

Evidence for Conserved Function of γ -Glutamyltranspeptidase in *Helicobacter* Genus

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Abstract

The confounding consequences of *Helicobacter bilis* infection in experimental mice populations are well recognized, but the role of this bacterium in human diseases is less known. Limited data are available on virulence determinants of this species. In *Helicobacter pylori*, γ -glutamyltranspeptidase (γ GT) contributes to the colonization of the gastric mucosa and to the pathogenesis of peptic ulcer. The role of γ GT in *H. bilis* infections remains unknown. The annotated genome sequence of *H. bilis* revealed two putative *ggt* genes and our aim was to characterize these *H. bilis* γ GT paralogues. We performed a phylogenetic analysis to understand the evolution of *Helicobacter* γ GTs and to predict functional activities of these two genes. In addition, both copies of *H. bilis* γ GTs were expressed as recombinant proteins and their biochemical characteristics were analysed. Functional complementation of *Escherichia coli* deficient in γ GT activity and deletion of γ GT in *H. bilis* were performed. Finally, the inhibitory effect of T-cell and gastric cell proliferation by *H. bilis* γ GT was assessed. Our results indicated that one gene is responsible for γ GT activity, while the other showed no γ GT activity due to lack of autoprocessing. Although both *H. bilis* and *H. pylori* γ GTs exhibited a similar affinity to L-Glutamine and γ -Glutamyl-p-nitroanilide, the *H. bilis* γ GT was significantly less active. Nevertheless, *H. bilis* γ GT inhibited T-cell proliferation at a similar level to that observed for *H. pylori*. Finally, we showed a similar suppressive influence of both *H. bilis* and *H. pylori* γ GTs on AGS cell proliferation mediated by an apoptosis-independent mechanism. Our data suggest a conserved function of γ GT in the *Helicobacter* genus. Since γ GT is present only in a few enterohepatic *Helicobacter* species, its expression appears not to be essential for colonization of the lower gastrointestinal tract, but it could provide metabolic advantages in colonization capability of different niches.

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Introduction

γ -Glutamyltranspeptidase (γ GT) is a threonine N-terminal nucleophile (Ntn) hydrolase that catalyses the transpeptidation and hydrolysis of the γ -glutamyl group of glutathione and related compounds [1]. γ GT is widely distributed in living organisms and is highly conserved, with mammalian and bacterial homologues often sharing more than 25% of sequence identity [2]. From the ~1000 of whole genome sequenced bacterial species available in MEROPS databases (<http://merops.sanger.ac.uk> [3]), 540 (~200 genera) possess γ GT-like proteins belonging to protease family T03. Moreover, several bacterial species carry multiple copies of genes annotated as γ GT, but the majority of these genes lack functional verification.

γ GT is found in all gastric *Helicobacter* species. However, among the 20 validly published enterohepatic *Helicobacter* species (EHS), only *H. aurati*, *H. bilis*, *H. canis*, *H. muridarum* and *H. trogontum* express this enzyme [4]. In *H. pylori*, γ GT represents an important virulence factor, playing a role in colonization [5,6] and pathogenesis [7–10]. It is constitutively expressed *in vivo* and *in vitro* [11] and enables the

bacterial cells to use extracellular glutamine and glutathione as a source of glutamate [9,12]. Purified Hp- γ GT was shown to be involved in upregulation of growth factors in MKN-28 gastric cells [7], to induce apoptosis in AGS cells [10], to inhibit T-cell proliferation [9] and to contribute to *H. pylori*-mediated H₂O₂ generation [8]. By contrast, the biochemical properties of EHS γ GT and its role in the colonization of the gut and in the pathogenesis of gastrointestinal and hepatobiliary diseases are completely unknown. EHS is a phenotypically and genotypically heterogeneous phylogroup within the *Helicobacter* genus [4], including species usually colonizing the intestinal tract and/or the liver of mammals and birds. Although EHS could be considered part of the normal microbiota of rodents, some species cause diseases in these animals [13]. In particular, *H. bilis*, an endemic EHS in most experimental mouse colonies, induces disease in susceptible animals and may substantially confound interpretations of some research studies [14]. Natural *H. bilis* infection in inbred [15] or outbred mice [16] has been associated with multifocal hepatitis. Moreover, *H. bilis* has been used experimentally to induce inflammatory bowel disease (IBD) in *mdr*^{-/-} and *IL-10*^{-/-} knock-out mice [16], typhlocolitis in the

C3H/HeN mice strain [17] and cholesterol gallstone formation in C57L mice [18]. *H. bilis* is able to infect and cause diseases in different animal hosts, showing one of the broadest host spectrums in the *Helicobacter* genus [19]. It was isolated from the aborted fetus of sheep and pig [19] and from chronic hepatobiliary diseases in hamsters [20]. *H. bilis* has been also isolated from human patients with chronic diarrhoea [21] and pyoderma gangrenosum-like ulcers [22]. In addition, several studies have reported an association of this species with chronic liver diseases [23,24] or biliary tract and gallbladder cancers [25,26] in human, using either PCR or serological tests. Limited data are available on virulence determinants of *H. bilis* [27–29], and no studies to date have described the biochemical and biological proprieties of *H. bilis* γ GT (Hb- γ GT).

In contrast to observations in gastric *Helicobacter* spp., the genome sequence of *H. bilis* ATCC 43879 revealed the presence of two *ggt* copies. In this study, we used a phylogenetic and a functional approach to analyse both *H. bilis* γ GT paralogues. Although both genes were phylogenetically related to other *Helicobacter* γ GTs, analysis of the recombinant proteins, western blot using specific antibodies, complementation of *E. coli* Δ *ggt* and mutation in *H. bilis* clearly showed that only one gene was responsible for *H. bilis* γ GT activity. The γ GT of *H. bilis* exhibited a similar affinity as *H. pylori* to γ -Glutamyl-p-nitroanilide and to L-Glutamine; however, it was significantly less active. Nevertheless, *H. bilis* γ GT inhibited T-cell and gastric cell proliferation at a similar level to that observed for *H. pylori* γ GT. The inhibition observed was mediated by an apoptosis-independent mechanism and suggested a conserved function of γ GT in *Helicobacter* genus.

Results

Sequence analysis revealed marked differences between two γ GT paralogues of *H. bilis* ATCC 43879

The *ggt* paralogues HRAG_01341 and HRAG_01828 of the human-associated *H. bilis* strain ATCC 43879 genome (NCBI ACDN0000000), were named *bgh1* (*H. bilis* *ggt* homologue 1) and

bgh2 (*H. bilis* *ggt* homologue 2), respectively. Nucleotide and amino acid similarity between the two *H. bilis* γ GT paralogues, and between each homologue and *H. pylori* γ GT (Hp- γ GT; HP1118) were analysed by pairwise global alignment. The two *H. bilis* γ GT paralogues showed 65.8% and 62.0% of nucleotide and amino acid identity, respectively. Moreover, in comparison with Hp- γ GT, Bgh2 showed 65.2% of amino acid identity, while Bgh1 showed 53.4%. To detect amino acid positions potentially involved in functional change, conserved sites of both Bgh1 and Bgh2 genes were evaluated on the basis of γ GT structural data available for *H. pylori* [30,31] and *E. coli* [32]. A multi-alignment including all *Helicobacter* γ GT sequences, other bacterial γ GTs and class IV cephalosporin acylase (CA) of *Pseudomonas* sp. strain SE82, was constructed (Figure 1). In Bgh2, all the functional sites described for *H. pylori* are conserved. In Bgh1, by contrast, amino acid substitutions in the 20 kDa subunit potentially involved in functional change were observed. The substitutions were as follows: Asp421Glu, Asp422Asn, Leu432Arg, Tyr433His, Ser452Thr and Gly473Ser (Hp- γ GT numeration).

To predict the localization of both *H. bilis* γ GT paralogues, the sequences were submitted to SignalP 3.0, using both Neural networks and Hidden Markov Models (HMM) methods [33], PSLpred (Hybrid Approach Based; [34]), PSORTb v3.0.2 [35] and CELLO v2.5 [36]. These tools were unable to uniformly predict the sub-cellular localization of both Bgh1 and Bgh2. To verify the presence of other potential start codons, we analysed the upstream region of both genes. We identified a second putative start codon 54 bp upstream of the annotated methionine of Bgh2, but no other potential start codons for Bgh1. Using the new predicted start codon, all of the analysis strongly predicted Bgh2 as a periplasmic protein, and SignalP detected the presence of *sec* signal peptide sequence with a potential cleavage site between positions 30 and 31 (VFA-AS). The same prediction was obtained for all γ GTs of *Helicobacter* spp. and *C. jejuni*. No twin-arginine signal peptide cleavage site was detected in any γ GTs of *Helicobacter* spp. and *C. jejuni*.

