Calibration-free assays on standard real-time PCR devices

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Quantitative Polymerase Chain Reaction (qPCR) is one of central techniques in molecular biology and important tool in medical diagnostics. While being a golden standard qPCR techniques depend on reference measurements and are susceptible to large errors caused by even small changes of reaction efficiency or conditions that are typically not marked by increased precision. Digital PCR (dPCR) technologies should alleviate the need for calibration by providing absolute quantitation using binary (yes/no) signals from partitions provided that the basic assumption of amplification a single target molecule into a positive signal is met. Still, the access to digital techniques is limited because they require new instruments. We show an analog-digital method that can be executed on standard (real-time) qPCR devices. It benefits from real-time readout, providing calibration-free assessment. The method combines advantages of qPCR and dPCR and bypasses their drawbacks. The protocols provide for small simplified partitioning that can be fitted within standard well plate format. We demonstrate that with the use of synergistic assay design standard qPCR devices are capable of absolute quantitation when normal qPCR protocols fail to provide accurate estimates. We list practical recipes how to design assays for required parameters, and how to analyze signals to estimate concentration.

Here we demonstrate a new type of assay that synergistically combines the advantages of digital and analogue quantitative PCR. The combination of the digital and analogue information provides for absolute quantitation with adjustable resolution. We also describe the procedure and derive a prescription for designing synergistic digital-analogue PCR assays and for analyzing the results. The assays that we describe in this report can be used to analyze difficult and exotic samples on standard qPCR (real-time) devices and to design new digital-analogue assays that maximize information gain from the partitions and thus reduce to minimum their number required to reach a particular level of precision in a given dynamic range of concentrations.

The analogue qPCR (in the format of Real-Time PCR1–4) was introduced in 1991 by Holland et al.5 (Fig. 1A). The technique traces the increase of intensity of fluorescence from a stained DNA product during PCR. Given that in each cycle the number of copies of the PCR product (the amplicons) increases q-folds, the concentration \( C_p \) of the product increases geometrically \( C_p \propto q^n \) in the cycle number n. If the product is fluorescent, one can assess the initial concentration \( C \) of the target strain by comparing the number of cycles needed to reach a threshold intensity of light between the sample and calibrated references.

qPCR can be used to determine the concentration of target DNA over a wide range of initial concentrations of the target. The fact that the estimate relies on comparison to a reference compromises the accuracy and precision of the measurement by variation in multiple experimental factors, such as quality of the reference samples, the type and quality of the buffer, the activity of the polymerase, the amount and specificity of primers, and the reproducibility of thermal cycling and/or of detection in the apparatus.

In spite of these limitations, qPCR remains the ‘golden standard’. The advantages include simple liquid handling protocols that do not require extensive partitioning of the sample, and straightforward mathematical procedures for obtaining the final result from the measurements on the sample and from the calibration curve.

Digital PCR

The idea of digital protocols root back to 1915 when McCrady6 introduced the limiting-dilution assay. He proposed the mathematical model—the most probable number method—for quantization of bacterial cells. Application of the digital protocols in quantitative assessment based on PCR was first proposed in 1992 by Sykes et al.4. The design proposed comprised 60 compartments grouped into six sets, each set being three-fold diluted to create a geometric sequence of the number of DNA copies from the sample. This idea was further developed...
in 1999 by Vogelstein and Kinzler. They proposed using identical partitions yielding either a positive ($s=1$) or negative ($s=0$) signal if the partition contained at least one target molecule ($m \geq 1$) or not ($m = 0$). As the assessment of the concentration by digital assays relies solely on the presence of a signal, and not on its analogue value, the resulting estimate of concentration is absolute i.e. not relying on comparison to any reference experiment (Fig. 1A). dPCR assays are typically also highly precise and sensitive \cite{3,7–10}. The digital schemes require strong amplification of the signal (i.e. number of molecules of the analyte), hence the digital scheme is applicable predominantly to qPCR\cite{7–10} and quantitative immuno-assays\cite{11,12}.

