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The *Neisseria gonorrhoeae* Obg protein is an essential ribosome-associated GTPase and a potential drug target

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Abstract

Background: *Neisseria gonorrhoeae* (GC) is a Gram-negative pathogen that most commonly infects mucosal surfaces, causing sexually transmitted urethritis in men and endocervicitis in women. Serious complications associated with these infections are frequent and include pelvic inflammatory disease, ectopic pregnancy, and infertility. The incidence of gonorrhea cases remains high globally while antibiotic treatment options, the sole counter measures against gonorrhea, are declining due to the remarkable ability of GC to acquire resistance. Evaluating of potential drug targets is essential to provide opportunities for developing antimicrobials with new mechanisms of action. We propose the GC Obg protein, belonging to the Obg/CgtA GTPase subfamily, as a potential target for the development of therapeutic interventions against gonorrhea, and in this study perform its initial functional and biochemical characterization.

Results: We report that NGO1990 encodes Obg protein, which is an essential factor for GC viability, associates predominantly with the large 50S ribosomal subunit, and is stably expressed under conditions relevant to infection of the human host. The anti-Obg antisera cross-reacts with a panel of contemporary GC clinical isolates, demonstrating the ubiquitous nature of Obg. The cellular levels of Obg reach a maximum in the early logarithmic phase and remain constant throughout bacterial growth. The in vitro binding and hydrolysis of the fluorescent guanine nucleotide analogs mant-GTP and mant-GDP by recombinant wild type and T192AT193A mutated variants of Obg are also assessed.

Conclusions: Characterization of the GC Obg at the molecular and functional levels presented herein may facilitate the future targeting of this protein with small molecule inhibitors and the evaluation of identified lead compounds for bactericidal activity against GC and other drug-resistant bacteria.

Keywords: Neisseria gonorrhoeae, Drug resistance, Obg proteins, GTPase, Drug target, Mant guanine nucleotides

Background

Neisseria gonorrhoeae (GC) is a Gram-negative bacterium and a human-specific pathogen that causes gonorrhea. This sexually transmitted disease remains a global health burden. The World Health Organization estimated 106.1 million new cases in adults in 2008, which was a 21 % increase compared to 2005 [1]. The disease usually manifests as cervicitis, urethritis, proctitis, conjunctivis, or pharyngitis. A significant proportion of women (\geq 50 %) and some men (\leq 10 %) undergo asymptomatic infections and

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Department of Pharmaceutical Sciences, College of Pharmacy, Oregon State University, 433 Weniger Hall, 103 SW Memorial PI, Corvallis, OR 97330, USA therefore many cases remain undiagnosed [2]. Untreated or inadequately treated gonorrhea often has serious longterm health consequences including endometritis, pelvic inflammatory disease, ectopic pregnancy, epididymitis, and infertility [2–4]. The serious sequelae of gonorrhea are exacerbated by a significant increase of the risk of HIV acquisition [5]. Pharmaceutical interventions against GC infections are limited to antibiotic regimens, as a preventive anti-gonorrhea vaccine does not exist. Antibiotic therapies, however, have been continually challenged by the remarkable ability of the bacteria to acquire and retain resistance [6]. Treatment failures associated with the current emergence of GC with decreased susceptibility to the last effective treatment



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option, third-generation cephalosporins, are concerning and emphasize the pressing need for the development of alternative antimicrobial strategies to combat drugresistant gonorrhea [1, 6-14].

Here, we focus on biochemical and functional characterization of the GC homolog of conserved bacterial Obg GTPases, NGO1990 (henceforth Obg_{GC}), as a target for the discovery of anti-gonorrhea compounds. Obg proteins (also recognized as YhbZ or CgtA) belong to the OBG-HflX superfamily within the TRAFAC (translational factors) class of P-loop (phosphate-binding loop) GTPases (reviewed in [15, 16]). The OBG family is comprised of four subfamilies: Obg, Nog1, DRG, and YchF. The family exists in all three domains of life with bacteria possessing Obg and YchF, archea having two Obg proteins and YchF, and eukaryotes commonly encoding four Obg proteins and YchF [15-18]. Structurally, the bacterial Obg proteins contain highly conserved N-terminal- and central-domains, and C-terminal domain that can vary in length and sequence, or as in Chlamydia, may even be absent [16, 19]. The N-terminal domain is glycinerich and has a unique fold, the Obg fold [20]. The signature GTP-binding domain shares overall topology with the small Ras-like GTPases, but the biochemical features of the Obg proteins are distinct from those of eukaryotic Ras-like proteins [21-26].

