**How well do laboratories adhere to recommended guidelines for cardiac biomarkers management in Europe? The CArdiac MARker Guideline Uptake in Europe (CAMARGUE) Study of the EFLM Task Group on Cardiac Markers.**

Running head Troponin utilisation survey

Paul Collinson1, Janne Suvisaari2, Kristin M. Aakre3,4, Hannsjörg Baum5, Christopher J. Duff6,7, Damien Gruson8, Angelika Hammerer-Lercher9, Kari Pulkki2, Sanja Stankovic10, Michel R. Langlois11,12 , Fred S. Apple13 and Päivi Laitinen2 for the EFLM Task Group on Cardiac Markers.

1 Department of Cardiology, St George's University Hospitals NHS Foundation Trust and St George’s University of London, London, UK.

2 Clinical Chemistry, HUS Diagnostic Center, University of Helsinki and Helsinki University Hospital , Helsinki, Finland.

3 Department of Medical Biochemistry and Pharmacology, Haukeland University Hospital, Bergen, Norway.

4 Department of Clinical Science, University of Bergen, Bergen, Norway.

5 Institute for Laboratory Medicine and Transfusion Medicine, Regionale Kliniken Holding RKH GmbH, Ludwigsburg, Germany.

6 Department of Clinical Biochemistry, University Hospitals of North Midlands NHS Trust, Stoke-on-Trent, UK.

7 School of Primary, Community and Social Care, Keele University, Stoke-on-Trent, UK.

8 Department of Clinical Biochemistry, Cliniques Universitaires St-Luc and Université Catholique de Louvain, Brussels, Belgium.

**9** Institute of Laboratory Medicine, Cantonal Hospital Aarau AG, Switzerland.

10 Center for Medical Biochemistry, Clinical Center of Serbia, Belgrade, Serbia.

11 Department of Laboratory Medicine, AZ St. Jan Brugge-Oostende AV, Brugge, Belgium.

12 Ghent University, Faculty of Medicine and Health Sciences, Ghent, Belgium.

13Department of Laboratory Medicine and Pathology, Hennepin Healthcare/HCMC, Minneapolis, MN 55415, USA and Department of Laboratory Medicine and Pathology, University of Minnesota, Minneapolis, MN, USA

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Correspondence to

Correspondence to: Professor Paul Collinson. Department of Clinical Blood Sciences, St George’s University Hospitals NHS Foundation Trust, Cranmer Terrace, London SW17 0QT, UK. Tel 0208 725 5923/5934 Fax 0208 725 5868

Email: [pcollins@sgul.ac.uk](mailto:pcollins@sgul.ac.uk) ; [paul.collinson@stgeorges.nhs.uk](mailto:paul.collinson@stgeorges.nhs.uk); [paul.collinson@ntlworld.com](mailto:paul.collinson@ntlworld.com)

**Abstract**

**Introduction.** The CARdiac MARker Guideline Uptake in Europe **(CAMARGUE)** program is a multi-country audit of the use of cardiac biomarkers in routine clinical practice.

**Methods.** An email link to a web-based questionnaire of 30 multiple-choice questions was distributed via the professional societies in Europe.

**Results.** 374 questionnaires were returned from 39 countries, the majority of which were in northern Europe with a response rate of 8.2-42.0%. The majority of the respondents were from hospital with proportionately more responses from central hospitals than district hospitals. Cardiac troponin was the preferred cardiac biomarker, evenly split between cardiac troponin T (cTnT) and cardiac troponin I (cTnI). Aspartate transaminase and lactate dehydrogenase are no longer offered as cardiac biomarkers. Creatine kinase, creatine kinase MB isoenzyme and myoglobin continue to be offered as part of the cardiac biomarker profile in approximately on 50% of respondents. There is widespread utilisation of high sensitivity (hs) troponin assays. The majority of cTnT users measure hs cTnT. 29.5% of laboratories measure cTnI by a non-hs method but there has been substantial conversion to hs cTnI. The majority of respondents used ng/L and use the 99th percentile as the upper reference limit (71.9% of respondents). A range of diagnostic protocols are in use.

**Conclusions.** There is widespread utilisation of hs troponin methods. A significant minority do not use the 99th percentile as recommended and there is, as yet, little uptake of very rapid diagnostic strategies. Education of laboratory professionals and clinicians remains a priority.