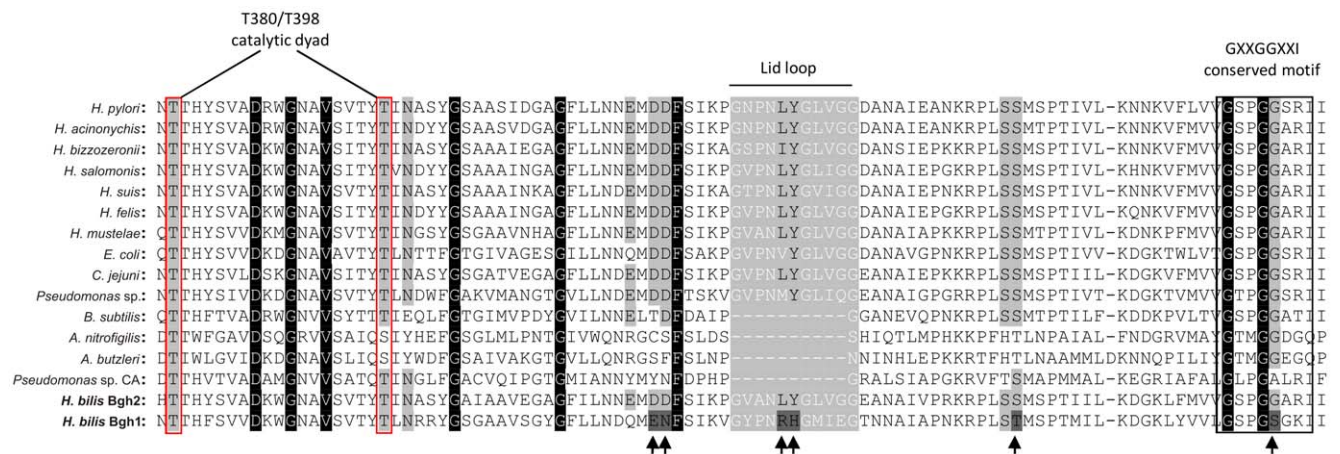


Figure 1. Multi-alignment of amino acid sequences of different bacterial γ GTs, *Helicobacter bilis* Bgh1 and Bgh2, and class IV Cephalosporin Acylase (CA). Multi-alignment between residues 379 and 477 (*Helicobacter pylori* γ GT numeration) is shown. Sequences of γ GTs of *H. pylori* (26695; HP1118), *H. acinonychis* (Sheeba; Hac_0598), *H. bizzozeronii* (CIII-1; HBZC1_08080), *H. salomonis* (O6A; EMBL FR821684), *H. suis* (HS1; HSUHS1_0265), *H. felis* (ATCC 49179; Hfelis_06880), *H. mustelae* (12198; HMU08020), *Escherichia coli* (K12; Swiss-Prot P18956), *Campylobacter jejuni* (81–176; CJ81176_0067), *Pseudomonas* sp. (A14; Swiss-Prot P36267), *Bacillus subtilis* (168; Swiss-Prot P54422), *Arcobacter nitrofigilis* (DSM 7299; Arnit_0203), *Arcobacter butzleri* (RM4018; Abu_0961), *H. bilis* (Bgh1=HRAG_01341; Bgh2=HRAG_01828) as well as class IV CA of *Pseudomonas* sp. (SE82; Swiss-Prot P15557) are shown. Residues completely conserved among the sequences are indicated with a black background. The catalytic dyad is highlighted with a red box, and the conserved motif GXXGGXXI is enclosed in a black box. The Lid loop consists of the residues G428 to G438 of *H. pylori* γ GT and is indicated in grey. Residues involved in the substrate recognition and the catalytic centre are highlighted in grey. Amino acid substitutions in Bgh1 potentially involved in functional change are indicated by arrows below the sequences. doi:10.1371/journal.pone.0030543.g001

Phylogenetic analysis of *Helicobacter* γ GTs

Phylogenetic relationships among epsilon proteobacteria and other bacterial γ GTs and *Pseudomonas* spp. CA were analysed (Figure 2). The Minimum Evolution tree based on amino acid sequence alignment showed that both *H. bilis* γ GT paralogues cluster with *Helicobacter* and *C. jejuni* γ GTs, forming, however, two distinct branches: Bgh2 and *C. jejuni* γ GT belonged to a single clade, while Bgh1 formed a separate branch.

To examine the phylogenetic relationships among *Helicobacter* γ GTs in more detail and to identify other *Helicobacter* species potentially carrying multiple γ GT sequences, consensus degenerate hybrid oligonucleotide primers were designed on the basis of highly conserved amino acid motifs (Table 1). A fragment of about 1300 bp, corresponding to an almost complete *ggt* sequence, was successfully amplified from *H. canis*, and two fragments, corresponding to an almost complete sequence of two distinct *ggt* copies, were amplified from *H. trogonutum* and *H. bilis* genomospecies FL56. From *H. aurati* and *H. muridarum*, amplification of only the N-terminal part of a single *ggt* copy was possible. An unrooted Minimum Evolution tree was built on the basis of an almost complete amino acid sequence of the γ GT pro-enzyme of *Helicobacter* spp. and *C. jejuni* (Figure 3a). Phylogenetic analysis showed that the two *ggt* copies amplified from *H. trogonutum* and *H. bilis* genomospecies FL56 corresponded to orthologues of *bgh1* and *bgh2*, whereas the sequences obtained from *H. canis*, *H. aurati* and *H. muridarum* (Figure 3a,b) clustered with other *Helicobacter* *ggt* and *bgh2*. In Figure 3, the phylogeny of the pro-enzyme was compared with those of the single sub-units. The trees showed differences in both the topology and the bootstrap supporting nodes, suggesting

different evolution of the two parts. The same results were obtained comparing the maximum likelihood trees of the nucleotide sequences (data not shown). Finally, to provide a measure for the selective pressure that the gene pair *bgh1* and *bgh2* was subjected to, the ratio between non-synonymous/synonymous substitution (Ka:Ks) was evaluated by sliding window analysis (window size 50 bp) using SWAKK [37]. A Ka:Ks ratio smaller than one was observed in all positions analysed. The average Ka:Ks ratio was calculated to be 0.175, indicating that the genes are under purifying selection. A similar Ka:Ks ratio was observed also for the *bgh1* and *bgh2* pairs of *H. trogonutum* and *H. bilis* genomospecies FL56 (data not shown).

Frequency of *bgh1* and *bgh2* genes in *H. bilis* strains

To determine the frequency of *bgh1* and *bgh2* in *H. bilis*, 33 *H. bilis* strains from our collection [38], including *H. bilis* type strain ATCC 51630^T, *H. bilis* ATCC 49314 and *H. bilis* ATCC 49320, were subjected to a specific *BclI* RFLP-PCR for *bgh1* and a specific PCR for *bgh2*. All strains were positive for the presence of both genes. Furthermore, *bgh1* orthologues of *H. bilis* type strain ATCC 51630^T and three canine strains [38] were sequenced and uncorrected distance matrices were constructed on the basis of both nucleotide and amino acid sequences. The overall nucleotide sequence identity observed varied from 94.7% to 97.3%, while the amino acid identity ranged from 97.5% to 100%.

Transcription of *bgh1* and *bgh2* by *H. bilis* CCUG 23435

The transcription of both *bgh1* and *bgh2* was evaluated in *H. bilis* CCUG 23435 at different time points. Bacterial growth was

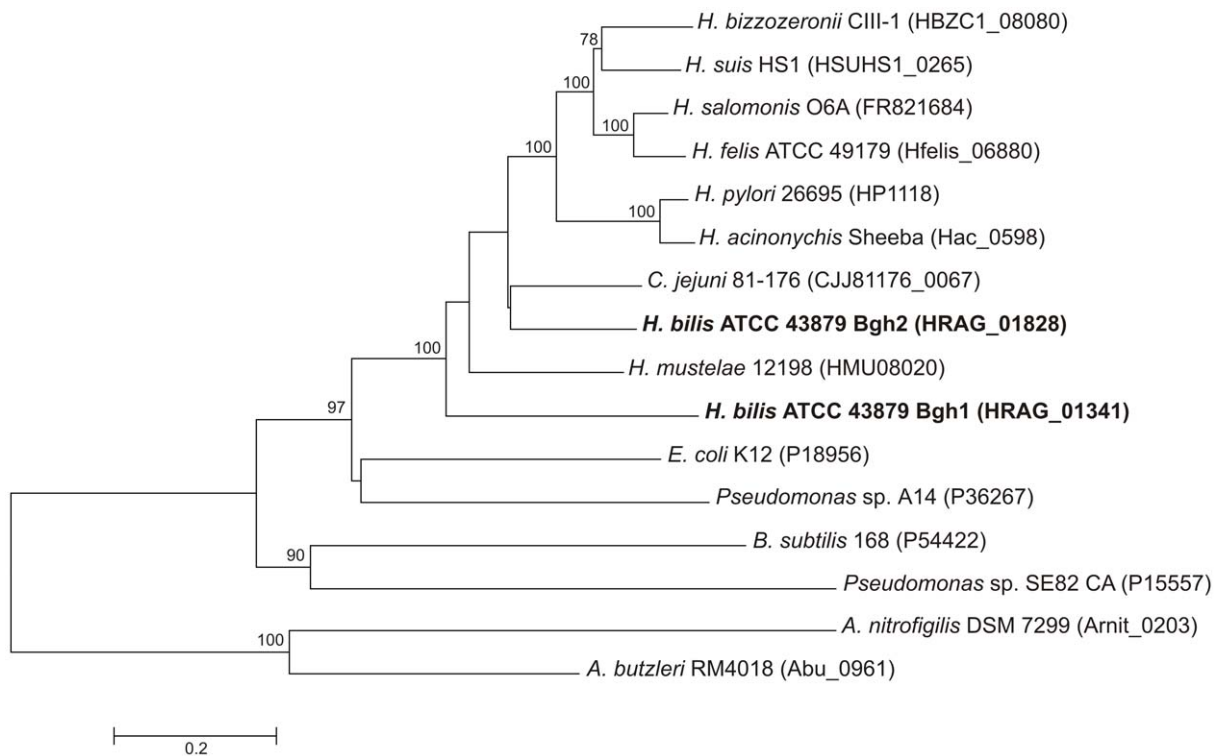


Figure 2. Unrooted tree based on complete amino acid sequences of different bacterial γ GTs, *Helicobacter bilis* Bgh1 and Bgh2, and *Pseudomonas* Cephalosporin Acylase (CA). The evolutionary history was inferred using the Minimum Evolution method and the evolutionary distances were computed using the Dayhoff matrix-based method. Bar indicates amino acid substitutions per position. Numbers at the nodes indicate support for the internal branches within the tree obtained by bootstrap analysis ($\geq 70\%$; percentages of 500 bootstraps). doi:10.1371/journal.pone.0030543.g002

Table 1. Oligonucleotides used in this study.

