Accurate Quantification of Cardiovascular Biomarkers in Serum Using Protein Standard Absolute Quantification (PSAQ™) and Selected Reaction Monitoring*

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Introduction of new diagnostic assays in the clinical setting requires an operating pipeline to efficiently translate putative biomarkers into validated biomarkers. Despite the discovery platforms’ capacity to generate well populated lists of candidate biomarkers, very few proteins reach the patient bedside as fully fledged “FDA-approved” biomarkers. This is largely because of divergences between analytical needs and performances of the techniques available for candidate biomarker evaluation (1, 2).

Candidate biomarker evaluation is a major process of the biomarker pipeline, positioned downstream of the biomarker discovery phase and necessary before clinical validation. Candidate evaluation aims to select, among hundreds of putative biomarkers, those of clinical relevance. Evaluation phase combines two steps which respectively consist in: (1) confirming a difference between physiological and pathological concentrations in biofluids (the so-called “qualification phase”) and (2) assessing the specificity of candidate biomarkers (the so-called “verification phase”) (1). Currently, because of its high throughput and high sensitivity, quantitative ELISA is the preferred assay format for studies evaluating biomarkers. However, as most candidates are likely to fail as relevant biomarkers, developing ELISA tests (with high quality antibodies) for all candidates is a financial burden for the diagnostics industry (3).

Thus, there exists an urgent need to develop analytical methods capable of reliable candidate evaluation, at high throughput and reasonable cost. Selected Reaction Monitoring (SRM)1 mass spectrometry combined with stable isotope dilution (SID-SRM) has shown promise as a solution to this

1 The abbreviations used are: CKB, Creatine kinase B chain; CKM, Creatine kinase M chain; CKBB, Creatine kinase BB homodimer; CKMB, Creatine kinase MB heterodimer; CKMM, Creatine kinase MM homodimer; LDH-B, Lactate dehydrogenase B; PCI, Primary percutaneous coronary intervention; SID, Stable isotope dilution; SRM, Selected reaction monitoring; PSAQ, Protein standard absolute quantification; PSAQ™ for accurate biomarker quantification.
Recently, in an effort to demonstrate the potential of SID-SRM for candidate biomarker evaluation, a multilaboratory study was set up to assess its analytical performances and potential transferability (7). Exogenous proteins, seven in all, were added to unfraccionated plasma samples. The spiked samples were analyzed by eight independent laboratories using SRM and isotope-labeled peptides as standards. The results obtained clearly demonstrated the capacity of SID-SRM to specifically and precisely quantify protein biomarkers in plasma. However, the results also revealed that the protein digestion rate was highly variable between laboratories. This variability had a significant effect on peptide recovery and on the accuracy of protein quantification. As suggested by the authors, this type of bias could be avoided if properly folded isotope-labeled protein standards were used as quantification standards (7, 8).

In 2007, we developed the PSAQ™ (Protein Standard Absolute Quantitation) method, which uses full-length isotope-labeled proteins as internal standards for absolute quantitative MS analysis. We demonstrated that, in contrast with peptide standards, adding isotope-labeled proteins before sample digestion enables accurate protein quantification, even for proteins resistant to trypsin digestion (9, 10). In addition, we, and others, have shown that this type of protein standard ("PSAQ standard") also corrects for protein losses that may occur during sample handling prior to trypsin digestion and liquid chromatography (LC)-MS analysis (11–17). This latter feature is a particular advantage for MS analysis of blood biomarkers. Indeed, as plasma/serum are highly complex matrices and display a huge dynamic range, sample prefractionation must be performed to detect low-abundance protein biomarkers (4).

In this study, we have tested a combination of the PSAQ strategy with SRM (PSAQ-SRM) for quantification of cardiovascular biomarkers in serum samples. Selected biomarkers include LDH-B, CKMB, myoglobin, and troponin I. For some of these validated biomarkers, a comparison of PSAQ-SRM data and ELISA results was performed on samples from patients having suffered myocardial infarction.

