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Molecular Cloning and Sequencing of Three Granulocytic *Ehrlichia* Genes Encoding High-Molecular-Weight Immunoreactive Proteins

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Granulocytic *Ehrlichia* was isolated from canine blood obtained from animals challenged with field-collected *Ixodes scapularis* and propagated in HL60 cells. PCR primers specific for the 16S ribosomal DNA (rDNA) of the *Ehrlichia* genogroup comprising *E. equi*, *E. phagocytophila*, and the agent of human granulocytic ehrlichiosis (HGE) amplified DNA from extracts of these cells. Sequence analysis of this amplified DNA revealed that it is identical to the 16S rDNA sequence of the HGE agent. A genomic library was constructed with DNA from granulocytic *Ehrlichia* and screened with pooled sera from tick-challenged, granulocytic *Ehrlichia*-infected dogs. Several clones were isolated and sequenced. Three complete genes encoding proteins with apparent molecular masses of 100, 130, and 160 kDa were found. The recombinant proteins reacted with convalescent-phase sera from dogs and human patients recovering from HGE. This approach will be useful for identifying candidate diagnostic and vaccine antigens for granulocytic ehrlichiosis and aid in the classification of genogroup members.

Members of the genus *Ehrlichia* include species which have a tropism for mononuclear phagocytes (*E. canis*, *E. chaffeensis*, *E. muris*, *E. sennetsu*, and *E. risticii*) and those which infect granulocytes (*E. ewingii*, *E. phagocytophila*, and *E. equi* [14, 26], and the recently discovered agent of human granulocytic ehrlichiosis [HGE] [3, 7]). Disease caused by granulocytic *Ehrlichia* (GE) is manifested by fever, lethargy, thrombocytopenia, and death, and many species from diverse geographical locations have shown evidence of natural infection, including horses (25, 27, 29, 38), dogs (24, 34, 35), small mammals (39, 40), and humans (4, 15).

The similar host range and near identity of the 16S rRNA genes of *E. phagocytophila*, *E. equi*, and the HGE agent (7) have raised the possibility that these organisms represent a single species (2, 28). In addition, Dumler et al. (9) have shown that they share significant antigenicity by immunofluorescence and immunoblot assays. Objective methods for species classification, e.g., molecular genetic analysis, have not been readily available, primarily because of an inability to culture these ehrlichiae in vitro. However, we (reference 43 and unpublished data) and others (13) have recently demonstrated successful cultivation of GE isolates from dogs and humans, respectively.

In this paper, we describe the use of purified GE obtained from in vitro culture of infected HL60 cells, a promyelocytic human cell line, to generate a genomic DNA library for expression screening with sera obtained from dogs experimentally infected with GE. The screening resulted in the isolation of recombinant clones containing complete genes encoding three putative proteins of GE, GE 160, GE 130, and GE 100 (named for apparent molecular mass in kilodaltons). One of these proteins, the 100-kDa protein, is similar in both glutamic acid content and repeated amino acid structure to an immunodominant 120-kDa *E. chaffeensis* protein (45). Both the 100- and 130-kDa granulocytic *Ehrlichia* proteins share some amino acid sequence homology to the 120-kDa *E. chaffeensis* protein.

**MATERIALS AND METHODS**

Isolation of GE in cell culture. The GE agent (strain USG3) was isolated as described previously (45). Briefly, blood from a dog experimentally infected by adult *Ixodes scapularis* ticks collected from both Westchester County, N.Y., and Montgomery County, Pa., was inoculated into a suspension culture of the human promyelocytic cell line HL60 (ATCC CCL 240). The cells were cultured in RPMI 1640 medium supplemented with 20% heat-inactivated fetal bovine serum, 2 mM L-glutamine, 1 mM sodium pyruvate, and 0.1 mM nonessential amino acids and maintained at 37°C and 6% CO2 in a humidified chamber. The fetal bovine serum was reduced to 10% once the culture was established. Ehrlichial infection was indicated by lysis of the HL60 cells. Cultures of USG3 (passages 7 to 21) were grown for 5 days prior to GE purification.