Oligonucleotides	Sequence
ggtCODEHOPfw-b	GATGAAGCGCGGAATGCTathgaycngc (conserved a.a. IDAA)
ggtCODEHOPrw-k	CAATTCTAATTTTCATCAGGTAGC caytgcattrg (conserved a.a. HMQW)
ggtCODEHOPfw-g	ATCCATATTATGGCTGAAGCTatgmngncargc (conserved a.a. MRQA)
ggtCODEHOPrw-g	ACGCTTCTATCAGCATAAGCTtgnckcatngc (conserved a.a. MRQA)
ggt_seqCODEHOPfw-b	GATGAAGCGCGGAATGCT
ggt_seqCODEHOPrw-k	CAATTCTAATTTTCATCAGGTAGC
ggt_seqCODEHOPfw-g	ATCCATATTATGGCTGAAGC
ggt_seqCODEHOPrw-g	ACGCTTCTATCAGCATAAGCT
1341fw	AAGTCGCACCAAAGGCAGCTAGTC
1341rw	ATCGCTTGCTGTCGCCCTATTTGTC
1828fw	AAGGTGGGGATACACTCG
1828rw	GGTGATTCTACCGCTTTTGC
1341fw-RFLP	GCAAAAAGAAGGCGAGAGTG
1341rw-RFLP	TGAAAGCGTGGAGATTCTAC
HRAG1341pBADKpnl	AGCAGGTACCCTATTATCATATCTAAAACCGCTAG
HRAG1341FwNdeI	ACGCCATATGATGCTATTATCATATCTAAAACCGCTAG
HRAG1341RwHindIII	CCCAAGCTTTTATATCTTCTCTAGGGTCTAATGTG
HRAG1828pBADKpnl	AGCAGGTACCACAGGGTTTTTAAGCGTGAGTG
HRAG1828FwNheI	CTAGCTAGCATGACAGGGTTTTTAAGCGTGAG
HRAG1828RwHindIII	CCCAAGCTTTTAAAACCTTTTTCTGGATCATTACTC
HRAG1828up5PstI	ATTCTGCAGATGACAGGGTTTTTAAGCGTGAG
HRAG1828up3XbaI	ATTCTAGAGTGATTACCCTTTTGTGATTTATAGGTTGTCC
HRAG1828dw5Kpnl	ATGGTACCGCAAGTTATGGTGCATAGCAGAGGTTGAAG
HRAG1828dw3EcoRI	ATGAATTCTAAAACCTTTTTCTGGATCATTACTC
U1catF2	ATTCTAGACGGCGGTGTTCTTTCCAAG
U1catR	ATGGTACCCGCCCTTAGTTCTAAAGGG
up1828	CGTCAGTTAAATTACTTGCAGCC
dw1828	CTTAAAAGGGGAGAGTTTATTACCTG
CatR	CCCTTATCGATTCAAGTGCATCATG
CatL	TAGTGGTCGAAATACTCTTTTCGTG

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monitored by measuring OD₆₀₀ up to 24 h, corresponding approximately to the end of the exponential phase. At 8, 12 and 24 h both genes were transcribed (data not shown).

Expression, purification and autoprocessing of recombinant Bgh1 and Bgh2 proteins

To further analyse the biochemical characteristics of both *H. bilis* γ GT paralogues, we expressed recombinant His-tagged Bgh1 and Bgh2 proteins in *E. coli*. Bgh2 was expressed without the signal peptide, while for Bgh1 the entire sequence was used. Recombinant expression of Bgh2 resulted in a soluble protein that was directly purified by Ni-affinity chromatography. By contrast, Bgh1 was expressed as an insoluble protein of ~70 kDa and was purified after refolding. The purity of the proteins on SDS-PAGE was >90% (Figure 4a). Bgh2 showed a catalytic activity for the substrate analogue L- γ -glutamyl-p-nitroanilide (gGpNA) and was synthesized as a pro-form of ~60 kDa, which undergoes autocatalytic processing to generate two subunits of ~40 and ~20 kDa (Figure 4a). By contrast, after solubilisation and refolding, recombinant Bgh1 showed no significant maturation after 24 h of incubation at 37°C and no activity for gGpNA

(Figure 4b). To estimate the secondary structure of Bgh1 and to verify the absence of a random folding, a Circular Dichroism (CD) spectrum of the recombinant protein was calculated. The spectrum minimum calculated for recombinant Bgh1 was 206 nm, indicating the absence of random coil. The secondary structure of Bgh1 was predicted by K2D3 [39] to contain 30.23% of alpha helix and 16.82% of beta sheet. The CD spectrum for Bgh1 resulted similar to the spectrum predicted by K2D3 and to those for properly folded proteins, e.g. lysozyme [40], indicating that recombinant Bgh1 has a defined secondary structure (Figure S1).

Autoprocessing of Bgh1 and Bgh2 in *H. bilis*

In order to evaluate the maturation of both Bgh1 and Bgh2 in *H. bilis*, antisera against both recombinant proteins were produced in mice and western blot analysis on *H. bilis* whole lysate was performed. The antisera showed high specificity for the corresponding *H. bilis* γ GT paralogue in ELISA test at 1:900 dilution selected for the subsequent western blots (data not shown). The western blot performed on *H. bilis* whole lysate using the antiserum against Bgh1 clearly showed a single band without indication of autoprocessing (Figure 5a). On the contrary, the western blot

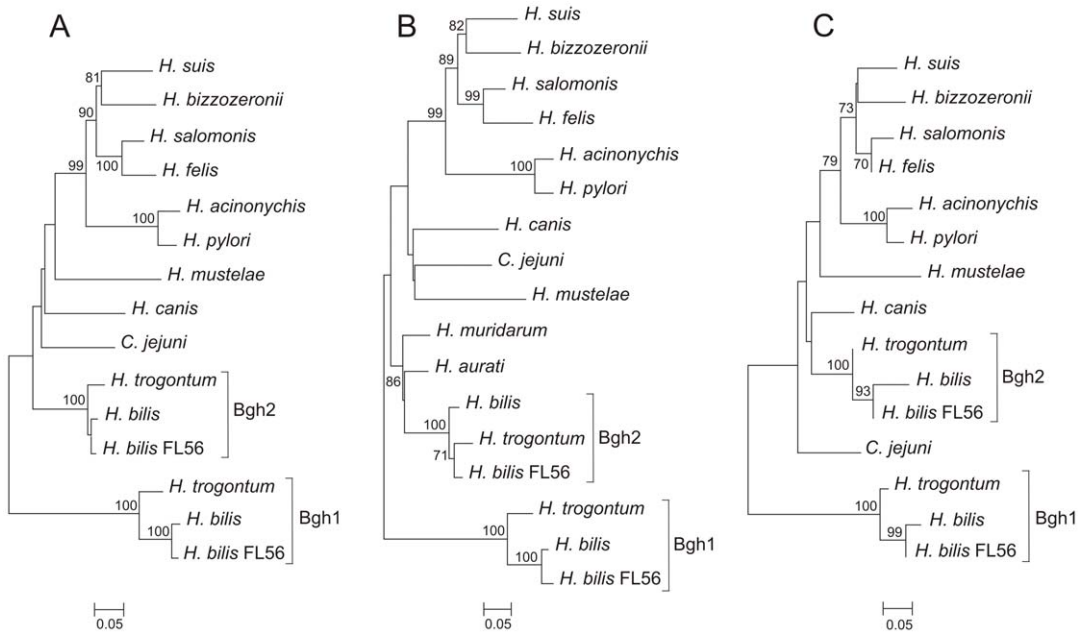


Figure 3. Unrooted tree based on the amino acid sequences of: *Helicobacter* spp. γ GTs, *Campylobacter jejuni* γ GTs and Bgh1 and Bgh2 homologues of *Helicobacter trogontum* and *Helicobacter bilis*. The evolutionary history was inferred using the Minimum Evolution method, and the evolutionary distances were computed using the Dayhoff matrix-based method. Bars indicate amino acid substitutions per position. Numbers at the nodes indicate support for the internal branches within the tree obtained by bootstrap analysis ($\geq 70\%$; percentages of 500 bootstraps). (A) Phylogeny of almost complete pro-enzyme sequences (447 AA). (B) Phylogeny of almost complete N-terminal γ GT sequences (Heavy chain; 333 AA). (C) Phylogeny of the almost complete C-terminal γ GT sequences (Light chain; 190 AA). doi:10.1371/journal.pone.0030543.g003

carried out using antiserum against Bgh2 revealed the expression of the pro-form and its corresponding two subunits (Figure 5b). These results confirmed that both proteins are expressed in *H. bilis* and that only Bgh2 undergoes autocatalytic processing.