**Introduction.**

The CARdiac MARker Guideline Uptake in Europe **(CAMARGUE)** program is a multi-country audit of the use of cardiac biomarkers in routine clinical practice undertaken by the European Federation of Laboratory Medicine cardiac markers task group (EFLM TG-CM). The objective is to benchmark clinical laboratory practice at the time of audit compared with the guidelines in use at the same time point. It therefore aims to describe the evolution of cardiac biomarker practice and to document how well laboratories remain aligned to clinical and laboratory guidelines.

The first audit focussed on the introduction of cardiac troponin (cTn) and natriuretic peptides into routine clinical use (1;2). Subsequent audits covered the evolution of cardiac biomarker testing as the evidence base and assay methodologies underwent progressive improvement (3-6). The most recent audit covered cardiac biomarkers and lipids(7). The major change since the last survey was undertaken in 2015 has been widespread endorsement of high sensitivity cTn (hs cTn) assays and their increasing availability from manufacturers. This has been reflected in a number of documents published since 2015. The International Federation of Clinical Chemists (IFCC) task force (now committee) on clinical applications of cardiac biomarkers, (IFCC C-CB) made a number of recommendations on implementing high sensitivity cTn (hs-cTn) assays in clinical practice. This covered the 99th percentile upper reference limit and calculating serial changes (8). Specifically, it distinguished between hs- cTn assays (assays which would detect cTn in 50% or more of healthy subjects with the analytical imprecision at the 99th percentile <10%) and conventional (contemporary) sensitive assays which would achieve this imprecision goal but not the detection of cTn in 50% or more of healthy subjects. The European Society of Cardiology (ESC) updated the guidelines on diagnosis of non-ST elevation myocardial infarction (NSTEMI) specifically recommending the use of hs cTn assays and that these could be used with measurements taken on admission and three hours from admission. In addition, they recommended for consideration rapid diagnostic algorithms based on admission measurement alone or sampling over short time intervals (measurement on admission and one hour from admission) (9). These changes are summarised in table 1. Finally, the fourth universal definition of myocardial infarction has further updated concepts with commentary on the analytical issues affecting cTn assays and specifically recommending hs cTn assays are used for routine diagnosis (10) as has the combined IFCC and Academy of the American Association of Clinical Chemists(11). This report concentrates on the use of cTn in routine clinical practice with particular emphasis on the use and implementation of high sensitivity assays.

**Methods**

The methodology used has been described in full in previous paper(1-4). Briefly a web-based questionnaire was developed by the EFLM TG-CM. The questions focused on recommendations of the IFCC C-CB (8), the ESC recommendations on NSTEMI (9) and the universal definition of myocardial infarction (10), for reasons discussed in the introduction. This questionnaire included 30 multiple-choice questions regarding cardiac biomarker testing and covered the analytical methods and manufacturer used, measurement units, decision threshold, and the use of decision-enhancing comments (supplementary Table 1). Structured questions were used with the option of adding free text. The questionnaire was implemented using a web-based survey system consisting of a HTML-AJAX interface and a CGI program storing the results in XML files on a database server. Results from raw data XML files were tabulated and the numbers of different answers for each question were calculated for further analysis using Microsoft Excel then the data extracted into a Microsoft Access database. The questionnaire was compiled with experience from the two previous questionnaires but also incorporated some modifications in design to allow more in-depth analysis of certain responses. A link to the online questionnaire was sent on March 1, 2019 to EFLM National Societies from 40 European member countries. The questionnaire was open till August 31, 2019. Statistical analysis was performed using the Analyse It (Analyse-It.com) add-in for Excel. Nonparametric statistics were used throughout.

**Results**

*Demographics*

A total of 374 questionnaires were returned from 39 countries, less than the previous survey (493 responses). Not all questionnaires were fully completed and where a null return (no response to the question) occurred, this is indicated in the subsequent analysis. Where questions are only relevant to hospital practice, non-hospital responders have been excluded from the analysis and this is indicated in the text. Responses were obtained from the majority of European countries (96% of all responses) but there was under-representation of the southern European countries (7.2% of responses) with the exception of Portugal. 84.8% of responses were from hospital laboratories with the breakdown shown in figure 1 (no response 6.4%). Proportionately, there were more central hospitals than district hospitals (203 versus 106) which was statistically higher than in the previous survey (219 versus 174, p = 0.009). 68.7% of laboratories reported that they were accredited (no response 0.5%). The majority of accreditation (76%) was to ISO 15189. Based on countries where reliable information was available from quality assurance scheme participant numbers, response rate was 8.2-42.0% with a median of 22%.