Purification of GE. USG3 cultures at approximately 80% cell lysis (monitored microscopically) were centrifuged at 840 × g for 15 min at 4°C to remove host HL60 cell debris. The supernatant was filtered through a Poretics (Livermore, Calif.) 5-μm-pore-diameter polycarbonate membrane, 47 mm in diameter, followed by a Poretics 3-μm-pore-diameter filter under negative pressure. The USG3 filtrate was centrifuged at 9460 × g in a Sorvall centrifuge for 30 min at 4°C. Following centrifugation, the GE pellet was resuspended in 5 ml of 25 mM Tris (pH 8.0)–10 mM MgCl2–0.19% NaCl. DNase I (Life Technologies, Gaithersburg, Md.) was added to a final concentration of 9 μg per ml, and the solution was incubated for 15 min at 37°C. Following incubation, the DNase was inactivated by the addition of 0.5 ml of 0.5 M EDTA, and the GE was pelleted at 14,000 × g in a Sorvall centrifuge for 30 min at 4°C.

PCR amplification and cloning of GE 16S ribosomal DNA. Universal eubacterial primers for the 16S rRNA gene (41) were modified to include restriction enzyme recognition sites as follows: forward primer, 5′ CTGACAGGTGTTGATC CTGG 3′ (PstI site); reverse primer, 5′ GGATCCTACCTTGTTACGACTT 3′ (BamHI site). These primers (0.5 μM) were added to a 100-μl reaction mixture containing 1× PCR buffer II (Perkin-Elmer Corp.); 1.5 mM MgCl2 (Perkin-Elmer Corp.); 200 μM (each) dATP, dCTP, dGTP, and dTTP; 2.5 U of AmpliTaq DNA polymerase; and 20 μl of USG3 DNA. Amplification was performed under the following conditions: 35 cycles at 94°C for 1 min, 53°C for 1 min, and 72°C for 2 min, followed by a 72°C incubation for 10 min. The amplified 1,500-bp fragment was digested with PstI and BamHI and ligated to pUC19 linearized with the same enzymes. The resulting clone, pUCHGE16S, was sequenced as described below.
Construction of a GE genomic library. Genomic DNA was isolated from purified GE with the QIAamp Genomic DNA kit (Qiagen, Chatsworth, Calif.) for library preparation (Stratagene, La Jolla, Calif.). The DNA was mechanically sheared to a 4- to 10-kb size range and ligated to EcoRI linkers. Linkered fragments were ligated into the EcoRI site of Lambda Zap II, and the library was amplified in Escherichia coli XL1-Blue MRF to a titler of 10^9 PFU/ml.

**Expression screening of the genomic library.** Bacteriophage were plated with XL1-Blue MRF^®^ and induced to express protein with 10 mM isopropyl-β-D-thiogalactopyranoside (IPTG) (Sigma, St. Louis, Mo.). Proteins were transferred to nitrocellulose filters, and the filters were washed with Tris-buffered saline (TBS: 25 mM Tris HCl [pH 7.5], 0.5 M NaCl). Washed filters were blocked in TBS containing 0.1% polyoxyethylene 20 cetyl ether (Brij 58) and incubated with rabbit anti-dog horse-radish peroxidase-conjugated immunoglobulin (Ig) antibody, washed, and developed with 4-chloronaphthol. Positive plaques were isolated, replated, and screened again. Plasmid DNA containing the putative recombinant clones was obtained by plasmid rescue (Stratagene).

**DNA sequencing and sequence analysis.** DNA sequencing of recombinant clones was performed by the primer walking method and with an ABI 373A DNA sequencer (ACGT, Northbrook, Ill.; Lark Technologies, Houston, Tex.; and Sequeneng, Shrewsbury, Mass.). Sequences were analyzed by the MacVector (Oxford Molecular Group) sequence analysis program, version 6.0. The BLAST algorithm (Altschul, Missouri) was used to search for homologous nucleic acid and protein sequences available on the National Center for Biotechnology Information (NCBI) server.

