A gonococcal homologue of meningococcal \( \gamma \)-glutamyl transpeptidase gene is a new type of bacterial pseudogene that is transcriptionally active but phenotypically silent

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Abstract

Background: It has been speculated that the \( \gamma \)-glutamyl transpeptidase (\( \text{ggt} \)) gene is present only in \textit{Neisseria meningitidis} and not among related species such as \textit{Neisseria gonorrhoeae} and \textit{Neisseria lactamica}, because \textit{N. meningitidis} is the only bacterium with GGT activity. However, nucleotide sequences highly homologous to the meningococcal \( \text{ggt} \) gene were found in the genomes of \textit{N. gonorrhoeae} isolates.

Results: The gonococcal homologue (\( \text{ggh} \)) was analyzed. The nucleotide sequence of the \( \text{ggh} \) gene was approximately 95\% identical to that of the meningococcal \( \text{ggt} \) gene. An open reading frame in the \( \text{ggh} \) gene was disrupted by an ochre mutation and frameshift mutations induced by a 7-base deletion, but the amino acid sequences deduced from the artificially corrected \( \text{ggh} \) nucleotide sequences were approximately 97\% identical to that of the meningococcal \( \text{ggt} \) gene. The analyses of the sequences flanking the \( \text{ggt} \) and \( \text{ggh} \) genes revealed that both genes were localized in a common DNA region containing the \( \text{fbp-ggt-ggh-glyA-opcA-dedA-abcZ} \) gene cluster. The expression of the \( \text{ggh} \) RNA could be detected by dot blot, RT-PCR and primer extension analyses. Moreover, the truncated form of \( \text{ggh} \)-translational product was also found in some of the gonococcal isolates.

Conclusion: This study has shown that the gonococcal \( \text{ggh} \) gene is a pseudogene of the meningococcal \( \text{ggt} \) gene, which can also be designated as \( \Psi \text{ggt} \). The gonococcal \( \text{ggh} \) (\( \Psi \text{ggt} \)) gene is the first identified bacterial pseudogene that is transcriptionally active but phenotypically silent.

Background

Two members of the gram-negative diplococci, \textit{Neisseria meningitidis} and \textit{Neisseria gonorrhoeae}, are particularly associated with pathological infections. \textit{N. meningitidis} is specialized for the mucosa of the nasopharynx and causes meningitis and septicemia. \textit{N. gonorrhoeae} is adapted for the mucosa of the urogenital tract and causes gonorrhoea and pelvic inflammatory diseases. Both species colonize only humans and share a great deal of relatedness at the nucleotide level [1]. This high degree of relatedness is reflected in the many common genetic, biochemical and antigenic features of the two bacteria.

\( \gamma \)-Glutamyl transpeptidase (also called \( \gamma \)-glutamyl aminopeptidase) (EC2.3.2.2; GGT) catalyzes the hydrolysis of \( \gamma \)-glutamyl compounds, and is found in a variety of
bacteria such as *Escherichia coli* [2] and *Helicobacter pylori* [3,4]. To distinguish *N. meningitidis* from *N. gonorrhoeae*, GGT activity is used as one of the identification markers for *N. meningitidis* because *N. meningitidis* is positive for this activity but *N. gonorrhoeae* and related species, e.g., *Neisseria lactamica*, are not [5]. In fact, the detection of GGT activity is applied for the identification of *N. meningitidis* in the Gonocheck II enzymatic identification system (E-Y Laboratories Inc., U.S.A.) [6-9]. From these empirical facts, it was believed that the gene encoding for GGT should exist only in *N. meningitidis*, but this has not been proven yet [3].

Recent remarkable progress in the sequencing of various genomes has led to the detection of nucleotide sequences that appear to be phenotypically silent, termed pseudogenes. The pseudogenes are defined as DNA sequences of formerly functional genes rendered nonfunctional by mutations and usually identified by their disrupted open reading frames (ORFs). Pseudogenes have been identified in a variety of eukaryotes, including insects [10], plants [11], and particularly vertebrates [10,12], but are relatively few in the bacterial genomes. Notable exceptions are intracellular bacterial parasites such as *Rickettsia prowazekii* and *Mycobacterium leprae* [13], which seem to have lost many genes due to obtaining nutritional supplies from the host cells. Cryptic genes such as the *cel* operon in *E. coli* [14-16] and the flagellar master operon in the genus *Shigella* [17-19] seem to be a kind of pseudogenes, but are different from pseudogenes because the cryptic genes completely retain intact ORFs, which can be occasionally activated by rare genetic events such as mutation, recombination, insertion of elements. As a whole, compared to the pseudogenes in eukaryotes, relatively few pseudogenes have been reported in bacterial genomes [20].

In this study, a gonococcal *ggh* gene, which is highly homologous to the meningococcal *ggt* gene, was found to be pseudogene. Sequence analyses of the flanking regions of both the *ggt* and *ggh* genes suggest that both genes were derived from a gene in a common ancestor, and subsequently diversified.

