Neutrophil elastase and cathepsin G protein and messenger RNA expression in bone marrow from a patient with Chediak–Higashi syndrome

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Abstract

**Aims**—To determine whether neutrophil elastase and cathepsin G are expressed, at transcriptional or translational levels, in the bone marrow from a patient with Chediak–Higashi syndrome.

**Methods**—Blood neutrophils were isolated from three patients with Chediak–Higashi disease and bone marrow was collected from one. Cell lysates were analysed for neutrophil elastase and cathepsin G activity by enzyme linked immunosorbent assay and western immunoblotting. Northern blotting was used to detect messenger RNA (mRNA) for cathepsin G, elastase and β-actin in bone marrow extracts, and immunohistochemistry was used to localise the enzymes in marrow myeloid cells.

**Results**—Elastase and cathepsin G were not detected in blood neutrophils from the patients with Chediak–Higashi disease, but were present in bone marrow cells, although immunohistochemistry showed they were not within cytoplasmic granules. The concentrations of elastase and cathepsin G in Chediak–Higashi bone marrow were about 25 and 15%, respectively, of those in normal marrow. Quantitative scanning of northern blots showed that elastase and cathepsin G mRNA, corrected for β-actin mRNA, were expressed equally in normal marrow.

**Conclusions**—Transcription of elastase and cathepsin G mRNA in promyelocytes of patients with Chediak–Higashi disease is normal, but the protein products are deficient in these cells and absent in mature neutrophils. This suggests that the translated proteins are not packaged into azurophil granules but are degraded or secreted from the cells.

Keys words: Chediak–Higashi syndrome, elastase, cathepsin G.

Chediak–Higashi syndrome is an autosomal recessive disorder characterised by abnormal cytoplasmic granules in a variety of cells. A major sequela of Chediak–Higashi syndrome is recurrent infection, which is thought to result from inefficient microbial killing by leucocytes, despite normal phagocytic activity and a normal respiratory burst. This defect therefore appears to be associated with a deficiency in killing of bacteria by microbialic proteins in phagolysosomes. Blood neutrophils of patients with Chediak–Higashi syndrome are deficient in the azurophil granule enzymes cathepsin G and elastase. Myeloperoxidase and the defensins, however, are present in normal or slightly reduced amounts, myeloperoxidase having been localised to the granules of the cells.

Cathepsin G and elastase are also deficient in blood neutrophils of the Chediak–Higashi animal model, the beige mouse. The bone marrow of the beige mouse, however, contains about half the elastase and cathepsin G content of normal mouse marrow, within neutrophil precursors but not mature cells. These results suggest that in this animal model the cathepsin G and elastase genes are transcribed and protein translation does occur, but the enzymes do not survive to the stage of mature cells. It is not known whether a similar situation exists in human Chediak–Higashi disease, or indeed whether transcription of these proteins is normal. The purpose of the present study was to analyse the messenger RNA (mRNA) and protein content of cathepsin G and elastase in the bone marrow of patients with Chediak–Higashi syndrome.

**Methods**

Three patients with Chediak–Higashi syndrome were studied. Diagnosis was based on a history of recurrent infection, depigmented areas of skin, hair or retina, and characteristic giant granules in leucocytes. None had evidence of neuropathy.

Patient JF is a 25-year-old man who was diagnosed when aged 13 years. He has suffered from recurrent gingivitis, eventually requiring extraction of all his teeth, and multiple skin infections which have been treated with antibiotics and occasional surgical drainage. He has had no major chest or urinary tract infections.

Patient GH is a 10-year-old boy, diagnosed when five years old. He has had a history of frequent respiratory tract and throat infections, oral ulceration, paronychia, and skin bruising following minor trauma. Relatively minor abrasions to the skin were delayed in healing and often became septic.

Patient AE is a four year old girl with a history of recurrent infections of the urinary tract, respiratory tract and throat from the age of six weeks. She has recently suffered from mouth ulcers and paronychia.