Complementation of *E. coli* Δ ggt

To evaluate the capability of *H. bilis* γ GT paralogues to functionally complement an *E. coli* γ GT-deficient strain, Bgh1 or Bgh2 (carried on plasmid pMRg3 and pMRg5, respectively) were introduced in *E. coli* CY128 [41]. *E. coli* DH5 α grown in LB overnight at 25°C was used as positive control in γ GT assay. Only Bgh2 was found to successfully complement *E. coli* CY128, while Bgh1 was unable to restore γ GT activity (data not shown). The induction of pMRg3 in *E. coli* CY128 resulted in a clear inhibition of growth. This finding was consistent with formation of inclusion

bodies, as a consequence of the overexpression of Bgh1. However, when the protein was expressed overnight at a lower temperature (15°C) with low amount of inducer (0.002% instead of 0.2%), there was no inhibition of the growth rate of *E. coli*, suggesting a low accumulation of insoluble proteins. The successful expression of Bgh1 under improved conditions was confirmed by SDS-PAGE (data not shown). Nevertheless, Bgh1 was not able to complement *E. coli* Δ ggt, while Bgh2, expressed under the same conditions, restored γ GT activity in the mutant strain.

Construction of a *H. bilis* mutant deficient in γ GT activity

To confirm that Bgh2 is responsible of the γ GT activity in *H. bilis*, a mutant of *bgh2* was constructed in CCUG 23435 strain. The *bgh2* gene was disrupted by insertion of a chloramphenicol cassette between positions 1182 and 1246, in order to delete the threonine-threonine catalytic dyad. The successful insertion of the chloramphenicol resistance cassette and the deletion of *bgh2* were confirmed by PCR. RT-PCR and western blot analysis showed the expression of *bgh1* in both wild-type and mutant strains (data not shown), indicating that the replacement of *bgh2* had no effect on the expression of the *bgh1* paralogue gene. Since complementation approaches are currently unavailable for *H. bilis*, another independent Δ *bgh2* mutant was used as a control for secondary mutations. Neither of the *H. bilis* mutants were able to hydrolyse the substrate analogue gGpNA. Moreover, in the supernatant of *H. bilis* wild-type γ GT activity was determined with a gGpNA turnover of 0.48 μ M/min at a concentration of 50 μ g/mL of total protein. No activity was detected in the supernatant of the mutant strain MR9 (Figure S2). These results indicate that only *bgh2* encodes a functional γ GT in *H. bilis* CCUG 23435 (Hb- γ GT). The mutants grew normally *in vitro*, as described for *H. pylori* γ GT-deficient strains [9], indicating that γ GT is not essential for survival and growth of both *H. pylori* and *H. bilis in vitro*.

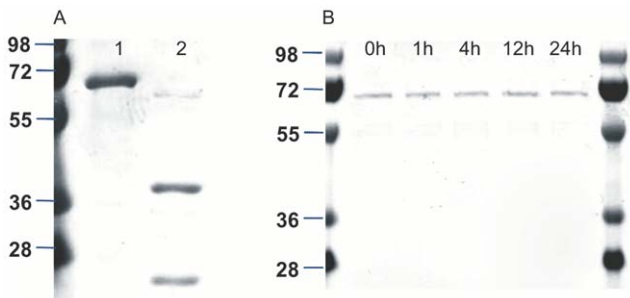


Figure 4. Purity and autoprocessing of recombinant Bgh1 and Bgh2. (A) SDS-page of the purified proteins after gel filtration: Bgh1 (Lane 1) and fully autoprocessed Bgh2 (Lane 2); (B) time-line for autoprocessing of Bgh1. doi:10.1371/journal.pone.0030543.g004

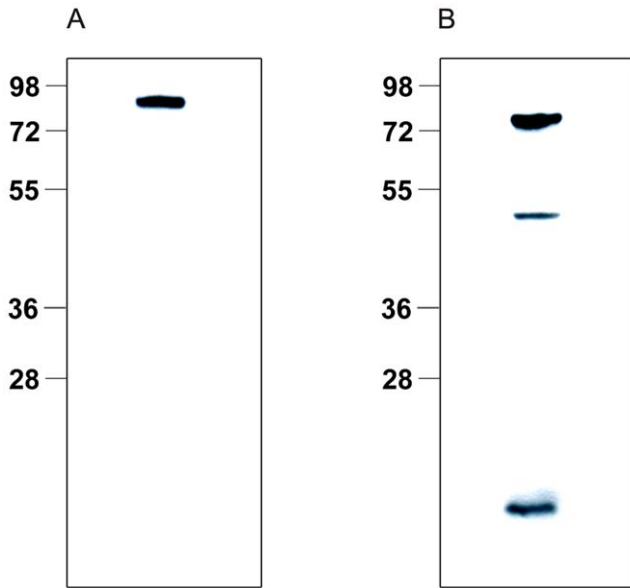


Figure 5. Western Blot on whole *H. bilis* lysate using specific antisera against Bgh1 and Bgh2. Western blot of whole *H. bilis* lysate using antisera against (A) Bgh1 and (B) Bgh2. doi:10.1371/journal.pone.0030543.g005

Biochemical characterization of recombinant Bgh2 (Hb- γ GT) and comparison with *H. pylori* γ GT (Hp- γ GT)

The kinetic parameters of recombinant Bgh2 (Hb- γ GT) were measured. The initial experiments were performed at pH 8.0. The K_M was determined by titration of the gGpNA concentrations between 1 and 1000 μ M. The processed Hb- γ GT had a K_M of $7.7 \pm 1.2 \mu$ M and a k_{cat} of $1.12 \pm 0.03 \cdot 10^2 / \text{sec}$. Compared with K_M and k_{cat} values calculated for Hp- γ GT ($K_M = 9.8 \pm 1.5 \mu$ M and $k_{cat} = 17.7 \pm 0.5 \cdot 10^2 / \text{sec}$), the apparent K_M value for Hb- γ GT was 20% lower, while the k_{cat} value was >10-fold reduced (Table 2). The pH dependence of the Hb- γ GT activity was analysed and compared to Hp- γ GT (Figure 6). The pH profile of Hp- γ GT activity was similar to reported data [30] (data not shown), whereas the activity optimum of Hb- γ GT was shifted to a more acidic pH between 6.0 and 7.0. By contrast, the K_M was increased more than 2-fold at this pH range. Initial experiments for comparing the substrate specificity of Hp- γ GT and Hb- γ GT for glutathione, D-glutamine, L-glutamine and L-glutamic acid were performed by a competition assay. A better binding affinity to the active γ GT and therefore a competitive inhibition of the gGpNA-reaction would be expected for its native substrates. As previously described [9,12], Hp- γ GT substrate inhibition was observed for both by glutathione and L-glutamine, whereas Hb- γ GT was only inhibited by L-glutamine (Figure S3).

Table 2. Comparison of the kinetic constants for *Helicobacter bilis* γ GT (Hb- γ GT) and *Helicobacter pylori* γ GT (Hp- γ GT).

	$k_{cat} [10^2 \text{ min}^{-1}]$	$K_M^{gGpNA} [\mu\text{M}]$
Hp-γGT	17.7 ± 0.5	9.8 ± 1.5
Hb-γGT	1.12 ± 0.03	7.7 ± 1.2

doi:10.1371/journal.pone.0030543.t002

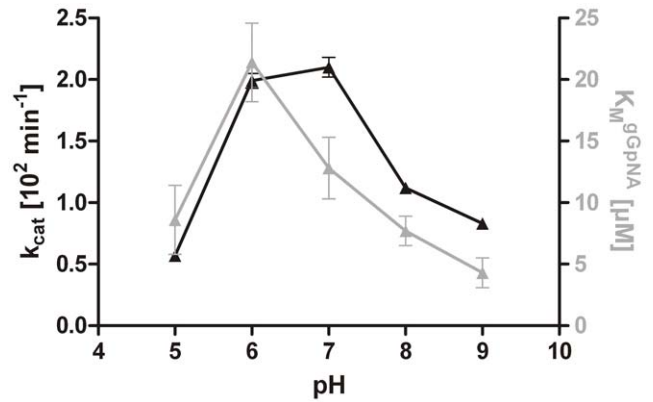


Figure 6. Comparison of the kinetic constants for *Helicobacter bilis* γ GT (Hb- γ GT) at different pH. Error bars in the graphs were calculated as SEM. The analysis was performed using Prism4 v4.03 (GraphPad Software, San Diego, CA USA). At indicated pH values, kinetic constants (k_{cat} in black and K_M in grey) for the hydrolysis of L- γ -glutamyl-p-nitroanilide (gGpNA) were determined. doi:10.1371/journal.pone.0030543.g006

Inhibition of human T-cell and AGS cell proliferation

To evaluate biological properties of recombinant Hb- γ GT and compare these with effects already observed for Hp- γ GT, the ability of Hb- γ GT to inhibit human T-cell and AGS cell proliferation was assessed. Recombinant Hb- γ GT efficiently inhibited the proliferation of Jurkat cells in a dose-dependent manner (Figure 7a). Concentrations of Hb- γ GT as low as 0.1 μ g/mL induced inhibition of T-cell proliferation. Comparison of the inhibitory potential of supernatants of wild-type and Δ ggt strains confirmed the results obtained with the recombinant proteins. The supernatants of *H. bilis* wild type reduced T-cell proliferation up to 60%, while the supernatant of Δ ggt strain had no significant inhibitory activity. When Jurkat cells were infected with acivicin pre-treated supernatant of *H. bilis* wild type, no significant reduction in proliferation of T-cells was observed, indicating that only Hb- γ GT is responsible for the inhibition of T-cell proliferation in our system (Figure 7b).