**Results**

**The gonococcal ggh gene was highly homologous to the meningococcal ggt gene**

Since GGT activity was detected only in *N. meningitidis* among the related species, it was speculated that the corresponding gene also existed only in *N. meningitidis*. However, by BLAST search, the nucleotide sequences highly homologous to the meningococcal *ggt* gene were found in the genome of *N. gonorrhoeae* FA1090 [GenBank:NC_002946]. The overall nucleotide sequence of the meningococcal *ggt* homologue was approximately 95% identical to that of the meningococcal *ggt* gene (data not shown and additional file). Eleven *N. gonorrhoeae* clinical strains were analyzed by PCR, and the corresponding DNA fragments were amplified in all of these strains (Figure 1B), indicating that this gene was generally present in *N. gonorrhoeae*. To analyze whether *ggt* homologues existed in the genomes of the other neisserial strains, Southern blotting was performed (Figure 1C). DNA fragments that hybridized with the meningococcal *ggt* gene were found in the meningococcal and gonococcal genomes (Figure 1C lanes 1–3) but not in the other neisserial genomes (Figure 1C lanes 4–11). These results suggested that the meningococcal *ggt* homologue was present only in *N. gonorrhoeae* among the neisserial species examined. The putative gene in *N. gonorrhoeae* was named *ggh* (*ggt* gonococcal homologue).

**Variations of the nucleotide sequences of ggh genes among clinical isolates**

To characterize the gonococcal *ggh* gene, the *ggh* genes were amplified from the chromosomal DNA of 11 *N. gonorrhoeae* strains and sequenced. The nucleotide sequences of the *ggh* genes from 7 *N. meningitidis* strains and of the *ggh* genes from the 11 *N. gonorrhoeae* strains were aligned, and the distance matrix calculated from these data was displayed as a phylogenetic tree (Figure 2A). The results revealed that the nucleotide sequences of the gonococcal *ggh* genes were more divergent than those of the meningococcal *ggt* genes (Figure 2A). Alignment of the nucleotide sequences of the 11 gonococcal *ggh* genes also showed that the mutations in the *ggh* gene consisted of the following four polymorphisms: 1) a 6-base insertion (named Type I), 2) an ochre mutation (Type II), 3) a 7-base deletion (Type III), 4) a 46-base insertion (Type IV) (Figure 2B and Table 2). In addition, all of the 11 *ggh* genes had one-nucleotide substitutions compared to the *ggt* genes in the same 25 sites (Figure 2B and additional file), with only 2 exceptions: a one-nucleotide variation in NIID109 (48th base A to G) and another in NIID105 (213th base G to A) (Table 2). The one-base substitutions in the common sites of the *ggh* genes strongly suggested that reconstruction of the *ggh* gene would have occurred at an early stage after speciation (See Discussion).

**Putative amino acid sequences in hypothetical coding region of the ggh genes**

Due to the ochre (Type II) and 7-base deletion (Type III) mutations, the ORF in each *ggh* gene was completely disrupted by the formation of 8 or 20 stop codons (Figure 3A). In fact, none of the gonococcal isolates showed any GGT activity (data not shown and [5,9]), indicating that there was no expression of functional GGT-like protein in *N. gonorrhoeae*. All of these results showed that the gonococcal *ggh* gene was a pseudogene of the functional *ggt* gene in *N. meningitidis*. On the other hand, if the two types of mutations (Types II and III) in the *ggh* genes were
The presence of a meningococcal \textit{ggt} gene homologue in \textit{N. gonorrhoeae}. A. A schematic diagram showing the position of the set of primers used for the detection of \textit{ggt} gene homologues. The black bar shows the region of the DNA probe used for Southern blotting in panel C. B. Amplification of gonococcal \textit{ggh} gene by PCR. The genomic DNAs of neisserial species used for PCR were as follows: lane 1, H44/76 (\textit{N. meningitidis} \textit{ggt}+); lane 2, NIID113 (\textit{N. meningitidis} \textit{ggt}::IS) [49]; lane 3–13, ATCC49226, NIID54, NIID102, NIID103, NIID104, NIID105, NIID106, NIID107, NIID108, NIID109, NIID111 (\textit{N. gonor- rhoeae}). C. Southern blotting using the meningococcal \textit{ggt} gene as a probe. Two micrograms of purified chromosomal DNA digested with \textit{ClaI} were subjected to this analysis. Lane 1, H44/76 (\textit{N. meningitidis} \textit{ggt}+); lane 2, NIID113 (\textit{N. meningitidis} \textit{ggt}::IS) [49]; lane 3, NIID54 (\textit{N. gonorrhoeae}); lane 4, ATCC23970 (\textit{N. lactamica}); lane 5, ATCC13120 (\textit{N. flavescens}); lane 6, ATCC14686 (\textit{N. denitrificans}); lane 7, ATCC25295 (\textit{N. elongata}); lane 8, ATCC14687 (\textit{N. canis}); lane 9, ATCC14685 (\textit{N. cinerea}); lane 10, NIID16 (\textit{N. mucosa}); lane 11, NIID17 (\textit{N. sicca}).