The digital assays are hoped to replace the standard analogue procedures. They provide absolute quantification without calibration of the experimental set-up. Still, it has to be ensured that no false positive or false negative signals are present due to the contamination of the sample or the limited amplification efficiency and specificity. They may also benefit from simplified execution of the amplification and detection protocols for end-point measurements (which may be simpler compared to real-time tracking of the signal). Yet the spread of the digital techniques in research and in diagnostics is slow due to the cost and complexity of the devices and protocols, as they require partitioning of the sample into large (or sometimes astronomically large) number of test volumes (typically thousands to millions partitions)\cite{13–15}. Furthermore, the precision and the dynamic range of a classic digital assay comprising identical partitions of the sample cannot be tuned independently. It is not, for example, possible to obtain high precision (low standard deviation of the estimate of concentration) in a narrow range of concentrations while using a small number of compartments, or vice-versa, to obtain an imprecise estimate over a wide dynamic range. We have recently shown\cite{16} a method for optimization of information gain from digital signals and for an optimum (rational) design of assays that minimizes the number of compartments required to obtain a given precision over a requested dynamic range. The protocol offers dramatic reduction (by orders of magnitude in comparison to classic digital methods) of the number of partitions. These end-point digital assays could be run on qPCR devices that use the strips (32) or (96, 192 or 384) well plates.

**The potential for synergy**

The existing methods of quantitative assessment based on PCR (dPCR and qPCR) present complementary features that have not yet been combined. The qPCR techniques offer relatively facile handling of samples. The sample is typically mixed with the PCR kit into one well. The increase of the real-time fluorescent signal is then compared with a calibration curve obtained from a reference sample. The method offers high information gain via analogue resolution of the signal yet relies heavily on calibration - an effect that may introduce significant errors in accuracy of the result. The digital assays alleviate the need for calibration, yet due to the intrinsically low (binary) information content of the signal from any single partition, the classic digital assays that use identical compartments require large numbers of them.

While the number of required partitions of the sample in a digital assay can be minimized\cite{7–10,16} while using only binary (yes/no) signals, here we show further reduction of the number of partitions of the sample or, alternatively, an enhancement of the precision of the assessment, obtained via a synergistic analysis of both the digital and analogue signals recorded on a qPCR instrument. The synergy combines the absolute character stemming from the digital analysis with precision and dynamic range delivered by the analogue (Real-Time) signals. The
synergistic assay combines the information from the digital and from the analogue signals to improve the precision of the estimate (Fig. 1B). Below we derive the synergistic assay step by step. Say that the sample contains the target analyte at (unknown) concentration C. A digital test, conducted on any isolated portion of the sample, provides information whether the partition, characterized by a volume \( V_d \) and a dilution factor \( d \), contained a number \( m_t \) of molecules \( C^*_m \), which depends solely on the volume and dilution of a test-volume \( V_d \), and a threshold number of molecules \( m_t \) (i.e. for \( m_t = 1 \), \( C^*_m = \log(2)/(dV_d) \)). Measurement of the real-time signal. In the qPCR, the level of signal (i.e. fluorescence) that is proportional to the number of copies of the amplicons in the compartment is measured at predefined time intervals, preferably at the end of every PCR cycle. The procedure returns an approximation (or estimation) of the real (non-integer) number of the cycle at which the signal from a given compartment crosses the pre-set threshold. Such identified threshold cycle number corresponds to a given, fixed (unknown, but same for all positive compartments) number of copies of the amplicons in that compartment. This allows using simple calculations to determine the ratios of initial numbers of molecules in these compartments as a function of their threshold cycle number \( c_t \).