Moreover, both recombinant Hb- γ GT and Hp- γ GT showed inhibitory effect on AGS cell proliferation in a dose-dependent manner as well, although the minimum concentration of protein required was 25 time higher (2.5 μ g/mL) compared to that able to inhibit the proliferation of human T-cells (Figure 8). No statistically significant differences were observed between the two proteins.

Apoptosis in AGS cell line

The ability of recombinant Hp- γ GT and Hb- γ GT to induce apoptosis in AGS cells was evaluated using AnnexinV-Propidium Iodide double staining and FACS analysis. After 24 h of incubation, concentrations of up to 5 μ g/mL of both recombinant proteins did not induce apoptosis in AGS cells (Figure S4). The average of percentage of apoptosis observed after treatment with Hp- γ GT and Hb- γ GT was $6.9 \pm 1.2\%$ and $6.8 \pm 0.9\%$, respectively. Those values were not statistically different from the percentage of apoptosis measured in the untreated cell ($7.3 \pm 1.9\%$).

Discussion

Two *Helicobacter* γ GT genes, identified in the shotgun genome sequence of the human strain *H. bilis* ATCC 43879 on the basis of sequence homology, were subjected to functional analysis to

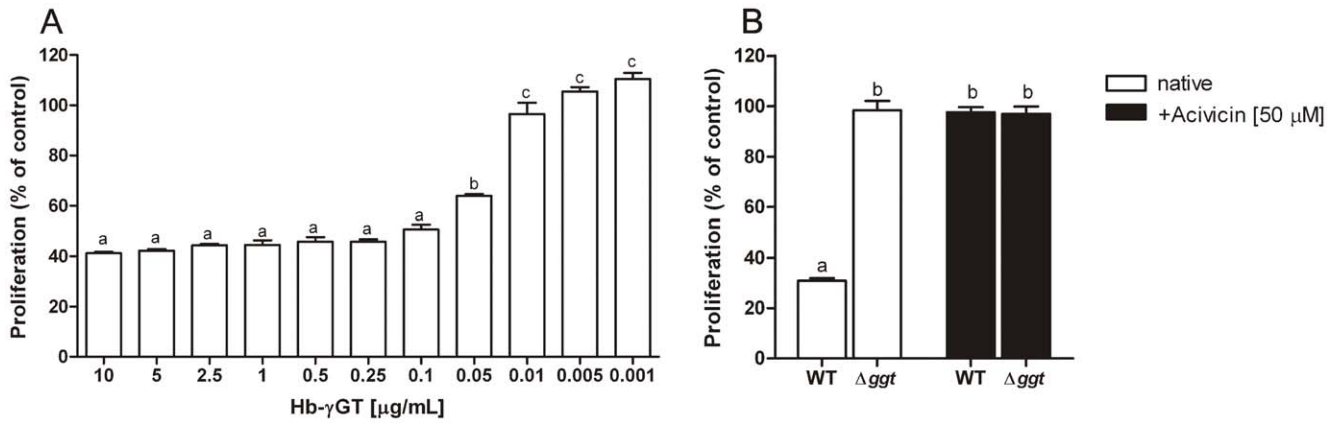


Figure 7. Inhibitory effect on T-cell proliferation (Jurkats) by *Helicobacter bilis* γ GT (Hb- γ GT). Error bars in the graphs were calculated as SEM. The analysis was performed using Prism4 v4.03 (GraphPad Software). Different letters on the bars indicate significant differences at $P < 0.05$. (A) Inhibitory effect with various amounts of recombinant Hb- γ GT (Bgh2) protein; statistical analysis was performed using one-way ANOVA, followed by the Bonferroni test. (B) Inhibitory effect of the culture supernatant of *H. bilis* wild-type CCUG 23435 (WT) and *H. bilis* Δ ggt MR9 (Δ ggt) with or without pre-treatment with Acivicin (Sigma-Aldrich); statistical analysis was performed using unpaired t -test. doi:10.1371/journal.pone.0030543.g007

confirm their annotations. Although the two genes *bgh1* and *bgh2* (*H. bilis* *ggt* homologue 1 and 2) showed significant homology and both are classified as T03.01 peptidases in MEROPS, the evaluation of conserved amino acid residues indicated a marked difference between these two paralogues, especially in the substrate-binding pocket. In fact, all critical functional residues are present in Bgh2, as well as in all the other *Helicobacter* γ GTs, but not in Bgh1. The amino acid substitutions observed in Bgh1 could potentially affect the enzyme maturation, catalytic activity and substrate specificity, suggesting different biochemical properties. Moreover, the absence of a signal peptide in Bgh1, in contrast to observations for Bgh2 and for the other *Helicobacter* γ GTs, suggests a different sub-cellular localization of this protein.

We first evaluated the phylogeny of *Helicobacter* γ GTs in relation with T03 proteases of other bacterial species. The Minimum Evolution tree clearly showed that both *H. bilis* T03.01 peptidase paralogues were phylogenetically associated with other ϵ -proteobacteria γ GTs. Nevertheless, Bgh1 formed an independent branch, indicating a different evolution. Moreover, among

Helicobacter species, only *H. trogonum* showed two distinct γ GT paralogues, as observed in *H. bilis*. In addition, the data suggested that both genes are not restricted to *H. bilis* ATCC 43879, but are common features among *H. bilis* genomospecies and universally shared by all members of the taxon.

A more detailed phylogenetic analysis further revealed that the two subunits of γ GTs have followed different evolutionary paths. The tree based on the C-terminal part resembled the topology of the tree of 16S rRNA and other housekeeping genes and includes *H. bilis* in the same branch as other enterohepatic *Helicobacter* spp. [38]. On the contrary the phylogenetic analysis of the N-terminal part separates the γ GTs of *H. bilis* and related species (*H. aurati*, *H. muridarum* and *H. trogonum*) from others enterohepatic *Helicobacter* species. On the basis of our phylogenetic analysis we can hypothesise that differing phylogeny of the subunits reflects a possible divergence of substrate specificity between the γ GTs of *H. bilis* and related species and the other *Helicobacter* or *Campylobacter* γ GTs. More studies are needed to verify this hypothesis.

In the trees based on the sequences of both subunits, the position of the clade including the orthologues of *bgh1* indicated that the paralogues potentially evolved by gene duplication from a common ancestral sequence probably before the speciation event of *H. bilis* and related species. The hypothesis of gene duplication is well supported also by the low $K_a:K_s$ ratio between the two paralogues (0,175), indicating that the two genes are under strong purifying selection [42].

Both sequence and phylogenetic analyses reveal clear differences between the two *H. bilis* T03.01 peptidases paralogues. To compare their biochemical properties, both genes were overexpressed in *E. coli* and the recombinant proteins were purified by affinity chromatography. The overexpression of Bgh1 resulted in formation of inclusion bodies due to the accumulation of an insoluble protein of ~ 70 KDa. After solubilisation and refolding, recombinant Bgh1 showed no significant maturation into two subunits and, subsequently, no γ GT activity. In contrast, recombinant Bgh2 was expressed in soluble form as a pro-enzyme of ~ 60 KDa, which undergoes autocatalytic processing to generate two subunits of ~ 40 and ~ 20 KDa and showed γ GT activity. No clear molecular evidence explains the absence of an autocatalytic process for Bgh1 since all of the amino acid residues critical for Hp- γ GT [2,43] and Ec- γ GT [44,45] autoprocessing

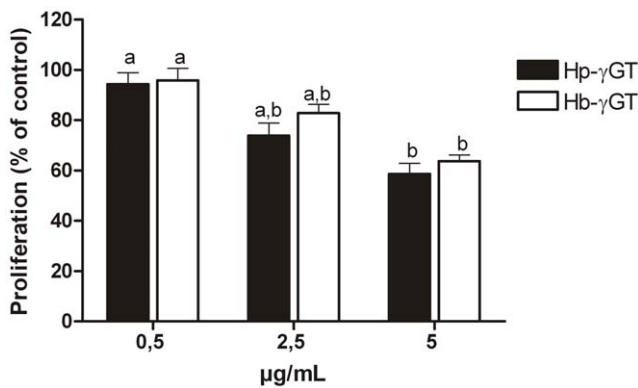


Figure 8. Inhibitory effect on AGS proliferation by *Helicobacter bilis* γ GT (Hb- γ GT). Error bars in the graphs were calculated as SEM. The analysis was performed using Prism4 v4.03 (GraphPad Software). Statistical analysis was performed using one-way ANOVA, followed by the Bonferroni test. Different letters on the bars indicate significant differences at $P < 0.05$. doi:10.1371/journal.pone.0030543.g008

are also conserved in Bgh1. We, therefore, cannot exclude that the absence of maturation could be due to misfolding *in vitro*. However, the CD spectrum of purified Bgh1 showed that the protein has a defined set of secondary structures indicating a non-random folding of the recombinant protein. Moreover, Bgh1 was unable to restore γ GT activity after complementation of *E. coli* Δ ggt mutant. Furthermore, the *H. bilis* strain in which *bgh2* was disrupted but still able to transcribe *bgh1* resulted in deficient γ GT activity. This data clearly indicate *bgh1* does not encode a functional *H. bilis* γ GT. As reported previously, the single amino acid substitution D433N converts Ec- γ GT to a class IV cephalosporin acylase [46]. Since in Bgh1 the same substitution is present, we decided to check the potential cephalosporin acylase activity of *H. bilis* CCUG 23435, which expresses both γ GT paralogues, as previously described for *E. coli* [46]. *H. bilis* did not show any cephalosporin acylase activity (data not shown), indicating that glutaryl-7-aminocephalosporanic acid is unlikely to be a substrate for Bgh1. To elucidate whether the lack of enzyme activity of Bgh1 is caused by the absence of autocatalytic maturation, western blot analysis on *H. bilis* whole lysate using specific antisera for both γ GT paralogues were performed. The results showed that *H. bilis* expresses Bgh1 which does not undergo autocatalytic processing, confirming the results obtained with the recombinant protein. However, further studies are needed to determinate potential substrates for Bgh1 and to investigate its role in the metabolism of *H. bilis*.

