Chlamydia pneumoniae Expresses Genes Required for DNA Replication but Not Cytokinesis during Persistent Infection of HEp-2 Cells

Gerald I. Byrne,1* Scot P. Ouellette,1 Zhao Wang,2 J. P. Rao,2 Lin Lu,2 Wandy L. Beatty,3 and Alan P. Hudson2

Department of Medical Microbiology and Immunology, University of Wisconsin School of Medicine, Madison, Wisconsin 53706; Department of Immunology and Microbiology, Wayne State University School of Medicine, Detroit, Michigan 48201; and Department of Molecular Microbiology, Washington University School of Medicine, St. Louis, Missouri 63110

Received 4 April 2001/Returned for modification 30 May 2001/Accepted 19 June 2001

Chlamydia pneumoniae causes community-acquired pneumonia and is associated with several chronic diseases, including asthma and atherosclerosis. The intracellular growth rate of C. pneumoniae slows dramatically during chronic infection, and such persistence leads to attenuated production of new elementary bodies, appearance of morphologically aberrant reticulate bodies, and altered expression of several chlamydial genes. We used an in vitro system to further characterize persistent C. pneumoniae infection, employing both ultrastructural and transcriptional activity measurements. HEp-2 cells were infected with C. pneumoniae (TW-183) at a multiplicity of infection of 3:1, and at 2 h postinfection gamma interferon (IFN-γ) was added to the medium at 0.15 or 0.50 ng/ml. Treated and untreated cultures were harvested at several times postinfection. RNA was isolated and reverse transcribed, and reverse transcription (RT)-PCR analyses targeting primary transcripts from chlamydial rRNA operons as well as dnaA, polA, mutS, minD, ftsK, and ftsW mRNA were done. Some cultures were fixed and stained for electron microscopic analysis, and a real-time PCR assay was used to assess relative chlamydial chromosome accumulation under each culture condition. The latter assays showed that bacterial chromosome copies accumulated severalfold during IFN-γ treatment of infected HEp-2 cells, although less accumulation was observed in cells treated with the higher dose. Electron microscopy demonstrated that high-dose IFN-γ treatment elicited aberrant forms of the bacterium. RT-PCR showed that chlamydial primary rRNA transcripts were present in all IFN-γ-treated and untreated cell cultures, indicating bacterial metabolic activity. Transcripts from dnaA, polA, mutS, and minD, all of which encode products for bacterial chromosome replication and partition, were expressed in IFN-γ-treated and untreated cells. In contrast, ftsK and ftsW, encoding products for bacterial cell division, were expressed in untreated cells, but expression was attenuated in cells treated with low-dose IFN-γ and absent in cells given the high dose of cytokine. Thus, the development of persistence included production of transcripts for DNA replication-related, but not cell division-related, genes. These results provide new insight regarding molecular activities that accompany persistence of C. pneumoniae, as well as suggesting requirements for reactivation from persistent to productive growth.

Chlamydia pneumoniae and Chlamydia trachomatis are intracellular bacterial pathogens that function as etiologic agents of important human diseases. The former, for example, causes community-acquired pneumonia and has recently been associated with various chronic diseases such as asthma and atherosclerosis (9, 16). C. trachomatis remains a significant cause of infectious, preventable blindness (trachoma) in the developing world (26) and is a sexually transmitted pathogen known to cause fertility deficits in women (26), as well as chronic arthritis in both sexes (for a review, see reference 14). Infections with each organism show high rates of recurrence (3, 8), but currently available information usually does not allow unequivocal differentiation between recurrences due primarily to reinfec-
been characterized (1). Interestingly, in one such model system ultrastructural analysis suggested that organisms of this species made persistent in culture by treatment with gamma interferon (IFN-\(\gamma\)) continue to replicate and partition their genomes, even in the absence of subsequent cell division (2). This provides at least a partial explanation for the lack of production of new infectious EB by persistent \(C.\) \textit{trachomatis}; however, no congruent information is available as yet for persistent \(C.\) \textit{pneumoniae}. The present study was undertaken to investigate similar aspects of the molecular genetic behavior of persistent \(C.\) \textit{pneumoniae}, using a well-described cell culture model system.