\textbf{Figure 1}
A. Split graph showing the relationships among ggt genes in 7 meningococcal strains and ggh genes in 11 gonococcal strains. The sequence data have been submitted to the DDBJ/EMBL/GenBank Databases under the following Accession Numbers:

- N. meningitidis strains H44/76 [DDBJ:AB089320], H114/90 [DDBJ:AB177989], 2996 [DDBJ:AB177990], NIID68 [DDBJ:AB177991], NIID76 [DDBJ:AB177992], NIID413 [DDBJ:AB177993] and NIID414 [DDBJ:AB177994];
- N. gonorrhoeae strains ATCC49226 [DDBJ:AB175023], NIID54 [DDBJ:AB175024], NIID102 [DDBJ:AB193248], NIID103 [DDBJ:AB175025], NIID104 [DDBJ:AB175026], NIID105 [DDBJ:AB193249], NIID106 [DDBJ:AB175027], NIID107 [DDBJ:AB175028], NIID108 [DDBJ:AB193250], NIID109 [DDBJ:AB194328], NIID11 [DDBJ:AB193251].

The scale bar represents uncorrected distances, and a fit parameter is also shown.

B. Alignment of the nucleotide sequences containing 4 kinds of differences in the ggt and ggh genes of N. meningitidis strains H44/76 and H114/90; N. gonorrhoeae strains ATCC49226, NIID54, NIID103 and NIID106, respectively. Sequence identity is represented as *, polymorphism among the sequences of the 6 strains is indicated by the appropriate letter, and the absence of a base is shown by a hyphen (-). The nucleotide substitution for an ochre (Type II) mutation is shown in bold. Boxes at the Type I and Type III mutations indicate the tandem repeat. The tetranucleotide repeat in type I is also shown as a gray box. The newly deduced start codon of the meningococcal GGT is shown in underlined bold.

Figure 2

A. Split graph showing the relationships among ggt genes in 7 meningococcal strains and ggh genes in 11 gonococcal strains. The sequence data have been submitted to the DDBJ/EMBL/GenBank Databases under the following Accession Numbers: N. meningitidis strains H44/76 [DDBJ:AB089320], H114/90 [DDBJ:AB177989], 2996 [DDBJ:AB177990], NIID68 [DDBJ:AB177991], NIID76 [DDBJ:AB177992], NIID413 [DDBJ:AB177993] and NIID414 [DDBJ:AB177994]; N. gonorrhoeae strains ATCC49226 [DDBJ:AB175023], NIID54 [DDBJ:AB175024], NIID102 [DDBJ:AB193248], NIID103 [DDBJ:AB175025], NIID104 [DDBJ:AB175026], NIID105 [DDBJ:AB193249], NIID106 [DDBJ:AB175027], NIID107 [DDBJ:AB175028], NIID108 [DDBJ:AB193250], NIID109 [DDBJ:AB194328], NIID11 [DDBJ:AB193251]. The scale bar represents uncorrected distances, and a fit parameter is also shown. B. Alignment of the nucleotide sequences containing 4 kinds of differences in the ggt and ggh genes of N. meningitidis strains H44/76 and H114/90; N. gonorrhoeae strains ATCC49226, NIID54, NIID103 and NIID106, respectively. Sequence identity is represented as *, polymorphism among the sequences of the 6 strains is indicated by the appropriate letter, and the absence of a base is shown by a hyphen (-). The nucleotide substitution for an ochre (Type II) mutation is shown in bold. Boxes at the Type I and Type III mutations indicate the tandem repeat. The tetranucleotide repeat in type I is also shown as a gray box. The newly deduced start codon of the meningococcal GGT is shown in underlined bold.
artificially corrected, the hypothetical amino acid sequences were approximately 97% identical to those of the meningococcal ggt genes and were highly conserved among the gonococcal ggh genes (Figure 3B). This result also supported the idea that the ggh and ggt genes were derived from a common ancestral gene and that the translational inactivation of the gonococcal ggh gene was solely due to the ochre (Type II) and the frame-shift mutations caused by the 7-base deletion (Type III).