Analysis of the recombinant protein, western blot on *H. bilis* whole lysate, complementation of *E. coli* Δ ggt and deletion of the gene in *H. bilis* CCUG 23435 clearly showed that only *bgh2* encodes the functional *H. bilis* γ GT. Hb- γ GT exhibited similar affinity to the substrate analogue gGpNA at pH 8.0 compared with that of Hp- γ GT; however, it was significantly less proficient, having more than 10-fold less k_{cat} . Moreover, in contrast to observations for Hp- γ GT [30], the affinity of Hb- γ GT to gGpNA varied in relation to the pH, being stronger at acidic and alkaline conditions and decreasing significantly at pH 6.0 and 7.0. In addition, Hb- γ GT showed the highest activity at pH 6.0, which is lower compared to previous reports for *H. pylori* and other bacterial γ GTs [30]. The results of the competition assay suggested that Hb- γ GT and Hp- γ GT have similar substrate specificity, utilizing not only the substrate analogue gGpNA but also L-glutamine. Nevertheless, Hb- γ GT differs from Hp- γ GT in its inability to bind glutathione as substrate. Therefore, we can speculate that *H. bilis* γ GT lost its ability to interfere the host redox system. However, the enzyme enables the bacterial cells to use extracellular glutamine as the source of glutamate. This hypothesis is supported by the presence in the *H. bilis* genome of a putative glutamate transporter gene (HRAG_00091; *gltS*) showing ~60% of amino acid identity with *H. pylori* GltS homologue [47]. Glutamate transporters *gltS* as well as *ggt* genes are missing in the EHS *H. hepaticus* genomes [47]. Both *H. hepaticus* and *H. bilis* are urease-positive *Helicobacter* spp., commensal bacterial species and/or opportunistic pathogens of mice, sharing the same niches in the lower gastrointestinal tract and causing similar diseases [48]. However, *H. bilis*, but not *H. hepaticus*, was also isolated from the stomach of different hosts [19]. Therefore, the expression of γ GT activity by *Helicobacter* spp. does not appear to be essential for colonization of the lower gastrointestinal tract of mice, but it could provide metabolic advantages in colonization capability of other tissues or in the adaptation of different hosts.

Although the importance of γ GT for physiology and pathobiology of mammalian tissues has been known for many years, its function in prokaryotic metabolism and its role in interaction with the host have been studied for only a few bacterial species

[6,9,10,12,47,49–51]. Recently it has been showed that γ GT has an important role in the interaction between *H. pylori* and the host by inhibiting T-cell proliferation [9], inducing apoptosis in gastric cells [10] and contributing to *H. pylori*-mediated H₂O₂ generation [8]. Here, we showed that *H. bilis* γ GT inhibited T-cell proliferation similar to what was observed for *H. pylori* [9,10]; the inhibitory effect of T-cell proliferation was completely abolished in the γ GT-deficient mutants of *H. bilis*. Our data demonstrate that this activity is not limited to *H. pylori* but is conserved among the genus, confirming the potential immune suppressive role of secreted *Helicobacter* γ GTs. The conserved function of this enzyme among *Helicobacter* spp. with different niches and host specificity opens a new scenario on the role of γ GT in the pathogenesis of *Helicobacter* associated diseases.

We detected AGS cell proliferation inhibition after treatment with both recombinant γ GTs, as observed for T-cells, but only when a high concentration was used (2.5 μ g/mL). However, in contrast to what was previously described [10], we showed that the suppressive influence of both Hb- γ GT and Hp- γ GT on AGS cells was mediated by an apoptosis-independent mechanism. The discrepancy between our results and those of Shibayama and colleagues [10] could attribute to different methodologies applied, in particular to the use of serum starvation by these authors. An increase of apoptosis was verified in AGS cells treated with both Hp- γ GT and Hb- γ GT under the same conditions used by these authors (data not shown), apparently confirming the apoptosis inducing potential of these enzymes. However, if this phenomenon is only detectable upon stress conditions (i.e. serum starvation), we cannot exclude that the observed apoptosis is a consequence of a nonspecific activity of γ GT. Further studies are needed to establish the real role of γ GT in the apoptosis of epithelial cells.

Methods

Bacterial strains, cell line, culture condition and DNA extraction

The bacterial strains and plasmids used in this study are listed in Table 3 and 4. All *Helicobacter* strains were routinely grown on HP medium (LabM Limited, Lancashire, UK) containing 5% bovine blood at 37°C in a microaerobic atmosphere (10% CO₂; 5% O₂) (Thermo Forma, Series II Water Jacketed Incubator; Thermo Fisher Scientific, Waltham, MA USA). *E. coli* strains were cultivated on Luria-Bertani (LB) agar or broth supplemented with 100 mg/L of ampicillin, 15 mg/L of kanamycin or 20 mg/L of chloramphenicol when needed. Jurkat T-cells (DSMZ, ACC 282, Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, Braunschweig, Germany) were cultured in RPMI 1640 (Invitrogen Corporation, Carlsbad, CA, USA) with 10% of foetal bovine serum (FBS; Gibco, Invitrogen). AGS cells (CLS, 300408; Cell Lines Service, Eppelheim, Germany) were grown in Dulbecco's Modified Eagle's growth medium (DMEM high glucose, 11971, Invitrogen) supplemented with 10% FBS and 200 U/mL of penicillin and 0.2 mg/mL of streptomycin (Pen Strep, Gibco, Invitrogen). Cell lines were incubated at 37°C with 5% CO₂. Bacterial DNA extraction was performed as described earlier [19].

PCR, cloning and sequencing

Consensus DEgenerate Hybrid Oligonucleotide Primers [iCO-DEHOP [52]] were designed on the basis of highly conserved amino acid sequences of available *Helicobacter* and *C. jejuni* γ GTs. Standard PCR was done in a 50 μ L reaction mixture containing 50 ng of DNA template, 25 pmol of each primer and 1.25 U of DyeNAzymeTM (Finnzymes Oy, Espoo, Finland). The 5' consensus

Table 3. *Helicobacter* species used in this study.

Species	Strain designation and genotype [ggt locus tag or accession number and/or bgh1 homologue accession number]	Hosts	Reference
Gastric <i>Helicobacter</i> spp.			
<i>H. acinonychis</i>	Sheeba [ggt, Hac_0598]	Cheetah; Big Felines	[60]
<i>H. bizzozeronii</i>	CIII-1 [ggt, HBZC1_08080]	Dog; Human	[61]
<i>H. felis</i>	ATCC 49179 [ggt, Hfelis_06880]	Cat	[62]
<i>H. mustelae</i>	12198 [ggt, Hmu08020]	Ferret	[63]
<i>H. pylori</i>	26695 [ggt, HP1118]	Human	[64]
<i>H. salomonis</i>	O6A [ggt, EMBL FR821684]	Dog	[38]
<i>H. suis</i>	HS1 [ggt, HSUHS1_0265]	Pig	[65]
Enterohepatic <i>Helicobacter</i> spp.			
<i>H. aurati</i>	ATCC BAA-1 ^T [ggt EMBL FR821682]	Hamster	[38]
<i>H. bilis</i>	CCUG 23435 (= ATCC 43879)[ggt, HRAG_01828; bgh1, HRAG_01341] MR9 (= CCUG 23435 Δ bgh2) ATCC 51630 ^T [bgh1 homologue EMBL FR821680] ATCC 49314 ATCC 49320 KO220 [bgh1 homologue EMBL FR821686] KO214 [bgh1 homologue EMBL FR821687] KO794 [bgh1 homologue EMBL FR821688] FL56 [ggt EMBL FR821679] [bgh1 homologue EMBL FR821676]	Mouse; Human; Dog; Cat; Pig; Sheep	[19]
<i>H. canis</i>	NCTC 12739 ^T [ggt EMBL FR821681]	Dog; Cat; Human	[66]
<i>H. muridarum</i>	CCUG 29262 ^T (= ATCC 49282 ^T) [ggt EMBL FR821683]	Mouse	[38]
<i>H. trogonatum</i>	ATCC 700114 ^T [ggt EMBL FR821678] [bgh1 homologue EMBL FR821677]	Rat; Mouse; Pig	[38]

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Table 4. *E. coli* strains and plasmids used in this study.

