Comparison of microfluidic digital PCR and conventional quantitative PCR for measuring copy number variation

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Received January 5, 2011; Revised and Accepted February 14, 2012

ABSTRACT

One of the benefits of Digital PCR (dPCR) is the potential for unparalleled precision enabling smaller fold change measurements. An example of an assessment that could benefit from such improved precision is the measurement of tumour-associated copy number variation (CNV) in the cell free DNA (cfDNA) fraction of patient blood plasma. To investigate the potential precision of dPCR and compare it with the established technique of quantitative PCR (qPCR), we used breast cancer cell lines to investigate HER2 gene amplification and modelled a range of different CNVs. We showed that, with equal experimental replication, dPCR could measure a smaller CNV than qPCR. As dPCR precision is directly dependent upon both the number of replicate measurements and the template concentration, we also developed a method to assist the design of dPCR experiments for measuring CNV. Using an existing model (based on Poisson and binomial distributions) to derive an expression for the variance inherent in dPCR, we produced a power calculation to define the experimental size required to reliably detect a given fold change at a given template concentration. This work will facilitate any future translation of dPCR to key diagnostic applications, such as cancer diagnostics and analysis of cfDNA.

INTRODUCTION

A key measurement challenge in diagnostic research involves identifying small changes in gene dosage or nucleic acid sequence that are commonly associated with genetic diseases. Copy number variations (CNVs) are changes in the genomic DNA leading to an abnormal number of copies of a DNA sequence (usually two copies per diploid genome). CNVs are caused by deletions, duplications or structural rearrangements of the genome. CNVs are involved in a large number of complex human diseases such as Down’s syndrome (trisomy 21) and many cancers, for example HER2 gene amplification in breast cancer (BC) (1–5). CNV measurements are used for routine screening in clinical diagnostics and their analysis can assist in subsequent prognostic monitoring (6,7). Clinical diagnostic methods currently include fluorescence in situ hybridization (FISH), comparative genome hybridisation (CGH), single nucleotide polymorphism (SNP) arrays, deep sequencing and real-time quantitative PCR (qPCR) (8–10).

Quantitative PCR (qPCR) is currently the most sensitive approach able to resolve ~1.5-fold changes (11,12). The discovery of cell free DNA (cfDNA) in blood plasma has provided a simple source of genetic material for pre-natal and tumour diagnosis (13–18) that could potentially enable routine minimally invasive sampling for subsequent CNV analysis. qPCR has recently identify amplified HER2 molecules in breast cancer patients with good correlation between the levels of amplification detected in the primary tumour and cfDNA (19). However, as only a proportion of the cfDNA is derived from the embryo or tumour, identification of an associated CNV is more challenging as the target DNA is effectively ‘diluted’ in a background of normal DNA. Consequently, a tumour-associated 5-fold increase in CNV becomes a 1.2-fold increase if only 5% of the cfDNA sample is derived from the tumour; this magnitude of CNV would be undetectable by current approaches.

One method that has shown promise for improving the limit of detection for nucleic acid quantification is digital PCR (dPCR) with a number of reports highlighting the
superior accuracy of dPCR for CNV analysis (20,21). dPCR has been reported to detect a 1.25-fold difference in copy number (21); however, to our knowledge, no direct comparison between qPCR and dPCR has been performed to ascertain if the latter is more sensitive. Another aspect that is not extensively addressed in the literature is that the ability of dPCR to measure small CNVs is directly dependent upon template concentration. This is a crucial consideration when considering cDNA as a template as its concentration can range considerably; from 2 to 30 ng/ml plasma in healthy individuals to 180–600 ng/ml plasma both during pregnancy and in cancer patients (13,16,18,22–24).

This study aimed to investigate this further using an in vitro BC gene amplification model to identify when a CNV is too small to be measured by qPCR and dPCR. Subsequently, we developed a model, based on the Poisson and binomial distributions, to determine the variance inherent in dPCR, and used this to perform power calculations which demonstrate the effects of the DNA-template concentration on the sensitivity of CNV measurements.

REAL-TIME QUANTITATIVE PCR

Real-time quantitative PCR was performed in accordance with the MIQE guidelines (Supplementary Table S1 and Supplementary Figure S1) (25). Ten microlitre reactions contained 1× TaqMan® gene expression mastermix (ABI 4369016), 1× gene-specific assay and 2 μl target DNA. qPCR was performed using the Prism 7900HT Real Time PCR system (ABI). Thermocycling conditions were 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 60 s.

