Identification of patients with triple antiphospholipid antibody positivity is platform and method independent

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KEY WORDS
antiphospholipid antibodies, antiphospholipid syndrome, chemiluminescent immunoassay, enzyme-linked immunosorbent assay, triple positivity

ABSTRACT
INTRODUCTION The risk of clinical complications in antiphospholipid syndrome (APS) increases when a patient is positive for all 3 types of antiphospholipid (aPL) antibodies. However, there is a considerable disagreement between various platforms for aCL and anti-β2-glycoprotein I (anti-β2GPI) measurement, which leads to discrepancies between these platforms in assessing aPL antibody positivity.

OBJECTIVES The aim of this retrospective cross-sectional study was to assess whether 2 different platforms, the QUANTA Lite® enzyme-linked immunosorbent assay and the QUANTA Flash® chemiluminescent immunoassay, identify the same subjects as triple positive in a group of patients with APS and comorbid autoimmune diseases.

PATIENTS AND METHODS The study included 220 patients with systemic autoimmune diseases (74 with primary APS; 47 with secondary APS; and 99 with systemic lupus erythematosus without APS). All patients were tested for IgG and IgM aCL and anti-β2GPI antibodies using both platforms.

RESULTS The agreement between the positive results for individual antibodies obtained using both platforms was not full, ranging from 81.8% to 90.9% in a pair-wise comparison. However, the number of patients with triple aPL antibody positivity was similar (80 by QUANTA Lite® and 86 by QUANTA Flash®); the agreement between the 2 platforms for the identification of patients with triple antibody positivity was 95.5% (Cohen’s kappa coefficient = 0.90). This resulted in a similar risk for APS-related clinical complications: an odds ratio of 24.9 for QUANTA Lite® and of 24.7 for QUANTA Flash®.

CONCLUSIONS Our results confirm a strong association between triple aPL antibody positivity and APS and indicate that the identification of patients with triple antibody positivity is platform independent. When aPL antibody profiles are assessed, the agreement between various methods is much higher than that for individual antibodies.

INTRODUCTION The international consensus statement on the classification criteria for definite antiphospholipid syndrome (APS) specifies anticardiolipin (aCL) and anti-β2-glycoprotein I (anti-β2GPI) antibodies of the immunoglobulin (Ig) G or IgM isotype in a medium or high titer as one of the laboratory criteria.¹ However, the agreement between various methods for aCL and anti-β2GPI antibody measurement is known to be suboptimal owing to discrepancies in the cutoff value, calibration, and technology.²⁻⁵ Consequently, the method that is used to assess the antiphospholipid antibody (aPL) status of a patient has a major impact on the consistency in diagnosis across different institutions.

The consensus statement (published in 2006) advises investigators to classify patients with APS into groups based on the number of laboratory criteria present: I) more than 1 laboratory criterion present (any combination); IIa) lupus anticoagulant (LAC) present alone; IIb) aCL antibody present alone; and IIc) anti-β2GPI antibody present alone.¹ Since then, a large number of publications have demonstrated that aPL antibody...
profiling is useful not only as a diagnostic tool but also for assessing thrombotic risk.\textsuperscript{5,6}

The risk for the first thrombosis in asymptomatic aPL antibody carriers was strongly associated with a triple positive antibody profile in several independent studies, where triple positivity was defined as a positive result in 3 different aPL antibody assays.\textsuperscript{7–9} Moreover, the cumulative incidence of thromboembolic events during a 10-year follow-up of patients with symptomatic APS was 44.2% in those with 3 positive aPL antibody results.\textsuperscript{10} Pregnancy failure during conventional therapies was also independently associated with triple aPL antibody positivity.\textsuperscript{11} It has been demonstrated that when initially positive aPL testing was repeated after 12 weeks according to the recommendations of the classification criteria, 98% of the subjects with triple antibody positivity had their aPL antibody status confirmed, in contrast with only 40% of single-positive subjects.\textsuperscript{12} These results suggest that triple positivity allows to diagnose APS with high reliability.

