A single blood test to rule out acute coronary syndrome

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Strategies to improve the assessment of patients with suspected acute coronary syndrome continue to evolve, in recognition that fewer than 20% of those attending the Emergency Department with chest pain receive a diagnosis of myocardial infarction. Identifying patients without myocardial infarction at an earlier stage has the potential to reduce hospital admissions for serial cardiac biomarker testing, and facilitate appropriate investigation for alternative causes. However, such strategies are only helpful if it can be demonstrated that they do not compromise patient safety.

In 2016, the National Institute for Health and Care Excellence (NICE) updated their guidance on the evaluation of patients with suspected acute coronary syndrome. For the first time, they recommended clinicians consider ruling out myocardial infarction if a patient has very low concentrations of cardiac troponin at presentation when measured using a high-sensitivity assay.1 This guidance could lead to a significant reduction in the proportion of patients who require serial testing, and may tempt clinicians to consider upgrading their infrastructure to facilitate implementation. In the UK, two high-sensitivity assays are recommended by NICE for use in clinical practice, the Roche Elecsys high-sensitivity cardiac troponin I assay (hs-cTnI) and the Abbott ARCHITECT_{STAT} high-sensitivity cardiac troponin, and there are important differences in the normal reference range, diagnostic thresholds, levels of imprecision and in the lowest absolute concentrations which can be reliably detected, also known as the limit of detection (LoD) (table 1).

NICE recommend clinicians apply the LoD as a threshold below which myocardial infarction can be safely ruled out at presentation. Such a strategy is only recommended for patients deemed to be at low risk of myocardial infarction 'as indicated by a validated tool'. During their appraisal, NICE considered evidence from studies including both the Thrombolysis in Myocardial Infarction (TIMI) score and the Global Registry of Acute Coronary Events score. Both scores were derived and validated in patients with confirmed myocardial infarction to confer prognosis, but over time, these scores have been implemented for risk stratification in patients with suspected, not confirmed, myocardial infarction. Importantly, cardiac troponin concentrations are embedded in both risk scores. NICE ultimately recommend the TIMI score, which has been previously validated in patients with suspected acute coronary syndrome alongside a contemporary troponin assay and serial testing, 2 but not with a high-sensitivity assay and the LoD at presentation alone.

Carlton *et al* provide the first validation of the NICE guidance in a pooled study of over 5000 patients, in five observational cohorts across two continents, with varying prevalence of major adverse cardiovascular events (4.8% to 15.6%).3 They found when an hs-cTnT of <5 ng/L (LoD) was applied alongside a TIMI score of o and a non-ischaemic ECG, the sensitivity and negative predictive value (NPV) were extremely high, at 99.5% (95% CI 98.1% to 99.9%) and 99.6% (95% CI 98.7% to 100%), respectively. They derived a metaestimate for sensitivity of 98.7% (95% CI 96.5% to 99.6%), with low heterogeneity observed between cohorts (I_2 15.3). For the hs-cTnI, using the LoD (<2 ng/L) and a TIMI score of o alongside a non-ischaemic ECG, the sensitivity was 98.9% (95%CI 97.4% to 99.6%) and NPV was 99.5% (95%CI 98.8% to 99.8%). The meta-estimate for sensitivity was similar (98.5%, 95% CI 95.4% to 99.5%) but the heterogeneity was high (I_2 73.7). The reason for the observed heterogeneity is unclear, but may reflect differences in the assay used for diagnostic adjudication and testing between cohorts. These strategies would identify between 17.9% (95% CI 16.6% to 19.3%) and 21.0% (95% CI 19.9%

to 22.2%) of patients as low risk, respectively. The authors evaluate several additional approaches not included in the recommendations of NICE, including the use of thresholds above the LoD in combination with the TIMI score, and their data suggest that higher thresholds (such as <7 ng/L on the hs-cTnT assay, or <5 ng/L on the hs-cTnI assay) could increase in the proportion identified as low risk without compromising sensitivity or NPV.