EXPERIMENTAL PROCEDURES

Biomarkers and Clinical Samples—Human LDH-B, creatine kinase MB heterodimer (CKMB), and myoglobin were purchased from Applichem (Darmstadt, Germany). Human troponin I and healthy human serum were obtained from Sigma-Aldrich (Saint Quentin Fallavier, France). Serum samples from five patients were provided by the Plateforme de Ressources Biologiques (Groupe Hospitalier Henri Mondor, Créteil, France). Five patients undergoing primary percutaneous coronary intervention (PCI) for ST-elevation myocardial infarction were included in the study. The protocol was approved by the hospital's institutional review board and all patients provided written informed consent for participation. Blood samples were taken at hospital admission and at regular intervals following PCI. Samples were collected in nontreated tubes (BD Biosciences, le Pont de Claix, France). All blood samples were centrifuged at 2200 × g for 10 min to obtain serum supernatants. These were immediately aliquoted and frozen at −80 °C. Additional blood samples were analyzed at the clinical chemistry laboratory for standard cardiac biomarker evaluation (Total CK activity, troponin I, and myoglobin) as described below. In this study, only serum samples collected at hospital admission and at 3 or 8 days after PCI were analyzed using the PSAQ-SRM approach.

Production of Full-length Stable-Isotope-Labeled Proteins (PSAQ Standards)—PSAQ standards were synthesized as previously described (9). Briefly, LDH-B, creatine kinase B chain (CKB), creatine kinase M chain (CKM), myoglobin, and troponin I genes were amplified by PCR using a cardiac cDNA library as template (Biochain Institute, Hayward, CA) (see supplemental Table S1 for primer sequences and PCR reaction conditions). Genes were cloned into the pIVEX 2.4d expression vector (5 Prime, Hamburg, Germany) using the In-Fusion™ PCR cloning system (Clontech, Saint Germain en Laye, France). The pIVEX 2.4d vector provides a N-terminal hexahistidine purification tag. Plasmids were cloned into XL1-Blue cells (Agilent Technologies, Massy, France), purified and sequenced (Cogenics, Meylan, France). Cell-free protein expression and isotope-labeling was performed using the RTS 500 Protemaster E. coli HY kit (5 Prime) in the presence of [15N] l-lysine and [13C6, 15N] l-arginine (Eurolab, Saint-Aubin, France). PSAQ standards were purified on a nickel affinity column (Qi Sepharose 6 Fast Flow resin, GE Healthcare, Orsay, France) using an imidazole gradient. PSAQ standards were checked for purity on SDS-PAGE using Coomassie staining (>95% purity). N-terminal hexahistidine purification tags were not removed as they are not expected to significantly modify PSAQ biochemical properties. Isotope-labeled proteins were quantified by amino acid analysis (MScan SA, Plan les Ouates, Switzerland). Isotope incorporation was verified by LC-MS and LC-SRM analysis and was found to be greater than 99% (see supplemental Fig. S1).

Depletion of Serum Samples—Serum samples (14 μl) from healthy donors and patients were spiked with defined amounts of PSAQ standards. To set up a titration experiment, samples from healthy donors were spiked with defined amounts of exogenous, unlabeled LDH-B, CKMB, myoglobin, or troponin I (see supplemental LC-SRM data). Serum samples were depleted of the six most abundant proteins using the Multiple Affinity Removal Spin cartrige (MARS 6) (Agilent Technologies) according to the manufacturer's instructions. The flow-through was concentrated to 15 μl using a 5000 Da cutoff ultrafiltration device (Vivascience, Hannover, Germany). Laemml buffer (10 μl) was added before SDS-PAGE analysis.

Troponin I Antibody Biotinylation—Troponin I antibody (Santa Cruz Biotechnology, Heidelberg, Germany) was dialyzed against phosphate-buffered saline (PBS) buffer. The antibody was then incubated with NHS-PEG2-Biotin reagent (Pierce/Thermo Fischer Scientific, Bребиёres, France) for 3 h at room temperature (NHS-PEG2-Biotin/antibody ratio: 20/1). After biotinylation, the antibody was once again dialyzed in PBS buffer. Biotinylation was checked by dot-blot analysis with neutravidin-HRP on a polyvinylidene difluoride membrane.