The name Obg originates from spoOB-associated GTPbinding protein of *Bacillus subtilis*, in which the *obg* gene was identified as a part of the *spo0B* operon [27]. Since its identification in 1989, Obg homologs have been demonstrated to be essential for viability not only in B. subtilis, but also Streptomyces coelicolor, Staphylococcus pneumoniae, S. aureus, Haemophilus influenzae, Caulobacter crescentus, Escherichia coli, Vibrio harveyi and V. cholerae [27–33]. These findings strongly suggest that Obg proteins are crucial for the survival of both gram-positive and gram-negative bacteria. The depletion of cellular Obg levels results in species-specific pleiotropic effects on bacterial physiology, including alterations of ribosome maturation and DNA synthesis; cell division and morphology; and induction of general, as well as (p)ppGpp-mediated stringent stress responses [15, 16, 34]. Growing lines of evidence support the link between Obg and ribosome function. Obg predominantly associates with the 50S ribosomal particles, and long-term Obg depletion results in reduced levels of 70S monosomes, ribosomal proteins S1, S14, S21 and L10, as well as a perturbed polyribosome profile [17, 22, 32, 35–37]. Recent studies suggest that in E. coli cultured under standard laboratory growth conditions, Obg acts as a checkpoint in the final steps of 50S subunit assembly and, via an interplay with (p)ppGpp, might modulate the production of large ribosomal particle in response to environmental cues [34]. Introduction of the temperature-sensitive variant of obg, G80E, in C.

crescentus caused a decline in both the growth rate and the amount of 50S subunits, even under permissive conditions [38]. Additional ribosome defects were not observed in non-permissive temperatures; however, the bacteria rapidly halted cell cycle progression and lost viability. Thus, the essential nature of Obg likely does not result directly from its function in the late stages of 50S subunit assembly [15, 38]. Obg might provide a key molecular nexus between different metabolic pathways to regulate cellular processes in response to the energy status of the cell [16]. All Obg proteins characterized to date bind GDP and GTP, and display relatively slow GTP hydrolysis, which can be moderately stimulated in the presence of purified 50S ribosomal particles [22, 24-26, 34, 39]. The mechanistic insights into how Obg participates in different metabolic paths and stress responses remain to be elucidated. Nevertheless, the Obg proteins appear to be promising molecular targets for the development of broadspectrum antibiotics against drug-resistant bacterial infections because of their essential nature, conservation, and strong link with pivotal physiological processes.

Results and discussion

Local gene context and Obg_{GC} domain architecture

In virtually all bacteria the *obg* gene has been reported to be physically linked to *rplUI* and *rpmA*, encoding 50S ribosomal proteins L21 and L27, respectively [15]. The inspection of the genetic organization of the *obg* region in available completed genome sequences of *Neisseriaceae*, however, revealed that this arrangement is not followed. In GC strain FA1090, the putative protein NGO1990, annotated as Obg, is the last ORF in a cluster comprised of NGO1989 and NGO1988. NGO1989 is a hypothetical protein present also in GC DGI18 and PID24-1, whereas NGO1988, encoding a homolog of the rRNA small subunit methyltransferase I, is located upstream of *obg* in other *Neisseriaceae*.

Analysis of the predicted amino acid sequence of NGO1990 from FA1090 revealed a typical structure of Obg GTPases (Fig. 1) and significant similarities to other Obg proteins (Table 1). The N-terminal domain of Obg_{GC} (amino acids 3-158) contains 26 glycine residues, similar to the *B. subtilis* Obg [20]. The central, GTP-binding domain (residues 160-348), includes five conserved G motifs (G1-G5) and two switch elements (switch I and switch II) that determine the active or inactive state of the G protein [15, 16]. As expected, the C-terminal domain shows the lowest conservation when compared to Obg homologs from different bacterial species (Additional file 1: Figure S1). Nevertheless, this region of Obg_{GC} contains clusters of acidic residues, a feature characteristic of Obg proteins [17, 33]. This charged C-terminus has been shown to be important for Obg



association with 50S ribosomal particles in *C. crescentus* as well as GTP and GDP binding in *V. harveyi* [17, 26].

Purification of Obg variants and evaluation of anti-Obg $_{\rm GC}$ antisera

To begin the characterization of Obg_{GC} , N- and Cterminally His-tagged versions of the wild type NGO1990, N-His-Obg_{GC} and C-His-Obg_{GC} (respectively), were overexpressed in *E. coli* BL21(DE3), and purified. The purified recombinant N-His-Obg_{GC} was subsequently used to obtain polyclonal rabbit anti-Obg_{GC} antisera. The antisera specifically recognized both the native and recombinant versions of Obg_{GC} (Fig. 2a). The purified proteins migrated in SDS-PAGE more slowly than the native protein and accordingly with the deduced molecular mass of Obg_{GC} (41.998 kDa) with the addition of the histidine epitope. Further, the antibodies cross-reacted with Obg homologs in the *N. meningitidis* serogroup B strains MC58 and NZ98/254 but failed to recognize Obg from *N. weaveri* and *E. coli* despite their 85 and 56 % identity to Obg_{GC} , respectively (Fig. 2b and Table 1). It is possible that the anti- Obg_{GC} antisera bind to the highly variable C-terminal domain of Obg_{GC} .