Given the importance of the triple positive aPL antibody status, our goal was to assess whether 2 different measurement methods, an enzyme-linked immunosorbent assay (ELISA) and a novel microparticle chemiluminescent immunoassay (CIA), identify the same subjects with APS and comorbid autoimmune diseases as triple positive.

**PATIENTS AND METHODS**  
**Patients** This was a retrospective cross-sectional study including 220 patients. Samples were collected at the Immunological Outpatient Clinic, Department of Internal Medicine, Jagiellonian University Medical College (Kraków, Poland) from patients referred to the clinic with a suspicion of systemic lupus erythematosus (SLE) or APS (or both). Of 220 patients, 74 had the diagnosis of primary APS; 47, of secondary APS; and 99, of SLE without APS. In the group of patients with secondary APS, 46 patients were diagnosed with SLE and 1 with mixed connective tissue disease. Data on the history of venous thrombosis, arterial thrombosis, and obstetric complications were available for all subjects. Thrombotic complications were confirmed in 111 patients (81 episodes of venous thrombosis and 43 episodes of arterial thrombosis), and pregnancy morbidity — in 30 of 97 women ever pregnant. A diagnosis of APS was made on the basis of the updated APS criteria.\textsuperscript{1} SLE was diagnosed according to the updated criteria of the American College of Rheumatology, whenever at least 4 criteria were fulfilled.\textsuperscript{13,14} The study was approved by a local ethics committee, and informed consent was obtained from all patients according to the Declaration of Helsinki.

**Sample preparation** Blood samples for the measurement of aCL and anti-β\textsubscript{2}GPI antibodies were collected in serum separation tubes and spun for 10 minutes at 3500 rpm at room temperature within 2 hours from sampling. Serum samples were then stored at −80°C until further analysis. For the detection of LAC, blood was drawn in sodium citrate tubes (3.2% [0.109 mol/l]; 1 part sodium citrate to 9 parts venous blood). Platelet-poor plasma was prepared by double centrifugation within 2 hours (first: 10 min/3500 rpm and second: 10 min/14000 rpm) and stored at −80°C for further analysis.

**Measurement of autoantibodies** All samples were tested for aCL and anti-β\textsubscript{2}GPI antibodies of IgG and IgM isotypes with QUANTA Flash\textsuperscript{®} CIA and QUANTA Lite\textsuperscript{®} ELISA (Inova Diagnostics Inc., San Diego, California, United States) according to the manufacturer’s instructions. The presence of LAC was also tested.

QUANTA Flash\textsuperscript{®} aCL and anti-β\textsubscript{2}GPI are semi-quantitative immunoassays that are run on the fully automated BIO-FLASH\textsuperscript{®} instrument (Biokit S.A., Barcelona, Spain). Results are expressed in (arbitrary) chemiluminescent units.

QUANTA Lite\textsuperscript{®} aCL and anti-β\textsubscript{2}GPI are standard ELISAs for the semiquantitative determination of aCL and anti-β\textsubscript{2}GPI antibodies. The QUANTA Lite\textsuperscript{®} aCL assays report results in GPL and MFL units, and the QUANTA Lite\textsuperscript{®} β\textsubscript{2}GPI assays — in (arbitrary) standard IgG and IgM units.

For the purposes of this study and in accordance with the international guidelines on aCL and anti-β\textsubscript{2}GPI antibody testing, only the values above the 99th percentile of the results obtained from 120 sex- and age-matched reference subjects were defined as positive.

LAC was determined in a 3-step procedure according to the guidelines of the International Society on Thrombosis and Haemostasis.\textsuperscript{15} Diluted Russell’s viper venom time (LA1-screen; Siemens, Erlangen, Germany) and a sensitive activated partial thromboplastin time (Diagnostica Stago, Gennevilliers, France) were used as screening tests, and LA2-confirm (Siemens) and Staclot LA (Diagnostica Stago)—as confirmatory tests. A reference value was established using the 99th percentile value obtained in the healthy population. A triple aPL antibody positive profile was defined as positivity for LAC and for aCL and anti-β\textsubscript{2}GPI antibodies of the same isotype by the same method.