Blood neutrophils were purified as described by Jepsen and Skottum. Venous blood was collected, from the three patients and from a
healthy volunteer, into lithium heparin anticoagulant and each blood sample was diluted with an equal volume of 0·15 mol/l NaCl solution. The diluted blood (5 ml) was layered onto two layers of Percoll (Pharmacia AB, Uppsala, Sweden); the upper layer was 2 ml 1·075 g/ml Percoll, the lower 3 ml 1·096 g/ml Percoll. The tubes were centrifuged at 2000 × g for 20 minutes. The neutrophils were harvested from the interface between the Percoll layers, washed three times in TRIS buffered RPMI 1640 medium (Flow Laboratories, Rickmansworth, UK), counted and resuspended in RPMI 1640 medium. The polymorphonuclear leucocytes (PMN) were >96% pure and >98% viable (assessed by exclusion of trypan blue). The neutrophil preparations from each subject were separated into two samples, centrifuged and the cell pellets retained. One sample from each subject (for enzyme activity assays) was lysed by the addition of 0·5 M Tris/HCl, pH 7·8, containing 1·0 M NaCl and 0·1% (v/v) Triton X-100 and sonicated (6 × 20 W) using a Lucas Dawe Ultrasones Sonprobe. The second sample (for western immunoblotting and enzyme linked immunosorbent assay (ELISA)) was lysed by sonication with 0·5 M Tris/HCl, pH 7·8 containing 1·0 M NaCl and 0·1% (v/v) Triton X-100, after the addition of protease inhibitors; 25·5 µl 100 mM diisopropyl fluorophosphosphate dissolved in dimethyl sulphoxide, 0·1 ml 0·1 M 1,10 phenanthroline in methanol and 0·1 ml 0·1 M iodoacetate in water. The neutrophil lysates were stored at −70°C until analysed.

Bone marrow (5 ml) was aspirated from the superior iliac spine of one of the patients (JF) after obtaining informed consent. After removal of a sample for routine haematological analysis, the remainder was dispensed into 30 ml RPMI 1640 culture medium, containing 10% (v/v) fetal calf serum and 2000 units of heparin anticoagulant. A portion of bone marrow was also retrieved from a haematologically normal bone marrow donor. The marrow samples were centrifuged at 200 × g for 20 minutes through 5 ml 1·096 g/ml Percoll to remove erythrocytes and erythroid precursors. The cells remaining above the Percoll layer were collected, washed in RPMI 1640 medium and cytospin preparations made for differential cell counts and immunochemistry. The remainder of the cells were counted and lysates were obtained for western blotting and enzyme assays as for blood neutrophils. Preparations of mRNA were also obtained from bone marrow for northern blotting. The mRNA was prepared from the bone marrow using a Promega PolyATtract System 1000 kit (Promega, Southampton, UK) and finally solubilised and stored as described by Chomczynski.9

Three complementary DNA (cDNA) probes were used. The elastase probe10 was a 516 base pair partial cDNA (supplied by Dr G Salvesen, University of Georgia, Athens, USA) subcloned into the EcoR1 site of pBluescript KS(−/−). The cathepsin G cDNA11 (supplied by Dr T Ley, Jewish Hospital, St Louis, USA) was 1·0 kilobase subcloned into the EcoR1 site of Bluescript. A β-actin probe12 (also supplied by Dr T Ley) was a 1·9 kilobase cDNA in the BamH1 site of Puc9.

Northern blotting was performed essentially as described by Maniatis et al.13 The mRNA preparations (about 10 µg) from normal and Chediak–Higashi bone marrow samples, suspended in denaturing buffer (MOPS containing formamide and formaldehyde), were subjected to electrophoresis in 1% (w/v) agarose gels in 0·2 M MOPS buffer (pH 7·0) containing 2·2 M formaldehyde, 50 mM sodium acetate and 1 mM EDTA. Following electrophoresis, the RNA samples were transferred by capillary blotting to Genescreen TM (NEN Research, Boston, USA). The Genescreen was prehybridised for 12 hours at 42°C before the addition of the cathepsin G probe or the elastase probe (100 ng), radiolabelled with [32P]dCTP to 10⁹ dpm/mg DNA using a commercial random prime labelling kit (Boehringer Mannheim, Lewes, UK). Hybridisation took place for 18 hours at 42°C in the presence of 50% formamide. The screens were then washed four times with 2 × standard sodium citrate (SSC) at 20°C and twice with 0·2 × SSC at 65°C (1 × SSC = 0·15 M NaCl, 0·015 M sodium citrate, pH 7·0). The screens were autoradiographed at −70°C and that which had been probed with the elastase cDNA was scanned with LKB Wallach 1205 β plate flat bed counter in the scintillation mode. When the radioactivity of the elastase probe scanned had decayed to undetectable levels, it was re-probed with radiolabelled β-actin cDNA, autoradiographed and scanned again. The ratios of radioactivity caused by hybridisation of elastase mRNA and β-actin mRNA in normal and Chediak–Higashi bone marrow samples were calculated from the scans.