Strains or plasmids	Relevant characteristics or genotype	Reference
<i>Escherichia coli</i> strains		
DH5 α	F ⁻ endA1 glnV44 thi-1 recA1 relA1 gyrA96 deoR nupG Φ 80d <i>lacZ</i> Δ M15 Δ (<i>lacZYA-argF</i>)U169, hsdR17(<i>r_K⁻ m_K⁺</i>), λ -	[67]
BL21 Rosetta 2	F ⁻ <i>ompT</i> <i>hsdS_B</i> (<i>r_B⁻ m_B⁻</i>) <i>gal dcm</i> (DE3) pLysSRARE2 (Cam ^R)	Novagen
JM109	endA1 glnV44 thi-1 relA1 gyrA96 recA1 mcrB ⁺ Δ (<i>lac-proAB</i>) e14- [F' traD36 proAB ⁺ <i>lacI^q</i> <i>lacZ</i> Δ M15] hsdR17(<i>r_K⁻ m_K⁺</i>)	Promega
CY128	DH5 α but Δ <i>ggt</i> Δ <i>ampC</i>	[41]
Plasmids		
pGEM [®] -T		Promega
pet28b(+)		Novagen
pBAD24	pMB1/M13 replicon, Ap ⁺ , AraC ⁺ , terminator (<i>rrnB</i>), promoter ParaBAD	[57]
pUC119	pMB1/M13 replicon, Ap ⁺ , <i>lacZα</i> ,	[68]
pUOA14	pMB1/M13 replicon, pIP1445 replicon, <i>lacZ'</i> , Cm ⁺ , Ap ⁺ , Km ⁺	[58]
pMRg1	pet28b(+), His ₆ -Bgp1 ⁺	this study
pMRg2	pet28b(+), His ₆ -Bgp2 ⁺	this study
pMRg3	pBAD24, Bgp1 ⁺	this study
pMRg5	pBAD24, Bgp2 ⁺	this study
pMRg9	pUC119, <i>bgp2::cam</i> ⁺	this study

doi:10.1371/journal.pone.0030543.t004

clamp region contains the sequencing primers used for direct sequencing of the PCR products. If secondary peaks or multiple sequences were present, the PCR products were inserted into pGEM[®]-T and sequenced on both strands using M13 primers. The area corresponding to nucleotides 702–1569 (amino acids 234–523) of *bgh1* was used to design specific primers for RFLP-PCR protocol. The PCR was done in a 50 μ L reaction mixture, as described above, and then the products were digested by *BclI* (NEB, New England Biolabs, Ipswich, MA, USA). *BclI* specifically cuts the PCR fragment in two pieces of 606 and 260 bp. EMBL accession numbers are listed in Table 3. All oligonucleotides used in this study are listed in Table 1.

Phylogenetic analysis

Phylogenetic analyses of γ GT amino acid sequences were conducted in MEGA4 [53] using the Minimum Evolution (ME) method. The amino acid sequences were aligned using MAFFT [54]. The evolutionary distances were computed using the Dayhoff matrix-based method, and the ME tree was searched using the Close-Neighbor-Interchange (CNI) algorithm (level 1). All positions containing alignment gaps and missing data were eliminated in pairwise sequence comparisons [53]. A multi-alignment of *ggt* sequences of *Helicobacter* spp and *C. jejuni* was built in MEGA4 on the basis of the amino acid sequence alignment. Maximum likelihood (ML) trees were constructed using PHYML [55] with a nucleotide evolution model selected by FindModel (<http://www.hiv.lanl.gov/>). Trees were visualized by iTOL [56]. Non-synonymous/synonymous substitution (Ka:Ks) was evaluated by sliding window analysis (window size of 50 bp) using SWAKK [37].

Expression of Bgh1 and Bgh2 and purification of recombinant proteins

Primers pairs were designed to amplify complete sequences of *bgh1* and *bgh2* genes using Phusion[®] High-Fidelity DNA Polymerase (Finnzymes). Bgh1 and Bgh2 were expressed as 6 \times His-tagged proteins by inserting the genes respectively in *NdeI* and *HindIII* or *NheI* and *HindIII* restriction sites of pet28b+ vector (Novagen[®], Merck KGaA, Darmstadt, Germany). The 6 \times His-tag was fused in the N-terminal part of the proteins. The resulting expression constructs pMRg1 and pMRg2 were sequenced and confirmed to be identical to *bgh1* and *bgh2*, respectively. Both expression constructs were used to transform *E. coli* Rosetta 2 (Novagen[®]). Expression of recombinant protein was induced with 0.5 mM IPTG for 4 h at 27°C. After centrifugation, pellets were solubilized in ice-cold binding buffer (50 mM Tris-HCl, 500 mM NaCl, 10 mM imidazole, pH 7.4) containing protease inhibitors and then lysed by subsequent sonication (5 times for 30 s) on ice. The supernatant was loaded on the HisTrap[™] column (GE Healthcare Biosciences, Pittsburgh, PA, USA) at 1 mL/min at 4°C, and bound protein was eluted with an imidazole-gradient (0–100 mM imidazole) using the elution buffer (50 mM Tris-HCl, 500 mM NaCl, 1000 mM imidazole, pH 7.4). Eluates were collected and then tested for purity by SDS-PAGE. Since purification of a soluble form of Bgh1 was not possible, the sonicated cells of *E. coli* (4 times for 1 min) were homogenized, solubilized in 6 M Gua-HCl, 1 mM DTT, 10% Glucose pH 8.5 and stirred for 1 h at 4°C. Following centrifugation, the supernatant containing the unfolded protein was loaded on a HisTrapHP[™] column (GE Healthcare) at 4°C. The bound protein was refolded with a gradient of 100 mM HEPES, 5% Glucose, 250 mM NaCl, 10 mM MgCl₂, 1 mM DTT pH 8.5 (0–100% of HEPES buffer). After the refolding, the column was washed extensively with 5 column volumes of HEPES buffer, and

bound protein was eluted with an imidazole-gradient (0–1000 mM) using HEPES buffer with 1000 mM imidazole. Eluate was collected and then tested for purity by SDS-PAGE. For further purification, fractions with the recombinant proteins from Ni-Sepharose affinity chromatography were pooled, dialysed overnight against 100 nM HEPES, 140 mM NaCl, 2.5% Sucrose pH 7.0, at 4°C and processed to the second and final purification step. The dialysed samples were loaded on a Superdex[™]75 column (GE Healthcare) and equilibrated with 100 nM HEPES, 140 mM NaCl, 2.5% Sucrose pH 7.0. All fractions collected were analysed by SDS-PAGE for the presence of recombinant proteins. Fractions containing the protein were pooled, aliquoted and stored at –80°C until further use.

Circular Dichroism (CD) spectrum of the recombinant Bgh1

To estimate the secondary structure of Bgh1 a CD spectrum of the recombinant protein was calculated. The measurements were performed on a JASCO J-810 spectropolarimeter equipped (Jasco Labor- und- Datentechnik GmbH, Gross-Umstadt, Germany) using a 0.2-mm-pathlength quartz-glass cuvette 1-mm. CD spectra were recorded with 50 μ L of the sample at final concentration of 15 μ M in 20 mM KH₂PO₄ pH 7.5, 50 mM K₂SO₄. All CD measurements were corrected by subtracting the buffer spectrum. The following parameters were used: data pitch, 0.1 nm; continuous scanning mode at 100 nm/min; response, 4 sec; band width, 1 nm; accumulation, 16. Smoothing was made by the method Means-Movement. Data were collected using Spectra manager v1.54.03 (Jasco).

Immunization of mice and Western-Blot on whole *H. bilis* lysate

Balb/c mice (3 mice per group) were immunized i.p. with 30 μ g of antigen and 10 μ g Cholera toxin (CT) (Sigma-Aldrich) as adjuvant at days 0, 7 and 14 (control group with CT only). Sera were taken 21 days after the last immunization and stored at –20°C. Animal experiments were performed according to the guidelines of the Bavarian Ministry of Animal Wealth Fare. For evaluation of Bgh-specificity, ELISA plates were coated with 2 μ g/mL recombinant Bgh1 and Bgh2 proteins in PBS pH 7.4 overnight at 4°C. After washing (4 \times 0.01% Tween-20 in PBS) and blocking (1% BSA in PBS), different dilutions of sera (in blocking buffer) were incubated for 1 h at 37°C. HRP-conjugated anti-mouse IgG antibody (1:3000) (Promega Corporation, Fitchburg, WI, USA) was incubated (1 h at 37°C) and detected with ELISA Opt EIATM (BD, Becton, Dickinson and Co., NJ, USA) according to the manufacturer's instructions.

For the western blot, *H. bilis* overnight culture was resuspended in 1 mL BHI (BD). Following lysis (sonication 4 \times 30 sec), the protein-concentration in cleared lysates was determined by BCA-Assay and adjusted to 1 mg/mL. Per lane 5 μ g of total protein were loaded. After blocking with 5% BSA in Tris-buffered saline (TBS), the Whatman Protran[®] Nitrocellulose membranes (GE Healthcare) were incubated for 1 h at RT with the sera of the immunized mice (1:800 diluted in TBS). Following washing, HRP-conjugated secondary anti-mouse IgG antibody (Promega) (diluted 1:3000 in TBS) was added and detected by the acridan-based ECL-system (Thermo Fisher Scientific).