For the hospitals, median bed number was 600 (range 3.0-3810, interquartile range 360-1000). Angiography was offered by 56.5% of the participants. 14.2% of laboratories were unaware as to whether or not their hospital offered angiography. 75.9% of central hospitals and 52.6% of district hospitals offered angiography. Overall, 319/374 (85.3%) offered a 24-hour service including 4 clinics and 3 freestanding laboratories (1.6% no response). When the analysis was confined to hospital affiliated laboratories, 95.6% offered a 24-hour service (0.8% no response) and in hospitals providing angiography, 99.4%.

*Biomarker preferences*

The preferred cardiac biomarker was cTn, offered overall by 95.2% of all laboratories surveyed, 99.1% of hospital laboratories (although it was not the first line test in 1.3% of those offering cTn) and 100% of those providing angiography, similar to the data from the 2013 survey (98.6% offered cTn). For hospitals not providing cTn, the reasons stated were financial. The median number of cTn measurements annually was 13,500 for hospital laboratories.

The trend of decline of aspartate transaminase (AST), lactate dehydrogenase or its isoenzymes (LD) and creatine kinase (CK) measurement was maintained (figure 2 upper panel). Measurement of CK and its CK-MB isoenzyme has remained stable from 2013 to 2019 (figure 2 lower panel) even when only hospital laboratories were examined (data not shown). Overall 93.6% of respondents used cTn as a front-line test in suspected acute coronary syndromes (ACS), 97.8% in hospitals, similar to 2013, but 66.6% continue to offer additional cardiac biomarkers to cTn as a routine. 356 laboratories currently measured cTn, 181 cardiac troponin T (cTnT, 50.8%,179 high sensitivity cTnT, hs-cTnT) and 173 cardiac troponin I (cTnI, 48.5%,122 high sensitivity cTnI, hs-cTnI) with 2 unspecified. The major change since 2013 has been the introduction of hs cTn methods. Currently 84.6% (301/356) of laboratories measure cTn with a high sensitivity method (59.5% hs-cTnT, 34.3% hs-cTnI). The impact of introducing hs cTn on the laboratory is shown in figure 3.

*Recent changes in cTn assays*

In the past two years 74.4% of those surveyed had not changed cTn method. Of those who measured cTnT 147/179 (82.1%) were already using hs-cTnT. Only 2 existing users, both measuring cTnT by point of care (POC), continued to use a non-hs assay. This probably represents service provision restricted to POC testing. Only 10 laboratories switched to hs-cTnT in the previous two years, the remaining 22 switched from cTnI or POC (1 laboratory) to hs-cTnT. Roche originally offered a conventional cTnI assay but did not develop an hs-cTnI assay.

Unlike cTnT, where 98.9% utilised hs-cTnT currently, 51/173 (29.5%) of laboratories currently measuring cTnI utilise a non-hs assay. In total 67/173 (38.7%) of laboratories measuring hs-cTnI reported no change in method. As would be expected this varied from 95.8% for Abbott to 0% (Roche cTnI). For Roche conversion to hs-cTnI measurement would have required changing manufacturer. However, some Roche cTnI users have changed to hs-cTnT. A detailed analysis of the data is available in the supplementary table 2. The current distribution of methods utilised is shown in supplementary figure 1. Utilisation by country was variable (supplementary figure 2) reflecting the different manufacturers domestic market position.