Neutrophil and bone marrow lysates were subjected to sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE), as described by Laemmli,14 on an LKB "Midi-" system (Pharmacia LKB Biotechnology, Milton Keynes, UK) using a 5% polyacrylamide stacking gel and 12·5% running gel. Each sample loaded contained lysate from 10⁵ cells. Following electrophoresis, the proteins were transferred to nitrocellulose paper (Hybond-C; Amersham International) by electroblotting. The nitrocellulose sheets were incubated with either sheep antineutrophil elastase15 or sheep anticathepsin G16 followed by donkey ant sheep IgG labelled with horseradish peroxidase (Binding Site, Birmingham, UK). Binding of peroxidase labelled antibody was visualised using 3-amino-9-ethylcarbazole as the substrate.

Cathepsin G was measured in blood neutrophil and bone marrow lysates using a direct capture ELISA. Nunc Immuno 96 well plates were loaded with 0·2 ml sheep antihuman cathepsin G, 20 µl/ml in phosphate buffered saline (PBS) (pH 7·2) and incubated overnight at 4°C. The wells were washed three times with washing buffer (PBS containing 0·1% Tween-20 and 1% (v/v) Haemaccel (Hoechst)) before the addition of 0·2 ml of each cell lysate diluted in washing buffer with 0·5 M NaCl. Reference
standards consisted of dilutions of cathepsin G, purified from an empyema as described by Martodam et al. The plates were incubated for two hours at room temperature before washing, followed by the addition of a horseradish peroxidase conjugated sheep antiacathepsin G. After a further two hours of incubation, the wells were washed and substrate added; 0·2 ml/well 1 mg/ml 0-phenylenediamine dihydrochloride (Sigma) in 0·1 M citrate/phosphate buffer (pH 5·0) with 1 μl/ml hydrogen peroxide. The colour reaction was stopped with 50 μl 0·5 M citric acid/well and the absorbance measured with a Dynatech MR 5000 plate reader at 490 nm. Cathepsin G concentrations in the cell lysates were calculated by interpolation from the reference values obtained with pure enzyme. The lower limit of detection was 3 ng/ml.

Neutrophil elastase in cell lysates was measured using an indirect competition ELISA. Nunc Immuno 96 well plates were loaded with 0·2 ml elastase, 1 μg/ml in PBS, pH 7·2, with 0·5 M NaCl and incubated overnight at 4°C. The cell lysates and dilutions of elastase, purified from an empyema, were each mixed with equal volumes of 25 μg/ml sheep antielastase, which were also incubated overnight at 4°C. The ELISA plates were washed and the lysates and reference mixtures (0·2 ml/well) added and incubated for two hours at room temperature. After further washing, the plate wells were loaded with 0·2 ml horseradish peroxidase conjugated donkey antisheep IgG (Binding Site) and incubated for two hours at room temperature, before the addition of substrate solution and measurement of the absorbance at 490 nm, as described earlier. The elastase concentrations in the cell lysates were calculated by interpolation from the inhibition binding reference line obtained with pure elastase, the lower limit of detection being 3 ng/ml. Results for both enzymes were expressed as pg protein/myeloid cell.

Cathepsin G and elastase activities in blood neutrophil and bone marrow lysates were measured in microtitre plate wells using fluorimetric substrates, Succ-Ala-Ala-Pro-Phe-paranitroanilide and Succ-Ala-Ala-Ala-paranitroanilide, respectively (Bachem Feinchemikalien, Bündorf, Switzerland). Samples (40 μl) of the cell lysates and dilutions of calibration standards of pure cathepsin G or elastase were mixed with 140 μl substrate buffer (0·05 mol/l TRIS/HCl buffer containing 0·5 mol/l NaCl and 1% (v/v) Triton X-100 and 1 mg/ml of the appropriate substrate). The plates were incubated at room temperature until colour change was sufficient and the absorbance of each well was measured at 410 nm with a Dynatech MR 5000 plate reader. Cathepsin G and elastase concentrations in cell lysates were calculated by interpolation from the calibration curves obtained with pure enzymes. The lower limits of detection for cathepsin G and elastase activity were 190 and 150 ng/ml, respectively. Results were expressed as activity representing pg enzyme/myeloid cell.

The cytocentrifuge preparations from the bone marrow samples were stained immuno-histochemically for cathepsin G and elastase as described previously.