**PCR analysis of USG3 and IL60 DNA.** PCR primer sets were designed based on the sequences of each of the three GE clones and are as follows: S2 (forward, 5'-GCGTCTCCAGAACCGA-3'; reverse, 5'-CTCTGTTGCCATAGG-3'); S7 (forward, 5'-GATGGTCCTCAGCATACGCCGC-3'; reverse, 5'-CAGAGATTACTCTTITGCGG-3'); and S22 (forward, 5'-CAGGCCTCTCTCTAC-3'; reverse, 5'-CCCTGTTGCTTATTGACG-3'). Each 50-μl reaction mixture contained 0.5 μM each primer, 1× PCR Supermix (Life Technologies, Gaithersburg, Md.) and either 100 ng of USG3 DNA, 100 ng of HL60 DNA, or 200 ng of plasmid DNA. PCR amplification was performed under the following conditions: 94°C for 30 s, 55°C for 30 s, and 72°C for 1 min. After 30 cycles, a single 10-min extension at 72°C was done. PCR products were analyzed on 4% NuSieve 3:1 agarose gels (FMC Bioproducts, Rockland, Me.).

**Western blot analysis.** Individual recombinant plasmid-containing cultures were induced to express protein with 5 mM IPTG. Bacterial cells were pelleted by centrifugation and resuspended in 5× Laemmli buffer (12% glycerol, 0.2 M Tris HCl [pH 6.8], 5% sodium dodecyl sulfate [SDS], 5% β-mercaptoethanol) at 200 μl per optical density unit of culture. Samples were boiled for 5 min, and 10 μl of each was analyzed by SDS-polyacrylamide gel electrophoresis (PAGE) (for dog sera) or NuPage (NoveX, San Diego, Calif.) electrophoresis (for human sera). Proteins were transferred to nitrocellulose filters, the filters were blocked in TBS-Brj 58, and the blots were probed with either the pooled dog sera referenced above or 1:1,000 dilutions of human sera. The study was done on pooled convalescent-phase serum samples from patients (no. 2 and 3 from New York, kindly provided by M. Aguero-Rosenfeld) and one sample from an individual in Wisconsin who was part of a seroprevalence study (no. 1, kindly provided by J. Bakken). Blots were washed and incubated with biotin-labeled goat anti-dog IgG (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, Md.) followed by peroxidase-labeled streptavidin (Kirkegaard & Perry Laboratories, Inc.) or horseradish peroxidase-conjugated antihuman IgG (Bio-Rad, Hercules, Calif.). After several additional washes, the dog sera were probed with biotinylated 4-chloronaphthol (Bio-Rad), and the human serum blots were detected by enhanced chemiluminescence (Pierce, Rockford, Ill.) and viewed by autoradiography.

**Nucleotide sequence accession number.** The nucleotide sequences of the GE genes described here have been assigned the following GenBank accession numbers: ank (GE 160), AF020521; rec (GE 130), AF020522; and gru (GE 100), AF020523.

**RESULTS**

**Cloning and sequencing of the USG3 16S rRNA gene.** Sources of GE used to generate a cell culture isolate have included blood from either infected humans (13) or dogs (32, 43). To assess the relatedness of the USG3 isolate to other GE isolates, including the HGE agent, the 16S rRNA gene was amplified by PCR with universal eubacterial primers (41). A 1,500-bp DNA fragment was isolated and cloned into pUC19 (see Materials and Methods), and the insert DNA was sequenced. The USG3 16S rRNA gene sequence was found to be identical to the GenBank sequence of the HGE agent (accession no. U02521). The sequences of other 16S ribosomal DNA PCR fragments generated with USG3 DNA as the template were also identical to the HGE agent sequence (reference 43 and data not shown).

**Isolation of recombinant clones identified with GE-positive dog sera.** Using pooled sera from adult I. scapularis-challenged, GE-infected dogs, positive clones were identified and purified as single plaques. pBluescript plasmids were rescued according to the Stratagene protocol, and DNA was purified with Qiagen plasmid purification kits. A number of restriction digests were performed with each clone to assess their relatedness (data not shown). Based on these digests, restriction maps were generated which showed that all of the clones represented just three different GE genes. One clone from each of the three gene groups (clones S2, S7, and S22) was chosen for further analysis.