Quantification was performed with the standard curve method using five standard dilutions, in triplicate, of normal female gDNA ranging from 16.50 to 0.33 ng (5000 to 100 haploid genome copies) per reaction. Due to the differences in molecular weight between the BC cell line gDNA, triplicate reactions were performed on a range of gDNA concentrations (16.50 to 0.33 ng per reaction) for HER2 and RNase P assays. This allowed identification of the gDNA concentration that fell within the range of the qPCR standard curves for both HER2 and RNase P assays. Subsequently, eight HER2 and eight RNase P reactions (16 reactions in total) were performed on the optimum weight of gDNA for each cell line (normal female: 0.71 ng, MCF-7: 0.60 ng, SK-BR-3: 0.94 ng and T-47D: 1.45 ng). The SDS software v2.4 (ABI) was used to calculate the quantification cycle (Cq) value, that is defined as the number of cycles at which the fluorescence signal is significant above the threshold, which was converted to copy number using the relevant standard curve. Replicate HER2:RNase P ratios were calculated by randomly pairing HER2 and RNase P copy numbers and calculating the 95% confidence intervals (CI) from the standard error of the mean and the two-tailed Student’s t-test. No template control (NTC) reactions were performed using water with no template; in all cases, no amplification occurred (Supplementary Figure S1c and S1d).

DIGITAL PCR

About 4.5 μl reactions containing 1× TaqMan® gene expression mastermix, 2× GE sample loading reagent (Fluidigm 85000746), 1× gene-specific assay and 1.35 μl target gDNA was pipetted into each loading inlet of a 48.770 digital PCR array (Fluidigm). The BioMark IFC controller MX (Fluidigm, San Francisco, CA) was used to uniformly partition the reaction from the loading inlet into the 770 × 0.84 nl chambers. dPCR was performed using the BioMark System for Genetic Analysis (Fluidigm). Thermocycling conditions were set as for qPCR. For BC cell line gDNA analysis, reactions were performed in quadruplicate panels for HER2 and RNase P assays (eight panels in total) on 0.6 ng BC cell line gDNA, estimated using A260 measurements. The Digital PCR Analysis software (Fluidigm) was used to set the Cq threshold and range (Supplementary Figure S2), and to count the number of positive chambers (H) out of the total number chambers (C) from which the Poisson distribution was used to estimate the average number of molecules per chamber (λ) so that λ = −ln(1 − H/C) (26). HER2:RNase P ratio (λ/λ0) and 95% CI were calculated as described in this publication (Table 1). NTC reactions were performed using water with no template; in all cases,
Table 1. Summary of equations derived in this study

<table>
<thead>
<tr>
<th>Description</th>
<th>Symbol</th>
<th>Equation</th>
<th>Equation in MS Excel</th>
<th>Worked example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of chambers analysed</td>
<td>C</td>
<td></td>
<td></td>
<td>770</td>
</tr>
<tr>
<td>Number of positive chambers for reference</td>
<td>H_r</td>
<td>-ln (1 - H_r/C)</td>
<td>-ln(1 - (H_r/C))</td>
<td>0.201</td>
</tr>
<tr>
<td>Number of positive chambers for target</td>
<td>H_t</td>
<td>-ln (1 - H_t/C)</td>
<td>-ln(1 - (H_t/C))</td>
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</tr>
<tr>
<td>Number of reference molecules per chamber</td>
<td>λ_r</td>
<td>ln (λ_r/λ_a)</td>
<td>ln (λ_r/λ_a)</td>
<td>0.283</td>
</tr>
<tr>
<td>Number of target molecules per chamber</td>
<td>λ_t</td>
<td>ln (λ_t/λ_a)</td>
<td>ln (λ_t/λ_a)</td>
<td>0.283</td>
</tr>
<tr>
<td>Log ratio estimate</td>
<td>R</td>
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<td>1 - e^{-λ_t/λ_a}</td>
<td>0.002</td>
</tr>
<tr>
<td>Variance for Log ratio estimate</td>
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<td>1 - e^{-λ_t/λ_a}</td>
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</tr>
<tr>
<td>Standard deviation for Log ratio estimate</td>
<td>σ_R</td>
<td>SQRT(σ_R^2)</td>
<td>SQRT(σ_R^2)</td>
<td>0.361</td>
</tr>
</tbody>
</table>

For equations in MS Excel, an ‘equals’ sign must be inserted before the formula and symbols should be replaced with the experimental values. A worked example is provided, with the values used to generate the variance model and to ensure correct equations are transferred.