**Single versus triple antibody positivity** Single aPL antibody positivity was defined as positivity for one of the following: LAC, aCL antibody, or anti-β\textsubscript{2}GPI antibody. Triple antibody positivity was defined as a positive result in 3 different aPL antibody assays, namely, LAC and aCL and anti-β\textsubscript{2}GPI antibody of the same isotype.\textsuperscript{7,8}

**Statistical analysis** A statistical analysis was performed using the Analyse-It Software (Version 4.00.1, Leeds, United Kingdom). The Spearman’s correlation and Cohen’s kappa coefficient tests were performed to assess the quantitative correlation between unit values and qualitative agreement between portions, and P values of less than
Agreement between QUANTA Lite® and QUANTA Flash® antibody results was good, ranging from 86.8% to 90.9% for IgG assays and from 81.8% to 90.9% for IgM assays, with Cohen’s kappa coefficients between 0.56 to 0.81, suggesting a moderate to substantial agreement. Most discrepant results between the 2 platforms were identified for the aCL IgM and anti-β2GPI IgG assays (TABLE 2).

Clinical performance of triple antibody positive status Despite significant differences between the clinical performance of individual QUANTA Lite® and QUANTA Flash® assays, when patients were classified as having single, double, and triple positivity, the number of triple-positive patients as detected by the 2 platforms was very similar (80 by QUANTA Lite® and 86 by QUANTA Flash®), while there was a significant discrepancy in the number of single-positive patients (40 by QUANTA Lite® and 21 by QUANTA Flash®). Altogether, 74 of 80 (92.5%) and 79 of 86 (91.9%) triple-positive patients had APS based on a historical...
Prevalence of triple positivity has the potential as a strong diagnostic and risk stratification tool. Over the last decade, a significant progress has been made in the diagnosis and management of APS. Beside the identification of new and promising aPL antibody specificities (antiphosphatidylserine/prothrombin and anti-β2GPI Domain 1 antibodies), several new concepts have been introduced, such as risk stratification, antibody profiling, and establishment of aPL antibody scores. New technologies and platforms have also emerged with favorable analytical and clinical characteristics.

One of the most important recent findings is the identification of triple antibody positivity as a strong diagnostic and risk stratification tool. Triple positivity has the potential of becoming an excellent laboratory tool for diagnosing APS and predicting thrombosis, but its usefulness may be hindered by the variable performance of different aPL antibody assays. Discrepancies between methods may lead to identifying different patients as triple positive depending on a platform or assay.

At the level of individual aPL antibodies, specifically for aCL IgM and anti-β2GPI IgG, significant differences were identified between the QUANTA Lite® and QUANTA Flash® assays in clinical sensitivity and specificity, together with a high level of disagreement in the classification of patients as antibody positive or negative. However, we have shown that a similar number of triple-positive patients were identified by the 2 methods: 80 with QUANTA Lite® and 86 with QUANTA Flash®. In addition, we have confirmed a strong association between triple antibody positivity and APS, regardless of the method. The OR for APS-related symptoms was 24.9 based on QUANTA Lite® results and 24.7 based on QUANTA Flash® results. This value is similar to the OR of 33.3 reported by Pengo et al in a large Italian cohort. Moreover, despite the differences between individual assays, we observed a total agreement of 95.5% for triple positivity between QUANTA Lite® and QUANTA Flash® platforms, demonstrating that the 2 methods identified the same patients as triple-positive. This finding has important implications for the future use of aPL antibody results because it suggests that the agreement between various platforms is better when assessing aPL antibody profiles than when assessing individual antibodies. In conclusion, our results not only confirm that triple aPL positivity has a strong association with APS, but also demonstrate that the identification of patients with triple antibody positivity is platform independent.

**Comparison of triple antibody positive status according to platforms** Almost a perfect agreement was observed between the triple positive antibody status identified by QUANTA Lite® and QUANTA Flash® platforms, with a total agreement of 95.5% (95% confidence interval [CI], 91.8%–97.5%) and a Cohen’s kappa of 0.90 (95% CI, 0.85–0.96). The results are presented in Table 3.

