Activation of the alternative complement pathway: clinical application of a new technique to measure fragment Ba

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SUMMARY A new laser nephelometric technique for the measurement of the alternative complement pathway fragment Ba has been developed. Activation of the alternative complement pathway was assessed in 16 patients with Gram negative bacteraemia, six with Gram positive bacteraemia, 20 with rheumatoid arthritis, and 18 healthy subjects. Patients with Gram negative bacteraemia had significantly higher values of Ba (median 14.8%) than controls (9.3%) (p < 0.01), while patients with Gram positive bacteraemia and rheumatoid arthritis had values similar to those of controls (10.1% and 9.5%). The technique proved sensitive and precise, and is suitable for the routine laboratory evaluation of complement activation through the alternative pathway.

Activation of the complement system has an important role in the pathogenesis of numerous immunologically mediated disorders.¹-³ The complement cascade is activated through two major pathways (fig 1). The alternative pathway is activated by complex polysaccharides of Gram negative bacteria, while the classical pathway is triggered by immune complexes.⁴ Both pathways converge at the level of C3 to progress in a common sequence leading to the formation of the membrane attack complex. Alternative pathway activation results in the splitting of factor B into a large Bb molecule (molecular weight 60,000) and a smaller fragment Ba (molecular weight 33,000). Bb contributes to the formation of the alternative pathway C3 convertase (C3bBb); Ba is physiologically inactive and takes no further part in the complement cascade.⁵ Increased concentrations of Ba thus provide a specific index of ongoing activation of the alternative complement pathway. In Gram negative bacterial infections complement is activated by direct contact with the cell wall of micro-organisms.⁶ Quantitation of complement activation via the alternative pathway may provide a useful tool for evaluating the progression and severity of Gram negative infections.⁷ In rheumatoid arthritis, a typical immune complex disease,⁸ complement is thought to be activated through the classical pathway,⁹ though activation through the alternative route has been reported.¹⁰

In this study we show how Ba can be separated from the larger factor B related molecules by precipitation through the alternative pathway.
itation with polyethylene glycol. Ba can then be measured by a laser nephelometric technique, using an antiserum reacting with determinants present on the whole molecule of factor B. We analysed activation of the alternative complement pathway in groups of patients with Gram negative bacteraemia (GNB), Gram positive bacteraemia (GPB), rheumatoid arthritis and in healthy subjects.

Material and methods

CLINICAL SAMPLES
Blood collected by venepuncture into a final edetic acid concentration of 10 mmol/l was immediately separated by centrifugation at 1000 g at 4°C for 15 minutes and the plasma stored at -70°C. Samples were obtained from 16 patients (nine men, median age 46 years, range 17–72) with blood samples positive on culture for Gram negative bacteria; six patients (three men, median age 34 years, range 1–70) with blood samples positive on culture for Gram positive bacteria, 20 patients (five men, median age 52 years, range 27–71) with “classical” or “definite” rheumatoid arthritis according to the criteria of the American Rheumatism Association, and 18 healthy subjects (nine men, median age 32, range 27–47). The degree of disease activity in patients with rheumatoid arthritis was assessed by measuring the erythrocyte sedimentation rate (ESR) (mm/first hour).

FRAGMENT Ba STANDARD
Normal human serum was incubated at 37°C for 60 minutes with inulin (BDH Biochemicals, Poole, Dorset, England) at a concentration of 3 mg/ml to activate exhaustively complement through the alternative pathway. After centrifugation at 1500 g for 30 minutes the supernatant was frozen in aliquots and stored at -70°C. This Ba enriched preparation was used throughout the study as the source of the Ba standard.

To separate fragment Ba from native factor B and fragment Bb 100 µl of Ba standard were added to 100 µl of polyethylene glycol 6000 (PEG) (Sigma Chemicals, Poole, Dorset, England) in Dulbecco's phosphate buffered saline (PBS), pH 7.2, at different final concentrations ranging from 2–24%, incubated for 90 minutes at 4°C, and centrifuged at 1500 g at 4°C for 30 minutes. This aims to precipitate the larger factor B and Bb, leaving Ba in the supernatant. The efficacy of various PEG concentrations in separating Ba was assessed by a double dimension immunodiffusion performed in 1% agarose 15 (BDH Biochemicals) in PBS containing 2% PEG.

To establish the working conditions of the assay, a checker board titration was performed. Supernatant from Ba standard treated with PEG was serially diluted in PBS and incubated at room temperature with different concentrations of anti-factor B antiserum (Dakopatts, High Wycombe, Buckinghamshire, England) in PBS. Readings were taken in a Behring laser nephelometer at 15, 30, 45, 60, 90, 120, 150 and 180 minutes and referred to a reference curve prepared from Ba standard.

Statistical analysis was carried out using the Wilcoxon's rank sum test and the Spearman's rank regression analysis.

Results

Ba was successfully separated from larger factor B-related molecules by precipitation in a final concentration of PEG of 18% (fig 2). At this concentration of PEG larger factor B related molecules were precipitated and the smaller fragment Ba remained in the supernatant. Higher concentrations of PEG tended to precipitate Ba, while lower concentrations did not completely remove larger factor B related molecules from the supernatant.

