Identification of an *Entamoeba histolytica* Serine-, Threonine-, and Isoleucine-Rich Protein with Roles in Adhesion and Cytotoxicity

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*Entamoeba histolytica* is a protozoan parasite that causes significant morbidity and mortality in the developing world (48). The life cycle is composed of two stages: the transmissible cyst that is released in the feces of infected humans, and the motile trophozoite, which resides in the large bowel. The infection begins when cysts are ingested, usually through contaminated food or water. Excystation occurs in the small bowel, and the motile trophozoites eventually colonize the large bowel.

The majority of individuals with *E. histolytica* infection remain asymptomatic; however, in some cases the parasites penetrate the colon wall causing dysentery and colitis and occasionally disseminate hematogenously to cause liver abscesses. The potential reasons for why so many individuals remain asymptomatic while others develop serious disease are not well understood. We have previously used expression profiling to identify *Entamoeba* genes whose expressions were strictly associated with virulent strains (R. C. MacFarlane and U. Singh, Infect. Immun. 74:340–351, 2006). One gene, which we have named EhSTIRP (*Entamoeba histolytica* serine-, threonine-, and isoleucine-rich protein), was exclusively expressed in virulent but not in nonvirulent *Entamoeba* strains. EhSTIRP is predicted to be a transmembrane protein and is encoded by a multigene family. In order to characterize its function in amebic biology, we used a double-stranded RNA-based approach and were able to selectively down-regulate expression of this gene family. Upon EhSTIRP down-regulation, we were able to ascribe cytotoxic and adhesive properties to the protein family using lactate dehydrogenase release and Chinese hamster ovary cell adhesion assays. EhSTIRP thus likely represents a novel determinant of virulence in *Entamoeba histolytica*. This work validates the fact that genes expressed exclusively in virulent strains may represent virulence determinants and highlights the need for further functional analyses of other genes with similar expression profiles.

*Entamoeba histolytica* is a leading cause of parasitic death globally. However, the molecular framework regulating pathogenesis is poorly understood. We have previously used expression profiling to identify *Entamoeba* genes whose expressions were strictly associated with virulent strains (R. C. MacFarlane and U. Singh, Infect. Immun. 74:340–351, 2006). One gene, which we have named EhSTIRP (*Entamoeba histolytica* serine-, threonine-, and isoleucine-rich protein), was exclusively expressed in virulent but not in nonvirulent *Entamoeba* strains. EhSTIRP is predicted to be a transmembrane protein and is encoded by a multigene family. In order to characterize its function in amebic biology, we used a double-stranded RNA-based approach and were able to selectively down-regulate expression of this gene family. Upon EhSTIRP down-regulation, we were able to ascribe cytotoxic and adhesive properties to the protein family using lactate dehydrogenase release and Chinese hamster ovary cell adhesion assays. EhSTIRP thus likely represents a novel determinant of virulence in *Entamoeba histolytica*. This work validates the fact that genes expressed exclusively in virulent strains may represent virulence determinants and highlights the need for further functional analyses of other genes with similar expression profiles.
lytica and E. dispar compared to that in E. histolytica HM-1:IMSS (33). Davis et al. used a microarray containing 70-mer oligonucleotides and reported 152 genes with significantly higher expression in E. histolytica HM-1:IMSS than in E. histolytica Rahman (18). Using an E. histolytica microarray based on the Affymetrix platform, Ehrenkauffer et al. identified 141 genes with significantly higher expression in E. histolytica HM-1:IMSS than in E. histolytica Rahman (24). Of these 141 genes, approximately 52 were not expressed in E. histolytica Rahman whereas the remainder were expressed but at lower levels than in E. histolytica HM-1:IMSS. The expression profile of E. dispar cannot be reliably determined using the oligonucleotide microarrays.