Data from the chlamydial genome sequence have identified orthologs for \textit{dnaA}, \textit{polA}, and \textit{mutS}, each of which encodes a gene product required for DNA replication and repair (4) in \(C.\) \textit{pneumoniae}; a \textit{minD} ortholog has also been identified, the product of which is required for chromosome segregation (24). Chlamydiae apparently do not have an \textit{fsZ} gene (24) but do possess orthologs for \textit{fsK} and \textit{fsW}, each of which encodes a product required for cell division (5). In the work described here, we provide information regarding differential expression of these \(C.\) \textit{pneumoniae} DNA replication- and cytokinesis-related genes under conditions of both persistent and productive growth. Our results indicate that mRNA from \(C.\) \textit{pneumoniae} genes encoding products required for chromosome replication, repair, and segregation are synthesized regardless of whether the organism is in a persistent or a productive growth state. Organisms undergoing productive infection express cytokinesis-related transcripts as expected, whereas persistent organisms do not. These observations provide new information concerning the basic biology of chlamydial persistence and may be useful in the design of improved treatment regimens for chronic chlamydial infections.

**MATERIALS AND METHODS**

**Cell culture and growth of \(C.\) \textit{pneumoniae}**. The human bronchial epithelial cell line HEP-2 was used for growth and propagation of \(C.\) \textit{pneumoniae}. HEP-2 cells were grown in Iscove’s modified Dulbecco’s medium (BioWhittaker, Walkersville, Md.) supplemented with 10% heat-inactivated fetal bovine serum (HyClone, Logan, Utah), 50 \(\mu\)g of vancomycin (Sigma, St. Louis, Mo.) per ml, and 10 \(\mu\)g of gentamicin (Life Technologies, Gaithersburg, Md.) per ml. Cells were routinely maintained at 35°C in 7% CO\(_2\) in a water-jacketed CO\(_2\) incubator and routinely maintained at 35°C in 7% CO\(_2\) in a water-jacketed CO\(_2\) incubator and were passaged two or three times per week in 162-cm\(^2\) cell culture flasks (Corning Costar, Cambridge, Mass.). \(C.\) \textit{pneumoniae} TW-183 was purchased from the American Type Culture Collection (Rockville, Md.) and propagated for 3 days in HEP-2 cells treated with 2 \(\mu\)g of cycloheximide (Sigma) per ml. Chlamydiae were collected from infected-cell sonicates by differential centrifugation, partially purified by centrifugation through a 30% Renografin (Braico Diagnostics, Princeton, N.J.) cushion, and resuspended in sucrose-phosphate buffer (SPB; 0.22 M sucrose, 0.2 M Na\(_2\)HPO\(_4\), 0.2 M NaH\(_2\)PO\(_4\), 5 mM glutamic acid [pH 7.4]), as described previously (15). Stock titers were determined, and stocks were stored at 10°C and resuspended in SPB. Chlamydial titers from each sample were determined as described for infectivity (15).

**Infectivity assays**. Monolayers of HEP-2 cells were infected with \(C.\) \textit{pneumoniae} at a multiplicity of infection of 1 and subsequently treated with IFN-\(\gamma\) at various doses (2 \(\mu\)g postinfection or left untreated. We adapted a highly quantitative real-time PCR assay system described and tested by others (21). Briefly, the chlamydia-directed primers target the two copies of the 16S RNA gene on the bacterial chromosome, while assay input is normalized simultaneously to host 18S rRNA gene sequences. The \(C.\) \textit{pneumoniae}-directed primers for the assay were designed using software supplied for this purpose by PE Biosystems (Foster City, Calif.); these primers were 5'-GTTGTTATTTAGTGCCGGA-3' and 5'-CCACCACTAA-GCTGATA-3'. Extensive control studies confirmed that only a single amplification product is generated by this primer system and that the product is the appropriate segment of the \(C.\) \textit{pneumoniae} 16S rRNA coding sequence. The human 18S-directed primer system used for normalization was purchased from PE Biosystems and was designed for this use. All assays were done several times, each in triplicate, using a PE Biosystems model 7700 sequence detector with the SYBR Green I real-time PCR assay as calculated using sequence detection software, version 1.7, from PE Biosystems.