**Information content of a binary signal**

The fact that the test volume (partition) contained a threshold number of molecules of analyte (or not) conveys a probabilistic information about the initial concentration \( C \). For example, under the assumption that the initial number of molecules \( m_t \) in the partition is given by a Poissonian distribution \( p(m_t) = e^{-Cd}/m_t! \), the positive signal can be translated into a density of probability that a given initial concentration \( C \) has caused the recording of a positive signal from the partition: \( \rho(C| m_t \geq m_{\tau}) = 1 - e^{-Cd}/m_t! \). Similarly, a negative signal can be translated into \( \rho(C| m_t < m_{\tau}) = e^{-Cd}/m_t! \). For example, for \( m_{\tau} = 1 \), \( \rho(C| m_t \geq m_{\tau}) = 1 - e^{-Cd}/m_t! \). The resulting sigmoidal probability functions are centered (i.e. present the value of 1/2) at a concentration \( C^* \). This concentration is characteristic of the partition only, i.e. in the example of \( m_{\tau} = 1 \), at \( C^* = \frac{\ln(2)}{V_d} \) (Fig. 2A) and does not depend on the (stochastic) value of the actual concentration of the analyte in the sample. The measure of the information gained from a single signal (Bernoulli trial) in a digital assay can be extracted with tools derived from the information theory. One such measure is the Shannon entropy \( H = \sum p_i \log(p_i) \), where \( i \) counts the two (positive/
negative) possible outcomes (Fig. 2B). In the dPCR, the efficiency of amplification of the presence of even a single-molecule of the analyte in the given partition is assumed to be 100%. Under this assumption, $H$ achieves maximum exactly at concentration $C^* = \ln(2)/s$. The information gain from the measurement of the state of a test-volume is highest at concentrations near $C^*$, for which $p(s = 1|C = C^*) = 1/2$ (Fig. 2A), and low for much smaller or much larger concentrations.

**Information content of digital assay**

In a purely digital assay (that uses only the end-point positive/negative signals) the estimate of $C$ is given by the product of all the probability densities obtained from individual compartments: $P(s_0, s_1, \ldots, s_{N-1}|C) = \prod_{i=0}^{N-1} p(s_i|C)$. The resulting distribution yields the estimate of the initial concentration of the analyte. This information can be expressed with the Bayesian formalism as the probability $p(C)$ that a given value of $C$ has caused the recorded outcome. The lack of prior knowledge of $C$ can be encoded with an initially constant density of probability $p(C) = 1/C_{\text{min}}$ of finding any particular value of $C$ between $C = 0$ and an arbitrary upper bound $C_{\text{max}}$. Then the information about $C$ gained from a single signal ($s = 1$ or $s = 0$) is

$$p(C|s = 1) = 1 - e^{-C_{\text{max}}/C_{\text{min}}} \text{ and } p(C|s = 0) = d\cdot e^{-C_{\text{max}}/C_{\text{min}}} \text{ (Fig. 2A).}$$

Since the sigmoidal contributions from the compartments $p(s_i|C)$ are centered on the concentrations ($C^*_i = \ln(2)/d\cdot v_i$) that depend solely on the properties of the compartment and do not depend on the actual concentration of molecules in the sample, many signals contribute little to the result of the assay\(^{16}\). In particular, the signals from compartments that contain either a high ($m \gg 1$) or very small expected number of molecules ($m \ll 1$), do not contribute much to the result of the assay\(^{16}\). This is because in the range of concentrations similar to the actual concentration $C$ of the analyte in the sample, these functions are closely equal to either unity or zero and are almost constant for $C^*_i \ll C$ or $C^*_i \gg C$, respectively.

**Optimized purely digital assays**

We have shown recently that it is possible to minimize the number of compartments needed to estimate the concentration of DNA with a requested precision within a requested range\(^ {16}\). In essence, we spread the values of $C$ of each of the compartments in the assay uniformly on the logarithmic scale. This procedure builds an analogue of a positional system, in which the role of digits is played by physical compartments of the examined sample. The positive (1's) and negative (0's) signals all have their well-defined positions in writing up the 'number' - i.e. the estimate of concentration of the analyte.

**Results**

**Combining digital and analogue information.**

Here we build on the digital assay and supplement it with the information from the real-time (analogue) signals. In our method we use the information on $c_{\omega}$ from the positive compartments to refine the estimate of concentration in the digital scheme. The real-time monitoring of compartments yields the number of cycle ($ct_{\omega}$) at which the intensity of fluorescence (which, in this example, is proportional to the number of DNA copies) exceeds a threshold value. Recording of the cycle numbers $ct_{\omega}$ and $ct_i$ from two separate compartments, allows us to estimate the ratio of initial numbers of molecules in them as: $m_{\omega}/m_i = q^{ct_{\omega}-ct_i}$. Quite interestingly, our method allows us to determine the amplification factor $q$ without any standardized calibration samples using only a known sequence of $d\cdot v$ and the 'analogue' values $ct_{\omega}$ [please see ESI for details].

