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# Trends in Microbiology

### Forum

## SARS-CoV-2 viral RNA levels are not 'viral load'

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Ct values are commonly used as proxies of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) 'viral load'. Since coronaviruses are positive singlestranded RNA [(+)ssRNA] viruses, current reverse transcription (RT)qPCR target amplification does not distinguish replicative from transcriptional RNA. Although analyses of Ct values remain informative, equating them with viral load may lead to flawed conclusions as it is presently unknown whether (and to what extent) variation in Ct reflects variation in viral load or in gene expression.

The SARS-CoV-2 pandemic has prompted an unprecedented and large-scale use of diagnostic tests: serological tests detecting antibodies, antigenic tests detecting viral proteins, or RT-qPCR tests detecting viral RNA [1]. Because the genetic information in coronaviruses is carried by RNA molecules, the first step in PCR-based tests includes RT of the viral RNA to DNA, which is subsequently amplified and quantified through gPCR. DNA guantification is typically achieved by measuring the fluorescence emitted by certain molecules bound to the amplified double-stranded DNA. The outcome is a numeric value, commonly called 'Ct' for 'cycle threshold' or 'Cq' for 'quantification cycle', corresponding to the amplification cycle at which the detected fluorescence exceeds baseline levels.

Thus, larger amounts of viral RNA in a sample lead to larger amounts of retrotranscribed DNA and lower Ct values (see e.g., [2] for a rapid presentation of RT-gPCR and a review of quantitative analysis methods). A relatively large number of PCR-based tests to detect infection by SARS-CoV-2 have been developed, most of them targeting several locations in the viral genome. A PCR test is considered positive if the Ct value is below a predefined threshold for at least one of the targets, that is, the genomic nucleotide sequence amplified by the test. The number of genomic targets below the threshold and the precise values of the thresholds vary across manufacturers and tests. The simultaneous amplification of multiple genomic locations by some tests was originally conceived to introduce robustness and increase specificity, but serendipity turned it into a way to detect 'variants of concern' as mutations in the target sequence prevented PCR amplification and led to negative results for specific variants [3].

Mass testing has resulted in the generation of extensive data consisting of Ct values corresponding to different viral targets per sample. Most often they serve diagnostic purposes, and their use in this context raises no conceptual concern. Several studies, however, have used these Ct values as proxies of viral load<sup>†</sup>, which is understandable, not only because these values were available anyway, but also because alternative quantification methods (e.g., plaque assays) are labor-intensive and are still not well standardized.

Unfortunately, an important aspect of the biology of coronaviruses is neglected when using Ct values as a proxy of viral load.

Given that they are (+)ssRNA viruses, newly synthesized (+) strand RNAs can be used either for replication or transcription. This makes it unclear to differentiate the process, namely, replication or transcription, that is quantified by RT-qPCR.

To make matters more complicated, coronaviruses produce two kinds of mRNA molecules, genomic (i.e., full-size) and a variety of subgenomic (sgmRNA) seqments [4]. All genomic and sgmRNAs contain the genomic 5' leader sequence as well as the 3' polyadenylated end. All sgmRNAs are nested into the 3' end of the genomic mRNA: the smallest samRNA contains only the open reading frame (ORF) at the 3' end of the genome (in SARS-CoV-2, the ORF located the closest to the 3' end, and whose corresponding sgmRNA has hitherto been amplified, is called N [5,6]); the second smallest contains the two ORFs lying at the 3' end of the full-size RNA (N and 8), and so forth up to the largest sgmRNA, which carries all viral ORFs except 1a and 1b. Only the genomic mRNA carries all viral ORFs. Thus, the ORF at the 3' end of the SARS-CoV-2 genome, that is, N, is carried by all viral mRNAs, the one after it, that is, 8, is carried by all viral mRNAs, but those carrying only N, while the ORFs at the 5' end, that is, 1a and 1b, are carried only by full-size, genomic mRNA (Figure 1). Upon translation and processing, the 1a and 1ab polyproteins form the RNA polymerase, while ORFs present in sgmRNAs encode structural and accessory proteins. If all viral genomic and sgmRNA types occurred at equal frequency (which is not the case; see next paragraph), the RNA sequences of the different ORFs would occur at different frequencies because the more they are located towards the 3' end, the larger the number of sgmRNA types carrying them.

Coronaviruses have evolved mechanisms to regulate gene expression through translational and, more to the point of this article, transcriptional regulation [4]. Finkel and colleagues [6] showed, using RNA sequencing techniques on cell cultures, that different SARS-CoV-2 transcripts occur at different

<sup>&</sup>lt;sup>†</sup> Pretty much every study referring to SARS-CoV-2 viral load uses Ct as its proxy. Rather than singling out one or several randomly, we opted to not cite any specific reference.