**RT-PCR analyses**. RT was done as previously described (6) with 5 \(\mu\)g of total RNA from each preparation, random hexamers as primers, and murine leukemia virus reverse transcriptase (Life Technologies). cDNA from each reaction was treated with RNase A, RNase T1, and RNase H, extracted several times with phenol-chloroform (24:1), and then collected as an ethanol precipitate (6). The genes targeted in RT-PCR are given in Table 1, along with the primer sequences. Primer sequences were designed using the GeneRunner program (Hastings Software, Hastings, N.Y.) based on published sequence information (http://stdgen.lanl.gov). Each primer set was tested to confirm that chlamydial but not host cell sequences were amplified; control experiments established that all assay systems had approximately equivalent sensitivity and were able to identify transcripts from the \(C.\) \textit{pneumoniae}. Amplification conditions for the first round of nested reactions were 4 min at 95°C, 35 cycles of 40 s at 95°C, 40 s at the annealing temperature, and 40 s at 72°C, and 10 min at 72°C. Annealing temperatures varied somewhat among the several primer sets. The second nested amplification round was done using similar cycling parameters and 10% of the first-round reaction mixture. The positive control for chlamydial transcriptional activity was demonstration of primary transcripts from the bacterial rRNA operons, as previously described (7). Amplifications were done using AmpliTaq DNA polymerase (Perkin Elmer) in a thermocycler (model PTC-100; MJ Research, Watertown, Mass.). Products were visualized by ethidium bromide staining of standard agarose electro-
TABLE 1. Primers used for RT-PCR assays targeting C. pneumoniae mRNA

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>adt1</td>
<td>5'-GCGGATACATACATCAGGCTAAAGAAG-3' (outer) 5'-GCTGATCGTGACATGACAAATGAAGG-3' (inner)</td>
</tr>
<tr>
<td>dnaA</td>
<td>5'-GACCTTCTGGGATCCCATGTCATTGGTG-3' (outer) 5'-GAGGATCGTCGAATGATGATGATGATGATGATG-3' (inner)</td>
</tr>
<tr>
<td>polA</td>
<td>5'-GCTGGACCGACGAAATAGGGGGGAGG-3' (outer) 5'-GAGGATCGTCGAATGATGATGATGATGATGATG-3' (inner)</td>
</tr>
<tr>
<td>mutS</td>
<td>5'-CAGTGGTGCATGGGGATGGGATTGGG-3' (outer) 5'-GAGGATCGTCGAATGATGATGATGATGATGATG-3' (inner)</td>
</tr>
<tr>
<td>ftsK</td>
<td>5'-GCTGACGTTGAACTGTTGAACTGTT-3' (outer) 5'-GAGGATCGTCGAATGATGATGATGATGATGATG-3' (inner)</td>
</tr>
<tr>
<td>ftsW</td>
<td>5'-GAGCATTGGAACGCTGTTGAACTGTT-3' (outer) 5'-GAGGATCGTCGAATGATGATGATGATGATGATG-3' (inner)</td>
</tr>
</tbody>
</table>