![Fig 2 Ba separation assessed by double dimension immunodiffusion. Central well: neat anti-factor B antiserum. Well A: edetic acid plasma; well B: normal human serum (NHS); well C: Ba standard (NHS treated with inulin). Wells D, E, F: supernatants recovered after PEG treatment of Ba standard at final concentrations of 16%, 18%, and 20%, respectively.](http://jcp.bmj.com)
The intra-assay coefficient of variation derived from estimating the same sample 30 times was 4.6%. This interassay coefficient of variation obtained by repeating the evaluation of the same sample on 10 different occasions was 13.8%.

Fig 4 shows the values in 16 patients with GNB, six with GPB, 20 with rheumatoid arthritis and in 18 healthy subjects. Ba values are expressed as a percentage of the top standard, which is regarded as 100%. Low yet measurable Ba values were present in all the normal subjects tested. Patients with GNB had significantly higher values of Ba (median 14.8%, range 7.0–38.3%) compared with controls (9.3%, 6.5–14.4%; p < 0.01). Nine (56%) had Ba values in excess of the upper limit of normal. Patients with GPB and with rheumatoid arthritis showed Ba values (10.1%, 6.1–12.7% and 9.5%, 4–17.4%, respectively) not significantly different from those of controls and significantly lower than those in patients with GNB (p < 0.01). Four patients with rheumatoid arthritis had Ba values in excess of the upper limit of normal. These patients had active disease (ESR > 40 mm first hour), and in patients with rheumatoid arthritis a
positive correlation was observed between the ESR and Ba values ($r = 0.49$, $p < 0.03$).

Discussion

To assess complement activation via the alternative pathway we established a laser nephelometric technique that measures fragment Ba. Significantly increased values of Ba were found in patients with Gram negative bacteraemia in which the alternative pathway is known to be activated. In contrast, Ba values were found to be normal in most of the patients with Gram positive bacteraemia and with rheumatoid arthritis. Our data show that when infection is suspected the finding of increased values of Ba implicates Gram negative micro-organisms.

Activation of the alternative pathway has previously been assessed by showing a reduction in the amount of the intact factor B molecule. This provides only indirect information concerning activation as extensive factor B breakdown would be required to produce a measurable reduction in the factor B molecule. Ba is generated by the enzymatic cleavage of factor B which occurs only when the alternative complement pathway is activated. The plasma concentration of Ba, therefore, provides a specific indication of the state of complement activation via this pathway. To measure Ba directly this fragment must first be separated from larger factor B related molecules, then quantitated by the use of anti-factor B antiserum. The separation step is required as the polyclonal anti-factor B antiserum recognises antigenic determinants present both on the parent molecule and the split product.

An essay exploiting PEG separation of Ba has been described previously. The quantitation of the fragment was, however, performed using radial immunodiffusion, which requires 48 hours to produce results and is insensitive. Evidence of activation through the alternative pathway has also been provided by electrophoretic techniques. These techniques, however, are insensitive, give only qualitative results, and are unsuitable for analysis of large batches of samples. In contrast, laser nephelometry is sufficiently sensitive to detect Ba in healthy subjects and permits the automated analysis of large sample batches in a short time.

In 56% of patients with Gram negative bacteraemia, Ba values were significantly increased, confirming the role of the alternative pathway. In a previous study 31% of patients showed a reduction in the values of factor B and properdin, when measured by a radial immunodiffusion technique. The greater number of patients showing evidence of activation of the alternative pathway in our study was probably a reflection of the higher sensitivity of our technique. Detection of increased amounts of the Ba fragment in patients with Gram negative bacteraemia suggests that this technique may be useful in the diagnosis and management of septicaemia. It has the advantage of providing results within five hours; blood cultures require at least 24 hours' incubation. The assay may be particularly useful in groups prone to infections but in whom the clinical diagnosis is difficult, such as neonates, the elderly, and the immunosuppressed.

The results obtained in Gram positive bacteraemia suggest that the alternative pathway is not activated. In contrast to Gram negative bacteria, the cell walls of Gram positive bacteria contain less lipopolysaccharide and are therefore poor activators of the alternative pathway.

Our results indicate that in some patients with rheumatoid arthritis there is increased breakdown of factor B. Complement activation in rheumatoid arthritis is known to proceed mainly along the classical pathway. The triggering of the alternative pathway is influenced by the concentrations of C3b. It is thus conceivable that high concentrations of C3b produced in rheumatoid arthritis through the classical route initiate the activation of the alternative pathway, ultimately leading to an increased breakdown of factor B and generation of Ba (fig 1). Interestingly, the extent of activation of the complement alternative pathway correlated significantly with the degree of disease activity, as assessed by the ESR.

The technique that we have developed may be useful in the management of high Ba values in patients with Gram negative bacteraemia, and in other conditions in which the alternative complement pathway has a role in the pathogenesis.

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