From the set of (positive) compartments that provided the recording of $ct_{\omega}$, we select the one (from now on indexed $\omega$) that provided the largest value of the cycle number $ct_{\omega}$, or equivalently, the compartment that contained initially the smallest number of molecules. We know that the reference ($\omega$) compartment contained at least a threshold number ($m_{\omega}$) of molecules (because it did yield the positive signal). We also know the set of measurements $ct_i$ from $\omega$ and from a set of other compartments. We can thus treat $\omega$ as a reference and calculate that each compartment $i$ contained at least $m_i = m_{\omega}q^{ct_{\omega}-ct_i}$ molecules of analyte before amplification. In essence, the set of "analogue" values $ct_i$ allow use to modify (i.e. increase the information content) of the probability density function contributed by each positive compartment.

In essence, instead of using probabilities that the number of molecules in a compartment is larger than one (standard digital signal), i.e. $m_i \geq 1$ we can use $m_i \geq d\cdot v_i\cdot e^{\Delta ct_{\omega}}$, where $\Delta ct_{\omega} = ct_{\omega} - ct_i$. Therefore, the positive signal can be translated into $p(\Delta ct_{\omega}|C) = e^{-Cdv_i\cdot \sum_{j=0}^{m-1} (Cd\cdot v_i)^j}/d\cdot v_i^m$. The resulting functions $p(\Delta ct_{\omega}|C)$ qualitatively preserve sigmoidal shape; however, they tend to be steeper and approach (with increasing $m$) the shape of the Heaviside step function. Moreover, they are shifted towards the actual concentration $C$ in the sample and therefore increase the information gain, which leads to an improved precision and accuracy of the result (Fig. 3).

**The architecture of the synergistic assay.**

In order to design the assay (the number and type of partitions of the sample), similarly to the method for rational design of a purely digital assay\(^ {16}\), we start with the concept of an active stripe, i.e. a set of compartments that contribute information to the estimate of $C$. As before, we start with the observation, that in a set of non-identical compartments, each of the compartments is centered (i.e. the probability $p(s(C) = 0.5$, where the information gain is maximal) at a different value of concentration depending solely on its size and/or dilution. Therefore, only a subset of aforementioned compartments really contribute information to the estimate of the actual concentration; other are too far and therefore are always positive or always negative, bringing no useful information (probability functions shown in Fig. 3A). In other words, very small (or highly diluted) compartments are not useful for probing small concentrations (they are always negative), while big compartments are not useful for probing high concentrations (they are always positive). The subset of compartments that contribute
to the information on concentration is called the active stripe. It is necessary to determine its characteristics – the number of compartments it comprises – to design a synergistic assay and test our method.

In order to determine the number of compartments that participate in the assessment of the concentration, we set an arbitrary input concentration \( C_{\text{input}} \) and build an assay. We start with a single compartment ‘centred’ on the concentration \( C_{\text{input}} \), i.e.

\[
d_{i}v_{0} = \ln(2)/C_{\text{input}}^{0}.
\]

Then, we add symmetrically \( \Delta N \) compartments on each ‘side’, that form geometric sequence:

\[
d_{i}v_{0}x^{i}, \quad i \in (-\Delta N, \Delta N).
\]

For any sequence (i.e. for any value of the progression factor \( x \), the increase of the number \( 2\Delta N + 1 \) of partitions in the active stripe results in an improved assessment of \( C \). However, at some value of \( \Delta N \) the precision of the assessment saturates and adding more compartments does not any more effectively improve the information content.