Establishment and analysis of copy number variation ratios

The in vitro gene amplification model was established by spiking T-47D gDNA into normal female gDNA at various percentages to generate a theoretical range of HER2:RNase P ratios between 1.00 and 2.00 based on the dPCR analysis (Supplementary Table S2). All ratios were diluted to give approximately 720 RNase P copies/μl. For qPCR, four replicates were performed per ratio for both HER2 and RNase P assays in two independent experiments (16 reactions in total). All CNV ratios were calculated by conversion of Cq values to copy numbers of HER2 and RNase P using a standard curve generated from five dilutions, in triplicate, of normal female gDNA ranging from 1,000 to 20 haploid genome copies/μl (Supplementary Figure S1c and S1d). For dPCR, four panels were performed per ratio for both HER2 and RNase P assays in two independent experiments (16 panels in total) to give approximately 155 RNase P copies per panel (λ_t = 0.2) where one panel on a 48.770 dPCR array contains 0.195 μl of template. CNV ratios were calculated using the equations described in this paper (Table 1).

To investigate the number of dPCR panels (containing 770 chambers each) needed to detect small ratio differences, a pseudo-random number generator (the ‘Math.random’ method in JavaScript) was used to assign a number to each HER2 and RNase P panel. In each case, the required numbers of panels were selected from the eight panel data set.

Statistical analysis and power calculations

Statistical analysis was performed using the MS Office Excel software (2003). Statistical comparisons to establish CNV limit of measurement were performed using the one-way analysis of variance test to compare the HER2:RNase P counts (dPCR) or copy number (qPCR) between the samples (in vitro gene amplification model ratio). Two-way analysis of variance was used to test for differences in the mean copy number between the respective replicate qPCR experiments. The two-tailed Student’s t-test was used to analyse the difference in HER2 counts or copy number between the sample and the calibrator (normal female gDNA). Power and associated calculations reported in this paper were carried out using the R statistical programming language (version 2.13, http://www.r-project.org). All scripts were written and run on a standard desktop personal computer (Optiplex, Dell Corporation). Further details of the methods and theory are given in the Statistical Supplementary Information.

RESULTS

CVN measurement by dPCR and comparison with qPCR

In order to investigate the accuracy of dPCR for CNV measurement, we used three BC cell lines with different HER2 gene copy number as a model of gene amplification. The RNase P assay was used as the reference gene for the diploid control. Assay optimization was performed using qPCR for HER2 and RNase P assays (Supplementary Figure S1). For dPCR, absolute quantification of HER2 and RNase P molecules were calculated from the number of positive counts per panel based on the Poisson distribution for the number of molecules in each chamber (Figure 1a).

dPCR analysis of normal female gDNA had a HER2:RNase P ratio of 1.03 and was not significantly different from qPCR analysis that had a HER2:RNase P ratio of 1.00 (P = 0.39; Figure 1b). Analysis of MCF-7 gDNA, which has a single HER2 allele deletion, gave a ratio of 0.44 by both dPCR and qPCR. T-47D gDNA, which has low-HER2 copy gain had a HER2:RNase P ratio of 2.00 and 1.96 for dPCR and qPCR, respectively. There was no significant difference in HER2:RNase P ratios between the two techniques when measuring

no positive chambers were observed (Supplementary Figure S2).

RNase P counts (dPCR) or copy number (qPCR) between the samples (in vitro gene amplification model ratio). Two-way analysis of variance was used to test for differences in the mean copy number between the respective replicate qPCR experiments. The two-tailed Student’s t-test was used to analyse the difference in HER2 counts or copy number between the sample and the calibrator (normal female gDNA). Power and associated calculations reported in this paper were carried out using the R statistical programming language (version 2.13, http://www.r-project.org). All scripts were written and run on a standard desktop personal computer (Optiplex, Dell Corporation). Further details of the methods and theory are given in the Statistical Supplementary Information.