Some genes involved in amebic pathogenesis are expressed at lower levels in E. histolytica Rahman, including cysteine proteases (EhCP6, EhCP4, EhCP7), gene products important in bacterial killing (AIG1, lysozyme) and stress response (70-kDa heat shock protein, peroxiredoxin) (18, 19), and the light subunit of the Gal/GalNAc lectin (3). Additionally, the proteophosphoglycans coating the surface of E. histolytica Rahman have truncated glucan side chains compared to those of the virulent E. histolytica HM-1:IMSS (35). The authors hypothesized that the glycans may protect parasite adhesion molecules from proteolysis or that the proteophosphoglycans may regulate the ability of parasite surface molecules to interact with host cell receptors. Two genes with lower expression in E. histolytica Rahman have been genetically linked to virulence: overexpression of the peroxiredoxin gene in E. histolytica Rahman imparted increased resistance to oxidative stress (19), and antisense inhibition of the light lectin subunit in E. histolytica HM-1:IMSS resulted in decreased cytotoxicity and fewer amebic liver abscesses in hamsters (3).

We previously identified a protein family rich in serine, threonine, and isoleucine residues which we have named EhSTIRP (Entamoeba histolytica serine-, threonine-, and isoleucine-rich protein) (33). We originally identified EhSTIRP as a putative virulence determinant based on its expression profile. EhSTIRP is highly expressed in two virulent strains (E. histolytica HM-1:IMSS and 200:NH) as well as in parasites causing disease in a mouse model of colitis (24, 27). Conversely, EhSTIRP was not expressed in the nonviralent E. histolytica Rahman or E. dispar (33). Additionally, EhSTIRP had low expression in the cyst stage of the parasite, which is noninvasive (24).

In order to determine if EhSTIRP is involved in parasite virulence, we down-regulated EhSTIRP expression in the virulent E. histolytica HM-1:IMSS via episcopal production of double-stranded RNA (dsRNA). Upon EhSTIRP down-regulation, parasites exhibited decreased levels of cytotoxicity and reduced ability to adhere Chinese hamster ovary (CHO) cells, implicating a role in amebic pathogenesis. This work validates our hypothesis that genes expressed exclusively in virulent strains may represent novel virulence pathways and increases the repertoire of genes that have been genetically linked to amebic virulence.

MATERIALS AND METHODS

Parasite strains and culture. E. histolytica strains HM-1:IMSS and Rahman were obtained from the American Type Culture Collection (ATCC) (http://www.atcc.org/). The parasites were grown under axenic conditions in Trypticase-yeast extract-iron-serum medium (TY1S-S3) with 15% adult bovine serum (Sigma), supplemented with penicillin (100 U/ml), streptomycin (100 µg/ml) (Gibco BRL), and 1× Diamond’s vitamins (Biosource International) in 15 ml glass culture tubes at 37°C as previously described (21). Parastate genotypes were tested using restriction fragment length polymorphism analysis of the EbsREHP gene using previously described methods (6, 15). The SREHP PCR products were digested with Aul, and results for all PCR products were compared to banding patterns from previously published studies (15). Growth rates were determined for parasites transfected with the control or dsRNA construct. Parasites (50,000) were inoculated into glass tubes at time point zero and grown under standard culture conditions, and cell counts were determined using a hemacytometer at 12- to 18-h intervals.

Sequence analysis. Genome sequences were downloaded from The Institute for Genomic Research (TIGR) (http://www.tigr.org/tdb/e2k1/eula1/) and the Sanger Institute Entamoeba histolytica GeneDB (http://www.genedb.org/GENDB/chistolytica/). We performed an identity alignment of nucleotide and amino acid sequences for all gene family members. Signal peptides were predicted using the SignalP algorithm (http://www.cbs.dtu.dk/services/SignalP/); transmembrane sites were predicted using the TIGR website.