Obg_{GC} binds GTP and GDP

GTPases cycle between being turned "on" in the GTPbound state and turned "off" in the GDP-bound state (Fig. 3a). In each state, G proteins undergo conformational changes and downstream effectors sense the GTP-bound protein complexes. Switch-off involves the exchange of GTP for GDP or hydrolysis of the γ -phosphate of GTP. The fluorescent N-methyl-3'-O-anthranoyl (mant) guanine nucleotide analogs, mant-GTP and mant-GDP, have been widely utilized for examining the nucleotide binding and GTP hydrolysis of various G-proteins including Obg homologs. The highly environmentally sensitive fluorescence of the mant group enables detection of nucleotide-protein interaction [22, 24, 26, 38, 40–43]. The binding of GTP to Obg requires the presence of physiological Mg²⁺ concentrations in *C. crescentus, E. coli*, and *V. harveyi*, whereas

Table 1 Comparison of the amino acid sequences of the N. gonorrhoeae Obg protein with Obg homologs

Organism	Accession number	Protein length	Region aligned	% identity	% similarity
Neisseria gonorrhoeae NCCP11945	B4RQP4	384	1-384	100	100
Neisseria meningitidis MC58	Q9JXE5	384	1-384	98	98
Neisseria lactamica 020-06	E4ZAV0	384	1-384	97	97
Neisseria weaveri LMG 5135	G2DJV3	384	1-384	85	91
Escherichia coli K12	P42641	390	1-344	56	71
Caulobacter crescentus NA1000/CB15N	B8GYI7	354	1-320	52	67
Bacillus subtilis 168	P20964	428	2-328	49	68
Chlamydia trachomatis D/UW-3/Cx	O84423	335	2-335	43	62
Homo sapiens	Q9H4K7*	406	72-364	40	58
Homo sapiens	A4D1E9**	308	77-293	35	52
Saccharomyces cerevisiae 204508	P38860	518	295-488	40	61

*ObgH1 (GTP binding protein 5, GTBP5; mitochondrial ribosome-associated GTPase 2, MTG2)

**ObgH2 (GTP binding protein 10, GTPBP10)

Α

T192AT193A **FA1090** P_{lac}::obg_{GC} Obg wt α-Obg α-MtrE α-Nq-MIP B N. meningitidis NZ98/254 N. meningitidis MC58 N. weaveri 1032 E. coli MC1061 =A1090 α-Obq Fig. 2 Validation of polyclonal rabbit anti-Obg_{GC} antisera. (a) The polyclonal rabbit anti-Obg_{GC} antibodies were used to probe the whole-cell lysates derived from wild type FA1090 and isogenic Plac::obg_{GC} as well as purified variants of Obg_{GC}. The bacteria were harvested following 2 h of growth in GCBL with (+) and without (–) 100 μ M IPTG, and the samples were matched by equivalent OD₆₀₀ units. Purified recombinant proteins (40 ng) include wild type Obg_{GC} with N-terminal 6 × His tag, (N-His-Obg_{GC}) and N-His-Obg_{GC} with T192AT193A substitutions. (b) Samples of whole-cell lysates derived from various Neisseria species, as indicated, were harvested from GCB and matched by equivalent OD₆₀₀ units. All samples were separated in 4-20 % Mini-PROTEAN TGX precast gels, the proteins were transferred onto the nitrocellulose membrane and probed with polyclonal rabbit anti-Obg_{GC} antisera raised against N-His-Obg_{GC}

Purified:

Obg-GDP complexes form over a wide range of Mg^{2+} concentrations [22, 24, 26]. The total intracellular Mg^{2+} content is about 100 mM in *E. coli* and *B. subtilis* and includes bound and free Mg^{2+} , with the latter ranging from 1–5 mM [44–46].

To determine whether Obg_{GC} requires Mg^{2+} to optimally bind mant-nucleotides, N-His- Obg_{GC} was incubated

with increasing concentrations of Mg^{2+} and either mant-GTP or mant-GDP. The Obg_{GC} binding profiles obtained for both nucleotides differed noticeably similarly to that observed for other Obg family members [22, 24, 26]. The optimal formation of mant-GTP-Obg_{GC} complexes occurred between 5 and 10 mM Mg²⁺, as indicated by maximal fluorescence (Fig. 3b, red circles). The binding of mant-GDP to Obg_{GC} did not require Mg²⁺ and was inhibited at above 1 mM concentrations (Fig. 3b, blue squares).