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Limit of detection for analysing copy number variations

To determine the limit of detection for analysis of CNVs by dPCR, an in vitro gene-amplification model was used, whereby T-47D gDNA was spiked into normal female gDNA to generate a theoretical range of HER2:RNase P ratios between 1.00 and 2.00 at low copy number (2.1 ng/μl) for analysis using dPCR and qPCR (Supplementary Table S2). Using dPCR, a ratio of 1.17 or more was significantly different from the experimentally derived normal female gDNA ratio of 1.03 (P < 0.0003) when eight panels where used (Figure 2a). There was good linear correlation between the expected and the observed ratios when measuring a CNV of ≥1.17 (R² = 0.9974) and this linear correlation was maintained when the line was extrapolated through the observed ratio for normal female gDNA (Figure 2a; dashed line). Furthermore, the slope and intercept of the linear correlation were measured as 1.05 and 0.05, respectively, demonstrating the accuracy in the measured ratios, and that no bias was introduced. An expected ratio of 1.12 was not significantly different from normal female gDNA when using eight panels (P = 0.67; Figure 2a). In all cases, the RNase P counts observed for each measurement were not significantly different from one another (P > 0.75). Analysing the in vitro gene-amplification model with qPCR demonstrated that ratios of 1.27 or more were significantly different from
normal female gDNA \((P < 0.0005)\) and maintained a linear correlation \((R^2 = 0.99;\) Figure 2b; dashed line). As was observed with the dPCR analysis, the observed ratios were accurate with no introduced bias as shown by the slope \((1.01)\) and intercept \((-0.0001)\) of the linear correlation. Ratios of \(\leq 1.22\) did not differ significantly from female gDNA \((P > 0.05;\) Figure 2b). As with dPCR, in all cases, the \(RNase\ P\) counts observed for each measurement were not significantly different from one another \((P \geq 0.27)\). Furthermore, no statistically significant inter-run variability was observed for either the \(RNase\ P\) or the \(HER2\) assays \((P \geq 0.99\) and \(P \geq 0.15,\) respectively).

**Determination of the number of dPCR panels needed to confidently detect small changes in copy number variation**

From our data, when the CNV to be measured is >1.5-fold, then four dPCR panels were sufficient while CNVs < 1.5-fold could be detected with up to eight dPCR panels (Figures 1b and 2a). As CNVs of <1.17 were not measurable using eight dPCR panels, it would be useful to be able to predict how many dPCR panels would be needed to detect such small CNVs. For such a prediction, we would need to assess discriminating power, and therefore, the true copy number difference at which we would reliably judge two materials to be different. This was achieved with a statistical power calculation, which required a test statistic and associated distribution.

The domain of interest in this study involved the reference \(RNase\ P\) assay adjusted to give an observed number of positive chambers in the region of 100–200 per panel \((2.1\text{ ng/\mu l})\) mimic the low concentration observed in cfDNA samples \((18)\). Based on the Poisson distribution, this can be used to estimate the number of molecules per chamber \((\lambda)\), and in this case, \(\lambda\) was approximately 0.2 in a 48,770 dPCR array (Table 1). At this concentration, the probability \(P\) that a chamber will give a positive signal is \(1 - e^{-\lambda}\) and is the same for every chamber \((26)\). The CNV is the ratio of \(\lambda\) estimates for the two groups being compared and as such, it is highly non-Normal. By taking the logarithm of the ratio for these two groups \((R)\), the data is transformed to produce a variable whose distribution is very close to Normal (Supplementary Figure S3a) and is given by the equation:

\[
R = \ln \frac{\lambda_t}{\lambda_e}
\]

where \(\lambda_t\) and \(\lambda_e\) correspond to the \(\lambda\) values of the target and reference genes, respectively, and \(\lambda = -\ln \left(1 - \frac{H}{C}\right)\) [Table 1; \(26\)]. Consequently, power calculations can be based on a \(t\)-test for a difference in the logarithm of the observed ratio of target to reference gene from zero. To calculate power, we first need an expression for the variance \((\sigma^2_R)\) of the log ratio \(R\). This was derived using first-order error propagation (see ‘Statistical Supplementary Information’) and is given here as:

\[
\sigma^2_R \approx \frac{1 - e^{-\lambda_t}}{C\lambda_t e^{-\lambda_t}} + \frac{1 - e^{-\lambda_e}}{C\lambda_e e^{-\lambda_e}}
\]