Immunoenrichment of Serum Samples—Before immunoenrichment, serum samples (1 ml) were spiked with defined quantities of troponin I PSAQ standard. To generate the titration curve, a range of concentrations of exogenous unlabelled troponin I was added to healthy donor serum samples (see supplemental LC-SRM data). Biotinylated antibody (1 mg) was added to the samples before overnight incubation at 4 °C on a rotating wheel. Dynabeads M-280 Streptavi
were concentrated for 3 min on a 300-nm x 150-mm C18 column (Dionex) at a flow rate of 300 nl/min and using a 60-min gradient from 10% B to 40% B in 30 min and from 40% to 90% in 5 min. Data were acquired in a positive ion mode with an ion spray voltage of 2200 Volts, curtain gas at 12 p.s.i and an interface heater temperature set at 4.9 s (multiplex detection in depleted serum samples) or 2.1 s (troponin I detection in immunoenriched serum samples). LC-SRM Data Analysis — Data analysis was performed using MultiQuant software (version 1.1 Applied Biosystems/MDS Sciex). Unlabeled/labeled peak area ratios were calculated for each SRM transition after careful verification of coelution profiles. Ratios obtained from the different SRM transitions were used to calculate the corresponding average peptide ratio. Then, ratios obtained for the different proteotypic peptides were combined to calculate the protein ratio and determine biomarker concentration in serum (see supplemental LC-SRM data and supplemental Figs. S2 to S5). From the 787 calculated peak area ratios, 11 (1.4%) outlier values were excluded from analysis (see supplemental LC-SRM data). Most of these outliers were related to matrix interferences impairing labeled or unlabeled peptide transition signals.

LLOQ Determination — LLOQ corresponds to the lowest concentration of an analyte that can be determined with acceptable precision and accuracy. Various approaches for determining LLOQ can be used, including signal-to-noise analysis, statistical analysis based on blank sample variance determination (19) and least-squares linear regression analysis (20). In this study, blank samples were not available (non-spiked serum samples contained endogenous levels of biomarkers). Consequently, LLOQ was determined according to the Food and Drug Administration criteria described in the guidelines for bioanalytical method validation (www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances). LLOQ was established as the lowest concentration of the titration curve that was measured with precision (CV) inferior to 20% and accuracy comprised between 80 and 120%.

ELISA and Enzymatic Assays — Myoglobin concentration in patient serum samples was determined using the Human Myoglobin ELISA Kit (Alpco Diagnostics, Salem, NH) according to the manufacturer’s instructions. To investigate troponin I concentration in patients’ samples, the Access AccuTnI Troponin I Assay (Beckman Coulter, Roissy, France), which is not a highly sensitive troponin I assay, was used as described by the manufacturer. Total CK activity was measured by spectrophotometry using a COBAS system (Hoffman La Roche, Basel, Switzerland) at the local clinical chemistry laboratory. All assay parameters are presented in supplemental Table S2.

RESULTS

Selection of Proteotypic Peptides and SRM Transitions — Once expressed and purified, each PSAQ standard was submitted to SDS-PAGE, in-gel digestion with trypsin and LC-MS/MS analysis (see supplemental LC-MS/MS data). This allowed us to experimentally determine the signature peptides (so-called “proteotypic” peptides) to be monitored for each cardiovascular biomarker targeted. Proteotypic peptides were chosen for their sequence uniqueness, verified by BLAST search, and ease of detection with LC-MS/MS. Peptides containing methionine and cysteine residues were not excluded from the proteotypic peptide panel. Indeed, our group has recently demonstrated that sulfur-containing peptides can be used for quantification if the redox status of the target protein and its PSAQ analog are “equalized” by H2O2 treatment before trypsin digestion (article under preparation, see also supplemental Figs. S2 and S5 which show titration curves obtained with such modified peptides). Based on LC-MS/MS data analysis, a list of putative SRM transitions was estab-
PSAQ™ for Accurate Biomarker Quantification

lished, which included three to six SRM transitions per proteotypic peptide. Then, using prefractionated serum matrix spiked with PSAQ standards and digested with trypsin, we experimentally selected the "best" SRM transitions, i.e. those effectively detected in the matrix and presenting no signs of interference. In the final SRM methods used for titration curves and patient samples, only the "best" SRM transitions (two to four per proteotypic peptide) were retained (Table I). Importantly, for each selected SRM transition, 2 precursor/fragment ions pairs were actually monitored: one for the labeled form of the peptide and one for its unlabeled form. Hence, LDH-B was characterized by four proteotypic peptides. We also monitored peptide VIGSGCNLDSAR, which is shared between the three LDH isoforms (LDH-A, LDH-B and LDH-C). This additional peptide allowed us to measure total LDH content in serum samples. Quantification of myoglobin, which is a small protein (17 kDa), was based on only one proteotypic peptide. Troponin I levels were evaluated using three proteotypic peptides. For CKMB, heterodimer concentrations could not be specifically investigated. However, we could examine the levels of CKB chains (using two proteotypic peptides) and CKM chains (using three proteotypic peptides) independently. Notably, in serum samples, CKB and CKM chains, in addition to making up the CKMB heterodimer, may also form CKMM and CKBB homodimers.