Constructs for expressing EhSTIRP dsRNA. To down-regulate EhSTIRP, we utilized a methodology previously described in reference 30. Briefly, a 395-bp fragment from the 3′ end of the EhSTIRP1 gene (99% nucleotide similarity to EhSTIRP2 and EhSTIRP3) was amplified by PCR and subcloned into the TOPO TA 2.1 vector (Invitrogen). The gene fragment was subcloned in opposing orientations separated by a 500-bp nonspecific stuffer region in pBluescript SK+ and then subcloned into the E. histolytica expression construct pJST4 at the KpnI and BamHI sites (26). This construct would be expected to generate dsRNA specific to the EhSTIRP gene family. A control construct was created using a similar methodology, except that 395 bp from the 3′ end of the gene and 350 bp from the 5′ end of EhSTIRP2 were cloned close to each other in the 500-bp stuffer region. This construct would not be expected to generate dsRNA.

Transfection of E. histolytica. Entamoeba histolytica HM-1:IMSS trophozoites (2 × 107 to 4 × 108) were transformed using 6 to 20 µg of plasmid DNA and 15 to 30 µl of Superfect (Qiagen) in each 35-mm petri dishes or 1.5-mL tubes according to the manufacturer’s instructions and as published previously (40). Stable transfectants were initially selected with G418 at 3 to 6 µg/ml, and cultures were maintained at 48 µg/ml G418.

Northern blot analysis. Northern blots were performed as previously described (32). Briefly, total parasite RNA was isolated using TRIZOL reagent (Invitrogen) according to the manufacturer’s instructions. RNA (10 to 20 µg) was separated by denaturing 12% agarose gel electrophoresis, transferred to Hybond-N membrane filters (Amersham-Pharmacia), and hybridized with radioactive probes using the ExpressHyb (Chontech) hybridization buffer. PCR product probes were sequenced and primed with random primers (5′-3′). Oligonucleotide probes were labeled with [32P]dATP and T4 polynucleotide kinase (New England Biolabs). Blots were exposed to film or a phosphorimaging screen, scanned, and prepared for publication using Adobe Photoshop version 7; Adobe, San Jose, CA). Blots were stripped using 0.5% sodium dodecyl sulfate and reused for subsequent hybridizations per the manufacturer’s suggestions. For loading controls, a probe for Ehelnin (locus m100031) was PCR amplified from E. histolytica HM-1:IMSS genomic DNA and labeled as described previously; or RNA bands were stained with methylene blue, denatured with deHO, and scanned.

The following primers were used in this study: Ehelnin forward, 5′-GCTGTGATGGG TCCAAAGGAG-3′; Ehelnin reverse, 5′-TTTCTGTGGAACACTGACTGTCC-3′; EhSTIRP1 5′ forward, 5′-TCAACTCTTCACATGTTTCCTCA-3′; EhSTIRP1 reverse, 5′-ATTACATTCAACAGAAGCATTG-3′; EhSTIRP2 5′ forward, 5′-GCTATTTGTTGAAATGAGG-3′; EhSTIRP2 reverse, 5′-GAAACTTGTTGG CCATTCCTT-3′; EhSTIRP3 5′ forward, 5′-GCCTGATGACGCTGTTACCT-3′; EhSTIRP3 reverse, 5′-GAACTTGGTTTG CACCTTCCTT-3′. The constructs for expressing EhSTIRP dsRNA were generated using a T7 RNA polymerase promoter construct in pBluescript STIRP2 specific, 5′-TTAATTCATTTCCATCTTCTGTTTGACC-3′; EhSTIRP3 specific, 5′-TTAATTCATTTCCATCTTCTGTTTGACC-3′; 5′-RACE outer primer, 5′-GCTGTGATGGG TCCAAAGGAG-3′; 5′-RACE inner primer, 5′-GCCTGATGACGCTGTTACCT-3′; EhSTIRP1 5′ reverse outer, 5′-TGGTATGCTTGGTACCTGTGAG-3′; EhSTIRP1 5′ reverse inner, 5′-GGGGATGTCAGAATTG-3′; EhSTIRP2 5′ reverse outer, 5′-TGGTATGCTTGGTACCTGTGAG-3′; EhSTIRP2 5′ reverse inner, 5′-GGGGATGTCAGAATTG-3′; EhSTIRP3 5′ reverse outer, 5′-TGGTATGCTTGGTACCTGTGAG-3′; EhSTIRP3 5′ reverse inner, 5′-GGGGATGTCAGAATTG-3′; 5′-RACE outer primer, 5′-GCTGTGATGGG TCCAAAGGAG-3′; 5′-RACE inner primer, 5′-GCCTGATGACGCTGTTACCT-3′; EhSTIRP1 promoter forward, 5′-GCTCAAAATCAATTGATAGCAAC-3′; EhSTIRP2 promoter forward, 5′-GGGTGCCAAATCAATTGATAGCAAC-3′; and EhSTIRP3 promoter forward, 5′-GAATTTGAAAAGAAGTTCCTAAT-3′.
5’ RACE and sequencing. 5’ Rapid amplification of cDNA ends (RACE) was carried out using a First Choice RLM 5’ RACE kit (Ambion) according to manufacturer’s instructions. PCR products were cloned using the TOPO-TA cloning kit and sequenced with M13F or M13R primers. Promoter analysis for E. histolytica Rahman was performed by PCR amplifying ~500 bp upstream of the predicted start codon from each of the three full-length EhSTIRPs from E. histolytica Rahman genomic DNA and sequencing the PCR products using an ABI PRISM 310 genetic analyzer (PE Applied Biosystems).