Comparison of the theoretical standard deviation of \(R\) [estimated from Equation (2) as \(\sqrt{\sigma^2_R}\)] with the observed standard deviation of the experimental data (Figure 2a) demonstrated good concordance (Supplementary Figure S3b). With a valid estimate of the variance \(\sigma^2_R\) in the log ratio, the upper and lower 95% CI can be calculated (Table 1). Power calculations can also be derived to determine the number of chambers required for an experiment capable of detecting a log ratio \(R\) with a defined test power of \(1 - \beta\) and at the \(1 - \alpha\) confidence level, where \(\alpha\) and \(\beta\) are the false-positive and false-negative rates, respectively. Our experiment was one in which a positive \(R\) was the expected result (an increase in copy number, so \(\lambda_t > \lambda_e\)), so a one-tailed test was appropriate. Statistical significance was declared at a \(P\)-value of \(\alpha\) or lower \((<0.05)\) and the test power was equal to 95% \((\beta = 0.05;\) Supplementary Figure S4). Power calculations involving a test statistic which has a Normal distribution can only be carried out numerically, as there is no analytical solution and details of the method are given in the ‘Statistical Supplementary Information’.

Using the power curve, where \(\lambda_t = 0.2\), we found that with 95% power, a fold change of 1.2 was easily detectable with five 770-chamber panels per gene assay, while ratios of 1.1 and below required greatly increased numbers of panels (>15 panels; Figure 3a). Using the curve to compare the data from the \(HER2\) \(in vitro\) gene amplification model with the number of panels required demonstrates that a \(\geq 1.17\) ratio can be measured with eight or fewer panels (Figure 3a); this is shown experimentally (Figure 2a). However, from the curve, a ratio of 1.12 was predicted to need more than 10 panels (Figure 3a), which is supported by our inability to measure with confidence a 1.12 ratio with only eight panels (Figure 2a). Our power curve predicts that when \(\lambda_t\) is 0.2, the smallest CNV ratio that could be measured using eight dPCR panels is approximately 1.15 (Figure 3a). This lies between our two experimental data points and therefore confirms the fitness of our model for this template concentration.

To further test the power calculations, each \(HER2\) dPCR panel was randomly paired with an \(RNase\ P\) panel and the desired number of paired panels \((1–8)\) were selected and used to calculate their CNV ratio and associated 95% CI (Figure 3b). Our power curve suggests that approximately three panels are needed to detect a ratio of 1.27 when \(\lambda_t = 0.2\) (Figure 3a) which was experimentally measurable with four or more panels (Figure 3b). Analysis of a fold difference of 1.17 predicted that approximately six panels are needed (Figure 3a), which is concordant with our experimental data (Figure 3b).

Power calculations were also performed for qPCR using the Student’s \(t\)-statistic (see ‘Supplementary Statistical Information’ for details) and the resulting power curves demonstrated that ratios greater than 1.25 can be measured with 95% power and eight replicate qPCR wells (Supplementary Figure S7). This compared well with the experimental data generated from the \(in vitro\) gene amplification model where a ratio of 1.27 was measured with eight qPCR wells but small ratios were not (Figure 2b). From the power curve, it was shown that using gDNA at the experimental defined
concentration, an excess of 20 qPCR wells would be needed to measure ratios of 1.17 or fewer (Supplementary Figure S7).

**DISCUSSION**

We have investigated the measurement capabilities of dPCR when investigating changes in CNV, described the limit of detection for a given experiment and demonstrate that dPCR exhibits greater sensitivity than qPCR when investigating subtle fold-differences. The potential of cfDNA as a source of template in minimally invasive diagnostics is becoming increasingly apparent. The measurable resolution of qPCR in our gene-amplification model for cfDNA, which we describe here as a ratio of 1.27, offers a theoretical diagnostic potential, however, the reality is that smaller CNVs, where the tumour contribution to the cfDNA can be as little as 5% of the total cfDNA (16,17) are still technically out of reach using qPCR. Furthermore, one of the major differences between qPCR and dPCR is that technical variability of qPCR can be high between and within laboratories (27). dPCR is notably less variable between experiments (28,29), which offers the possibility of reproducibly more accurate results. In this study, a CNV ratio of 1.17 was significantly detected using eight panels and therefore, the ability of dPCR to detect incrementally smaller fold differences than qPCR demonstrates the potential of this method for future CNV clinical diagnostics.