Quantification of Cardiovascular Biomarkers in Depleted Serum Samples—A rapid sample preparation workflow, combining depletion of the six most abundant serum proteins with short SDS-PAGE migration, was developed (Fig. 1A). For each cardiovascular biomarker, a four-point titration experiment was performed by spiking 14 μl serum samples with defined amounts of both unlabeled protein (LDH-B, CKMB, myoglobin, or troponin I) and corresponding PSAQ standard. Zero samples were also constituted. The spiked LDH-B, CKMB, myoglobin, and troponin I quantities were calculated to generate titration curves covering physiological levels to highest pathological concentrations. All titration points were performed in full-process triplicates (see supplemental LC-SRM data). For CKMB heterodimer, we investigated the levels of CKM chains and CKB chains by simultaneously spiking CKM and CKB PSAQ standards. After in-gel trypsin digestion, the mix was analyzed by LC-SRM which revealed a 2/1 CKM/CKB stoichiometry. This indicates that the commercial CKMB heterodimer contained excess CKM chains, possibly originating from a contaminating CKMM homodimeric form. Commercial CKMB is purified from myocardium containing excess CKMM (75%), thus contamination of CKMB preparations would not be surprising. The underestimation of CKMB in serum may stem from CKB and CKM PSAQ standards and CKMB heterodimer behaving differently during prefractionation and/or digestion. This hypothesis was supported by native-PAGE analysis of CKB and CKM PSAQ standards. CKB PSAQ standard was found to be monomeric whereas CKM PSAQ standard was structured as a CKMM homodimeric (data not shown).

Quantification of Troponin I in Immunoenriched Serum Samples—Generally, LC-SRM detection of serum proteins present below 100 ng/ml is difficult when using depletion as unique prefractionation method (4). However, we hypothesized that the combination of depletion and SDS-PAGE might allow troponin I detection, particularly for samples with concentrations in the range of 100 ng/ml. Indeed, myoglobin could be detected at just 40 ng/ml using this procedure. However, troponin I could not be detected, even at the highest concentration tested (272.5 ng/ml). Most likely, this is because of its interaction with abundant proteins retained on the depletion cartridge (19). Therefore, a prefractionation method based on immunocapture coupled to SDS-PAGE was developed to validate the PSAQ-SRM approach for this specific biomarker (Fig. 1A). To perform titration experiments, healthy serum samples (1 ml) were spiked with defined amounts of troponin I and its corresponding PSAQ standard. An immunoenrichment protocol was optimized using a bio-
### Table I

Peptide sequences and SRM transitions. Peptides from PSAQ standards are mentioned with the C-terminal isotope-labelled amino acid in bold. Methionine dioxidation (ox2) or cystein trioxidation (ox3) modification states are indicated.