Therefore, the standard deviation \( \sigma \) of the estimate of the concentration \( E(C) \) initially decreases with increasing \( \Delta N \) but at some point saturates at the limit value \( \lim_{\Delta N \to \infty} \sigma (x) \) [ESI]. We determined the value of \( \Delta N \) at which \( \sigma (x) \) saturates as the optimum number of compartments \( \Delta N_{x} \) in the active stripe [ESI]. If we substitute the requested precision \( \sigma_{\text{max}} \) for \( \lim_{\Delta N \to \infty} \sigma (x) \) and invert it, we find that the common ratio \( x \) of the geometric sequence of compartments can be closely approximated as a simple algebraic function of \( \sigma_{\text{max}} \):

\[
\Delta N_{x} = \left[ \frac{\ln(2)}{C_{\text{input}}} \right],
\]

\[
\alpha = \sqrt{\frac{\sigma_{\text{max}}^{2} + \beta \sigma_{\text{max}} + \gamma}{\delta \sigma_{\text{max}}}},
\]

\[
N = 2\Delta N + \lfloor \log_{x} (1/\Omega) \rfloor.
\]

The value of the common ratio \( x \) provides the sequence \( d_{i}v_{0} \) of the consecutive compartments that provide assessment with a requested precision. The value of \( \Delta N \) provides ’margins’ of the compartments outside. Knowing that the geometric progression is self similar, it is enough to span the assay to provide the requested precision within the required dynamic range of concentrations \( C \in (C^{-}, C^{+}) \) with

\[
d_{i}v_{0}x^{i} = \ln(2)/C_{\text{input}}^{i},
\]

with \( \alpha, \beta, \gamma, \delta \) and \( \varepsilon \) being positive constants [ESI].
The precision of the assessment can be further improved by increasing the number of test volumes for Real-Time assays, which is usually smaller than for synergistic assays. However, one has also need a calibration line, which is usually calculated from 15–25 experiments with standardized samples. Still, the number of test volumes for Real-Time assays is usually smaller than for synergistic assays. However, one has also need a calibration line, which is usually calculated from 15–25 experiments with standardized samples. Still, the number of test volumes for Real-Time assays is usually smaller than for synergistic assays. However, one has also need a calibration line, which is usually calculated from 15–25 experiments with standardized samples. Still, the number of test volumes for Real-Time assays is usually smaller than for synergistic assays. However, one has also need a calibration line, which is usually calculated from 15–25 experiments with standardized samples. 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diluted in three elution buffers from DNA isolation kits (AE elution buffer QIAamp DNA Mini Kit (Qiagen), MBL5 NucleoMag Blood (MACHEREY-NAGEL) and MagJET Whole Blood Genomic DNA Kit (Thermo Scientific) and water to obtain model samples with 25 000 copies of the target DNA per mL. The absolute quantification of sample DNA molecules were calculated using 4 internal kit’s calibrators (from 10 to 10.000 copies per μl) without any dilutions and modifications. The qPCR analysis produced highly scattered results with the following averages: 96 514 [1/mL] for water, 79 368 [1/mL] for the AE buffer, 29 684 [1/mL] for the MBL5 buffer and 24 967 [1/mL] for the MagJET buffer. Thus the change of the buffer in the isolation protocol may heavily influence the accuracy of the assay changing the readout of the qPCR analysis even by a factor of 400% while not signalling the systematic error in the spread of the results (i.e. in reduced precision).

The reason why we observe such a variability in the estimate of concentration provided by qPCR can be the presence of PCR inhibitor in the buffer containing DNA standard. In the experiments we used standardized samples diluted 100 times. If the sample was diluted with water (Fig. 5A, blue point) the inhibiting agent practically disappeared, and therefore higher readout was observed. The agent in question is the ethylenediaminetetraacetic acid (EDTA), present in the buffer, which neutralizes DNases and reduces DNA degradation. The presence of EDTA also lowers the amount of free Mg²⁺ ions, which affects the specificity and efficiency of primers. If the sample is diluted 100 times, we eliminate EDTA and make PCR more effective, yielding higher signal. To verify our hypothesis, we repeated the experiment using DNA elution buffers for the dilution of the sample. This time, the results obtained were close to the real value. Hence, only by changing the elution of the clinical sample (either by water or elution buffer), we can accidentally change the estimated provided by qPCR by 400%. Therefore, it is vital to control very accurately the conditions of the reaction, because even a minor alteration significantly changes the final readout. Moreover, a reliable quantitative comparison of the samples purified using different methods may be challenging, or even impossible.