RESULTS

Accumulation of chlamydial DNA and expression of DNA replication- and cytokinesis-related genes during infection of HEp-2 cells. C. pneumoniae completes the standard developmental cycle within HEp-2 cells. However, growth of the organism in infected cell cultures is usually done in the presence of cycloheximide to increase the yield of infectious EB. To provide controls for the relative amount of chlamydial DNA produced under various treatment conditions, including treatment with rIFN-γ (see below), a highly quantitative real-time PCR assay targeting the C. pneumoniae 16S rRNA genes was used. The data shown in Fig. 1 indicate that when chlamydial DNA levels were compared as a function of incubation time in untreated and cycloheximide-treated HEp-2 cells, a time-dependent increase occurred, and as expected, greater amounts of chlamydial chromosome accumulated in cycloheximide-treated than untreated host cells. Part of this observed increase may have been due to less host 18S rRNA in cycloheximide-treated samples, but chlamydial growth does occur to a greater extent in host cells treated with cycloheximide. We found that at 96 h postinfection, the final time point examined in these experiments, the cycloheximide-treated cultures had accumulated in excess of 3.5-fold more chlamydial DNA than the untreated culture. In the treated cultures, accumulation of bacterial chromosome over 96 h was at the level of about 270-fold, indicating seven to eight cell divisions undertaken by each RB within the inclusions. Several repeats of these determinations yielded virtually identical results.

As a control for transcript-related experiments given below, the time course of expression was determined for a panel of C. pneumoniae genes whose products are involved in replication and partition of the bacterial chromosome, as well as the cell division process, in untreated and cycloheximide-treated HEp-2 cells. Figure 2 shows representative results from these nonquantitative RT-PCR studies. In both treated and untreated infected HEp-2 cells, primary transcripts from the RNA operons were apparent at 12 h postinfection, as was mRNA from adt1, which specifies an ATP-ADP exchange protein of the organism. Similarly, dnaA, polA, and ftsK all were expressed by 12 h after initiation of infection, the earliest time assayed in both treated and untreated cells; expression of each of these genes continued through 96 h postinfection. Similar results were obtained for mutS, minD, and ftsW (data not shown).

Effects of IFN-γ on C. pneumoniae development and ultrastructure in IFN-γ-treated HEp-2 cells. Treatment of HEp-2 cells with rIFN-γ 2 h after infection with C. pneumoniae caused a dose-dependent inhibition of infectivity recovery (Fig. 3). Transmission EM of infected HEp-2 cells demonstrated classic EB and RB developmental forms 48 h postinfection (data not shown). When cells were treated with subinhibitory doses of rIFN-γ, normal-appearing inclusions also were observed after 48 h of growth (Fig. 4A). In contrast, infected HEp-2 cells
treated with 0.5 ng of rIFN-γ per ml for 48 h demonstrated essentially universal development of enlarged persistent forms, as judged from EM (Fig. 4B). This form of the organism is indicative of noninfectious, poorly dividing, greatly enlarged, and aberrantly shaped RB and is known to be a result of IFN-γ-mediated induction of the tryptophan-decyclizing enzyme indoleamine-2,3-dioxygenase, which limits availability of the required amino acid tryptophan (1). The presence of these aberrant forms was used as a basis for comparison of selected transcriptional activities with organisms progressing through a productive infection in HEp-2 cells.

Viability of *C. pneumoniae* in untreated and rIFN-γ-treated HEp-2 cell cultures. To determine if the morphologically aberrant chlamydiae observed in EM studies exhibit metabolic activity, infected HEp-2 cells treated for 48 h with 0.15 or 0.5 ng of rIFN-γ per ml were assessed for the presence of primary transcripts from the chlamydial rRNA operons. These transcripts were detectable by RT-PCR by 12 h postinfection in chlamydia-infected cells not treated with rIFN-γ, and their expression continued through 96 h (Fig. 2); similarly, the chlamydial *adt1* gene was expressed from 12 to 96 h postinfection in untreated cells. Transcripts from these genes were not detectable in chlamydial EB but were demonstrable in infected HEp-2 cells treated with either high- or low-dose rIFN-γ (Fig. 5). These data support the contention that while *C. pneumoniae* organisms grown in the presence of rIFN-γ assume an aberrant morphologic form, they do remain viable, as judged by the presence of RNA species characteristic of metabolically active bacteria.

