Granulocyte Chemotaxis Measured in a Boyden Chamber Assay by Quantification of Neutrophil Elastase

By G. Pelz, Almut Schettler and H. Tschesche
Lehrstuhl für Biochemie, Fakultät für Chemie, Universität Bielefeld, Bielefeld, Germany

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Summary: The human polymorphonuclear leukocyte reaction in response to chemotactic or chemokinetic stimulation is often assayed using the Boyden chamber technique. We present a quick and reliable method for evaluating Boyden chamber experiments, which avoids time-consuming cell counting and does not require expensive equipment. This method is based on assaying human neutrophil elastase, a serine protease derived from polymorphonuclear leukocytes. We tested the method in different types of Boyden chambers equipped with two superimposed filters or a filter amnion membrane combination. The chambers were incubated with the cells for 2 h then dismantled and the elastase activity in supernatant, filters or membrane was assayed. The results were compared with the results obtained by cell counting, or measured by determination of myeloperoxidase. There was a good correlation between the cell count and elastase technique ($r = 0.90$), but the elastase method achieved higher intra- and inter-assay precision. Myeloperoxidase and elastase results also correlated well ($r = 0.94$) and showed comparable intra- and inter-assay precision. With the elastase method it was also possible to quantify polymorphonuclear leukocyte reactions on an amnion membrane surface. In amnion membrane assays the percentage of cells which reacted in response to formyl-peptide stimulation was not altered by varied cell concentrations, and polymorphonuclear leukocytes showed little unstimulated adherence or migration.

Introduction

Polymorphonuclear leukocytes are the primary effector cells at sites of acute inflammatory processes. Their ability to follow gradients of chemotactic reagents and migrate through vessel walls and tissue barriers is essential for their function in the host defence system. Several in vitro methods for detecting chemotactic activity of polymorphonuclear leukocytes have been developed, including the so-called Boyden chamber invented by Boyden in 1962 (1). The Boyden chamber consists of two (upper and lower) compartments separated by a microporous filter membrane. When a chemotactic substance is placed in the lower, and the cells are placed in the upper compartment of the chamber, polymorphonuclear leukocytes migrate into the filter along the resulting concentration gradient. The method mainly used to measure the chemotactic response is the cell counting technique. The distance which the cells travel into the filter (leading-front technique) can be determined after fixation and staining. Alternatively the number of cells which migrate a given distance into the filter is counted. Several other methods have been recommended to avoid time-consuming microscopic counting of the cells (for review see l.c. (2)). Gallin et al. (3) described a method in which cells were labelled with the radioactive tracer $^{51}$Cr. Two overlayed filters were used. The number of polymorphonuclear leukocytes which penetrated through the first filter was determined by measuring the radioactivity of the lower filter. Somersalo et al. (4) invented a method also using two filters, and measuring the content of human polymorphonuclear leukocyte myeloperoxidase in the lower filter. This method avoided working with radioactive isotopes and was easier to perform than cell-counting techniques.

We have developed a method similar to Somersalo's based on quantifying human polymorphonuclear leukocyte elastase. Human polymorphonuclear leukocyte elastase is easier to handle and to measure than myeloperoxidase, which makes elastase assays suitable for routine use. Moreover, elastase and myeloperoxidase assays may complement one another when experimental conditions do not permit the use of the myeloperoxidase method.

Materials and Methods

Isolation of polymorphonuclear leukocytes

Fresh blood was taken by venipuncture from healthy volunteers. Polymorphonuclear leukocytes were isolated from the heparinized blood according to the method of Boyum (5). Polymorphonuclear leukocytes were separated from the blood by sedimentation in a mixture of Ultravist/Methocel (18.2 ml/100 ml Ultravist, Schering, Berlin; 1.45 ml/100 ml Methocel, Sigma, München), followed by density gradient centrifugation with Histopaque-1077 (Sigma, München). Some remaining erythrocytes were removed by hypotonic lysis, and the polymorphonuclear leukocytes resuspended in a buffer solution containing 140 mmol/l NaCl, 2.7 mmol/l KCl, 8 mmol/l Na₃HPO₄·2H₂O, 1.5 mmol/l K₃PO₄ (pH 7.3), 0.5 mmol/l MgCl₂·6H₂O, 0.9 mmol/l CaCl₂·H₂O, 1 g/l glucose and 10 g/l human serum albumin (Institut Mérieux GmbH, Leimen). Viability of the polymorphonuclear leukocytes was >95% as determined by exclusion of trypan blue.

