Immunochemical and ultrastructural study of multiple myeloma with a heavy chain protein in the serum

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SUMMARY A patient with multiple myeloma had antigenically related monoclonal Fc-gamma fragments and complete IgG-kappa molecules in the serum. The urine contained only Fc-gamma fragments in the absence of Bence-Jones protein. The two distinct M-components in the serum showed electrophoretic identity but could be separated by chromatography. The simultaneous presence of complete monoclonal IgG molecules and Fc-gamma fragments, though difficult to detect, could be a frequent occurrence in multiple myeloma, and it could be defined as 'double paraproteinaemia'. A detailed ultrastructural study was performed in this case and showed fibril bundles being released from the malignant plasma cells; such fibrils could be the supramolecular organisation of the neosynthesised heavy chain fragments.

The association of a heavy-chain disease protein with a complete immunoglobulin in a monoclonal gammopathy has been described in less than 10 cases from 1968 to the present.1–7 It was observed in patients who, despite a dissimilar clinical picture, suffered mainly from a lymphoma-like disease. The clinical condition was constantly characterised by hepatomegaly and/or splenomegaly, an increased incidence of respiratory tract bacterial infections, and a complete absence of clinical and radiological lesions in the bones.

We present a case showing the clinical features of multiple myeloma (osteoporosis, bone marrow plasma cell infiltration, frequent respiratory tract bacterial infections), which had two monoclonal proteins in the serum. The first protein was a complete IgG-kappa M-component, and the second was composed of Fc-gamma dimers, which were also present in very low quantities in the urine. Despite the single narrow band on cellulose acetate electrophoresis, the second M-component was suggested by the presence of monoclonal Fc-gamma fragments in the urine concentrated up to 300-fold, proteinuria being less than 150 mg/24 hours.

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Material and methods

METHODS FOR PROTEIN STUDIES
Zone electrophoresis was performed on cellulose acetate strips with a Saitron 5003 apparatus and a Celloscan densitometer; quantitative determination of immunoglobulins was performed by radial immunodiffusion (Behring Partigen plates) according to Mancini8; immunoelectrophoresis with the Scheidegger9 micromethod against normal human serum and anti-human immunoglobulin (or Ig fragment) monospecific sera: anti-normal human (anti-NHS), anti-gamma chain, anti-alpha chain, anti-mu chain, anti-kappa and anti-lambda light chain, anti-IgG/Fab, anti-IgG/Fc, antiIgG/Fd serum (from Behring Institute, Marburg Lahn, West Germany).

The serum gammaglobulin fraction was obtained by ammonium sulphate precipitation according to Stanworth.10 Serum gel-chromatography11 was performed on a 2-5 × 110 cm Sephadex G-200 column; the pooled G-200 eluted fractions were rechromatographed on a 1·7 × 95 cm Sephadex G-100 column; in both cases we used 0-1 M phosphate buffer, pH 7·6. Urine gel-chromatography was performed on a 1·7 × 95 cm
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G-100 column; the elution buffer was 0.1 M acetate, pH 5-4. The fractions were read by an ISCO-UAS continuous-flow spectrophotometer (lambda = 280 nm UV) and collected by an LKB apparatus.

For SDS-polyacrylamide gel electrophoresis, we used bovine serum albumin (BSA) (68 000 MW) and carbonic anhydrase (45 000 MW) as markers; molecular weight (MW) was obtained on a logarithmic scale plotting RF (run length/total gel length) against MW values.

For the subclass characterisation of the isolated immunoglobulin and fragments, double diffusion according to Ouchterlony was performed in 2% agar gel in saline + 2% PEG 4000 with rabbit anti-human IgG1, IgG2, IgG3, IgG4 monospecific sera (from the Central Laboratory of the Netherlands Red Cross Blood Transfusion Service, Amsterdam, The Netherlands). Total protein values were obtained by Lowry's method.