Complementation of *E. coli* Δ ggt

bgh1 and *bgh2* were amplified as described above and were inserted between *KpnI* and *HindIII* restriction sites of pBAD24 expression vector [57], resulting in pMRg3 and pMRg5,

respectively. Both expression constructs were used to transform *E. coli* CY128 Δ ggt. The recombinant strains were grown in LB medium with ampicillin at 37°C to OD₆₀₀≈0.4, and the expression of the corresponding gene was induced with 0.2% L-arabinose for 4 h at 25°C. The basal expression of pMRg3 and pMRg5 vectors was inhibited by using 0.2% of glucose instead of L-arabinose. Expressions were verified by SDS-PAGE electrophoresis and Coomassie Blue stain. A total of 10⁹ cells were harvested by centrifugation and stored at -70°C before the γ GT assay.

Construction of *H. bilis* isogenic mutant

Chromosomal inactivation of *bgh2* was performed in *H. bilis* CCUG 23435. Deletion was introduced by allelic exchange using vector pUC119 in which ~1100 bp of the 5' -end and ~500 bp of the 3' -end of the target gene and the chloramphenicol resistance gene from pUOA14 [58] were cloned. The resultant plasmid, pMRg9, was constructed and amplified in *E. coli* DH5 α and used as a suicide plasmid in *H. bilis* CCUG 23435. *H. bilis* mutant was obtained by electroporation as described for *H. hepaticus* [59]. The mutant strain, *H. bilis* MR9 (*bgh2::cam*), was selected on an HP medium containing 5% bovine blood supplemented with chloramphenicol (20 mg/mL). The site of recombination was verified by PCR.

RNA extraction and RT-PCR

To analyse the expression of both *bgh1* and *bgh2* during growth, *H. bilis* wild-type CCUG 23435 was cultivated in Brain Heart Infusion (BHI; BD, NJ USA) containing 20% FBS and Vitox supplement (Oxoid Ltd., Cambridge, UK) at 37°C microaerobically by continuous shaking at 150 rpm. OD₆₀₀ was measured at 8, 16 and 24 h and the same amount of cells was treated with RNAProtect Bacteria Reagent (Qiagen GmbH, Hamburg Germany). RNA was extracted using RNeasy Mini Kit (Qiagen) and treated with TURBO DNaseTM (Applied Biosystems/Ambion, Austin, TX, USA). Moreover, *H. bilis* CCUG 23435 and the mutant MR9 were cultivated in YT×2 medium (16 g/L Tryptone, 10 g/L Yeast Extract, 5 g/L NaCl) containing 10% FBS at 37°C in a microaerobic atmosphere. After 24 h RNA was extracted as described above. cDNAs were synthesized from 1 μ g of total RNA using SuperScript[®] III Reverse Transcriptase (Invitrogen) using random hexamers (Finnzymes). RT-PCR reactions were carried out in a total volume of 50 μ L as described above, using 1 μ L of cDNA and 1 μ L of RNA as negative control and 20 pmol of specific primers for *bgh1* and *bgh2*. Thermocycling conditions were 94°C for 2 min followed 40 cycles of 94°C for 45 s, 58°C for 45 s and 72°C for 1 min. The PCR products were sequenced using the same primers.

T-cell and AGS proliferation assay

For measurement of the effect of *H. bilis* γ GT on T-cell proliferation, Jurkat cells were seeded at 5×10³ per well in a complete culture medium in 96-well microtiter plates and a variable amount of protein (0.001 to 10 μ g/mL) or sterile-filtrated culture supernatant (1 μ g/mL total protein) was added in a total volume of 100 μ L. As a control for the inhibition of T-cell proliferation, the recombinant γ GT from *H. pylori* 26695 (HP- γ GT) was used. The HP- γ GT was produced as previously described [9]. Additionally, supernatant were preincubated with 50 μ M Acivicin for 30 min at 37°C to inactivate the γ GT. Cells were incubated at 37°C for 48 h and analysed for proliferation as described previously [9]. Results are expressed as the mean±SEM from three experiments.

For the measurement of the effect of recombinant *H. bilis* and *H. pylori* γ GT on proliferation of AGS cells, 5×10³ cells per well

were seeded in culture medium in a 96-well plate and treated with 0.5–5 μ g/mL of protein in a total volume of 100 μ L. Cells were incubated at 37°C for 48 h and analysed for proliferation as described above.

Apoptosis in AGS cells

Apoptosis in gastric epithelial cells was determined by Fluorescence-activated cell sorting (FACS). AGS cells were seeded in a 6-well plate till they reached a confluence of 60–70%, and treated for 24 hours with 0.5–5 μ g/mL of purified recombinant γ GT proteins from *H. pylori* and *H. bilis* in DMEM with 10% of FBS. Untreated cells were used as a negative control and cells treated with 10 μ M of Staurosporine (LC laboratories, Woburn, MA, USA) as a positive control for apoptosis. Cells were harvested with 0.25% Trypsin-EDTA (Invitrogen) and 2×10⁵ cells were washed once with PBS. The pellets were resuspended in 10 μ g/mL of AnnexinV-FITC antibody (BD, Becton, Dickinson and Co., NJ USA) in FACS buffer and incubated for 20 minutes in dark at room temperature. Cell suspensions were then diluted with FACS binding buffer and 10 μ L of 50 μ g/mL of Propidium Iodide (Sigma-Aldrich, Inc, St Louis, MO, USA) was added before acquiring samples. Cell debris was excluded by scatter gating (forward vs. side scatter). Approximately 50,000 events were collected per sample using CyAn ADP FACS cytometer (Beckman Coulter, Inc., CA, USA) and data was analysed with FloJo software (Tree Star, Inc, Oregon, USA). The percentage of apoptotic cells is expressed as the double positive cell population (AnnexinV+PI) and pre-apoptotic cells stained only for AnnexinV.

γ GT enzymatic and competition assays

For determination of γ GT activity, ~10⁹ bacteria were suspended in 400 μ L of reaction buffer containing 1 mM of L- γ -glutamyl-p-nitroanilide (gGpNA) as described previously [6]. Additionally, culture supernatants, containing 50 μ g/mL protein were analysed by diluting 50 μ L of supernatant in 150 μ L of reaction buffer. The kinetic parameters for the enzymatic reaction of the recombinant proteins were measured by following the cleavage of gGpNA. To determine the exchange rate of the substrate to 4-p-nitroaniline, the reaction was monitored at 405 nm and calculated by using the reported extinction coefficient of 8800 M/cm. Enzymatic activity was measured in 0.1 M Tris HCl with a pH range from 5.0 to 9.0, 20 mM glycyglycine and gGpNA at a concentration range from 1 μ M to 2.5 mM. The experiments were carried out at 37°C for 30 min using a Mithras LB940 Well-Reader. The initial rates were determined at each gGpNA concentration and entered in the Michaelis-Menten equation. To investigate the affinity of the postulated γ GT-substrates (i.e. L-Glutamine, D-Glutamine, L-Glutamic acid and Glutathione), we made a competition experiment in which the hydrolysis and transpeptidation of gGpNA was analyzed by measuring the absorption at 405 nm in the presence of γ GT-substrates. For this assay we used 250 ng of the Hb- γ GT or Hp- γ GT per reaction with 100 μ M gGpNA, 10 mM GlyGly and the substrates in a concentration-range between 0–5 mM in 100 mM Tris/HCl pH 8.0. The measurement was performed for 15 min at 37°C in a total volume of 200 μ L.

Supporting Information

Figure S1 CD spectrum and photomultiplier voltage for recombinant Bgh1. (A) CD spectrum for recombinant Bgh1 (black line), the predicted spectrum for Bgh1 obtained by K2D3 (red line) and lysozyme (blue line). The spectrum for lysozyme was taken from the Protein Circular Dichroism Data Bank (<http://>

pcddb.cryst.bbk.ac.uk/). (B) Photomultiplier voltage of the spectrometer for recombinant Bgh1. (TIF)

Figure S2 Rate of gGpNA turnover of culture supernatant (50 μ g/mL total protein) of *Helicobacter bilis* wild-type CCUG 23435 (WT) and *H. bilis* Δ ggt MR9 (Δ ggt). The white bars show the results for untreated culture supernatants, while the black bars show the results after treatment of culture supernatants with 50 μ M of Acivicin. (TIF)

Figure S3 γ GT-activity competition with glutathione, L-/D-Glutamine and L-Glutamic Acid. Error bars in the graphs were calculated as SEM. The analysis was performed using Prism4 v4.03 (GraphPad Software). Inhibitory effect with various amounts of substrate analogues on (A) *Helicobacter bilis* γ GT (Hb- γ GT) and (B) *Helicobacter pylori* γ GT (Hp- γ GT). (TIF)

Figure S4 Apoptosis analysis of AGS cells after 24 hour treatment with γ GT recombinant proteins. AnnexinV and

PI staining of AGS cells analysed by Flow cytometer treated with 0.5, 0.25 and 5 μ g/mL of recombinant proteins from *Helicobacter bilis* (Hb- γ GT), and *Helicobacter pylori* γ GT (Hp- γ GT), compared to untreated cells and cells treated with staurosporin. (TIF)

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Author Contributions

Conceived and designed the experiments: MR CB MG JR. Performed the experiments: MR CB JR NE SJ FA. Analyzed the data: MR CB JR SJ. Contributed reagents/materials/analysis tools: MLH HH MG PV. Wrote the paper: MR CB MLH. Final approval of the version to be published: MR CB JR HH FA SJ NE PV MG MLH.

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