Cytotoxicity assay. Host cell cytotoxicity was measured using a lactate dehydrogenase (LDH) release assay (Bovision) as previously published (32). Colonic epithelial (Caco-2) cells were grown on 24-well culture dishes in Dulbecco’s modified Eagle’s media supplemented with 10% serum and penicillin/streptomycin until polarized. On the day of the experiment, Caco-2 cells were washed and the media changed to TYI-S-33 media. Amoebae were added at a 1:15 (ameba:Caco-2 cell) ratio and the cultures were incubated at 37°C. Samples of culture supernatant were taken at 3, 6, and 9 h, diluted 1:10 in 150 mM phosphate-buffered saline, mixed 1:1 with reaction mixture, and transferred to a 96-well plate. The absorbance at 490 nm was read using a Bio-Tek Instruments EL311SX plate reader. Caco-2 cells incubated in TYI-S-33 media alone or syringe lysed were used as controls.

Adhesion assay. Adhesion experiments were conducted as published previously (39). Briefly, E. histolytica cells were mixed with CHO cells in M199 media at a 1:20 ratio and incubated on ice for 90 min. The amebae were washed twice and samples counted on a hemacytometer. Amebae with three or more CHO cells attached were considered positive for adhesion in this assay.

RESULTS AND DISCUSSION

Characterization of the EhSTIRP gene family. In the current E. histolytica genome assembly and annotation, the EhSTIRP gene family consists of five members: EhSTIRP1 (GenBank accession no. XM_645112), EhSTIRP2 (XM_644280), EhSTIRP3 (XM_658084), EhSTIRP4 (XM_651335), and EhSTIRP5 (XM_647165). Three members (EhSTIRP1, EhSTIRP2, and EhSTIRP3) are approximately 8 kb in length, making them some of the largest annotated genes in the E. histolytica genome. The other two, EhSTIRP4 and EhSTIRP5, appear to be truncated forms (Fig. 1A). In the TIGR database, only EhSTIRP1 is annotated as having a signal peptide. In order to determine if the absence of a signal peptide in EhSTIRP2 and EhSTIRP3 may be a result of an annotation error, we determined the transcription start site by performing 5’ RACE for each full-length gene family member. Utilizing the first in-frame start codon downstream of the transcription initiation site, both EhSTIRP2 and EhSTIRP3 are predicted to have signal peptides with 86% and 93% probability, respectively, according to the SignalP 3.0 program (7) (data not shown). All gene family members are predicted to contain a single transmembrane domain. These data together with additional topology information (WoLF PSORTII [28]) indicate that the gene family members are likely transmembrane proteins with the vast majority of the protein facing the luminal side and only a short (34-amino-acid) cytoplasmic tail and are likely to be present on the plasma or vesicular membranes (data not shown). Future studies with antisera to EhSTIRPs will be important to definitively prove the localization of the protein products.