METHODS FOR CELLULAR STUDIES

Electron microscopy

For electron microscopy, buffy coat cells, prepared according to Russo, and sternal bone marrow aspirate were fixed (2 hours) in 2% glutaraldehyde in 0.1 M phosphate buffer, pH 7.4, then rinsed with the same buffer (2 hours), and post-fixed (2 hours) in 1:33% osmium tetroxide solution in the same buffer. After rapid dehydration with increasing ethanol concentrations, the specimens were embedded in an Epon-812 mixture. Ultrathin sections, prepared with a diamond knife and stained with uranyl acetate (20-40 seconds) and lead hydroxide (6 minutes), were observed with a Philips-300 electron microscope.

Results

CASE REPORT

An 82-year-old woman was admitted to Agostino Gemelli Hospital of the Catholic University, Rome, in January 1977 complaining of fatigue, back pain, loss of weight, and recent bronchopneumonia for which she had received antibiotic therapy. In the same year she had been referred to two other hospitals with similar respiratory tract infections.

Physical examination

Physical examination revealed a still evolving bronchopneumonia, spontaneous pain in the dorso-lumbar region, and acute pain on percussion of the sternum; in addition, we found hepatomegaly without splenomegaly in the absence of lymphadenopathy and palatal oedema.

Laboratory investigations

The routine laboratory test results were as follows: a total body skeletal x-ray showed a diffuse and severe osteoporosis without osteolysis; a liver biopsy showed a well-preserved structure without infiltration and/or sclerosis. The bone marrow aspirate (sternum) presented atypical, immature, polynucleated, and bizarre plasma cells (approximately 40%), whereas the other cells maintained a normal morphology but were slightly decreased in number. Other haematological findings were: ESR 54 mm/h, haemoglobin 8.6 g/dl; red cell count 3.4 x 10^{12}/l; white cell count 7.4 x 10^{9}/l; platelet count 700 x 10^{9}/l; total serum proteins 80 g/l; alkaline phosphatase 86 IU; differential white cell count in the normal range.

Immunological data

Serum electrophoresis on cellulose acetate showed hypoalbuminaemia (34.9 g/l) and a quite homogeneous and sharp migration band in the gammaglobulin region, which was 32-2% of the total proteins (Fig. 1). Quantitative determination of immunoglobulins showed: IgG 44.5 g/l (4450 mg/100 ml), IgA 0.9 g/l (90 mg/100 ml), and IgM 11.0 g/l (110 mg/100 ml).

Serum immunoelectrophoresis showed an anomalous precipitation arc with rabbit anti-human gamma heavy chain serum and a very faint precipitation arc with anti-kappa light chain serum.

Urine immunoelectrophoresis showed no demonstrable precipitation arcs against anti-NHS at a concentration of 100-fold, usually sufficient to demonstrate Bence-Jones proteinuria in most cases of multiple myeloma. When the urine was concentrated up to 300 times, immunoelectrophoresis

![Fig. 1 Serum electrophoresis showing a homogeneous migration band in the gammaglobulin region.](http://jcp.bmj.com/ on December 18, 2017 - Published by group.bmj.com)
revealed an anomalous precipitation arc against only anti-gamma heavy chain (or Fc-gamma) serum.

**Protein studies**

Serum gel-chromatography on Sephadex G-200 was performed (Fig. 2) and the isolated fractions (tubes 48 to 65) developed an immunoelectrophoretic precipitation arc abnormally shaped in the cathodal tract with anti-gamma heavy chain and anti-Fc gamma monospecific sera, while no precipitation arcs were found with anti-light chain (kappa or lambda) sera.

![Fig. 2 Chromatographic separation of serum on Sephadex G-200. The fractions from tubes 48 to 65 (shaded area) react only with anti-Fc gamma, and anti-gamma sera.](image)

A further separation of the pooled fractions (tubes 48 to 65) on Sephadex G-100 gave two different chromatographic peaks (Fig. 3). The first (tubes 45 to 68) on immunoelectrophoretic analysis showed an irregular precipitation arc with anti-Fab gamma (Fig. 4a), anti-Fc gamma (Fig. 4b), and anti-kappa (Fig. 4c) sera, respectively. This fraction contained a complete monoclonal IgG-kappa protein, further characterised as belonging to the G1 subclass by means of double diffusion performed with monospecific anti-IgG subclass sera. The second peak (tubes 70 to 88) showed an abnormal precipitation arc only with anti-gamma heavy chain and anti-Fc gamma sera, and not with anti-Fab gamma, anti-kappa, or anti-lambda sera (Fig. 5).