| Biomarker | UniProt accession number | Peptide sequence | SRM transitions | Collision energy (Volts) |
|-----------|--------------------------|-----------------|-----------------|-------------------------|
| LDH-B     | P07195                   | IVVVTAGVR       | 457.3 503.3     | 27.9                    |
|           |                          |                 | 457.3 602.4     | 27.9                    |
|           |                          |                 | 457.3 701.4     | 27.9                    |
|           |                          | IVVVTAGVR       | 462.3 513.3     | 27.9                    |
|           |                          |                 | 462.3 612.4     | 27.9                    |
|           |                          |                 | 462.3 711.4     | 27.9                    |
|           |                          | GLTSVINQK       | 480.3 502.3     | 29.0                    |
|           |                          |                 | 480.3 601.4     | 29.0                    |
|           |                          |                 | 480.3 688.4     | 29.0                    |
|           |                          | GLTSVINQK       | 484.3 510.3     | 29.0                    |
|           |                          |                 | 484.3 609.4     | 29.0                    |
|           |                          |                 | 484.3 696.4     | 29.0                    |
|           |                          | SLADELALVDVLEDK | 815.4 504.3     | 45.8                    |
|           |                          |                 | 815.4 718.4     | 45.8                    |
|           |                          |                 | 815.4 1001.6    | 45.8                    |
| Total LDH:| P00338                   | VIGSGC(ox3)NLDSAR| 620.3 333.2     | 36.0                    |
| LDH-A, LDH-B and LDH-C| P07195 |                | 620.3 448.2     | 36.0                    |
|           | P07864                   |                 | 620.3 675.3     | 36.0                    |
|           |                          | VIGSGC(ox3)NLDSAR| 625.3 343.2     | 36.0                    |
| Creatine kinase B | P12277 |            | 625.3 458.2     | 36.0                    |
|           |                          |                 | 625.3 685.3     | 36.0                    |
| Creatine kinase M | P06732 |            | 616.8 742.4     | 35.8                    |
|           |                          | DLFDPIIEDR      | 616.8 857.4     | 35.8                    |
|           |                          |                 | 621.8 752.4     | 35.8                    |
|           |                          | VLTPELYAELR     | 621.8 867.4     | 35.8                    |
|           |                          |                 | 652.4 990.5     | 37.6                    |
|           |                          | VLTPELYAELR     | 652.4 1091.6    | 37.6                    |
|           |                          |                 | 657.4 1000.5    | 37.6                    |
|           |                          |                 | 657.4 1101.6    | 37.6                    |
| Myoglobin | P02144                   | FEEILTR         | 454.3 502.3     | 27.7                    |
|           |                          |                 | 454.3 631.4     | 27.7                    |
|           |                          | FEEILTR         | 459.3 512.3     | 27.7                    |
|           |                          |                 | 459.3 641.4     | 27.7                    |
|           |                          | ELFDPIISDR      | 602.8 700.4     | 35.1                    |
|           |                          |                 | 602.8 962.5     | 35.1                    |
|           |                          | ELFDPIISDR      | 607.8 710.4     | 35.1                    |
|           |                          |                 | 607.8 972.5     | 35.1                    |
|           |                          | GGDDLDPNYVLSSR  | 754.4 935.5     | 42.7                    |
|           |                          |                 | 754.4 1050.5    | 42.7                    |
| Troponin I | P19429 |              | 545.0 553.3     | 31.2                    |
|           |                          | VEADPGHGWNSLVR | 545.0 702.9     | 31.2                    |
|           |                          |                 | 548.3 558.3     | 31.2                    |
|           |                          |                 | 548.3 707.9     | 31.2                    |
|           |                          | TLLQLIAK        | 450.3 685.5     | 27.5                    |
|           |                          |                 | 450.3 572.4     | 27.5                    |
|           |                          | TLLQLIAK        | 454.3 693.5     | 27.5                    |
|           |                          |                 | 454.3 580.4     | 27.5                    |
|           |                          | NITEIADLTQK     | 623.3 675.4     | 36.2                    |
tinylated antitroponin I antibody and streptavidin coated beads. Because of the strong biotin-streptavidin interaction, antitroponin I antibody was retained on streptavidin beads during elution, further improving sample decomplexification. However, as Laemmli buffer was used to elute troponin I, direct trypsin digestion was not possible. Therefore, samples were submitted to “stacking” SDS-PAGE before in-gel digestion (supplemental Fig. S5A). After trypsin digestion and peptide extraction samples were analyzed using LC-SRM analysis. With this serum prefractionation method, troponin I could be detected at 500 pg/ml of serum, which is slightly higher than physiological concentrations (~350 pg/ml). However, sensitive and accurate troponin I quantification was possible over the pathological concentration range (5.5 to 272.5 ng/ml) (Fig. 2D, supplemental Fig. S5). PSAQ-SRM assay analytical performances after troponin I immunoenrichment are presented in Table II.

Interestingly, the troponin I proteotypic peptide ISADAM(ox2)M(ox2)QALLGAR includes serine 149 which was previously reported to be phosphorylatable (21). In this study, all quantification results obtained from this peptide were consistent with those obtained from the two other proteotypic peptides (see supplemental Fig. S5 and supplemental LC-SRM data). Therefore, we can hypothesize that: (1) serine 149 was primarily not phosphorylated or (2) serine 149 phosphorylation was removed as troponin I was released in blood flow.