Comparisons of the intrafamily nucleotide and amino acid percent identities revealed nearly complete conservation at the 3’ end between EhSTIRP1, EhSTIRP2, and EhSTIRP3 and much less conservation at the 5’ end of the EhSTIRP gene family. We found that the promoter for EhSTIRP1 (defined as the region 500 bp upstream of the predicted start codon) was significantly different from EhSTIRP2 and EhSTIRP3 (47% and 45% identity, respectively) while those two gene promoters (EhSTIRP3 and EhSTIRP2) were quite similar to each other (98% identity).

EhSTIRP expression is restricted to Entamoeba strains and stages with virulence potential. Using gene-specific oligonucleotide probes from a region of low intrafamily nucleotide similarity (63% to 77% identity), Northern blot analysis was performed and indicated that each of the three full-length family members were expressed under standard axenic culture conditions in E. histolytica HM-1:1MSS; we did not detect any transcript from any of the gene family members in the non-virulent E. histolytica Rahman (Fig. 1B). EhSTIRP has an interesting expression profile: highly expressed in all strains with virulence potential (E. histolytica HM-1:1MSS, E. histolytica 200:NIH, and trophozoites during colonial invasion) and low or absent expression under all conditions with low virulence potential (E. histolytica Rahman, E. dispar, and E. histolytica cysts). Furthermore, although a number of genes have higher expression in virulent strains than in nonvirulent strains, the completely absent expression in E. histolytica Rahman and the high expression in E. histolytica HM-1:1MSS is an unusual expression profile for the E. histolytica transcriptome.

We previously reported that Entamoeba dispar does not contain a full-length intact copy of EhSTIRP according to the available genome sequence (35). We are currently investigating the mechanism by which EhSTIRP is silenced in E. histolytica Rahman. We have sequenced the majority of the EhSTIRP1 coding region in E. histolytica Rahman and found it to be nearly identical to that in E. histolytica HM-1:1MSS, indicating that E. histolytica Rahman most likely contains at least one full-length copy of the gene (33). The average promoter length in E. histolytica is ~350 bp (Jason Hackney, personal communication). We sequenced approximately 500 bp of the promoter for each of the three full-length gene EhSTIRP family members in E. histolytica Rahman and found them to be highly similar (>98%) to the promoter sequences for each gene in E. histolytica HM-1:1MSS. DNA methylation is known to silence expression of virulence genes in E. histolytica (1). We treated E. histolytica Rahman with 23 μM 5-azacytidine for 72 h and found no change in expression of EhSTIRPs, indicating that silencing via DNA methyltransferase is an unlikely mechanism of silencing in the Rahman strain (I. K. M. Ali and U. Singh, unpublished data). We also treated E. histolytica Rahman with trichostatin A (TSA) at 75 nM for 48 h and 150 nM and 300 nM for 24 h. TSA has been shown to be a potent inhibitor of histone deacetylase activity in E. histolytica (6, 38). TSA treatment modulates amebic gene expression (23) and has been shown to affect the expression level of other genes involved in amebic virulence (31). After treatment with TSA, we noted no change in EhSTIRP expression in E. histolytica Rahman by Northern blot analysis (data not shown), making it unlikely that deacetylation is responsible for the lack of EhSTIRP expression in E. histolytica Rahman. Transcriptional gene silencing has been previously documented in E. histolytica. Bracha et al. (11) showed that upon transfection with a plasmid containing the amoebapore gene and a portion of the short interspersed nuclear element EhSINE1, both chromosomal and episomal copies of the amoebapore gene were permanently silenced, and this silencing was stable even after removal of the plasmid (2). This permanent and stable epigenetic silencing mechanism is not completely understood, and whether a similar mechanism is functional in E. histolytica Rahman is not currently known. We previously reported that the
mobile elements EhLINE1 (long interspersed nuclear element) and EhLINE3 had lower expression in *E. histolytica* Rahman (33).