The genetic organization of the ggt- and ggh-flanking regions in the genomes of N. meningitidis and N. gonorrhoeae

By using the information in the database for N. meningitidis strain MC58 [21], N. gonorrhoeae strain FA1090 and N. lactamica (the neisserial species most closely related to the above two species) ST-640 strain, the flanking regions of the meningococcal ggt and the gonococcal ggh genes were further analyzed. The ggt and ggh genes were both localized in the identical gene cluster of fbp-ggt (or ggh)-glyA-opcA-dedA-abcZ in the genomes of N. meningitidis and N. gonorrhoeae, respectively (Figure 4). The fbp-glyA-dedA-

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Table 1: Oligonucleotides used in this study

| Oligonucleotide name | Position in sequence | Length (bp) | Sequence (5’-3’) | Reference |
|----------------------|----------------------|-------------|------------------|-----------|
| ggt-3                | *1265–1286           | 21          | GACTGCTGATGACATTAGCGG | [49]      |
| ggt-4                | *3250–3228           | 22          | GATTACTCAAAATTTCCCTCTA | [49]      |
| ggt-5                | *1791–1811           | 20          | CGATGCCTGCGACGCCCGAA | [25]      |
| ggt-6                | *2676–2685           | 23          | ATAGACATTGCCGCTTATCC | [25]      |
| ggt-7                | *2241–2262           | 22          | CAAGATTATCTGATTATCAAG | [25]      |
| ggt-9                | *2779–2800           | 21          | GGGCGAAACAGTGCGCAATCG | [25]      |
| ggt-10               | *2089–2068           | 21          | TGTAGCGGCACACCCATTCCG | [25]      |
| ggt-18               | *1554–1534           | 21          | CGGTAGCTCCGCTGATGTT | [25]      |

*Primer for RT-PCR

| Oligonucleotide name | Position in sequence | Length (bp) | Sequence (5’-3’) | Reference |
|----------------------|----------------------|-------------|------------------|-----------|
| ggt-29               | *1452–1475           | 24          | GGATGCTCAAGTCTCATCTGCAAT | This study |
| ggt-20               | *1682–1659           | 24          | TGTCGCTCTGCCCGCCACCCATTCCG | This study |
| ggt-31               | *1878–1901           | 24          | GTACGCGCTGTATCCCTCAAACCTG | This study |
| ggt-22               | *3079–3056           | 24          | CGCACATCGTCTTCTATACCCCAA | This study |

*Primer for primer extension

| Oligonucleotide name | Position in sequence | Length (bp) | Sequence (5’-3’) | Reference |
|----------------------|----------------------|-------------|------------------|-----------|
| primer-ext-2         | *1492–1467           | 26          | GTATTAACCTACCTTTGATGCGAT | This study |

(Biotin-labeled at the 5’-terminus)

*Numbers of positions indicate the position from the 5’-nucleotide of the ggt locus in N. meningitidis strain H44/76 [DDBJ:AB175033].

Table 2: Mutation types found in the ggh genes of 11 N. gonorrhoeae strains

| Strains       | Mutation type* |
|---------------|----------------|
|               | I (+6 bp) | II (ochre) | III (Δ7 bp) | IV (+46 bp) |
| ATCC49226     | -       | +        | -          | -          |
| NIID102       | -       | +        | -          | -          |
| NIID104       | -       | +        | -          | -          |
| NIID108       | -       | +        | -          | -          |
| NIID111       | -       | +        | -          | -          |
| NIID109 (48th A to G) | -       | +        | -          | -          |
| NIID103       | +       | +        | +          | +          |
| NIID54        | +       | +        | -          | -          |
| NIID106       | -       | +        | -          | -          |
| NIID107       | -       | +        | -          | -          |
| NIID105 (213th G to A) | -       | +        | -          | -          |

*“+” or “-” denotes the presence or absence of the mutation, respectively.
*Mutation type corresponds to the categories in Figure 2B.
A. Positions of stop codons in the hypothetical ORF in the \textit{ggh} gene. Black squares indicate the positions of stop codons and white squares indicate the ochre (Type II) mutation shown in Figure 2B. * indicates the position of the 7-bp deletion (Type III) mutation in the \textit{ggh} gene. B. Putative translated products from the corrected nucleotide sequences of the \textit{ggh} genes. The Type IV insertional mutation shown in Figure 2B is removed and the site of Type III deletion is replaced by the letter B. The site of the ochre (Type II) mutation is shown as X. The identical amino acid sequences between \textit{N. meningitidis} and \textit{N. gonorrhoeae} are shown in a black box and amino acid sequences common to only \textit{N. gonorrhoeae} are shown in a gray box.

Figure 3

A. Positions of stop codons in the hypothetical ORF in the \textit{ggh} gene. Black squares indicate the positions of stop codons and white squares indicate the ochre (Type II) mutation shown in Figure 2B. * indicates the position of the 7-bp deletion (Type III) mutation in the \textit{ggh} gene. B. Putative translated products from the corrected nucleotide sequences of the \textit{ggh} genes. The Type IV insertional mutation shown in Figure 2B is removed and the site of Type III deletion is replaced by the letter B. The site of the ochre (Type II) mutation is shown as X. The identical amino acid sequences between \textit{N. meningitidis} and \textit{N. gonorrhoeae} are shown in a black box and amino acid sequences common to only \textit{N. gonorrhoeae} are shown in a gray box.
abcZ locus was also found in the genome of *N. lactamica* but lacked the ggt and opcA homologues (Figure 4). This highly conserved genetic organization implied that a DNA island containing an original ggt gene was first incorporated into an ancestor's genome of the above three species and subsequently diversified after the speciation (see Discussion).