**Multiplexed PSAQ-SRM Analysis of Clinical Samples and Correlation with ELISA/Enzymatic Assay Results**—Once the prefractionation and quantification processes had been characterized for each biomarker, multiplexed LC-SRM detection was assessed on healthy donor serum samples. After spiking the samples with defined quantities of PSAQ standards, prefractionation and SDS-PAGE, gel bands were cut, pooled and proteins were digested with trypsin (Fig. 1A). LDH-B, myoglobin, CKM and CKB could be simultaneously detected in depleted serum samples without difficulty and without time-scheduled acquisition (Fig. 1B). We then applied this multiplex PSAQ-SRM method to the analysis of serum samples from patients with myocardial infarction. Serum samples from five patients were collected at two time-points: (1) immediately after hospital admission and (2) 3 to 8 days after PCI. All samples were analyzed using the PSAQ-SRM method after depletion (14 μl) or troponin I immunoenrichment (1 ml). LDH-B, myoglobin, CKM and CKB chains and troponin I were all detected in these samples (Table III, supplemental Figs. S6 and S7). For each patient, biomarker concentration changes between the two collection time-points were consistent with blood-release kinetics i.e. an early increase in troponin I, myoglobin, and CKMB (day 0) and a delayed LDH-B augmentation (days 3 or 8). As myoglobin, CKMB and troponin I are routinely used clinically to confirm myocardial injury, correlation between ELISA or enzymatic assays and the PSAQ-SRM approach could be assessed for these patient samples (Table III and Fig. 3). Total CK enzyme activity and CKB and CKM concentrations estimated by PSAQ-SRM correlated well, with a \( R^2 \) value of 0.89 (Fig. 3A). Excellent correlation between ELISA and PSAQ-SRM results \( R^2 = 0.95 \) were observed for myoglobin. Surprisingly however, the slope of the correlation curve was equal to 5.29, suggesting that PSAQ-SRM systematically quantifies five times more myoglobin than ELISA (Fig. 3B). Following this result, we compared the commercial myoglobin ELISA standard directly to our PSAQ standard by mixing them in equal amounts. After trypsin digestion and LC-SRM analysis of the 1/1 mix, the ELISA standard was found to be 5.39 times less concentrated than the PSAQ standard (see supplemental Fig. S8). From these results, we can surmise that either the concentration of the myoglobin ELISA standard was overestimated by the supplier or that it had been degraded during storage. For troponin I quantification, a correlation coefficient \( R^2 \) of 0.79 was found between ELISA and PSAQ-SRM with a correlation curve slope of 1.41 (Fig. 3C). Possibly, this lesser correlation originates from troponin I interactions with serum proteins. By masking epitopes, such interactions could differentially influence ELISA detection (based on two monoclonal antibodies) and PSAQ-SRM immunocapture (based on one monoclonal antibody).

**DISCUSSION**

In 2006, Rifai and co-workers described the biomarker pipeline and the promise MS held for biomarker research (1). In particular, they highlighted how SID-SRM analysis might solve the technological hurdle of biomarker evaluation. However, application of SID-SRM as part of biomarker develop-
ment requires key analytical performances to be attained, including specificity, sensitivity and confident quantification. In addition, to surpass ELISA it is crucial to offer multiplexing capabilities and antibody-free prefractionation. Recently, an interlaboratory study assessed the analytical features of a multiplexed SID-SRM assay and demonstrated the accelerated throughput and transferability of this type of analysis. This study, however, highlighted quantification accuracy as a limitation of the method, particularly when sample prefractionation was necessary (7, 8). The goal of our study was to demonstrate that the use of full-length isotopically-labeled proteins used as quantification standards (PSAQ standards) could significantly advance the performances of a SRM-based biomarker evaluation platform.

LDH-B, CKMB, myoglobin, and troponin I were chosen as model biomarkers to assess the performances of the PSAQ-
SRM method. Troponin I, CKMB, and myoglobin are currently used in hospital laboratories to rapidly confirm myocardial injury. Measurement of LDH-B levels was abandoned because its release in the bloodstream is delayed, reaching its maximal concentration 72 h after myocardial infarction. However, to evaluate the PSAQ-SRM method, the combination of these four biomarkers was of particular interest. First because they belong to different concentration classes, with LDH-B being the most abundant (µg/ml of serum) and troponin I requiring very sensitive assays (below 1 ng/ml in serum). Second, ELISA or enzymatic tests are available for troponin I, myoglobin and CKMB, making a comparison between ELISA and PSAQ-SRM assays possible.

