplification of only the bisulfite-converted DNA sequence, but not of the non-converted DNA template. To enforce that the primers efficiencies are methylation-independent, no CpG sites were included in the primer pairs. Two probes, one labeled with FAM and one with HEX, were designed to bind converted and non-converted DNA sequence, respectively. Two reactions with 8000 droplets or lower were excluded from further processing. Thermal cycling conditions were 95°C for 10 min, followed by 40 cycles at 94°C for 30 secs, 56°C for 60/90 secs with a final 10 min at 98°C (Table 1). After PCR amplification using the T100™ Thermal Cycler Bio-Rad, the positive droplets (HEX/FAM) were counted with QX200 droplet reader (Bio-Rad). The total number of copies/reaction was calculated using the Bio-Rad CFX manager software and data was analyzed using the QuantaSoft 1.7.4 software (version 3.1) using the Single Threshold mode. The percentage methylation was calculated as follows: % methylation = 100/(1 + 2^(ΔCtFAM-ΔCtHEX)). Reactions were performed in triplicate and the mean with 95% CI was calculated.

**Genome-wide methylation screening**

DNA was extracted from 2 slices (10 μm) from the FFPE samples (Qiagen), quantified using PicoGreen and checked for quality using the Illumina FFPE QC Kit. The DNA was then subjected to a Bisulfite conversion (EZ DNA Methylation Kit, Zymo Research). Next, 250 ng DNA was subjected to Illumina’s FFPE restoration protocol, per manufacturer’s protocol. Samples were run on Infinium HumanMethylation450 BeadChip (450K). Several selective filters (detection filter, bead-count filter, SNP-filter, cross-reactivity filter, XY-filter) were applied to the probe sets, as described in Nordlund et al [27], and Dedeua- waerden et al [28]. Within-sample β-values were normalized for probe-design using an adjusted version of the PBC algorithm [28] and corrected for batch-effects using surrogate variable analysis [29].

**Disclosure of potential conflicts of interest**

The authors report no conflict of interest.

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**ddPCR**

The ddPCR reaction mixture consisted of the 2 × ddPCR Supermix for probes (no dUTP) (Bio-Rad), forward and reverse primer (900 nM), probe (250 nM) and 2 to 5 μl bisulfite-converted DNA in a total reaction volume of 25 μl. Methylated and non-methylated controls and a no-template control (water) were added in every experiment. A volume of 20 μl of PCR mixture was added to the middle row of wells of the droplet generation cartridge (DG8) followed by addition of 70 μl of droplet generation oil to the left row. After completion of the droplet generation (average 15,000 droplets/well) in the QX200 droplet generator, the contents were transferred to a PCR plate. Reactions with 8000 droplets or lower were excluded from further processing. Thermal cycling conditions were 95°C for 10 min, followed by 45 cycles of 94°C for 30 secs, 60°C for 60/90 secs with a final 10 min at 98°C (Table 1). After PCR amplification using the T100™ Thermal Cycler Bio-Rad, the positive droplets (HEX/FAM) were counted with QX200 droplet reader (Bio-Rad) and data was analyzed using the QuantaSoft 1.7.4 software (Bio-Rad). The total number of copies/reaction was calculated and results were reported as the percentage methylation, i.e., the fractional abundance of methylated DNA alleles. Reactions on the control DNA dilutions were performed in triplicate and the mean with 95% CI was calculated. Clinical samples were analyzed in single reactions.

**qPCR**

The qPCR reaction mixture consists of 2x Epitect Methylight Master Mix (Qiagen), forward and reverse primer (400 nM) and probe (200 nM) and 2 μl bisulfite-converted DNA. Water is added to a total reaction volume of 20 μl. Thermal cycling conditions were 95°C for 5 min, followed by 45 cycles of 95°C for 15 sec, 60°C for 60 sec; The Bio-Rad CFX-96 was used. Ct values were calculated by the Bio-Rad CFX manager software (version 3.1) using the Single Threshold mode. The percentage methylation was calculated as follows: % methylation = 100/(1 + 2^(ΔCtFAM-ΔCtHEX)). Reactions were performed in triplicate and the mean with 95% CI was calculated.
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