**The expression of the ggh gene in *N. gonorrhoeae***

The hitherto identified bacterial pseudogenes are not expressed transcriptionally or translationally [20,22-24]. To examine the ggh transcriptional expression, dot blot analysis was first performed and RNA that hybridized with the ggt probe was detected in the total RNAs of 4 *N. gonorrhoeae* strains (Figure 5A). The transcriptional expression was also confirmed by RT-PCR, and the products were amplified with all 3 sets of primers from total RNA of all 4 *N. gonorrhoeae* strains tested (Figure 5B). These results strongly suggested that the full-length ggh gene transcript was expressed in *N. gonorrhoeae*. Primer extension analysis further revealed that the gonococcal ggh RNA was transcribed from the same starting point as the meningococcal ggt mRNA (Figure 5C and 5D). All of these results indicated that the gonococcal ggh gene was transcriptionally active though it was a pseudogene.

To further study the translational expression of the truncated GGT-like protein in *N. gonorrhoeae*, Western blotting was performed with anti-meningococcal GGT rabbit antiserum [25]. When the same amounts of the whole cell extracts were analyzed (Figure 6B), approximately 15-kDa bands were detected in the extracts of NIID103 and NIID106 *N. gonorrhoeae* strains (Figure 6A lanes 5 and 7). Because the 15-kDa protein was not observed in the Aggh background of NIID103 and NIID106 *N. gonorrhoeae* strains (Figure 6A lanes 6 and 8), the 15-kDa protein was thought to be the ggh gene product whose translation was terminated at the 145th codon (Figure 3B). However, the 15-kDa protein was not found in the extracts of the ATCC49226 and NIID54 *N. gonorrhoeae* strains (Figure 6A, lanes 3 and 4) and seemed
Transcriptional expression of the ggh genes. A. Dot blot analysis using the meningococcal ggt gene as a probe. One to 16 micrograms of RNAs isolated from H44/76, HT1089 (H44/76 Δggt::spc) [39], ATCC49226, NIID54, NIID103 and NIID106 were subjected to this analysis. B. RT-PCR to detect the transcripts of the gonococcal ggh genes. The schematic figure in the box depicts the position of the primers used in this experiment (see Table 1). RT-PCR was performed without reverse transcriptase (RTase) (lanes 1 to 5) or with RTase (lanes 6 to 10). Lanes 1 and 6, H44/76 (N. meningitidis); lanes 2 and 7, ATCC49226 (N. gonorrhoeae); lanes 3 and 8, NIID54 (N. gonorrhoeae); lanes 4 and 9, NIID103 (N. gonorrhoeae); lanes 5 and 10, NIID106 (N. gonorrhoeae). The marker in the left-most lane is φ X174 DNA digested with HaeIII. Primer sets used for RT-PCR are shown on the left side, and the corresponding PCR products are indicated by arrows on the right side. C. Primer extension analysis to detect the transcriptional start point of the ggt and ggh genes. Total RNA extracted from H44/76 (N. meningitidis), ATCC49226 and NIID106 (N. gonorrhoeae) was used for the primer extension with AMV reverse transcriptase XL and biotin-labeled oligonucleotide ggt-ext-2. The arrow on the right side indicates the transcriptional start site. D. Alignment of the nucleotide sequences of the upstream regions of the ggt and ggh genes. The sequence data have been deposited in the DDBJ/EMBL/GenBank Databases under the following Accession Numbers: N. meningitidis strains H44/76 [DDBJ:AB193252], H14/90 [DDBJ:AB193253], N. gonorrhoeae strains ATCC49226 [DDBJ:AB193254], NIID54 [DDBJ:AB193255], NIID103 [DDBJ:AB193256]. An identical nucleotide is represented as *. The transcriptional start site is shown in bold as +1. The putative -35, -10 elements and Shine-Dalgarno sequence (SD) are depicted in the box, and the ideal -35 and -10 nucleotide sequences are shown above the boxes. The previously predicted start codon (ATG) [25] and newly predicted start codon (GTG) of the meningococcal {lit ggt} gene are underlined. The amino acid sequence deduced from the putative start codon GTG (shown in bold) in the meningococcal {lit ggt} gene is also shown under the corresponding nucleotide sequences.
not to be expressed in any of the other gonococcal strains (see Discussion).