While NICE recommend use of the TIMI score, the true need for clinical risk scores in this setting is uncertain. A recent meta-analysis of 9269 patients found a normal ECG and an hs-cTnT result below the LoD provided excellent NPV (99.3%, 95% CI 97.3% to 99.8%) and sensitivity (98.7%, 95% CI 96.6% to 99.5%) for the diagnosis of myocardial infarction, without the need for additional risk scores.4 There were no deaths at 30 days in patients classified as low risk with the index test. This reflects our understanding that patients classically considered high risk (due to increasing age or cardiovascular risk factors such as diabetes, renal disease or prior ischaemic heart disease) have chronic elevation in high-sensitivity cardiac troponin concentrations (within the normal reference range) and are less likely to have low concentrations to support early discharge. Indeed, the European Society of Cardiology advocate use of the LoD at presentation in conjunction with the ECG, but do not recommend the addition of clinical risk scores.

However, there is undoubtedly an appetite for clinical risk scores in some settings, perhaps due to the additional perceived diagnostic confidence they provide. One of the most widely used risk scores, the HEART score, was developed and validated in a suspected acute coronary syndrome population. This score is based on clinical variables selected a priori (*H*istory, *E*CG, *Age*, *R*isk factors, cardiac *T*roponin) with arbitrary weighting chosen on a pragmatic basis. A recent meta-analysis of 11 217 patients demonstrated this score had a sensitivity of just 96.7% (95% CI 94.0% to 98.2%), below the threshold of 99% which most emergency department physicians deem acceptable.5 Whether use of this score offers additional benefit over risk

stratification with troponin alone is unclear. Comparative studies including risk stratification thresholds alone or in combination with risk scores are required to determine if improvements in safety can be obtained.

There are important limitations to the study findings of Carlton *et al.* It is notable that there was a significant proportion of missing data, with over 2000 patients excluded from the hs-cTnT cohort and 784 patients from the hs-cTnI cohort. This reflects the pragmatic basis on which the data were compiled for this study, but may have contributed to important differences in risk profile between cohorts. Patients in the hs-cTnT cohort were older, with more cardiovascular risk factors, and one may therefore anticipate higher baseline cardiac troponin concentrations and a reduced likelihood of a missed myocardial infarction compared with the hs-cTnI cohort.

So, is it time to adopt the NICE guidance? The evidence supporting implementation of strategies including low high-sensitivity cardiac troponin concentrations is strong, but some uncertainty remains. All studies on which the NICE recommendations are based were observational in nature (ie, no patients were discharged from hospital on the basis of a single troponin result), and the same applies to the vast majority of studies in this area. A multicentre randomised controlled trial of implementation of a rule-out strategy using a single hs-cTnI concentration (<5 ng/L) has completed recruitment in Scotland (NCT:03005158), and similarly, a randomised controlled trial of the hs-cTnT is being planned (LoDED study, personal communication Dr Edd Carlton). Both trials will give complementary insight into the safety and efficacy of this approach.

For the clinician who plans to implement this strategy prior to the outcomes of these randomised controlled trials, there are some important considerations. Patients who present early after onset of symptoms are challenging to recruit and therefore under-represented in all observational cohort studies. It is therefore recommended that serial testing is performed in all who present early after onset of symptoms. Similarly, any patient with myocardial ischaemia on the ECG should not be considered for early rule out and should undergo serial troponin testing. Of utmost importance is an awareness of the assay in use at your institution, the normal reference range and the appropriate diagnostic and risk stratification thresholds which are not equivalent. Where low concentrations are reported, it is important to ensure appropriate standards for clinical reporting can be met and maintained under routine working conditions. As noted by NICE, implementation of the proposed early rule-out strategy should include clinical audit, with attention paid to the time taken to rule out the diagnosis and on the clinical outcomes of patients with suspected acute coronary syndrome.

Clinicians should be confident that newer approaches using low concentrations of cardiac troponin are a magnitude safer than prior strategies using the 99th centile alone at presentation and 3 hours as recommended in the European Society of Cardiology guidelines.6 Clinicians should be prepared to restructure the assessment of patients with suspected acute coronary syndrome in their institution to harness the potential of high-sensitivity cardiac troponin testing, and to improve the efficiency and safety of healthcare delivery.