A similar detailed study was performed on the urine. The total 24-hour urine protein was about

![Fig. 3 Chromatographic separation of pooled fractions (tubes 38 to 65) on Sephadex G-100. No reaction is obtained with monospecific anti-light chain (kappa or lambda) sera from the fractions eluted with the second peak.](image)

100 mg/100 ml (Ig/l), but, also after routine concentration, it was still too low for electrophoretic studies; therefore, it was necessary to concentrate the urine sample up to 350-fold to find either a faint anodal band related to albumin or a monoclonal cathodal band. The latter showed, on immunoelectrophoresis, an abnormal precipitation arc with anti-Fc gamma (Fig. 6a) and anti-gamma heavy chain (Fig. 6b) sera, but no precipitation arcs with anti-Fab gamma, anti-kappa, or anti-lambda sera.

The fragments were further characterised as belonging to the G1 subclass, as previously described. SDS-polyacrylamide gel electrophoresis on serum and concentrated urine showed that the unique thick band of urine migrated in close relation with an analogous thick band present in the serum; their approximate molecular weight was estimated to be about 50 000 (Fig. 7). After mercaptoethanol reduction" of this gamma chain related protein, the resulting 24 000 MW fragments were considered as monomers.

**Light microscopy studies**

On light microscopy of marrow smears, 40% plasma cells were counted. Most of them appear irregular in shape, frequently polynucleated (2-6 nuclei), and containing one or more nucleoli.

**Electron microscopy studies**

Under the electron microscope, different morphologi-
Immunoelectrophoretic study of multiple myeloma

Fig. 5  Immunoelectrophoresis of serum fractions eluted with the second peak on Sephadex G-100 (tubes 70 to 88). An abnormal precipitation arc is obtained only with anti-gamma heavy chain and anti-Fc gamma sera.

Fig. 6  Immunoelectrophoresis of urine. An abnormal precipitation arc is obtained only with anti-Fc gamma (a) and anti-gamma (b) sera.

Cytological populations of plasma cells could be detected. The most frequent type is shown in Fig. 8, and its features are similar to those described in other myelomatous or heavy chain disease plasma cells. In particular, the rough endoplasmic reticulum (RER) is abundant, it frequently presents annulate lamellae, and its cisternae are full of a homogeneous, dense material (Fig. 9). The Golgi apparatus is also quite abundant (Fig. 9); in its maturing (concave) face different sized vesicles are seen, containing a very electron-dense material; some of the largest are deformed by their crystalline content, which is well recognized only at times, and at others can be deduced...
in the serum of a patient suffering from clinical multiple myeloma. The first M-component has been identified as an IgG1-kappa protein and the second as an Fc gamma I fragment (comparable to a gamma heavy chain protein). Despite the presence of a heavy chain protein in the serum, the clinical pattern was different from that of classical Franklin's disease.

Many of the signs that give a lymphoma-like pattern to Franklin's disease were absent, such as palatal oedema, erythema, as well as lymph node, liver, and spleen involvement. On the other hand, this case is much more like a multiple myeloma, as far as the severe osteoporosis and morphological plasma cell features are concerned; the increased respiratory tract infection, however, is a less specific sign present in both diseases, probably due to the relative humoral immunity deficiency.

On zone electrophoresis, as only one M-component was detectable, the complete IgG-kappa monoclonal protein and the gamma chain fragments could not be shown. The presence of the latter, successively isolated by means of gel-chromatography in the serum, was merely suggested by the finding of Fc gamma fragments in the urine.

Fc gamma dimers have already been found in extremely low quantities by other authors in the serum of normal people, whereas in our patient they represented about one-third of the total myelomatous protein, as shown by chromatographic separation on Sephadex G-100.