Recently, an error model relating to the 95% theoretical CI for copy number versus number of chambers was published (21). According to this model, approximately 1,200 chambers (equates to 1.5 panels) would be needed to detect a CNV ratio of 1.25. However, when the authors performed this in a real experiment, they were unable to realise this predicted precision, requiring twice the number of chambers/panels to measure the targeted difference (21). The authors explain that this was directly due to the fact that, for their model, they fixed the $\lambda_t$ at a value of 0.6, whereas the real experiment (where $\lambda_t$ values were 0.18 and 0.37 for their two samples) was below this value and so did not have sufficient power to resolve the given difference with the predicted number of chambers.

When we used this model to estimate the number of chambers needed for our experiments, we found that this was also underestimated and due to the smaller $\lambda_t$, which in the case of our experiment was about 0.2. Weaver and co-workers (21) used the point (in observed number of panels) at which calculated 95% CIs just overlap, assuming a Poisson distribution for individual counts. The point of theoretical CI overlap is an indicator of discriminating power, in that it increases as precision degrades and allows comparison between samples. However, basic statistical theory shows that a significant difference at the 95% confidence level would usually show substantially overlapping CIs for the two independent observations. In this case, statistical significance focuses on the error rate under the null hypothesis; the probability of wrongly declaring a result significant, or the false-positive rate. In contrast, we were interested in assessing discriminating power and therefore, the true copy number difference at which will we reliably judge two materials to be different, that is the false-negative rate. Therefore, we generated a power calculation based on the variance and false-negative rate of a given measurement of copy number ratio.

Our power calculation method takes the DNA-template concentration into consideration using the corresponding $\lambda_t$ and estimating the number of panels required. Validation of the theory is demonstrated with our experimental data (Figure 3b). Based on this, further theoretical power curves for a range of $\lambda_t$ values (0.1–0.8) were derived and are given in Supplementary Figure S5 for researchers to use for future experimental design. From these power curves, we predict that to investigate a 1.25-fold difference, where $\lambda_t$ is 0.2 or 0.4 (approximate values for $\lambda_t$ of 0.18 and 0.37) with 95% power requires a four- or two-panel experiment,
respectively (Supplementary Figure S5), which was consistent with Weaver and co-workers observations. The power calculations described in this publication are based on a one-tailed test as we were interested detecting HER2 amplification (where \( \lambda_i > \lambda_t \)) which has clinical and prognostic relevance (30,31). Further applications of this model would include detection of trisomy, e.g. in Down’s syndrome (32). However, a modification of this model to include detection of both amplification and deletion of gene copies would have a wider scope, e.g. in characterization of in vitro culture cells over time (33), detection of polyploidy in plants (34) or as an alternative to CGH analysis or FISH for cancer diagnostics. Therefore, we have expanded our model to incorporate a two-tail test (Supplementary Figure S6).

We also generated power curves for qPCR based on a one- or two-tailed test (Supplementary Figure S7). These curves demonstrate that while dPCR appears to detect only incrementally smaller fold differences than qPCR using a one-tailed test and eight replicate measurements (1.17 versus 1.27 in this study), an additional 12 qPCR wells per assay or more would be needed before qPCR could potentially measure a similarly small ratio to dPCR. This is dependent upon the standard deviation of the qPCR (in this case, was approximately 10% for both assays across all experiments) as the number of replicates needed will increase if precision is reduced. The binary nature of dPCR means that the precision is more independent of variation in assay amplification, making it easier to optimize and standardize between laboratories.

As conventional qPCR is currently approximately one 20th of the expense of an equivalent dPCR analysis, it does still offer a practical alternative method to measure smaller CNVs. However, the increased template required to perform the larger number of replicate measurements could be a disadvantage where sample is limiting. qPCR has the benefit of being able to measure lower plasma template concentration than the dPCR method used here as its larger reaction volume facilitates the addition of more template. qPCR is also scalable offering the option of adding even more template, where sample permits, by further increasing the volume of the reaction. This is not currently possible with the dPCR approach used here; however, with the development of higher volume and throughput dPCR methods this option will also be possible (35–37) with comparable accuracy (38).