Samples were fractionated using a decomplexification method based on the depletion of abundant proteins combined with SDS-PAGE. Using this straightforward sampling, only 14 µl of serum were necessary to simultaneously quantify 3 biomarkers at their physiological levels. Likely because of the high ion-current potential of its proteotypic peptide (22), myoglobin could be quantified down to 40 ng/ml (equivalent to 33 femtomoles in 14 µl). For troponin I quantification, however, this type of fractionation, applied to tiny volumes of serum, was not sufficient. Not only is troponin I a low-abundance biomarker, but it has also been shown to interact with abundant proteins retained by serum depletion devices (19). Therefore, we developed an immunocapture approach to

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**Fig. 2. Titration of cardiovascular biomarkers in serum samples.**

A, LDH-B titration curve. This curve was generated using four proteotypic peptides. Titration curves for each proteotypic peptide and related SRM transitions are presented in supplemental Fig. S2.

B, Myoglobin titration curve. These results were obtained using one proteotypic peptide. Two SRM transitions were monitored and the corresponding data are presented in supplemental Fig. S3.

C, CKM chain, CKB chain and CKMB titration curves. CKMB dimer was artificially spiked into samples. CKB and CKM chain concentrations were estimated independently (from 2 and 3 proteotypic peptides, respectively). The CKM+CKB titration curve represents a combination of the CKB and CKM titration curves. Further details on the different proteotypic peptides monitored and SRM transitions considered are available in supplemental Fig. S4.

D, Troponin I titration curve based on three proteotypic peptides. Titration curves for each proteotypic peptide and related SRM transitions are presented in supplemental Fig. S5. For all titration curves, error bars represent standard deviations obtained from 3 full-process replicates.
Measurement of this protein might also have been possible using depletion on larger serum volumes, combined with mild-detergents to improve elution from the depletion cartridge and by replacing SDS-PAGE with chromatographic methods. Indeed, Keshishian and coworkers have recently described an antibody-free sampling approach involving protein depletion, trypsin digestion and SCX peptide separation. Unlike PSAX-SRM standards, possibly because the standards did not exactly mimic the biochemical behavior of the CKMB heterodimer, quantification was less accurate for this protein than for the other proteins studied. Nevertheless, in this study, we also tried to quantitatively detect troponin T down to 5 ng/ml in 50 µl plasma samples. Another recent study detected 4 ng/ml of PSA in 100 µl serum samples using albumin depletion, trypsin digestion, SPE peptide separation, and LC-SRM analysis (23). Given that most proteins of clinical interest are present in the ng/ml range in serum or plasma, matrix decomplexification and compensate for digestion variability. Our results demonstrate that accurate and precise biomarker quantification can be achieved by spiking isotopically labeled proteins (PSAQ standards) into serum samples early in the sample preparation process. Provided that PSAQ standards behave exactly like the target proteins, quantification is accurate, even in cases where recovery may have been affected by extensive fractionation (depletion or immunocapture, and SDS-PAGE). Similarly, PSAQ standards also compensate for variable digestion yields (17). In this study, we also tried to quantify a protein heterodimer, CKMB, using both CKB and CKM standards. Possibly because the standards did not exactly mimic the biochemical behavior of the CKMB heterodimer, quantification was less accurate for this protein complex than for the other proteins studied. Nevertheless, in patient serum samples, increases in CKM and CKB chains measured by PSAQ-SRM were highly consistent with total CK activity results. Therefore, it is mandatory to successfully detect biomarker candidates (4). In this context, the use of isotope-labeled peptide standards, added at late stages of the analytical workflow, may not provide optimal quantification accuracy and reproducibility (7–9). Ideally, quantification standards should correct for losses occurring during sample decomplexification and compensate for digestion variability. Our accuracy comprised between 80 and 120%. * CKMB LLOQ could not be determined as quantification accuracy was lower than 80% (65%).