**Analysis of proteins encoded by GE clones.** Because recombinant clones were selected based on immunoreactivity, bacterial lysates of each clone were analyzed by SDS-PAGE and Western blotting to identify the immunoreactive proteins. The blots were probed with the same pooled canine sera used to screen the library. Figure 1A shows the reactivity of the anti-sera against purified USG3. Several immunodominant proteins with sizes of 36, 43, and 45 kDa were observed, and other less-immunoreactive proteins of various molecular masses are observed.
shown. Figure 1B shows the Western blot of the library clones, S2, S7, and S22. A single high-molecular-mass protein (indicated by arrows) was detected by the sera for each isolated clone. These proteins were not expressed in the absence of IPTG induction and were not detected with preimmune dog sera (data not shown). The approximate molecular masses of each protein are as follows: 160 (S2), 100 (S7), and 130 (S22) kDa. These proteins are not the major immunodominant antigens of the purified cell culture isolate USG3, two of which migrate at about 43 to 45 kDa (Fig. 1A), but they could be present among the higher-molecular-mass immunoreactive proteins.

The three GE proteins were also analyzed by Western blotting for their reactivity with human sera. Figure 2 shows three different human serum samples tested against purified USG3 antigen and the three GE recombinant proteins. Each serum sample reacted strongly with one of the three GE proteins: no. 1 with GE 160 and less strongly with GE 130, no. 2 with GE 100, and no. 3 with GE 130. All three serum samples reacted with several proteins in the purified USG3 lanes (GE lanes).
serum sample from a Rhode Island HGE patient recognized two of the three GE proteins, and other HGE patient sera tested also recognized one or two of the proteins (data not shown). None of the samples we have tested to date react strongly with all three recombinant GE proteins. Negative control human sera did not react with the GE proteins on Western blots (data not shown).

It is interesting to note that these human sera do not strongly recognize proteins with sizes of 100, 130, and 160 kDa with USG3 lysates as the antigen source. This may be due to underrepresentation of these proteins in the purified cell culture antigen, either because they are secreted or because they are expressed only after infection of a host species.

DNA sequencing and database analysis of recombinant clones. The inserts from each of the recombinant plasmids S2, S7, and S22 were completely sequenced by the primer walking method. Sequence analysis (MacVector 6.0; Oxford Molecular Group) showed that each clone contained a single large open reading frame encoded by the plus strand of the insert and that each one appeared to be a complete gene. The portions of the DNA sequence containing the open reading frame and corresponding amino acid sequence of each of the clones are shown as follows: Fig. 3, S2, GE 160; Fig. 4, S7, GE 100; and Fig. 5, S22, GE 130. A possible ribosome binding site sequence upstream from the initiating ATG codon is underlined in each figure.

The numbers of nucleotides sequenced upstream of each reading frame were 1,500 bp for the S2 insert, 500 bp for the S22 insert, and 200 bp for the S7 insert. These sequences were examined for likely ehrlichial promoter sequences based on homology to the E. coli -35 and -10 consensus sequences (16). Of the three clones, S2 had the closest homology to the E. coli promoter sequences, and these are underlined in Fig. 3. A number of possible promoter sequences were found for the S2 and S7 genes, but in the absence of direct experimental evidence, it is difficult to determine which sequence elements are necessary for promoter function. Protein expression in all three clones was IPTG dependent, and thus it is likely that the vector lacZ promoter is used for transcription of the Ehrlichia RNA in E. coli.

Amino acid sequence analysis of the DNA sequences of the three gene clones showed that all contain regions of repeated amino acids. A schematic version of these repeat structures is shown in Fig. 6 and 7. The 160-kDa protein (Fig. 6) has three groups of repeats. The first set consists of a number of ankyrin-like repeat units of 33 amino acids, the second consists of repeat units of 27 amino acids, and the third consists of repeat units of 11 amino acids. The ankyrin repeats were revealed by a BLAST database search for protein homologies. Ankyrin repeats occur in at least four consecutive copies and are present in yeast, plants, bacteria, and mammals (5, 6). Figure 6 shows a multiple alignment of the 160-kDa protein ankyrin repeats under a consensus sequence derived from the analysis.
of several hundred similar ankyrin-like motifs (6). The eighth repeat sequence holds to the consensus only through the first half of the repeat unit and may not represent a full ankyrin-like repeat.