**Discussion**

In this study, it was shown that the gonococcal ggh gene is a member of bacterial pseudogenes, and is transcribed but not properly translated so that active ggh protein product is not produced. 11Ph-mtTF/A [26], OsMu4-2 [27], NAB8-A [28], Makorin1-p1 [29], Dnm3a2 [30] and pseudoNOS [31] genes are known to be transcriptionally eukaryotic pseudogenes. In neisseriae, the gonococcal porA and ΨopcB genes have been reported as neisserial pseudogenes [1,32,33]. The porA pseudogene contains mutations in the promoter and the coding regions, and is not translated [24]. While some hypothetical bacterial pseudogenes with repetitive runs of A and T are speculated to be potentially expressed by transcriptional slippage [34], the expression, including the transcription, has not been proven yet. To our knowledge, the gonococcal ggh gene is the first identified bacterial pseudogene that is transcriptionally active.

The 15-kDa derivative of the putative ggh protein product is detected in the NIID103 and NIID106 N. gonorrhoeae strains but not in the ATCC49226 and NIID54 N. gonorrhoeae strains (Figure 6A). Since the predicted amino acid sequences of the putative 15-kDa proteins seem to be similar among the 4 gonococcal strains, the reason why the 15-kDa protein was not detected in ATCC49226 and NIID54 is not clear. The 15-kDa protein might be degraded in ATCC49226 and NIID54 backgrounds but not in NIID103 and NIID106 backgrounds. It seems unlikely that the 15-kDa protein encoded by the ggh gene has an essential function for N. gonorrhoeae because the 15-kDa protein is not always detected in any of gonococcal strains (Figure 6A).

Why does the gonococcal ggh gene still retain the transcriptional activity? There are some examples in which RNAs transcribed from pseudogenes have some biological functions: antisense RNA expressed from the pseudoNOS gene hybridizes with nNOS (nitric oxide synthase) mRNA, resulting in the suppression of the nNOS gene expression in the neurons of the snail *Lymnaea stagnalis* [31]. RNA of Makorin1-1p, a pseudogene of Makorin1, regulates the Makorin1 mRNA stability, which is important for the correct formation of the kidneys and bone in mice [29]. Some eukaryotic genes may be duplicated and one of the plural genes may be subsequently reconstructed due to its redundancy, resulting in a pseudogene. However, since a bacterial pseudogene generally does not have a functional counterpart (wild-type gene) in a single organism, the ggh RNA does not seem to have the same kind of biological function as the pseudoNOS and Makorin1-1p genes. In fact, we could not find any prominent phenotype for a Δggh gonococcal mutant (unpublished data). However, we cannot exclude the possibility that the ggh RNA has some biological function(s) in other milieus such as the urogenital tract and further analyses will be required to address this possibility.

The ggt and ggh genes are located in the fbp-ggt (ggh)-glyA-opca-dedA-abcZ common gene cluster in the genomes of both *N. meningitidis* and *N. gonorrhoeae* (Figure 4) [23]. The genome of *N. lactamica* lacks ggt and opcA homologues in the fbp-glyA-dedA-abcZ locus (Figure 4 and [23]). It would not be the result of chance that the two ggt (or ggh) and opcA genes are located in the same respective sites of the fbp-glyA-dedA-abcZ gene locus of both *N. meningitidis* and *N. gonorrhoeae* but not in that of *N. lactamica*. Moreover, it is also unlikely that a nonfunctional ggh gene was horizontally transferred into the gonococcal fbp-glyA-opca-dedA-abcZ gene cluster since such a pseudogene could not have been sustained due to the lack of selection [23]. Therefore, it seems more probable that the fbp-ggt-glyA-opca-dedA-abcZ gene cluster was present in an ancestor of the three neisserial species, and has been subsequently diversified independently among the three species, as shown for the opcA gene [23]. During the diversification, the meningococcal ggt gene has been maintained in an active state while the gonococcal ggh gene has been reconstructed by insertion, deletion and substitutions, resulting in the translational inactivation. In *N. lactamica*, the ggt and opcA homologues might have been lost because of their dispensability for *N. lactamica* (see below).