Chromosome replication in rIFN-γ-treated *C. pneumoniae*. The control experiments described above indicated that replication of the chlamydial chromosome was under way by 24 h postinfection in both untreated and cycloheximide-treated infected HEp-2 cells (Fig. 1). When the relative levels of C.
pneumoniae DNA were compared in untreated cells and those treated for 48 h with 0.15 or 0.5 ng of rIFN-γ per ml, evidence for bacterial chromosome replication in the IFN-γ-treated samples was observed (Fig. 6). Clearly, less bacterial DNA was produced in the cultures treated with 0.5 ng of IFN-γ per ml than in untreated cultures, but these data confirm the contention that persistent C. pneumoniae are metabolically active and that aberrant chlamydial morphology and growth did not cause complete inhibition of genome replication.

Expression of chlamydial DNA replication- and cytokinesis-related genes in untreated and rIFN-γ-treated infected HEp-2 cells. Control studies (Fig. 2) clearly indicated that the chlamydial dnaA, polA, ftsK, and ftsW genes are expressed as early as 12 h postinfection in HEp-2 cells, regardless of whether the cells are treated with cycloheximide. To determine whether

FIG. 4. Ultrastructural analysis by transmission EM of HEp-2 cells persistently infected for 48 h with C. pneumoniae. (A) Treatment with low levels of IFN-γ (0.15 ng/ml) resulted in the development of typical inclusions containing EB and RB. Normal RB are indicated by arrows. (B) Treatment with 0.50 ng of IFN-γ per ml to induce persistence engenders development of grossly enlarged RB (arrowheads). N, nucleus. Bar = 1 μm.

FIG. 5. Representative RT-PCR analyses targeting primary transcripts from the C. pneumoniae rRNA operons (A) and adt1 (B) in untreated infected HEp-2 cells and infected HEp-2 cells treated with low or high doses of rIFN-γ. Cells were infected with chlamydiae and harvested at 48 h posttreatment, and RNA was prepared from each harvested culture (see Material and Methods). RT-PCR analyses were performed using primers given in Table 1. Lanes: C+, positive PCR control for each primer set, using C. pneumoniae DNA as the amplification template. C−, negative RT-PCR control using cDNA from uninfected HEp-2 cells as the amplification template; RT−, negative RT-PCR control using cDNA from untreated infected HEp-2 cells as the amplification template; RT−, negative RT-PCR control showing the results of PCR with each primer set in the absence of RT of RNA preparations used; Act, amplification product from an RT-PCR assay targeting host β-actin mRNA; EB, amplification product from pure EB RNA. Sizes of the amplification products are given in the legend to Fig. 1.

FIG. 6. Results from quantitative real-time PCR assays to determine the relative level of accumulation of C. pneumoniae chromosomal DNA during infection of untreated infected HEp-2 cells and infected HEp-2 cells treated with either 0.15 or 0.50 ng of rIFN-γ per ml. Cells were infected with C. pneumoniae TW-183 as described in Materials and Methods. Both treated and untreated cultures were grown without cycloheximide and were harvested at 48 h postinfection or posttreatment. DNA was prepared, and a real-time PCR assay system was used to determine the relative level of Chlamydia DNA from each preparation. Input into each assay was normalized to the host 18S rRNA genes, as described in Materials and Methods. Data are triplicate mean values relative to the mean value obtained for untreated cells at 12 h postinfection (not shown). Standard errors are indicated.
and partition-related genes were expressed in rIFN-γ-treated cells, indicating that those genes are expressed during IFN-γ-induced persistent infection. In contrast, genes required for bacterial cell division either are not expressed or are expressed only at an extremely low level during persistence, thus essentially eliminating cytokinesis during this growth state. The functional result of expression of DNA replication and segregation genes by the bacterium should be accumulation of chromosomal DNA; the results presented here confirm that C. pneumoniae DNA does indeed accumulate during cytokine-induced persistence, although the level of accumulation is low.