The synergistic PCR algorithm should be insensitive to such problems (Fig. 5B), as it provides absolute quantification and does not require any calibration. To prove it, we performed two series of tests on the samples prepared with the two buffers (water and MagJET buffer) that produced the largest difference in the results from qPCR analysis. In each series we run 12 synergistic analogue-digital tests, using the same qPCR protocol. In a stark contrast to the classic qPCR analysis, the synergistic algorithm correctly estimates the concentration regardless of the method of dilution. The assay was designed to provide the estimates with a 60% standard deviation over a 4 log dynamic range. In each assay we diluted the sample into a sequence of 165 μL partitions and added to each 15 μL of the master mix. The PCR was conducted on the Applied Biosystems 7500 Fast Real-Time System according to the GeneProofs’ prescription. As shown in Fig. 5B, the results of the 24 experiments corroborate very well with the calculated expected distribution of the estimates of concentration. The average of the 12 estimates for the samples diluted with water was 23 355 copies/mL (6.6% off the reference) and the average for the 12 samples diluted with the MagJET buffer was 25 163 copies/mL (0.65% off the reference). The standard deviation of the results corroborates with the predicted precision of the assay span on only 16 independent wells.

Please note that while the 12 repetitions of the 16-well synergistic assay provide a correct average of the estimate of concentration of the target DNA, the precision of such a small (16-well) assay is limited (60%, provided that no inaccuracies from the signal readout or the preparation of the sample are present). If better precision is needed we suggest constructing a bigger assay (i.e. one that uses more wells on the plate) via the algorithms described in this report. For example using 96 wells, one can construct a synergistic assay that delivers an estimate with a precision of 25% over a 4-log dynamic range.

Finally, we tested a set of synergistic assays that could be easily implemented in current qPCR devices. The assays were designed to cover the dynamic range of Ω = 10⁵ with σ = 40% (N = 32, x = 0.54), and Ω = 10⁸ with σ = 52% (N = 16, x = 0.4) and Ω = 10⁴ with precision σ = 10% (N = 200, x = 0.9). The performance of the assays is shown in Fig. 6A (each blue line shows a trend drawn from 10,000 grand canonical Monte Carlo simulations for
each assay). The observed performance of the assays agrees with analytical predictions, i.e. the precision and dynamic range of each assay is as requested in the design.

The results of the performance tests of synergistic assays shown in Fig. 6 are a good starting point for a more detailed comparison with current state-of-art solutions: classic dPCR (i.e. end-point measurement using identical compartments) and previously published optimized dPCR (end-point measurement using non-identical compartments)\(^{16}\). For a fixed number of compartments, digital assays offer assessment within a fixed dynamic range with fixed precision, while multivolume or multidilution designs are more flexible and can be tuned for precision and dynamic range to meet user's requirements. Moreover, such assays gain a competitive advantage over state-of-the-art solutions (dPCR and qPCR) thanks both to their high performance taking into account relatively low technical requirements, and low cost.

**Discussion**

In conclusion, we have shown that the use of synergistic combination of digital and analogue techniques may bring a new tool in quantitative assaying of DNA load in various samples. The use of information from digital and analogue signals allows to design quantitative assays that provide an absolute (no calibration needed) and precise assessment of concentration of the analyte. Moreover, the technical requirements for running such assays are minimized thanks to the simplified partitioning of the sample (reduction of the number of compartments) and no need for calibration.

The model that we presented also allows to tune independently the dynamic range and precision of the assay and, consequently, construct assays that provide different required precision in different ranges of concentration.