Human amnion membrane

Normal term placentas were obtained from the hospital immediately after delivery and stored in phosphate-buffered saline containing penicillin and streptomycin (100 × 10³ IU/l and 1 mg/l, respectively; Sigma, München). The adherent chorioamniotic membrane was peeled away from the amnion, and the epithelial cell layer was scraped off after treatment with 40 g/l deoxycholate solution for 30 min (Sigma, München). The remaining basement membrane was washed carefully with the buffer solution used for resuspending polymorphonuclear leukocytes (see above).

Boyden chamber preparation

Chemotaxis chambers similar to that described by Russo et al. (6) were used. The two parts of the chemotaxis chamber were separated either by one 5 μm micropore filter (Millipore S. A., Molsheim), a two filter combination with one upper 5 μm or 8 μm filter and one lower 0.22 μm filter, or a human amnion membrane and an underlying 0.22 μm filter. Filters were soaked with buffer A containing 140 mmol/l NaCl, 2.7 mmol/l KCl, 8 mmol/l Na₃HPO₄·2H₂O, 1.5 mmol/l K₃PO₄, 0.5 mmol/l MgCl₂·6H₂O, 0.9 mmol/l CaCl₂·H₂O, 1 g/l glucose, 10 g/l human serum albumin, pH 7.3, for 30 min before use. The lower compartment of the chambers was filled with 2 ml buffer A alone or with various concentrations of N-formyl-methionyl-leucyl-phenylalanine in the buffer A to induce chemotaxis. Experiments were started by placing 300 μl of polymorphonuclear leukocyte suspension in the upper compartment. After 2 h incubation at 37 °C, (humidity 90%, 95% air, 5% CO₂), the remaining supernatant in the upper compartment was collected. Non-adherent cells were removed from the upper filter or the amnion membrane by washing 3 times with 300 μl of the buffer required for the subsequent enzymatic assay, and combined with the supernatant.

Polymorphonuclear leukocyte quantification

Cell count method

The polymorphonuclear leukocytes in the filters were fixed with aqueous ethanol (volume fraction 0.7) and stained with hematoxylin as described by Boyden (1). The filters were cleared with xylene and placed on a glass slide. Polymorphonuclear leukocytes in each filter were counted in five random, high power fields (×400) scanning through the whole depth of the filter. Total polymorphonuclear leukocytes for each filter were calculated from this, and the results expressed as a percentage of the number of cells added to each chamber.

Enzymatic assays

After collection of the supernatant, the Boyden chambers were dismantled, and each filter or amnion membrane disc was placed in 1 ml of elastase or myeloperoxidase test buffer containing 10 ml/l Triton X-100 (elastase test buffer: 0.2 mol/l triethyamine hydrochloride containing 1.0 mol/l NaCl and 2 g/l benzalkonium chloride, pH 8.0; myeloperoxidase test buffer: 83 mmol/l citric acid, 112 mmol/l Na₂HPO₄·2H₂O, pH 5.6). Cells and granules were disrupted by twofold sonication (Branson cell disrupter, B-15 microtip, step 4 for 10 s, after 20 min step 5, 15 s). The resulting cell lysates were cleared by high speed centrifugation (15 min, 15,000 g). Elastase activity was assayed at 25 °C and samples were measured in elastase test buffer using MeO-Suc-Ala-Ala-Pro-Val-4-nitranilide (MeO-Suc, methoxyxycinnyl) (1 mmol/l dissolved in dimethyl sulfoxide) in a final concentration of 0.05 mmol/l as substrate (7). For determination of the polymorphonuclear leukocyte mean elastase content 200 μl of the cell suspension were centrifuged (10 min, 400 g) and the resulting pellet treated in the same manner as the Boyden chamber samples. The test was standardized with human polymorphonuclear leukocyte elastase (Elastin Products Inc.). To determine spontaneous elastase release during migration, filters were placed in 1 ml of the buffer solution used for resuspending polymorphonuclear leukocytes (see above) and centrifuged at 400 g for 10 min, and elastase activity in the supernatant was assayed. For measurement of myeloperoxidase activity, samples were incubated at 37 °C for 45 min in myeloperoxidase test buffer (83 mmol/l citric acid, 112 mmol/l Na₂HPO₄·2H₂O, pH 5.6) using 0.91 mmol/l ABTS containing 125 μl/l H₂O₂ (300 g/kg) as substrate. Elastase and myeloperoxidase activities were monitored at 405 nm (Eppendorf photometer).