*Choice of reporting units*

Current IFCC guidelines recommend criteria for classification of cTn methods and for reporting hs-cTn in nanograms/L and non-high sensitivity assays in micrograms/L(12). The majority of laboratories (338) correctly characterised the methodology they were using but 17/356 (4.7%) were using an assay that did not meet high sensitivity criteria but reported the assay they were using as high sensitivity (no response 1). Of the 356 laboratories measuring cTn, hs-cTn was measured in 283, 69.6% (197/283) reported in ng/L and 14.5% pg/ml. 9.5 % reported in ng/ml and 4.9% µg/L. 1.4% of laboratories reported both units. In the 52 laboratories not utilising high sensitivity assays 48.1% (25/52) reported in ng/L with 26.9% in ng/ml and 13.5% in µg/L. 5.9% (21/356) did not respond

*Diagnostic cut offs*

There has been a significant shift in the choice of the 99th percentile as the URL for AMI. 71.9% (256/356) utilise the 99th percentile although the 8.1% reported not using any decision limit and no data was obtained in 9.8% (supplementary figure 3). This is significantly different from the previous surveys with the choice of the 99th percentile now dominant (figure 4).

A sex specific upper reference limit was used in only 23% (68/300) of laboratories utilising hs-cTn assays (10% non-responders). In addition to the 99th percentile being the dominant discriminant for AMI, there has been a switch to using the correct numeric value. For hs cTnT 74.4% now use the recommended value of 14 ng/L, a significant improvement from 2013 (50.2%, p = 0.0001).

For laboratories measuring cTn (n = 356), decision to use the AMI cut-off and choice of cut off was again dominated by the recommendations of the instruction for use (IFU), the package inset from the manufacturer. This confirmed the trend seen in the past three surveys and was statistically significant (p < 0.0001, supplementary figure 4). The choice of AMI cut-off was derived from the IFU in 56.7% (202/356) of responses and from expert recommendations or derived from the literature in 28.7% (non-response rate 14.6%). However, when the IFU was used as the source of data for selection of AMI cut-off (202), the 99th percentile was utilised in 85% (172/202), 5% continued to use the 10% CV with the other 10% using a variety of cut offs. More interestingly, the reason for selecting the 99th percentile as AMI cut-off (n = 256) was not the IFU alone. The IFU was the prime reason for selecting the 99th percentile in only 67.2% (172/256) of respondents with 28.5 % citing clinical or professional reasons (no response in 4.3%).

Analysis of alternative clinical use, diagnostic algorithms, sample timings, interpretive comments and demand management was confined to hospital laboratories measuring cardiac cTn only (314). 45.5% use cTn for other clinical diagnostic purposes than diagnosis of ACS of which the majority was monitoring after cardiac revascularisation procedures (supplementary figure 5). Only 1.9% reported using a different diagnostic discriminant when cTn was used for these purposes.

*Testing protocols*

48.7% used serial testing with 7.6% using admission rule out plus serial testing up to 3 hours and 10.8% the ESC 0/1 hour rule out algorithm. 32.5% did not specify a particular testing strategy. Similar results were seen if hs-cTn assays only were considered. Serial testing (n = 153) was most commonly 0-3 hour sampling (58.8%, 90/153) followed by 0-6 hour sampling (15.7%). Taking all the available data from which sample timings could be extracted (n = 174), diagnosis was expected to be complete by 3 hours in 68.0 % or 6 hours in 21.7% with other strategies in use in only 10.3%. 24% of laboratories provided interpretation as a computer-generated comment and 4% calculated the delta (although absolute or relative was not specified) but the remainder did not indicate that any additional information was provided. 55.4% of respondents documented a derivation of their diagnostic algorithm but 45.6% did not. The largest single source of derivation of a diagnostic algorithm was the ESC (25.5%). Although laboratories use admission testing plus 0-3 hour serial testing or 0-3 hour serial testing (n = 104), this strategy was derived from ESC recommendations in only 34.6%.

66% of all the hospital laboratories indicated that they have a procedure in place for regular review of the diagnostic pathway for suspected ACS patients. 46.1% reviewed on an annual basis and in total 84.5% reviewed within 3 years. 38.9% of respondents said test ordering was determined by clinician preference alone. However, 41.1% used structured ordering either an order set plus additional clinician ordering (22.6%), a timed set alone (1.0%) or a combination of a timed order set plus single orders (17.5%). 14.6% of respondents provided no data. Only 3.5% of participants utilised a demand management strategy for cTn testing by limiting the number of cTn orders. No data was available on the average number or timing of cTn orders per patient.