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Figure 1. Expression and replication of severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2). The SARS-CoV-2 genome is carried by a 29.9 kb single-stranded RNA molecule, directly translatable by the cellular machinery as a messenger RNA (hence the positive polarity). The open reading frames (ORFs) located at the 5'of the full-length genome are translated into two polyproteins post-translationally processed into several non-structural proteins (nsps) involved, along with other viral and host cell proteins, in the replication-transcription complex (RTC). Replication of the full genomic (+)RNA at the 3' end by the RTC generates the full-size (-)RNA, which serves as the template for genomic replication. Replication does not necessarily proceed as far as the 5' end of the full-length viral genome, and can produce instead nine kinds of (-) subgenomic mRNAs (sgmRNAs). For the sake of simplicity, only two of them are exemplified in the figure, one reaching the N ORF and another one reaching the S ORF. These (-)sgmRNAs can be transcribed into (+)sgmRNAs, whose genetic sequences are nested by increasing inclusion from the coding 3' extremity. Although full-length genomic mRNA and sgmRNAs can encode several ORFs, only the ORF located at the 5' end of the curresponding (+)mRNA molecule is actually translated, for example: only ORF1a and ORF1b are translated from the full-length genomic mRNA even if this molecule spans all other viral ORFs.

abundancies, with variation spanning one order of magnitude (see their Figure 2b, mRNA axis). The exception is the transcript coding the accessory protein 7b, which is approximately three orders of magnitude less abundant than the most common one, N, coding the nucleocapsid protein. Transcript abundances further vary through time during experimental infection (see Figure 4d,e in [6]) as well as between cell lines of different hosts infected by different SARS-CoV-2 isolates (see Figure 4d,f in [6] and Figure 3c in [5]). The ranking order of transcript RNA abundance is thus variable across experimental conditions and does not match the order of the ORFs in the genome. Finally, in bovine coronavirus it was shown that sgmRNAs can directly serve as templates for the synthesis of the corresponding (–)sgmRNAs as well as for the synthesis of shorter nested sgmRNAs [7], allowing for additional regulation of mRNA and protein abundance.

In SARS-CoV-2 the different targets of RT-qPCR are differentially affected by transcriptional activity because ORFs closer to the 3' end are present in more mRNA types than ORFs closer to the 5' end. Further, the relevance and biological meaning of the Ct values corresponding to the different targeted locations in an individual sample should be considered with a lot of caution, as gene expression depends on environmental, (viral) genetic, and host (epi)genetic factors, as well as on the natural history of the infection. As Sola and colleagues wrote in 2015, 'limited information is available on the temporal regulation of viral translation, replication, and transcription over the course of infection and on how switching between these processes occurs' [4]. This remains largely true, especially in the context of the ongoing pandemic. Interestingly, at least under some experimental conditions, quantification of infectious virions through plaque assays yielded statistically significant differences between SARS-CoV-2 genotypes while quantification through single-target RT-qPCR did not reveal differences for the same treatments (Figure 3b,c in [8]). Overall, it is unclear how good a proxy of viral load Ct values are, or what the observed differences in Ct values actually reflect. Their use in qualitative diagnostics is not problematic as long as the detection thresholds in these kits are properly established, but finer quantitative applications deserve more caution. Thus, using Ct values to infer whether an infection is progressing versus resolving within an individual or growing versus declining within a population should not be problematic, provided sufficient

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sampling and appropriate standardization. On the other hand, using Ct values to predict contagiousness may be riskier for several reasons: sample variability [9], Ct variation across qPCR targets [10], variation among patients in their physiological status [10,11], infection age [10,11], or viral variants [12], to name a few. It should be emphasized that, with the exception of sample variability, these sources of variation may result in changes in viral replication and/or viral gene expression though it is presently unknown whether and to what extent they would lead to relative increases or decreases.

Despite experimental biases associated with RT-qPCR protocols and differential robustness with respect to input sample quality [9], quantitative analyses of Ct values may be highly informative, for example, in allowing for the detection of patterns in 'levels of RNA' in patients with different characteristics (such as viral variant, host gender, age, clinical presentation and stage of the infection) or in allowing to correlate such patterns with epidemic properties in host populations. For example, given that *a priori* viral replication levels should be reflected equally by all RT-qPCR targets, differences in Ct values among targets lying in different viral ORFs should reflect different transcription levels. Such observations could help reveal interesting, and potentially epidemiologically significant, variations, for example, among SARS-CoV-2 variants.

In summary, since Ct values quantify 4. both viral replication and transcription, 5. the relevance of their use depends on how inferences may be affected by imprecise estimates of viral load. 7.

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#### **Declaration of interests**

There are no interests to declare.

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