**Results**

**NORTHERN BLOTTING**

Autoradiographs of northern blots obtained

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**Figure 1 Northern blotting with cDNA probes for cathepsin G (CATH G), elastase and β-actin in bone marrow.** The figure shows the results of blotting bone marrow myeloid cell mRNA from a normal donor (n) and a patient with Chediak-Higashi disease (ch). The transcripts were of different sizes but the blots are aligned for convenient comparison.
with mRNA from normal and Chediak-Higashi bone marrow samples using the elastase, cathepsin G and β-actin cDNA probes are shown in fig 1. Although we attempted to load similar amounts of RNA from normal and Chediak-Higashi marrow samples, it is evident, by comparing the intensities of the resulting β-actin autoradiographs, that the normal marrow samples contained more mRNA. Nevertheless, visual comparison of the autoradiographs suggested that the ratios of cathepsin G:β-actin and elastase:β-actin were similar in bone marrow samples from both subjects. This was confirmed by the quantitative scanning of the elastase and β-actin blots. The ratio of elastase:β-actin transcripts in the normal marrow was 0·34 and that for the Chediak-Higashi marrow was 0·40, suggesting that the two marrow lys-

Figure 2 Western blotting for elastase and cathepsin G in blood neutrophils. The figure shows western blotting for cathepsin G (A) and elastase (B) in blood neutrophil lysates. Lane 1, molecular weight markers; lane 2, neutrophils from the healthy subject; lane 3, neutrophils from a patient with Chediak-Higashi disease. Normal neutrophils contain both enzymes with apparent molecular weights of 29 500 (arrows).

Figure 3 Western blotting of bone marrow. The figure shows western blotting for cathepsin G (A) and elastase (B) in lysates from bone marrow cells. Lane 1, bone marrow from a patient with Chediak-Higashi disease; lane 2, normal marrow. The lysates from normal and Chediak-Higashi bone marrow show cathepsin G and elastase bands of molecular weights consistent with mature active protein (arrows). The Chediak-Higashi marrow lysates also show elastase and cathepsin G bands of smaller sizes.
Cathepsin G and elastase were detected, using ELISA, in the bone marrow from the Chediak-Higashi patient. Cathepsin G was measured at a concentration of 0·23 pg/myeloid cell, compared with a concentration in normal marrow of 0·97 pg/cell. Elastase was also present in the Chediak-Higashi marrow at a concentration of 0·37 pg/myeloid cell, compared with the normal marrow value of 2·3 pg/cell. Thus, Chediak-Higashi marrow myeloid cells contained 24 and 16% of the cathepsin G and elastase proteins, respectively, found in the normal marrow. Cathepsin G activity in the marrow cells represented 0·09 pg/cell in Chediak-Higashi marrow and 0·93 pg/cell in normal marrow. The elastase activity in Chediak-Higashi marrow was 0·15 pg/cell and was 1·2 pg/cell in normal marrow. Thus, cathepsin G and elastase activity in Chediak-Higashi marrow cells represented 9·8 and 15·8%, respectively, of that in normal marrow cells.
IMMUNOHISTOCHEMISTRY OF CYTOSPUN PREPARATIONS

Figure 4 shows the results of immunohistochemical staining of bone marrow, from the Chediak–Higashi patient and a normal donor, for cathepsin G. In the normal marrow cathepsin G could be seen within the granules of the cells, whereas the Chediak–Higashi promyelocyte granules showed no staining.

Discussion

In the present study we have confirmed the observation of Ganz et al.\(^8\) that elastase and cathepsin G are undetectable in mature blood neutrophils from patients with Chediak–Higashi disease. The fundamental cause of neutrophil cathepsin G and elastase deficiency in Chediak–Higashi syndrome has not, however, been established. Elastase, coded for on chromosome 19pter\(^18\) and cathepsin G, on chromosome 11q14-2,\(^11\) are both transcribed, synthesised and packaged within the promyelocyte during myeloid differentiation in the bone marrow,\(^11\) when the azurophil granules are formed;\(^22\) transcription and synthesis cease before the cells become mature neutrophils. Both enzymes are translated as pro-proteins and thence processed via enzymatically inactive pro-proteins to the mature active enzymes.\(^22\) Activation from the pro-enzyme appears to involve the proteolytic removal of an acidic N terminal dipeptide, Ser-Glu in the case of elastase and Gly-Glu for cathepsin G, and C terminal extensions of 20 and 11 amino acids in elastase and cathepsin G, respectively. The N terminal sequences of the mature proteins share considerable homology, suggesting common dipeptidyl peptidases are responsible for processing the pro-proteins.\(^22\) The C terminal extensions of the pro-proteins are not homologous. The mechanisms determining packaging to the azurophil granules are not known. The deficiency of these enzymes in Chediak–Higashi syndrome could potentially occur at any of these several stages of processing. The identification of the mature enzymes within bone marrow promyelocytes of the beige mouse suggested, however, that transcription of the genes for these enzymes and translation of the proteins do occur. Thus, the deficiency would be a consequence of post-translational events.