The presented methodology and design of quantitative assays may lead to the development of new, reliable and high-throughput devices for quantitative assaying and find use in the point-of-care applications. From the practical point of view, the reduction of the number of compartments per assay allows running multiple independent assessments using one standard multiwell plates on the standard qPCR machines. For example, one standard 96-well plate is enough to run (i) 3 independent assays that provide dynamic range within 5 orders of magnitude and precision \(\sigma = 50\%\), or (ii) 4 assays with \(\Omega = 10^4\) and \(\sigma = 55\%\), or (iii) 6 assays with \(\Omega = 10^3\) and \(\sigma = 60\%\). As synergistic PCR assays can readily be designed and executed on standard qPCR devices, they can be used to assess samples for which accuracy of the estimates is important and calibration is difficult or impossible, especially in research applications. In a further perspective, the combination of the new assays with simple detection schemes and high speed amplification systems can be used to introduce an inexpensive system offering precise and absolute quantization taking few minutes from the insertion of the sample, or constructing fast high-throughput systems for the rapid analysis of large number of samples.

**Methods**

**Digitalized RT-PCR assay is immune for initial sample buffer composition.** Materials. All experiment were prepared on IVD certificated PCR kit for Cytomegalovirus detection (GeneProof). Internal calibrator from the kit was used as a DNA template after 40 times diluted in water or 3 different elution buffers from commercially available DNA isolation kits (AE elution buffer from QIAamp DNA Mini Kit (Qiagen), MBL5 elution buffer from NucleoMag Blood (MACHEREY-NAGEL) and MagiET elution buffer from Whole Blood Genomic DNA Kit (Thermo Scientific) to obtain model samples with 25 000 copies of the target DNA per mL.

Methods. To compare the traditional qPCR with digital approach three-step amplification protocol was performed in 7500 Fast Real-Time System (Applied Biosystems) according to Cytomegalovirus PCR kit prescription: UGD decontamination 37 degrees for 2 min an initial denaturation at 95 degrees for 10 min. Subsequently, target
amplification involved 45 cycles of 5 s at 95 degrees, 40 s at 60 degrees for annealing, then extension for 20 s at 72 degrees. After amplification cycles, PCR products were evaluated for quality using melt curve analysis, which entailed 15 s at 95 degrees, 1 min at 70 degrees, 15 s at 95 degrees and 1 min at 55 degrees.

**Experimental Verification of the Synergistic Analogue-Digital Algorithm.** Materials. The reaction was performed in a volume of 20 μL, consisting of 4.5 μL of diluted plasmid DNA, 125 nM of forward and reverse primers (F: tcttgccctttctgcc, R: gatcggctcgagaatcattgcg) and 10 μL of SensiFAST SYBR No-ROX mix (Bioline).

Methods. We used the pJET1.2 plasmid with a fragment of LepA gene cloned from Mycobacterium smegmatis. The initial concentration of DNA was quantified with the use of a NanoDrop device. DNA used for all tests were stored in frozen aliquots.

A three-step amplification protocol was performed in 7500 Fast Real-Time System (Applied Biosystems); an initial denaturation was performed with one cycle at 95 degrees for 10 min. Subsequently, target amplification involved 50 cycles of 15 s at 95 degrees, 25 s at 62 degrees for annealing, then extension for 15 s at 72 degrees. After amplification cycles, PCR products were evaluated for quality using melt curve analysis, which entailed 15 s at 95 degrees, 1 min at 70 degrees, 15 s at 95 degrees and 1 min at 55 degrees.

14 different DNA concentrations were tested from 0.08 to 500,000 DNA molecules in first well (from 0.004 to 25,000 molecules/μL). The geometric sequences of the modification factors of compartments comprising tested assays were made via multi-dilution approach, i.e. the volume of all the compartments were same and the dilution factor changed geometrically.

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**Author Contributions**
P.R.D. and P.G. developed the algorithms, analyzed data and wrote the manuscript; K.G. and S.B. designed the experiments and, together with P.R.D. and P.G., the implementation of the algorithms for diagnostic assays; K.G. and S.B. executed the experiments.

**Additional Information**

**Supplementary information** accompanies this paper at http://www.nature.com/srep

**Competing Interests:** P.R.D. and P.G. declare competing financial interest in possession of shares of Curiosity Diagnostics Sp. z o.o. that seeks commercial implementation of the method.

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