Improvements to qPCR using similar technologies to those advancing dPCR will increase the throughput of qPCR, as has already been described (21), although this is also at the cost of smaller input volume which often requires pre-amplification (39); this is not routinely needed for dPCR. It should also be noted that for qPCR (conventional or high throughput) to measure such small fold changes, it is not only important that precise well-optimized assays are used, but it is also essential that some estimation of efficiency is made (25). This can be done by standard curves to estimate copy number, as described here, or by performing efficiency corrected \( \Delta \Delta Cq \) (40). However, this fact further complicates using qPCR for small fold change measurement.

Our model also raises the question of why variable \( \lambda_i \) should need to be considered at all when performing dPCR. The alternative would be to ensure the optimum concentration in the first place. However, this is only possible when the sample DNA is at or higher than this value, and if the concentration is lower than the optimum \( \lambda_i \) then more chambers would be needed, and therefore, our model would provide an idea of how many. This issue is applicable when using cfDNA as a template, which is by nature low in concentration; around 2–30 ng/ml blood plasma in normal healthy humans (13,16–18,22,23), which corresponds to approximately 600–9900 normal haploid genomes per ml. Measuring HER2 status using cfDNA offers considerable potential as a diagnostic and prognostic test (19,30,31).

Our power calculation provides a good foundation from which to design subsequent experiments; in our assessment, we have used a high-quality DNA template in our proof-of-principle experiments to model the cfDNA. Additional factors that will need to be considered, building on the findings of this study, include template integrity, PCR-assay efficiency and the impact of matrix effects as integral parts of subsequent translational research, if this approach is to be used in the context of HER2 and other clinical measurements. Recently, power calculations based on a multivolume dPCR assays were generated and demonstrated that different reaction volumes influence the dynamic range and precision of dPCR, could minimize the total number of dPCR reactions needed and separated the upper and lower limits of quantification. This would allow samples of differing concentration to be analyzed in parallel without compromising one sample over the other (37).

An additional function of this study was to directly compare microfluidic dPCR with conventional qPCR for CNV using the same gDNA template and reaction assays. Both methods gave similar HER2 CNV results using BC cell line gDNA with varying numbers of HER2 gene copies per diploid genome. Two BC cell lines were concurrent with the published data; MCF-7 gDNA, that has monosomy for chromosome 17 assigned a HER2:RNase P ratio of < 0.5 (one copy per diploid genome) (41,42) and T-47D gDNA gave a HER2:RNase P ratio of < 2 (3–4 copies per diploid genome) (43–46). Analysis of the SK-BR-3 gDNA identified high levels of HER2 amplification that corresponds with the published copy number range (14–24 copies per diploid genome) (41,43–45,47). However, there was a discrepancy between the HER2:RNase P ratio obtained by dPCR (7.15) and qPCR (9.43).

This difference could be attributed to the large number of HER2 gene amplifications occurring on the same molecule (concatamers of HER2 molecules inserted at the same point in the genome) that are observed in this cell line using FISH (48,49) but do not occur in the MCF-7 or T-47D cell lines. Unlike qPCR, dPCR may be unable to accurately quantify this type of gene amplification as linked genes cannot be separated into individual chambers. Qin and co-workers (20) have suggested that this problem can be overcome by a pre-amplification step using gene-specific primers to separate the linked
copies before performing dPCR. However, such a step requires careful validation because of the potential bias that can be introduced that may outweigh the necessary precision for detecting small CNVs (29).

In conclusion, our data suggests as microfluidic dPCR becomes more established, it will offer a new level of precision and the clinical benefits of measuring smaller CNVs in more challenging samples like cfDNA, will become possible. However, the pre-clinical and translational research necessary for this to be realized will need to consider the issues explored by this study. The model we describe here both provides a mechanism to facilitate this research and highlights the issues around ensuring the template concentration is included as a central consideration when preparing CNV studies using dPCR. This will better enable dPCR experiments to be designed, increasing the impact of future research.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–2, Supplementary Figures 1–7 and Supplementary Statistical Information.

ACKNOWLEDGEMENTS

The authors would like to acknowledge Dr Malcolm Burns and Dr Alison Devonshire for critical review of the manuscript.

FUNDING


Conflict of interest statement. None declared.

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