We did not attempt to quantitate relative transcript levels for the C. pneumoniae genes assayed in either untreated or IFN-γ-treated infected HEp-2 cells. Nonetheless, it is unlikely that the ftsK and ftsW mRNAs were missed in the analysis of the high-dose-cytokine-treated samples. Control studies not shown here indicated that the relative sensitivities of the RT-PCR assay systems employed are approximately equivalent, and each of the assays routinely identifies the targeted transcripts from 10 to 30 bacterial cells. In the untreated infected cells, amplified products from the dnaA, mutS, minD, polA, ftsK, and ftsW cDNA were demonstrable even after the first round of the nested PCR. In the low-dose-rIFN-γ-treated cells, products from the replication-related genes were easily identifiable after the two amplification rounds, as they were in cDNA from the high-dose-cytokine-treated cells. No products for the two cytokinesis-related genes could be identified even after the second, nested amplification in cells treated with 0.50 ng of cytokine per ml, although the nonnested assays for primary transcripts from the rRNA operons gave clear products from the same RNA-cDNA preparations. The important point is that we could identify no products derived from either of the cytokinesis-related mRNAs in the high-dose-rIFN-γ-treated infected HEp-2 cells, indicating that expression of the two cell division-related genes is severely attenuated, even if it is not completely abolished, during cytokine-induced persistent C. pneumoniae infection.

This study provides important new information regarding gene regulation patterns intrinsic to intracellular chlamydial growth, during both active and persistent infection. Clearly, differential gene expression does occur in these two states, almost certainly as a result of modulation of the host cell environment. One mechanism by which chlamydiae might sense host environmental changes is through a two-component signaling system, as has been described for a variety of other bacterial pathogens (13). For example, a histidine kinase (designated Cpn 0584; http://www.stdgen.lanl.gov) has been identified in the C. pneumoniae genome, and this enzyme shows possible two-component regulatory activity. Importantly, a recent study has shown that C. trachomatis exhibits differential, developmentally regulated expression of its three sigma fac-

FIG. 7. Representative RT-PCR analyses targeting transcripts from C. pneumoniae DNA replication- and cell division-related genes in untreated infected HEp-2 cells and infected HEp-2 cells treated with low or high doses of rIFN-γ. Cells were infected with chlamydiae and harvested at 48 h posttreatment, and RNA was prepared from each harvested culture (see Material and Methods). RT-PCR analyses were performed using primers given in Table 1. (A) primary rRNA transcripts; (B) dnaA; (C) polA; (D) ftsK. Lanes: C+, positive PCR control for each primer set, using C. pneumoniae DNA as the amplification template; C−, negative RT-PCR control, using cDNA from uninfected HEp-2 cells as the amplification template; RT−, negative control showing the results of PCR with each primer set in the absence of RT of RNA preparations used; ACT, amplification product from an RT-PCR assay targeting host β-actin mRNA. Input was normalized to the host β-actin mRNA. Sizes of the amplification products are given in the legend to Fig. 1.

these chlamydial DNA replication- and cytokinesis-related genes are expressed during persistent growth induced by cytokine treatment, we assessed the presence of mRNA encoding them in RNA prepared from infected HEp-2 cells treated for 48 h with 0.15 or 0.50 ng of rIFN-γ per ml. As indicated by representative assays (Fig. 7), chlamydial DNA replication-and partition-related genes were expressed in rIFN-γ-treated infected HEp-2 cultures regardless of the cytokine concentration in the growth medium. However, amplification products of the chlamydial ftsK and ftsW genes, whose products are required for cell division, were attenuated in cultures treated with 0.15 ng of rIFN-γ per ml, and they were not found in RNA or cDNA preparations from cells grown with the higher cytokine concentration.