We are currently investigating whether the silencing of EhSTIRP in *E. histolytica* Rahman is related to the LINE or SINE elements or small RNAs.

**Down-regulation of EhSTIRP expression.** In order to down-regulate the EhSTIRP gene family, we transfected *E. histolytica* HM-1:IMSS parasites with a construct coding for the production of dsRNA specific to the 3' region of the gene family, which should target all family members for silencing. As a control, we constructed a plasmid containing nonannealing ends (Fig. 2). This methodology was previously used to down-regulate expression of the formin homology gene EhDia1, as well as the kinesin family gene EhKlp5 (17, 30). After selection with G418 at 48 μg/ml, the expression levels of all EhSTIRP genes were drastically reduced as shown by reverse transcription-PCR (data not shown) and Northern blot analysis (Fig. 3). The PCR-amplified gene segment used as a probe in the

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**FIG. 1.** EhSTIRP gene family structure and expression. (A) The EhSTIRP gene family is made up of five members. The GenBank accession number is listed for each gene. EhSTIRP4 and EhSTIRP5 may represent truncated versions of the gene. The thinner bars at either end of the gene represent the 5' promoter region and presumptive 3' untranslated region. EhSTIRP2 and EhSTIRP3 have very similar 5' promoter regions (98% nucleotide identity), whereas the 5' promoter region from EhSTIRP1 was only 45% and 47% identical to EhSTIRP3 and EhSTIRP2, respectively. The gene family members are highly similar at both the nucleotide and amino acid levels towards the 3' end/C terminus (~99%) and less similar to one another at the 5' end/N terminus. The overall similarity is higher between EhSTIRP2 and EhSTIRP3 than between either gene and EhSTIRP1. A signal peptide ( ) is shown for the three full-length gene family members and a transmembrane domain ( ) is shown for each gene. Gene-specific oligonucleotide probes are denoted as follows: ( ). A PCR-generated probe capable of detecting all full-length EhSTIRPs and used in the dsRNA constructs is denoted as follows: ( ). (B) *E. histolytica* HM-1:IMSS expresses each of the EhSTIRP full-length members, whereas *E. histolytica* Rahman does not express any of the full-length gene family members. Separate blots were probed for EhSTIRP expression in *E. histolytica* HM-1:IMSS, whereas since no signal was observed in *E. histolytica* Rahman, the same blot was sequentially probed. rRNA stained with methylene blue is shown as a loading control.
Northern blot analysis (shown in Fig. 1A) would be predicted to hybridize with all gene family members. We also looked individually at each full-length EhSTIRP gene family member and found that each transcript was significantly reduced in parasites producing dsRNA specific to the EhSTIRP gene family (Fig. 3B). In order to determine whether the EhSTIRP dsRNA parasites had altered growth, we compared their growth rates to the growth of parasites transfected with a control plasmid maintained at the same drug selection. We found no significant difference in growth compared to amebae transfected with control plasmid during the first 20 h. At time points beyond 20 h and up to 72 h, the EhSTIRP dsRNA parasites grew slightly slower than the control strains (data not shown). All parasite strains were routinely passed every 48 h and all experiments for cell killing and adhesion were performed with equal numbers of parasites for short time points (<9 h), time points at which there were no detectable growth differences between EhSTIRP dsRNA and control parasite strains.

Interestingly, after nearly 1 year in cell culture, the EhSTIRP dsRNA-based down-regulation spontaneously reverted and EhSTIRP expression reverted to wild-type levels. The spontaneous loss of dsRNA-based gene down-regulation has not previously been reported in E. histolytica, and the underlying cause of this phenomenon is currently being investigated (R. C. MacFarlane and U. Singh, unpublished data). However, all phenotypic assays described in this work were performed at a time when the gene was clearly down-regulated by either reverse transcription-PCR or Northern blot analysis.