Ultrastructural observations enabled us to correlate the immunochemical results with the abnormal plasma cell features. Electron micrographs, in addition to the moderately electron-dense material located in the RER cisternae, show two main storage sites of organised material in pathological plasma cells. The first appears as granules of various sizes originating from the Golgi apparatus and consisting of dense material often in a crystalline form, similar to Russel's bodies found in classic myeloma plasma cells. The second is represented by numerous 10 nm fibril bundles. Such fibrils have been described frequently in myelomatous plasma cells, but their meaning has not yet been clarified.

Our ultrastructural findings and previously reported data on supramolecular organisation are consistent with those experimental models, which showed that intracytosolic microfilaments and microfibrils may originate from monomeric or dimeric globular proteins (45 000-60 000 MW) and polymerise as filaments. Subsequently, two such filaments give place to a supramolecular arrangement (eg, microfilament or microfibril). An analogous process could have taken place in our patient's

Fig. 7 SDS-polyacrylamide gel electrophoresis of serum and urine. The unique thick, homogeneous, and narrow band of urine migrates in close relation with an analogous band of serum (approximate MW = 50 000).

from the hexagonal or pentagonal vesicle perimeter (Fig. 10a, b, c).

In close relation to the Golgi apparatus and to the RER, numerous fibril bundles (9-10 nm diameter) are seen (Fig. 9). Despite the difficulty of identifying their origin, they seem to originate from Golgi apparatus dictosomes cisternae (Fig. 9) or from the open RER cisternae (Fig. 11a). In some cases we noted that part of the cytoplasm containing such bundles seems to come off the plasma cell to release its fibrillar and cytosolic content in the blood (Fig. 11b).

Treatment and course

The patient survived for one year, complaining during that time of increasing bone pain. Despite the administration of cyclophosphamide she died in August 1978. A necropsy confirmed the bone marrow aspirate and radiological findings.

Discussion

Two different monoclonal components were found
Fig. 8  Electron micrograph of a binucleated plasma cell (× 12 500).
Fig. 9  Detail of a Golgi zone presenting parallel membrane arrangement, numerous dense vesicles of various size, and fibril bundles, which seem to originate from the Golgi apparatus vesicles (arrow) ($\times$ 34 000).
Fig. 10 Various aspects of Russel’s bodies.
(a) Golgi apparatus with numerous dense vesicles, fibril bundles, and one large paracrystalline granule (× 36 000).
(b) Detail of a polygonal granule. The crystalline array is probably lost with the fixation and embedding procedures (× 72 000).
(c) In other plasma cells (nearer to the specimen surface) the paracrystalline array of the granules is well preserved (× 32 000).
Fig. 11  Morphodynamic appearance of the fibrils. 
(a) Microfibrils and their probable subunits near an open RER cisterna (× 81 000).  
(b) Clasmatosis with abundant microfibrils (× 65 000).
plasma cells if we assume that a globular dimer (Fc gamma fragment) might be produced by the endoplasmic reticulum and transferred to the Golgi apparatus.

Figures 9, 11a, and 11b show a close spatial relation between these organules and the fibrils. For such molecules this arrangement is probably consistent with the thermodynamic minimum-energy state conformation and with the intracytosolic ionic content. For this reason it is not likely that such fibrils are composed of the complete monoclonal immunoglobulin, which is normally found in Rssel's bodies, as described in classical multiple myeloma.

Finally, at the periphery of the plasma cell, the fibril bundles enveloped in a part of the cytoplasm, seemed to be released from the cell. The dimers found in the serum and urine could represent the depolymerisation product of the released fibrils, due to the different ionic strength and energy charge of the haematic environment, in comparison with the cytosolic milieu. These sequential phases support the hypothesis of a 'de novo' synthesis of such fragments.

Buxbaum and Alexander obtained from a plasma cell culture heavy chain fragments, presenting a molecular weight similar to that of our fragment, with no evidence of a larger molecule. According to Adlersberg, such fragments might be a synthetic product, being the mutation result of the malignant clone already producing the complete protein. However, we cannot exclude the possibility that serum Fc gamma fragments might result from the proteolytic cleavage of the complete immunoglobulin by physiological plasma endopeptidases.

References


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