**DISCUSSION** In this cross-sectional study, we have shown that 2 different platforms, an automated microparticle CIA and a traditional ELISA, identified the same patients as having triple aPL antibody positivity.

Over the last decade, a significant progress has been made in the diagnosis and management of APS. Beside the identification of new and promising aPL antibody specificities (antiphosphatidylserine/prothrombin and anti-β2GPI Domain 1 antibodies), several new concepts have been introduced, such as risk stratification, antibody profiling, and establishment of aPL antibody scores. New technologies and platforms have also emerged with favorable analytical and clinical characteristics.

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**Contribution statement** TI contributed to the concept and design of the study as well as the analysis and interpretation of the data. MK interpreted the data and conducted statistical analyses. MC-L was responsible for clinical evaluation of the patients. SP conducted the analysis. JM was responsible for critical writing, revising the manuscript for intellectual content, and the final approval of the manuscript.

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**Table 3** Contingency table showing agreement between triple positivity status as identified by QUANTA Lite® and QUANTA Flash® assays

| Assays | QUANTA Flash® chemiluminescence assay |
|--------|--------------------------------------|
| QUANTA Lite® enzyme-linked immunosorbent assay | no triple positivity | triple positivity | total |
| no triple positivity | 132 | 8 | 140 |
| triple positivity | 2 | 78 | 80 |
| total | 134 | 86 | 220 |
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Streszczenie

Wprowadzenie  Ryzyko powikłań klinicznych w zespole antyfosfolipidowym (antiphospholipid syndrome – APS) zwiększa się, gdy u chorego występują wszystkie trzy rodzaje przeciwiał antyfosfolipidowych (antiphospholipid – aPL). Niemniej jednak brak zgodności między różnymi metodami pomiaru przeciwiał antykardioliopinowych (anticardiolipin – aCL) oraz przeciw β₂-glikoproteinie I (anti-β₂-GPI) jest znaczący, co prowadzi do rozbieżności w ocenie dodatnich wyników pomiędzy tymi metodami.

Celem tego retrospektywnego badania przekrojowego było zbadanie, czy dwie różne metody pomiarowe: QUANTA Lite® ELISA (metoda immunoenzymatyczna) i QUANTA Flash® CIA (metoda chemiluminescencyjna) pozwalają zidentyfikować tych samych chorych z potrójnie dodatnim wynikiem oznaczania przeciwiał aPL w grupie pacjentów z APS i towarzyszącymi chorobami autoimmunologicznymi.

Pacjenci i metody  Do badania włączono 220 osób z ogólnoustrojowymi chorobami autoimmunologicznymi (74 badanych z pierwotnym APS, 47 badanych z wtórnym APS oraz 99 badanych z toczniem rumieniowatym układowym bez APS). U wszystkich pacjentów oznaczono aCL i anti-β₂-GPI w klasach IgG i IgM z wykorzystaniem obu metod pomiarowych.

 Wyniki  Zgodność dodatnich wyników dla pojedynczych przeciwiał uzyskanych obiema metodami nie była pełna, wahała się od 81,8% do 90,9% w poszczególnych parach. Liczba pacjentów z dodatnimi wynikami dla wszystkich trzech rodzajów przeciwiał aPL była natomiast podobna (80 dla QUANTA Lite® i 86 dla QUANTA Flash®); zgodność pomiędzy obiema metodami dla rozpoznania chorych z potrójnie dodatnim wynikiem wynosiła 95,5% (współczynnik kappa Cohena = 0,90). W rezultacie obserwowano zbliżone ryzyko powikłań klinicznych APS – iloraz szans wynosił 24,9 dla QUANTA Lite® i 24,7 dla QUANTA Flash®.

Wnioski  Uzyskane wyniki potwierdzają silny związek pomiędzy obecnością potrójnie dodatnich przeciwiał aPL a APS oraz wykazują, że rozpoznanie chorych z potrójnie dodatnimi przeciwialami nie zależy od metody oznaczenia. W sytuacji, kiedy oceniane są profile aPL, zgodność pomiędzy różnymi metodami jest znacznie większa niż zgodność dla pojedynczych przeciwiał.