### Table II

| Biomarker        | Sample prefractonation method | Serum volume | LLOQ* (ng/ml) | Range of tested concentrations (ng/ml) | Linearity (R²) | Accuracy (slope value) | Precision at LLOQ (CV in %) |
|------------------|-------------------------------|--------------|---------------|---------------------------------------|----------------|------------------------|-----------------------------|
| LDH-B            | Depletion + SDS-PAGE           | 14 µl        | 510           | 510 (endogenous) → 5500               | 0.99           | 1.18                   | 8                           |
| CKB chains       | Depletion + SDS-PAGE           | 14 µl        | ND            | 20 (endogenous) → 1125                | 0.99           | 0.20, 0.65 (CKMB)      | ND*                         |
| CKM chains       | Depletion + SDS-PAGE           | 14 µl        | ND            | 260 (endogenous) → 1125               | 1              | 0.45                   | ND*                         |
| Myoglobin        | Depletion + SDS-PAGE           | 14 µl        | 500           | 40 (endogenous) → 1000                | 1              | 1.14                   | 8                           |
| Troponin I       | Immuno-enrichment + SDS-PAGE   | 0.5 to 1 ml  | 5.5           | 0.29 (endogenous) → 272.5             | 1              | 0.95                   | 3                           |

* LLOQ was established as the lowest concentration of the average titration curve that was measured with precision (CV) inferior to 20% and accuracy comprised between 80 and 120%. * CKMB LLOQ could not be determined as quantification accuracy was lower than 80% (65%).

### Table III

PSAQ-SRM and ELISA quantification of LDH-B, CKB, CKM, myoglobin and troponin I in patient serum. Biomarker concentrations were determined using at least 1 proteotypic peptide and 2 SRM transitions except for values indicated with:

* 1 proteotypic peptide detected with 1 SRM transition

** 2 proteotypic peptides, each detected with 1 SRM transition

See supplemental Figure 7 which shows the corresponding SRM transition chromatograms.
ical samples has to be established. For this, a method that can accurately and precisely quantify biomarkers over both physiological and pathological concentration ranges would be of particular value. In addition to meeting this criterion, as PSAQ standards correct for variable recovery due to differences in sample handling and digestion efficiency, their use should significantly improve interassay and interlaboratory reproducibility.

Because of its selectivity, SRM analysis provides a high detection specificity (5). However, when working with highly complex samples such as serum, this selectivity might not be sufficient to avoid matrix interferences (24). In this context, the use of isotope dilution standards, such as labeled peptides or proteins, which coelute with the target improve the specificity of the analysis (12, 25). Finally, we have shown that PSAQ standards offer the largest coverage for quantification even making it possible to take cystein- and methionin-containing peptides into account after protein oxidation (18).

In our study, multiplexing was limited to the detection of four biomarkers in depleted serum samples. However, technological advances, including scheduled SRM (26, 27) and iSRM (28), have made it possible to monitor hundreds of proteins in a single experiment. Recently, Kuzyk and coworkers used isotope-labeled peptide standards and LC-SRM to quantify 45 proteins in plasma. Thirty-one of these proteins were potential biomarkers of cardiovascular diseases (25). The most frequent question about implementation of the PSAQ method concerns the availability of PSAQ standards. In this study, the cell-free expression and isotolabeling of LDH-B, CKB, CKM, myoglobin, and troponin I PSAQ standards were optimized in less than 2 months. These biomarkers are not post-translationally modified, which facilitates their expression in bacterial lysates. However, phosphorylated or glycosylated PSAQ standards can be produced using specific production systems (11, 13). Certainly, the throughput of PSAQ standard production could be further enhanced by avaling of cDNA libraries specifically developed for protein expression (29, 30).

In conclusion, this study clearly demonstrates the relevance of using isotope-labeled protein standards for multiplex and reliable quantification of biomarkers in prefractionated clinical samples. We are currently concentrating our efforts on the generation of isotope-labeled protein libraries to increase the availability of PSAQ standards and widen the use of the PSAQ-SRM analytical strategy. Applying PSAQ-SRM as part of patient samples and correlation with enzymatic or ELISA assays. A, Correlation between total CK enzymatic activity and CKMB concentration, as determined by PSAQ-SRM, in clinical serum samples after depletion of abundant proteins (five patients with myocardial infarction, two collection time-points). B, Correlation between ELISA and PSAQ-SRM results for quantification of myoglobin in clinical serum samples after depletion of abundant proteins (five patients with myocardial infarction, two collection time-points). C, Correlation between results for quantification of troponin I in clinical serum samples by ELISA or in immunoenriched samples by PSAQ-SRM (five patients with myocardial infarction, two collection time-points).
of the biomarker development pipeline should help bridge the gap between biomarker discovery and clinical applications.

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