In the present study we have shown that in bone marrow from a patient with Chediak–Higashi syndrome the levels of the mRNA transcripts for cathepsin G and elastase, when corrected for \(\beta\)-actin mRNA levels, were similar to those of normal marrow. Thus, if \(\beta\)-actin transcription is normal in Chediak–Higashi myeloid cells the results would suggest that transcription both of cathepsin G and elastase is also normal.

The bone marrow myeloid cells of the patients with Chediak–Higashi syndrome contained mature, active elastase and cathepsin G. As in the beige mouse, the concentrations of these enzymes were lower than those of marrow from a normal subject and undetectable in mature blood neutrophils. The Chediak–Higashi marrow protein concentrations were only about one fifth of those seen in normal marrow. This is less than the value for the beige mouse marrow, which was reported to be about 60% of normal.\(^6\)

Enzyme activity measurements in marrow samples were slightly lower than the protein concentrations estimated using ELISA. This difference may represent a technical discrepancy. Alternatively, it is possible that some inactive, unprocessed, pro-elastase and pro-cathepsin G were present within the marrow cells, although there was no evidence for this on the western blots. The western blots of elastase and cathepsin G from Chediak–Higashi bone marrow samples showed, in addition to molecular weight bands consistent with the mature proteins, smaller forms which were not observed in lysates of normal marrow or of blood neutrophils. These small forms could represent differences in glycosylation. Alternatively, they might represent proteolysed forms although proteinase inhibitors were added to the marrow and neutrophils before lysis to prevent proteolytic modification of proteins during the processing of samples for blotting. The presence of the “smaller” form of elastase and cathepsin G in Chediak–Higashi marrow suggests, therefore, that some proteolytic degradation may have occurred before sampling—that is, in vivo. This has relevance to the possible fate of these proteins in Chediak–Higashi marrow, but further studies would be necessary to establish the reasons for this observation. Nevertheless, the presence of active elastase and cathepsin G in the myeloid cells of the Chediak–Higashi marrow shows that these proteins had been synthesised and processed to mature, active enzymes. This indicates that a deficiency in proteinases involved in processing of the pro-enzymes is not responsible for the absence of elastase and cathepsin G in Chediak–Higashi neutrophils.

Takeuchi and Swank\(^3\) characterised inhibitors of elastase and cathepsin G, which formed covalent complexes with the enzymes, in mature neutrophils of beige mice; these inhibitors were absent from normal mouse cells and beige mouse pro-myelocytes. It was suggested that the absence of elastase and cathepsin G activities in beige mouse neutrophils was because of the presence of these inhibitors. This would not, however, explain the absence of immunoreactive elastase and cathepsin G in bone marrow neutrophils, either as native proteins or complexed with putative inhibitors. In addition, the presence of elastase and cathepsin G within marrow myeloid cells, but not apparently within the cells’ granules, suggests that these enzymes are absent from mature neutrophils in Chediak–Higashi syndrome because of a failure of transport to, or packaging within, the primary granules of the marrow promyelocytes.

Granule defects are a common feature of several cell types in Chediak–Higashi syndrome, including cells which do not express cathepsin G or elastase.\(^1\) This suggests that a fundamental defect in granule structure may
prevent the packaging of some, but not all, neutrophil azurophil proteins. Presumably elastase and cathepsin G, but not myelo-peroxidase or the defensins, share common requirements for packaging. Little is known, however, about the mechanisms responsible for intracellular transport of azurophil proteins. It is likely that failure to package cathepsin G or elastase successfully may result in these proteins being degraded intracellularly, resulting in their absence from mature neutrophils, following cessation of synthesis. This would explain the smaller molecular weight forms of these enzymes observed on the western blots from the Chediak–Higashi bone marrow. Should this hypothesis be correct, the detection of cathepsin G and elastase activity within Chediak–Higashi marrow cells would suggest that post-translational processing of these enzymes takes place before the proteins enter the azurophil granules. The site of processing events has, to date, been unclear.22

In conclusion, our results suggest that the absence of elastase and cathepsin G in Chediak–Higashi blood neutrophils results from a primary defect in azurophil granule structure which precludes the successful packaging of these enzymes, but permits that of other azurophil granule proteins. Further studies are indicated to identify the fundamental defect in Chediak–Higashi disease and to establish whether failure of phagolysosome formation or the deficiencies of specific granule proteins are responsible for defective bacterial killing by leucocytes in this disease.

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