*Internal and external quality assessment*

Overall, of those laboratories who measured cTn (n = 356) 81.2% indicated they used internal quality control, 2.5% that they did not and 16.3% provided no data. The most frequently used control was around the 99th percentile (66.8%) distribution of the controls are shown in supplementary figure 6. Only a minority of laboratories (1.7%) used a control close to the limit of detection of the assay. This was confined to hospital laboratories only. 75.3% participated in external quality assessment (proficiency testing) for cTn, 17.7% did not provide any data and 7% did not participate.

**Discussion**

There are six main findings from this study. First, the measurement of cTn is now the preferred cardiac biomarker and is provided by nearly all laboratories. The only reason for non-provision was financial. Second, CK, CK-MB and myoglobin continue to be measured with little change from 2013, with AST and LD no longer measured. Third, there has been widespread adoption of high sensitivity cTn measurement. Fourth, there has been widespread adoption of the 99th percentile URL as the diagnostic discriminant for AMI, with the main source of information the manufacturers IFU. Fifth, although there is a shift towards use of rapid diagnostic algorithms, they are not yet in use by the majority of hospitals. Sixth, the use of structured ordering protocols occurred in less than 50% although documentation and regular formal review of the diagnostic protocol in use was reported by the majority, but not all, hospitals.

The continued use of CK, CK-MB and myoglobin is similar to that seen in the previous survey(4). There are several possible reasons. Some respondents may have indicated that they retain these tests as part of a generalised test menu rather than specifically for cardiac disease. This would be consistent with the majority of laboratories reporting that cTn was the first line test. Continued measurement of CK is likely for detection of muscle pathology as it has a defined clinical role and might be combined with cTn measurement for differential diagnosis between cardiac and skeletal muscle injury. As such it might be considered by laboratories as part of the “cardiac biomarker panel”. Continued measurement of CK-MB and myoglobin may also be specifically used as part of a cardiac panel. However, it is difficult to justify the retention of myoglobin and CK-MB as “cardiac biomarkers”. The diagnostic sensitivity of myoglobin for early detection is superseded by conventional sensitive cTn (13) with hs-cTn assays offering superior diagnostic sensitivity. It is also difficult to justify continued measurement of CK-MB. The usual justifications given are the detection of reinfarction, although this is incorrect(14), and for monitoring after percutaneous invention (PCI). CK-MB is used as a marker in PCI trials. It is difficult to justify the retention of CK-MB measurement on financial grounds as cTn is diagnostically superior and now of comparable cost. Where information is available, the reason for retention of CK-MB is stated to be a clinician preference. It is to be hoped that recent recommendations from the ESC will see the retirement of CK-MB (15).

Measurement of cTn by a high sensitivity method is strongly encouraged although not mandated in current guidelines but has increasingly been adopted. For some methods (cTnT) this has been largely due to the manufacturer replacing a pre-existing assay by a better version with withdrawal of the previous assay version. Manufacturers of cTnI have progressively introduced a high sensitivity version of that assay in response to the perceived clinical need and commercial pressures. The only exception is Roche diagnostics who already had an hs cTnT assay. Laboratories are increasingly adopting hs cTnI with some manufacturers actively encouraging this transition. Where point of care testing is used, where hs cTn assays are not available, such a transition may be problematic, as was found in this survey, with the only option shifting to a laboratory-based assay.

In accordance with the universal definition, the 99th percentile is now the dominant decision limit for the diagnosis of MI. The main reason will be evolution of cTn assays. The shift from the Roche fourth-generation cTnT assay to the fifth-generation (high sensitivity assay or designated GEN 5 per the FDA in the USA) was accompanied by a significant improvement in assay sensitivity and imprecision. The conventional sensitive cTnI assays previously in use, for the most part, had analytical characteristics that supported the use of the 99th percentile as diagnostic discriminant. The major source of information on the choice of 99th percentile, as well as reference intervals comes from the manufacturers IFU. This trend is even more marked from the previous survey and highlights the need for independent validation of manufacturers claims as has been discussed previously(3;4).