The 130-kDa protein (Fig. 7) has a repeat unit of 26 to 34 amino acids which occurs eight times in the carboxy-terminal half of the protein. The sequence varies somewhat from repeat to repeat. A database search revealed that it is similar to the 120-kDa protein (45). An amino acid sequence alignment of a motif common to both proteins is shown in Fig. 7. This motif is represented by a bold line and occurs four times in an identical fashion in the E. chaffeensis 120-kDa protein (designated A) and eight times in the GE 130-kDa protein (a-1 to a-4). Sequence a-1 is repeated three times, related sequences a-2 and a-3 are each repeated twice, and related sequence a-4 is found once. The positions of these sequences in the proteins are indicated by the small bold lines. (B) Alignments of two different sequence motifs which occur in the E. chaffeensis 120-kDa protein (B-1 to B-3 and C-1) and the GE 100-kDa protein (b-1 and c-1). Bold and cross-hatched boxes indicate the positions of these sequences in the proteins. Identical amino acids are surrounded by boxes, and conserved amino acids are in capital letters.

Verification that clones are GE-derived with PCR analysis. Based on the DNA sequences of each clone, PCR primers were designed to amplify specific regions of each open reading frame (see Materials and Methods). Primer pairs specific for S22, S2, and S7 were used in separate PCRs to amplify three different templates: GE DNA (from USG3), HL60 DNA, or...
the 160-, 130-, and 100-kDa proteins. Aside from the 16S We also describe three previously unidentified proteins of GE, cern United States (3, 7). However, classification of GE has for disease in patients in the upper midwestern and northeast-

E. chaffeensis is responsible for a tick-borne fever of cattle, sheep, and goats and occurs primarily in Europe (10, 33), and E. equi is found in horses in North and South America (14, 26). The agent of HGE has been identified as the organism responsible through the proteins GE 100 and GE 130.

The presence of repeated regions in the 100-, 130-, and 160-kDa proteins may in part explain the differences between the expected molecular masses and the observed molecular masses on SDS-PAGE, which are much higher. This phenomenon has been reported for other proteins containing repeated regions, including rickettsial proteins (1, 36), malaria antigens (21), the fibronectin-binding protein from Staphylococcus au-

Investigators have speculated that high percentages of certain amino acids in these proteins or posttranslational modifications may contrib-

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A search of the major protein and nucleotide databases with the NCBI BLAST algorithm did not reveal any significant similarities of the GE proteins to any known proteins, other than the ankyrin repeats of the 160-kDa protein and the homology between the 100-kDa protein or 130-kDa protein and the 120-kDa protein of E. chaffeensis. The function of the E. chaffeensis 120-kDa protein is not known, but the hydrophilic nature of its repeat domain suggests that it may be located on the surface of E. chaffeensis (45). The GE 100-kDa protein is also hydrophilic in the repeat region (data not shown) and may also be surface associated. Although both the 100- and 130-

It is unclear whether the three GE proteins described in this report may be useful as potential diagnostic reagents or vac-


discussion

Ehrlichiae which have a tropism for peripheral blood gran-

The GE isolate we describe here is indistinguishable from the purified plasmid DNA of each clone. Figure 8 shows the results obtained for primers of S2 (lanes 2 to 4), S7 (lanes 5 to 7), and S22 (lanes 8 to 10). All three clones were specific to GE and were not present in HL60 DNA (lanes 2, 5, and 8). In each case, the size of the PCR product determined with genomic DNA as the template was the same as that generated by the purified plasmid DNA.

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creted into the culture medium or their expression is tied more closely to natural infection, they are clearly important immunologically. All three react with convalescent-phase sera from dogs, sheep (data not shown), and humans diagnosed with HGE. The E. chaffeensis 120-kDa protein, the counterpart of the 100-kDa GE protein, has shown sensitivity and specificity as a diagnostic reagent with a panel of E. chaffeensis patient sera (44). However, Western blots with convalescent-phase sera from HGE patients do not show a consistent response to the GE 100-kDa protein. It is not known whether there is any immunologic cross-reactivity between the GE 100-kDa protein and the E. chaffeensis 120-kDa protein or whether the two could be used to distinguish monocytic from granulocytic ehrlichiosis. Some human antisera have been reported to react with both E. chaffeensis and HGE antigens, but it is not known whether this represents cross-reactive epitopes or separate exposures to the two ehrlichial agents (30, 32).

In summary, the expression library approach for the isolation and characterization of GE proteins has resulted in the discovery of three previously unidentified genes which express high-molecular-weight, immunoreactive proteins. Further study of these proteins will aid in the classification of the members of the GE genogroup as corresponding genes are sequenced and compared and will be important in elucidating the role of these proteins in immunity and pathogenesis.

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