It is also interesting that the ggh gene has not been fully deleted from the gonococcal genome. The kinds and sites of mutations in the ggh genes are relatively few and highly conserved, respectively, among the gonococcal isolates (Figure 2B, Table 2 and additional file). It is also noted that, while in general the RNA polymerase-binding sites and SD regions of pseudogenes are highly degraded, there are also a few exceptions in species such as *Y. pestis* that could have emerged in recent evolutionary times [35]. Since the ribosome-binding sites (Shine-Dalgarno homologues) of the gonococcal ggh gene are identical to those of the meningococcal ggt gene and the RNA polymerase-binding sites are almost completely conserved (with a one-nucleotide difference) (Figure 5D), it is speculated that the reconstruction of the ggh gene may have occurred in relatively recent evolutionary times. From the evolutionary viewpoint, the drastic deletion of the approximately 2-kb DNA region containing the ggh gene in *N. gonorrhoeae* may not have been likely to occur in such a short period. Alternatively, deletion of the 2-kb DNA region may not be more advantageous for gonococcal evolution than reconstruction involving short deletions, insertions and substitutions.
Western blotting with anti-meningococcal GGT rabbit antiserum [25] (A) and Coomassie Brilliant Blue staining (B) of the whole cell extracts after SDS-PAGE. Bacterial whole cell extracts equivalent to 0.025 OD_{600} were analyzed. Lane 1, *N. meningitidis* strain H44/76; lane 2, HT1089 (H44/76 Δggt::spc); lane 3, ATCC49226 (*N. gonorrhoeae*); lane 4, NIID54 (*N. gonorrhoeae*); lane 5, NIID103 (*N. gonorrhoeae*); lane 6, HT1195 (NIID103 Δggh::spc); lane 7, NIID106 (*N. gonorrhoeae*); lane 8, HT1196 (NIID106 Δggh::spc). Black arrows show the bands corresponding to the processed small and large subunits of meningococcal GGT and the gray arrow indicates the 15-kDa band corresponding to the truncated protein product of the gonococcal ggh gene. M stands for molecular weight marker.
The maintenance of a functional ggt gene in *N. meningitidis* would have some advantages for its survival. *N. meningitidis* causes meningitis, which is due to the meningococcal invasion into the human central nervous system, including cerebrospinal fluid (CSF) [36-38]. It has been shown that meningococcal GGT has a physiological function of acquiring cysteine from environmental γ-glutamyl-cysteinyl peptides under cysteine-limited environments such as the CSF [39]. Almost all (98.8 %) meningococcal isolates from humans are positive for GGT activity [40]. All of these results suggest that the GGT activity is important for *N. meningitidis* but not for *N. gonorrhoeae*. The dispensability of GGT activity for *N. gonorrhoeae* seems to be consistent with the fact that a cysteine-limited milieu such as the CSF in humans is not a natural gonococcal habitat. However, it is not very likely that the milieu of CSF exerts selective pressure for an active ggt gene because human CSF is not a relevant milieu for human-to-human spread of meningococcus [41]. We believe that the meningococcal GGT must have some unknown essential function(s) for *N. meningitidis* and further studies will elucidate the function(s).

**Conclusion**

Our data on the *ggh* gene indicate that the *ggh* gene in *N. gonorrhoeae* is a pseudogene of the functional *ggt* gene in *N. meningitidis*. To our knowledge, the *ggh* gene is the first reported bacterial pseudogene that is transcriptionally active but phenotypically silent. Our findings may also contribute to understanding the speciation of *N. meningitidis* and *N. gonorrhoeae*.

**Methods**

**Bacterial strains and growth conditions**

Seven *N. meningitidis* strains (H44/76, H114/90, 2996, NIID68, NIID76, NIID413 and NIID414) were described in our previous reports [42,43]. Ten *N. gonorrhoeae* strains (NIID54, NIID102, NIID103, NIID104, NIID105, NIID106, NIID107, NIID108, NIID109 and NIID111), one *N. mucosa* strain (NIID16) and one *N. sicca* strain (NIID17) were clinical isolates donated by T. Kuroki and Y. Watanabe. All of the clinical strains were mutually independent: they were isolated in different periods and from different persons who lived in different areas of Japan. The following 7 neisserial strains were obtained from ATCC (species / ATCC no.): *N. gonorrhoeae* / ATCC49226; *N. lactamica* / ATCC23970; *N. flavescence* / ATCC13120; *N. denitrificans* / ATCC14686; *N. elongata* / ATCC25295; *N. canis* / ATCC14687; *N. cineria* / ATCC14685. All of the strains were stored by the gelatin disc method [42] and cultivated on GC agar (Becton-Dickinson) supplemented with 1 % IsoVitaleX enrichment (Becton-Dickinson) at 37°C in 5 % CO₂ or in GC broth (1.5 % proteose-peptone, 0.5 % NaCl, 0.05 % soluble starch, 0.1 % K₂HPO₄, 0.4 % KH₂PO₄, 1 % IsoVitaleX, 5 mM NaHCO₃ 10 mM MgCl₂) at 37°C with shaking.

**Isolation of chromosomal DNA, PCR, Southern blotting and dot blotting**

Isolation of chromosomal DNA, PCR, Southern blotting and dot blotting were performed as described in our previous report [43].

**Nucleotide sequence determination and analyses**

The ggt and ggh genes were amplified with a set of primers (ggt-3 and ggt-4) by PCR, and the resulting products were purified using High Pure PCR Product Purification Kit (Roche) as templates for sequencing. The sequencing was performed as described previously [25]. Primers used for sequencing are listed in Table 1. Raw data from the ABI sequencer were assembled with the program DNASIS ver. 3.2 (HITACHI, Japan). The sequence alignment was performed with GENETYX-MAC ver.11 (GENETYX, Japan). Phylogenetic analyses were performed by constructing a distance matrix of nucleotide mismatches using the web site of the Belozersky Institute at Moscow State University [44] and visualized by Split decomposition analysis with the program SPLISTREE, version 3.2 [45].