The advantage of high sensitivity assays is earlier diagnosis of AMI and in particular the ability to rule out in appropriately selected patients on a single measurement performed on admission to hospital. The clinical utility of rapid diagnosis based on hs-cTn assays is supported by an evidence-based review (16) by the National Institute of Health and Care Excellence (NICE), recently updated with a review of the most recent literature (17;18). Only a minority of laboratories are using very rapid rule out algorithms, the majority utilising 0-3 hour or 0-6 hour strategies. It is possible that the respondents did not understand the question as phrased or that the diagnostic protocols in use in the emergency department are not agreed with or involve the laboratory. But, the results are in agreement with another survey on utilisation of rapid diagnosis (19) and the experience of a number of the authors on discussing this subject with clinical colleagues. A possible obstacle is that patients with possible unstable angina pectoris (UAP) still need urgent (within days) diagnosis and follow-up. Hospitals implementing the rapid rule-out algorithms for NSTEMI therefore need to establish an out-patient clinical pathway for rapid follow-up of patients ruled-out for NSTEMI who might have UAP.

The survey continued to highlight deficiencies in communication between the laboratory and clinician users. Some laboratories were unaware whether their hospital offered angiography, a fundamental part of modern clinical cardiology and predicated by cTn measurement. Although there is documentation and review of a diagnostic protocol, there is a lack of structured ordering and demand management. This will contribute to the substantial inappropriate over requesting of cTn testing, degrading diagnostic efficiency(20) and therefore wasting healthcare resources.

Study limitations.

The study was largely confined to European laboratories with variable uptake by country with the best response from northern European countries although the project team aimed to achieve as wide a geographical representation as possible. The responses were consistent so the conclusions are likely to be valid, but will have been modulated by individual understanding of the questions as formulated. As English is the language of the EFLM the questionnaire was not translated which might also be a limitation.

In conclusion, the measurement of cTn by high sensitivity methods is now the predominant methodology in use certainly throughout northern Europe. The adoption of the main benefits of these assays, rapid diagnostic algorithms, appears to lag behind adoption of the methodology itself. There is clearly a need for closer interaction between laboratory and clinical staff to achieve the potential benefits.

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Table 1. Comparison of guidelines from the European Society Cardiology from 2011 to 2017.

|  |  |  |
| --- | --- | --- |
|  | ESC guideline 2011 | ESC guideline 2017 |
| First line biomarker | Cardiac troponin | Cardiac troponin |
| Assay | High sensitivity assay | High sensitivity assay |
| Diagnostic cut-off | 99th percentile upper reference limit (URL) of a normal reference population with 10% CV at the URL | Sex-specific URLs |
| Analytical quality | No methodological recommendations (standardization and analytical performance goals) | CV less than 10% mandatory for hs-Tn. Assays with CVs >20% at the 99th percentile should not be used |
| Point of care testing (POCT) | POCT recommended when lab TAT more than 60 min | POCT not recommended as unable to meet high sensitivity performance recommendations |
| Laboratory clinician interaction | No recommendations | “Thus, clinicians must learn about their local assay and should look for reliable information, for example, available on the International Federation of Clinical Chemistry and Laboratory Medicine (IFCC) website (http://www.ifcc.org/ executive-board-and-council/eb-task-forces/task-force-on-clinical-applications-ofcardiac-bio-markers-tf-cb/), when they have questions concerning analytical issues." |

Figure legends

Figure 1 Categories of respondent. Central = University/Teaching hospital or Regional centre/Secondary and Tertiary hospital; District = District General or Secondary Care hospital; Laboratory = Free standing laboratory; Clinic = Primary care or Ambulant care facility; Community = Community hospital (non-urgent care).

Figure 2. Proportion of laboratories using other biomarker as a percentage of total numbers. Upper panel laboratories using aspartate transaminase (AST) lactate dehydrogenase (LD) or hydroxybutyrate dehydrogenase (HBD). Lower pane laboratories using creatine kinase (CK) or its MB isoenzyme (CK-MB) and myoglobin.

Figure 3. Distribution of laboratories currently utilising or changing to a high sensitivity troponin method. Change to hs = change to a high sensitivity method supplied by the same company (supplier); hs cTn change in company = existing user of a high sensitivity method but changing to a different company (supplier); POCT to hs cTn = change from point of care testing to a high sensitivity troponin method; cs cTn instrument change the same company = continues to use a non high sensitivity cTn method from the same company (supplier).

Figure 4. Percentage utilisation of decision limits for acute myocardial infarction. Changes to the guidelines of the European Society of Cardiology (ESC) are indicated.