Critical appraisal to using relative or absolute cardiac troponins change for diagnosing acute myocardial infarction

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Both cardiac troponins T (cTnT) and I (cTnI) are universally considered the reference biomarkers for diagnosing acute myocardial infarction (AMI) (1). Irrespective of the guidelines or recommendations used for diagnosing AMI, the conventional approach entails a baseline measurement eventually followed (i.e., non-diagnostic values at the first measurement or suggestive electrocardiogram changes) by serial sampling at different times points, which are aimed to detect suggestive changes of cardiac troponins (cTn) reflective of an acute ischemic event (2,3). Two different strategies have been proposed for interpreting results of serial cTn testing, the former entailing an absolute variation of concentration from the baseline value (i.e., expressed in cTn concentration, ng/L), and the latter based on a relative change from the baseline value (i.e., expressed in percentage increase, %). The most commonly used cutoffs of cTn variation used for serial sampling encompass an absolute delta comprised between 5 and 10 ng/L or a ~20% variation, since these thresholds actually reflect the within-intra-individual biological variation of cTn in emergency department patients (4).

Though many studies have been published on the diagnostic efficiency of both strategies and currently there is general consensus that the use of the absolute variation may be more clinically useful for both diagnosing and ruling out AMI (5,6), absolute changes of cTn concentration may occur for non-biological causes, i.e., due to preanalytical or analytical issues (7). The latter aspect is especially significant. Although the new generation of high-sensitivity (HS) cTn immunoassays is characterized by considerably magnified analytical performance (including a lower analytical imprecision) (8,9), an improved analytical variation may impact the efficiency of diagnostic algorithms based on absolute cTn variation during serial sampling.

A pragmatic representation of this issue is given in Figure 1A, showing the confidence interval (CI) limits of Roche HS cTnT values based on a previously estimated analytical imprecision (intra-assay coefficient of variation; CV) of 8.4% (10), which is then applied to either an absolute increase of 5 ng/L (as suggested by Marjot et al.) (11) or the conventional 20% increase from the baseline cTnT value. The baseline value (straight black line) would never be included within the CI limits drawn according to the level of analytical imprecision of the method when the diagnostic threshold is based on the 20% increase of cTnT from baseline (dotted blue lines). On the other hand, for cTnT concentrations >50 ng/L, the baseline values would be included within the CI limits drawn according to the level of analytical imprecision of the method when the diagnostic threshold is based on a 5 ng/L increase of cTnT from baseline (dotted blue lines). Practically, this means that the window of analytical imprecision of the cTnT immunoassay may virtually impair the diagnostic performance of the absolute increase after a certain concentration (i.e., 50 ng/L for cTnT).

The absolute variation may exhibit better diagnostic performance when using HS troponin immunoassays with lower analytical imprecision. Figure 1B shows the CI limits of Abbott HS cTnI values based on a previously estimated analytical imprecision (mean intra-assay CV) of 3.7% (12), which is then applied to either an absolute increase of 6 ng/L (as suggested by Neumann et al.) (13) or the
conventional 20% increase from the baseline cTnI value. Like cTnT, the baseline value (straight black line) would never be included within the CI limits drawn according to the level of analytical imprecision of the method when the diagnostic threshold is based on the 20% increase of cTnI from baseline (dotted blue lines). However, for cTnI concentrations >150 ng/L the baseline values would be included within the CI limits drawn according to the level of analytical imprecision of the method when the diagnostic threshold is based on a 6 ng/L increase of cTnI from baseline (dotted blue lines). This means, again, that the window of analytical imprecision of the cTnI immunoassay may virtually impair the diagnostic performance of the absolute increase after a certain concentration (i.e., 150 ng/L for Abbott cTnI).

This straightforward concept can hence be easily deployed to all the commercially available cTnI and cTnT HS immunoassays, by knowing in advance the analytical imprecision and the optimal thresholds for both relative and absolute variations of the method. Although it can be argued that the threshold over which the CI limits of imprecision will include the baseline cTn values are relatively distant from the 99th upper reference limit (URL; 13 ng/L for cTnT and 27 ng/L for cTnI, respectively), cTn concentrations as high as 6.5-fold the URL are frequently observed in patients admitted to the emergency department with non-ischemic chest pain. These patients, who are finally diagnosed with myocardial injuries different from AMI, regularly undergo serial cTn monitoring for ruling out acute myocardial ischemia (11,14). In these conditions, values as high as 85 ng/L for cTnT and 176 for cTnI can be observed, thus exceeding, in both cases, the thresholds after which the imprecision of the cTn immunoassays would make it unadvisable using the absolute variation (i.e., 50 and 150 ng/L, respectively).

Taken together, the data emerged from our analysis seemingly shows that the relative change may outperform (and may be also considered a safer approach) the absolute variation in patients presenting with intermediate or high cTn values (Figure 1). On the other hand, this strategy may be less efficient for diagnosing AMI in patients presenting with low cTn values or when using narrow serial testing (i.e.,

Figure 1 Impact of the analytical variability using the absolute or relative cardiac troponin variation for diagnosing acute myocardial infarction (AMI). The straight black line identifies the baseline value, whilst the range comprised between the dotted blue lines and between the dotted red lines designate the imprecision window of the 20% cardiac troponin increase from baseline and the imprecision window of the absolute cardiac troponin increase from baseline, respectively.
1- or even 2-hour protocols), so that the absolute change may generally appear more clinically and analytically reliable in these conditions (5). Pragmatic working solutions may thus entail the use of absolute cTn increases when the concentration is lower than the 99th URL, whilst the 20% variation may then be more analytically robust and reliable for cTn values above such limit, as shown in Figure 2. This may henceforth lead the proposal of a revised diagnostic algorithm for AMI, which will need to be tested in future studies, combining symptoms onset, analytical characteristics of cTn immunoassay (namely the limit of detection and the URL), as well as the absolute and relative cTn variation (see Figure 3).

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Figure 2 Strategy based on the combination of absolute and relative cardiac troponin variation for diagnosing acute myocardial infarction (AMI). The straight black line identifies the baseline value, whilst the range comprised between the dotted blue lines and between the dotted red lines designate the imprecision window of the 20% cardiac troponin increase from baseline and the imprecision window of the absolute cardiac troponin increase from baseline, respectively. URL, upper reference limit.

Figure 3 Algorithm based on the combination of absolute and relative cardiac troponin variation for diagnosing acute myocardial infarction (AMI). cTn, cardiac troponin; LoD, limit of detection; URL, upper reference limit.
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