DISCUSSION

Active ocular or urogenital infection with C. trachomatis represents a clinically significant event. However, it is now clear that long-term, persistent infection with this organism often engenders severe and difficult-to-treat sequelae. Similarly, the pneumonia resulting from respiratory infection by C. pneumoniae can be significant, but increasing evidence sup-

ports both the existence and the clinical importance of chronic infection with this organism. Evidence also argues that this latter state may be of the most import in terms of overall human health. A good deal of information is currently avail-

able concerning biochemical and molecular genetic character-

istics of persistent C. trachomatis. For example, morphologic

and microbiologic studies indicate that during persistence, cells of this organism are characterized by a reduced capacity for the enlarged and aberrantly shaped RB to undergo cell division but that these abnormal intracellular organisms continue to replicate their genomes (2). Failure to undergo cell division has been noted for IFN-γ-mediated induction of persistence in C. pneumoniae (22), and data in the present report provide an explanation for that lack of cytokinesis. Results of RT-PCR analyses, targeting C. pneumoniae genes whose products are required for chromosomal DNA replication and partition, indicate that those genes are expressed during IFN-γ-induced persistent infection. In contrast, genes required for bacterial cell division either are not expressed or are expressed only at an extremely low level during persistence, thus essentially elimin-

ating cytokinesis during this growth state. The functional result of expression of DNA replication and segregation genes by the bacterium should be accumulation of chromosomal DNA; the results presented here confirm that C. pneumoniae DNA does indeed accumulate during cytokine-induced persistence, although the level of accumulation is low.

We did not attempt to quantitate relative transcript levels for the C. pneumoniae genes assayed in either untreated or IFN-γ-treated infected HEp-2 cells. Nonetheless, it is unlikely that the ftsK and ftsW mRNAs were missed in the analysis of the high-dose-cytokine-treated samples. Control studies not shown here indicated that the relative sensitivities of the RT-PCR assay systems employed are approximately equivalent, and each of the assays routinely identifies the targeted transcripts from 10 to 30 bacterial cells. In the untreated infected cells, amplified products from the dnaA, mutS, minD, polA, ftsK, and ftsW cDNA were demonstrable even after the first round of the nested PCR. In the low-dose-rIFN-γ-treated cells, products from the replication-related genes were easily identifiable after the two amplification rounds, as they were in cDNA from the high-dose-cytokine-treated cells. No products for the two cytokinesis-related genes could be identified even after the second, nested amplification in cells treated with 0.50 ng of cytokine per ml, although the nonnested assays for primary transcripts from the rRNA operons gave clear products from the same RNA-cDNA preparations. The important point is that we could identify no products derived from either of the cytokinesis-related mRNAs in the high-dose-rIFN-γ-treated infected HEp-2 cells, indicating that expression of the two cell division-related genes is severely attenuated, even if it is not completely abolished, during cytokine-induced persistent C. pneumoniae infection.

This study provides important new information regarding gene regulation patterns intrinsic to intracellular chlamydial growth, during both active and persistent infection. Clearly, differential gene expression does occur in these two states, almost certainly as a result of modulation of the host cell environment. One mechanism by which chlamydiae might sense host environmental changes is through a two-component signaling system, as has been described for a variety of other bacterial pathogens (13). For example, a histidine kinase (designated Cpn 0584; http://www.stdgen.lanl.gov) has been identified in the C. pneumoniae genome, and this enzyme shows possible two-component regulatory activity. Importantly, a recent study has shown that C. trachomatis exhibits differential, developmentally regulated expression of its three sigma fac-
tors, and it has been postulated that this pattern of expression reflects a more global pattern of chlamydial gene expression (25). Congruent information is not yet available for transcription of C. pneumoniae sigma factor genes, but experiments are now under way to meet this need. The present study necessarily required selecting genes to study their transcriptional activity under various growth conditions. More complete analysis of C. pneumoniae and host cell transcription patterns will require microarray analyses. These studies are also in progress.

ACKNOWLEDGMENTS

This work was supported by NIH grants AR-42541 and AI-44055 (A.P.H.) and AI 19782 and AI 42790 (G.I.B.). G.I.B. and A.P.H. contributed equally to this work.

REFERENCES