EhSTIRP contributes to amebic cytotoxicity. We first investigated whether the EhSTIRP gene family was involved in cytotoxicity. In order to do this we compared the ability of control and EhSTIRP-down-regulated parasites to damage Caco-2 colonic epithelial cell monolayers. For these experiments we combined E. histolytica HM-1:IMSS parasites with differentiated Caco-2 cells at a 1:15 ratio. Monolayer

![FIG. 3. Northern blotting indicates a significant decrease in EhSTIRP gene expression when exposed to dsRNA targeting the 3' end of the EhSTIRP gene family. (A) Northern blot analysis shows substantial down-regulation of EhSTIRP expression in the presence of dsRNA targeted to the 3' end of the EhSTIRP gene family (parasites selected at 48 μg/ml G418). A PCR product from the 3' end of the gene family that should detect EhSTIRP1-3 was used as a probe. RNA probed with an actin probe and methylene blue-stained rRNA (grayscale) are shown as loading controls. (B) Gene-specific oligonucleotide probes were also used and indicated that the expression level of each full-length EhSTIRP was decreased in the dsRNA-expressing parasites. RNA probed with actin and methylene blue-stained rRNA (grayscale) are shown as loading controls.](image-url)
damage was assessed at 3, 6, and 9 h and compared to syringe-lysed monolayers. We found a significant difference between control and EhSTIRP-down-regulated parasites at all time points (P < 0.05) (Fig. 4). Overall, the EhSTIRP-down-regulated parasites showed a ~45% decrease in cytotoxicity across all time points examined compared to that of the control strains exposed to similar amounts of G418 drug selection. Importantly, EhSTIRP-down-regulated parasites had a decreased cytotoxic ability at both the early (3 h) and late time points (9 h) after host cell interaction. The cytotoxicity data are a significant finding, as the relative difficulty in manipulating gene expression in *E. histolytica* has limited the ability to associate particular genes with pathogenicity. To date, only a few amebic genes have been genetically shown to be involved in cytotoxicity (3, 11, 16, 25, 29, 43, 47).

**EhSTIRP has a role in amebic adhesion to host cells.** Adhesion is intimately linked to cell killing and is one of the first steps in pathogenesis on a cellular scale and a prerequisite for host cell destruction. Once a role in cytotoxicity had been established, we set out to determine whether the EhSTIRP-down-regulated parasites had a defect in adhesion to host cells. To test the adhesive ability of the parasites, we combined the control and EhSTIRP dsRNA parasites with CHO cells at a 1:20 ratio and incubated them on ice for 90 min as previously described (39). Amoebae with three or more CHO cells attached after washing scored positive for the assay. The average of five experiments is shown in Fig. 5. In these experiments we noticed a significant difference (~35% decrease in adhesion; P = 0.025) between control and EhSTIRP-down-regulated parasites in their ability to adhere CHO cells. Importantly, we did not observe complete abrogation of parasite adhesion to host cells. The primary adhesive molecule on the surface of *E. histolytica* is the Gal/GalNAc lectin. Inhibition of the amebic lectin with galactose or N-acetylgalactosamine inhibits cell adhesion and killing (22, 37). However, complete inhibition of amebic adhesion to host cells is not seen even at very high concentrations of galactose (100 to 500 mM) (22), indicating that additional adhesins may be involved. In addition to the Gal/GalNAc lectin, a 112-kDa protein complex (25) and KERP1 (42), a lysine- and glutamic-acid rich protein, have also been implicated in adhesion, although their roles are currently less well understood. Based on the observed phenotype when EhSTIRP is down-regulated, we believe that EhSTIRP facilitates adhesion and subsequent host cell destruction and functions in the molecular program of *E. histolytica* pathogenesis. Studies to elucidate the role of the EhSTIRP gene family in in vivo models of amebic colitis and liver abscess formation will be crucial future experiments.

**Conclusions.** As with most protozoan parasites, many of the genes in the *Entamoeba histolytica* genome have no homologues in other systems and are simply annotated as hypothet-
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