All assays were carried out in duplicate and intra-assay precisions for all assays were calculated from

\[
\frac{SD}{\bar{x}} \times 100 = CV% 
\]

SD is the mean of standard deviations for all samples, \(\bar{x}\) is the mean of all samples, and CV is the coefficient of variation.

Results

Elastase content of neutrophils

The total content of polymorphonuclear leukocyte elastase was determined in separate samples after disruption of the polymorphonuclear leukocytes by


donor: hydrogen-peroxide oxidoreductase (EC 1.1.1.7)
treatment with detergent and a twofold sonication procedure. This procedure ensured the complete release of elastase from the granules. The elastase content was calculated from the determined enzyme activity. The mean protein content was 14.90 ± 2.7 μg elastase for $5 \times 10^6$ polymorphonuclear leukocytes ($n = 18$, 13 donors). This was comparable to the values reported in literature (8). Therefore, we were able to reliably detect 10 000 cells per sample. Spontaneous release of elastase in the supernatant was <2%, as determined in the supernatant of the samples and the filters after low speed centrifugation.

Comparison of the elastase and cell counting methods

To compare the elastase and the cell counting methods, two experiments were carried out using parallel methods. The elastase and cell counting method were used to measure the number of cells which migrated in a 5 μm filter after chemotactic stimulation. For this purpose, polymorphonuclear leukocyte suspensions (cell count from $1.25 \times 10^9/l$ to $5.0 \times 10^9/l$) were placed in the upper compartment and chemotaxis was induced by $10^{-7}$ mol/l N-formyl-met-leu-phe in the lower compartment. Under these conditions most of the polymorphonuclear leukocytes migrated into the filter or adhered to the surface. Both methods gave similarly high results (fig. 1), whereby the cell count method showed about 8% more polymorphonuclear leukocytes in the filters (mean for all concentrations), and the mean of standard deviations for all concentrations was 14.2% for the cell count method and 8.9% for the elastase method. Higher polymorphonuclear leukocyte concentrations led to a slight decrease in cells in the filter; +11% in the elastase method, +13% in the cell count method.

Both methods were then tested in a two filter migration assay (fig. 2). A 0.22 μm filter was overlayed with an 8 μm filter, then chemotactically induced, and spontaneous migration into both filters was measured at two polymorphonuclear leukocyte concentrations. A high percentage of polymorphonuclear leukocytes (up to 78%, elastase method) adhered to the surface or migrated into the upper filter even without stimulation. N-Formyl-met-leu-phe stimulation led to an increase in the polymorphonuclear leukocyte content.
of the upper filter of up to 95% (elastase method). A higher cell concentration decreased the percentage of cells in the upper filter without stimulation, but N-formyl-met-leu-phe stimulation diminished this effect. Only a few polymorphonuclear leukocytes were found attached to the lower filter without N-formyl-met-leu-phe stimulation. The cell count technique often failed to show any cell in the randomly chosen high power fields. Chemotactic stimulation led to a distinct increase in migration. The elastase method in all cases resulted in higher migration rates into the lower filter. The correlation coefficient between the two methods was \( r = 0.90 \). In all cases the inter-assay standard deviations were higher with the cell count method. The intra-assay precision of the elastase and the cell count method was calculated from both experiments. The coefficients of variation in the upper (only) and lower filter were calculated separately. Results for the elastase method were 6.1% (n = 14) and 24.4% (n = 8), respectively. Results for the cell count method were 12.8% (n = 14) and 30.1% (n = 8), respectively.