**Construction of ∆ggh::spc N. gonorrhoeae mutants**

HT1195 (NIID103 ∆ggh::spc) and HT1196 (NIID106 ∆ggh::spc), in which a spectinomycin resistance gene (*spc*) was inserted into the *ggh* gene, were constructed as follows: A 2-kb fragment containing the *ggh* gene of NIID103 or NIID106 was amplified by PCR and cloned in the Smal site of pUC18 (Takara Bio) to construct pHT412 or pHT413, respectively. A blunted 1-kb fragment containing the *spc* gene [25] was inserted into the EcoRV sites of pHT412 and pHT413, respectively. The EcoRV sites are located at 277 bp and 1642 bp (NIID103 [DDBJ:AB175025]) and at 271 bp and 1590 bp (NIID106 [DDBJ:AB175027]) downstream from the transcriptional start point of each *ggh* gene, respectively (see Figure 5D). The resulting plasmids were named pHT414 and pHT415, respectively. Five hundred nanograms of the plasmids linearized by digestion with EcoRI were transformed into NIID103 and NIID106, respectively, as described previously [43]. Spectinomycin-resistant clones were selected on GC agar plates containing 75 μg/ml spectinomycin. The resulting mutants were isolated as ∆ggh::spc NIID103 (HT1195), and ∆ggh::spc NIID106 (HT1196), respectively. The allelic exchange was confirmed by PCR and Southern blotting.

**RT-PCR**

Bacteria grown on GC agar plates were suspended in 20 ml of GC broth to an OD₆₀₀ of 0.1 and continuously cultured to mid-log phase (OD₆₀₀ of ~ 0.6) at 37°C with shaking. The total RNA was isolated from the harvested cells using GenElute Yeast RNA Miniprep Kit (Sigma-Aldrich) according to the manufacturer’s instructions.
bacteria as previously described [46] with an additional
treatment with DNase I. RT-PCR was performed using one
step RT-PCR Kit Ver. 1.1 (Takara Bio, Japan) with approxi-
mately 2 µg of total RNA according to the manufacturer’s
instructions. The products were visualized by electro-
phoresis in a 2 % agarose gel followed by ethidium bro-
mide staining.

**Primer extension analysis**

Fifty micrograms of total RNA and 5 pmol of biotin-
labeled primer (ggt-ext-2) were hybridized in 20 µl of
buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 250 mM
KCl). The hybridized RNA-DNA probe was treated with
35 units of AMV reverse transcriptase (RTase) XL (Takara
Bio) in a reaction mixture (250 µM dNTPs, 1 × AMV
reverse transcriptase XL buffer) at 37°C for 30 min. The
ethanol-precipitated DNA product was dissolved in 20 µl
of formamide dye (80 % formamide, 10 mM NaOH, 1
mM EDTA, 0.025 % bromophenol blue, 0.025 % xylene
cyanol). Sequencing of the ggt and ggh genes with the ggt-
ext-2 primer was performed by using 47F polymerase
sequencing high -cycle- (TOYOBO, Japan). Aliquots of the
reaction products were analyzed by electrophoresis on 8
% acrylamide-7 M urea gel followed by capillary blotting
to Hybond-N+ (Amersham). The bands were visualized
with Imaging high (TOYOBO) according to the manufac-
turer’s protocol.

**SDS-PAGE and Western blotting**

SDS-PAGE and Western blotting were performed as
described previously [43] using 1 × 103-fold diluted anti-
GCL polyclonal rabbit antiserum [25] and 2 × 103-fold
diluted horseradish peroxidase-conjugated secondary
antibody (Amersham)

**Abbreviations**

GGL, γ-glutamyl transpeptidase; IS, insertional sequence;
ORF, open reading frame; RTase, reverse transcriptase

**Authors' contributions**

HT carried out all of the studies including molecular
genetic studies, sequence determination, sequence analy-
ses and drafting the manuscript. HW made a critical read-
ing of the manuscript and final approval of the version to
be published.

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**Additional material**

**Additional File 1**

Alignment of the nucleotide sequences within the ggt and ggh genes of N. meningitidis H44/76 [DDBJ:AB089320]; N. meningitidis H119/90 [DDBJ:AB211221]; N. gonorrhoeae ATCC49226 [DDBJ:AB175023]; N. gonorrhoeae NIID103 [DDBJ:AB175025] and N. gonorrhoeae NIID106 [DDBJ:AB175029] strains, respectively. Sequence identity is represented as *, polymorphism within the sequences of the 5 strains is indicated by the appropriate letter, and the absence of a base is shown with a hyphen (–). Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2180-5-56-S1.pdf]
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