**Comparison of the elastase and myeloperoxidase methods**

Both methods were applied to a two filter assay with an upper 5 \( \mu \text{m} \) filter and a lower 0.22 \( \mu \text{m} \) filter (fig. 3). The results of the elastase method for spontaneous and N-formyl-met-leu-phe-stimulated migration into the upper filter were clearly lower (35.8 = -20.5% and 74.6 = -16.6%, respectively), compared with previous results from 8 \( \mu \text{m} \) filters. Transmigration of polymorphonuclear leukocytes to the lower 0.22 \( \mu \text{m} \) filter was in accordance with previous results. The results of the myeloperoxidase method and elastase method results correlated with \( r = 0.94 \). Interassay standard deviations for both methods were comparable. The intraassay precision was calculated as above. The coefficients of variation for the elastase method in the upper and lower filter were 6.1% and 29.4% (n = 6). Results for the myeloperoxidase method were 5.8% and 25.2% (n = 6).

**Application of the elastase method in amnion membrane models**

All measurements were carried out using the elastase method. Chemotaxis chambers with a piece of human amnion membrane mounted on a 0.22 \( \mu \text{m} \) filter were assembled, and N-formyl-met-leu-phe concentrations ranging from \( 10^{-5} \text{ mol/l} \) to \( 10^{-11} \text{ mol/l} \) (plus controls without chemoattractant) were applied to the lower compartment of the chambers. An N-formyl-met-leu-phe concentration of \( 10^{-8} \text{ mol/l} \) induced maximum adherence to, and migration through the membrane (fig. 4). At higher N-formyl-met-leu-phe concentrations even fewer cells than in unstimulated controls were detected in/on the membrane. Examination of the underlying filter revealed that polymorphonuclear leukocytes migrated through the entire amnion membrane into the filter at only the most stimulatory N-formyl-met-leu-phe concentrations. Even with optimal N-formyl-met-leu-phe stimulation only a small percentage of the cells was detected on the filter: 0.5 ± 0.3% (5 × \( 10^{-9} \text{ mol/l} \) N-formyl-met-leu-phe) and 0.5 ± 0.4% (1 × \( 10^{-8} \text{ mol/l} \) N-formyl-met-leu-phe). Higher or lower N-formyl-met-leu-phe concentrations did not induce any detectable migration into the filter.

A second test showed that variation of the polymorphonuclear leukocyte concentration did not alter the percentage of polymorphonuclear leukocytes found.
Fig. 4. The effect of N-formyl-met-leu-phe concentration on the polymorphonuclear leukocyte response in chemotaxis chambers equipped with an amnion membrane. The two parts of the chemotaxis chamber were separated by a slice of human amnion membrane on an underlying 0.22 μm filter. The polymorphonuclear leukocyte concentration in the upper compartment was $10^{10}$/l. The results were expressed as the percentage of total polymorphonuclear leukocytes which migrated into, or adhered to the amnion membrane, detected by the elastase method. Means ± SD from 4 experiments.

Fig. 5. The effect of polymorphonuclear leukocyte concentration on the neutrophil response in chemotaxis chambers equipped with an amnion membrane, measured by the elastase method. The chemotaxis chambers were separated by a slice of human amnion membrane on an underlying 0.22 μm Millipore filter; 10^{-4} mol/l N-formyl-met-leu-phe was added to the lower compartment for stimulation. The results show the percentage of cells which migrated into, or adhered to the amnion membrane. Means ± SD from 6 experiments.

Discussion

This report describes a new method for assaying polymorphonuclear leukocytes in Boyden chambers operated conventionally or equipped with an amnion membrane. Measurement of elastase activity proved to be a quick and easy method for evaluating Boyden chamber experiments. Human leukocyte elastase is a serine protease primarily associated with polymorphonuclear leukocytes. However, monocytes also contain some human leukocyte elastase, and recently Lungarella et al. (9) reported that eosinophils also contain an elastase similar to the major isoenzyme of neutrophil elastase. But the enzyme content of these cell types is reported to be much lower than in neutrophils (the monocyte elastase content is approximately 6% of that in neutrophils (10)).

As our cell isolation procedure resulted in 99% polymorphonuclear leukocytes, the elastase content of contaminating cells could be assumed to be extremely low. The enzyme activity measured in our tests therefore derived almost entirely from neutrophils. As recently reported, adherent polymorphonuclear leukocytes release only very little (< 2%) of their elastase content even after formyl peptide treatment (11). This corresponded with our findings. As a result, the elastase method is not affected by elastase secreted into the medium.

Cell counts and elastase activity in the filters correlated closely in most cases. Although the correlation was adequate, the cell count and elastase results showed systematic differences. Cell counting results in the upper filter were always higher than those calculated from the elastase method. Close examination of the filters revealed that the areal distribution of polymorphonuclear leukocytes was not homogenous. In most cases polymorphonuclear leukocytes were more numerous on the periphery of the filter. Therefore, the results of the cell count method are strongly affected by the distribution of the high power fields on the filter. This effect can hardly be quantified, and any way of choosing the high power fields is more or less arbitrary. In contrast, cell count results for the lower filter led to lower migration rates than elastase activity. We decided to use a fine pore "catcher" filter as recommended by Keller et al. (12) in the lower position. This prevented polymorphonuclear leukocytes from migrating through both filters; they were blocked and remained attached to the surface of the second filter. But, weakly attached cells might have been washed off during the cell count staining procedure and therefore could not be counted. The elastase method avoids this problem.

in the amnion membrane after 2 h of N-formyl-met-leu-phe stimulation (fig. 5). This percentage was about 25 to 45% lower than in similar filter experiments; moreover, in contrast to the filter experiments, unstimulated adherence to or migration of the polymorphonuclear leukocytes into the amnion membrane was minor.
Since the results of each elastase test were calculated as a percentage of the total elastase content present in the same test, errors resulting from a different polymorphonuclear leukocyte number in the chambers were also eliminated.

Somersalo et al. (4) presented a modified Boyden chamber assay based on myeloperoxidase. We found that elastase and myeloperoxidase assays gave comparable results, but elastase had some clear advantages in routine use. The enzymatic assays could be carried out at nearly room temperature, and the samples could be stored for up to three days at 4 °C without much loss of activity. The elastase method may also be a preferable alternative to myeloperoxidase assays when experimental conditions influence the myeloperoxidase measurements.

Russo et al. (6) recommended the use of human amnion membrane for Boyden chamber investigations. The amnion membrane is a three-layered structure of epithelial cell layer, basement membrane and loose stroma tissue. When the epithelial cell layer is removed the basement membrane appears as a dense collagen network. We found that on this natural matrix barrier most polymorphonuclear leukocytes did not adhere or migrate without further stimulation. N-Formylmet-leu-phe stimulates about 40% of the cells to migrate through the membrane or adhere to the surface. This percentage was not altered by an 8-fold rise in cell concentration. Amnion membrane chambers, therefore, permit a sensitive assay widely unaffected by variations in cell concentration. They proved to be well suited for quantifying polymorphonuclear leukocyte migration following different types of stimulation. Of course, adherence to, or partial migration into, an amnion membrane cannot be distinguished by the elastase method, but the method provides useful information on leukocyte reactions on a natural basement membrane. However the percentage of polymorphonuclear leukocytes migrating through the whole membrane, even after optimal stimulation, was rather low.

The elastase method provided some advantages over other methods used to evaluate Boyden chamber investigations. It avoids time-consuming and laborious cell labelling with 51Cr, which could also lead to pre-stimulation of the sensitive cells, i.e. the unpolarized polymorphonuclear leukocytes. It is easy to perform without expensive equipment and much less tedious than the cell counting technique. In addition it can provide information on the percentage of cells able to respond to different stimuli and, in the two-filter assay, on the number of cells which migrated through the layer of the upper filter.

Thus, this novel enzymatic approach for measuring granulocyte chemotaxis by quantification of neutrophil elastase provides a simple standard procedure for routine assays.

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References


Prof. Dr. Harald Tschesche
Universität Bielefeld
Lehrstuhl für Biochemie
Fakultät für Chemie
Postfach 1001 31 »
D-33501 Bielefeld
Germany