Subsequently, binding of mant-GTP and mant-GDP was assessed for both N-and C-His-Obg_{GC}, as addition of a six-histidine epitope to the C-terminus of the V. harveyi Obg completely abolished interaction with GTP and resulted in a weak binding of GDP [26]. Likewise, the C. crescentus Obg containing influenza virus hemagglutynin tag demonstrated a reduction in protein function [17]. In contrast, the C-His-Obg_{GC} showed very similar properties to N-His-Obg_{GC} (Fig. 3c). Binding of either variants of Obg_{GC} to mant-GTP and mant-GDP led to 1.9- and 1.3-fold enhancement in mantnucleotide fluorescence, respectively. These results suggest that subtle perturbations to the C-terminus of Obg_{GC} are not detrimental to protein function. The C. abortus Obg naturally lacks the C-terminal domain, yet the protein is a functional GTPase, with similar activity to other Obg proteins, and binds to the 50S large ribosomal particle [19].

mantGTP hydrolysis by Obg_{GC}

We next examined the GTPase activity of purified N-His-Obg_{GC} and C-His-Obg_{GC} by monitoring the decrease in fluorescence that is associated with the single-turnover conversion of bound mant-GTP to bound mant-GDP (Fig. 3d). The peak of fluorescence was recorded for 3 h at 1 min intervals. The reduction in fluorescence was fitted to a single exponential decay with a firstorder rate constant, k_h , of 2.3×10^{-4} s⁻¹ and 2.1×10^{-4} s^{-1} , or half life (T_{1/2}) of 49.1 min and 53.2 min, for N- $\mathsf{His}\text{-}\mathsf{Obg}_{\mathsf{GC}}$ and $\mathsf{C}\text{-}\mathsf{His}\text{-}\mathsf{Obg}_{\mathsf{GC}}\text{,}$ respectively. Therefore, the GTP hydrolysis rates of both Obg_{GC} variants are very similar and are approximately twenty times slower than that of the V. harveyi Obg and two-fold slower than C. crescentus and E. coli Obg proteins, respectively [22, 24, 26]. These differences may reflect distinct Obg control or function in distantly related bacterial species.

Alteration of switch I element of Obg_{GC} abolishes GTP but not GDP binding abilities

The two adjacent threonine residues, T192 and T193, which coordinate Mg^{2+} , are ubiquitously present within the G2 domain of Obg proteins (Fig. 1). However, their function has been assessed only in *C. crescentus* [47]. To address their importance for guanine nucleotide binding in Obg_{GC}, the double T192AT193A mutant protein with N-terminal-His epitope was constructed and purified.



Compared with the wild type Obg_{GC} , the mutated protein exhibited completely impaired mant-GTP binding, whereas a 1.3-fold increase in fluorescence was observed in the presence of mant-GDP, indicating unaffected formation of Obg-GDP complexes (Fig. 3c). A similar effect was observed in the *C. crescentus* Obg, and T193 was identified as the pivotal residue. The *obg* T193A allele was not able to support *C. crescentus* growth, which demonstrated that the Obg GTPase activity was a prerequisite for cell viability [47].

Depletion of Obg_{GC} has deleterious effect on GC survival

To examine whether Obg_{GC} plays a critical function in GC physiology, we used an allelic exchange approach and placed NGO1990 under the control of the isopropyl- β -D-thiogalactoside (IPTG)-inducible promoter, P_{lac} , in its native chromosomal locus in GC FA1090. The resulting conditional knockout strain, FA1090 P_{lac} :: obg_{GC} , failed to grow when inoculated directly from the freezer stocks onto the gonococcal base agar solid medium (GCB) lacking IPTG, whereas robust bacterial growth was observed in the presence of the inducer (Fig. 4a).

Subsequently, to examine the effect of Obg_{GC} depletion on GC viability over time, non-piliated and translucent colonies of FA1090 P_{lac} :: obg_{GC} , harvested from GCB supplemented with 100 μ M IPTG, were washed, suspended to the same OD₆₀₀ of 0.1, divided, and cultured in gonococcal base liquid (GCBL) medium in the presence or absence of IPTG. After 2 h under repressive conditions, Obg_{GC} was still detectable by immunoblotting (Fig. 2a) and the bacterial proliferation rate was indistinguishable from the permissive condition (Fig. 4b). Similarly, depletion of *C. crescentus* Obg using the P_{xyl} promoter and repressive growth conditions (glucose instead of xylose) resulted in much lower but detectable levels of Obg even 12 h after a carbon shift [17].

Prolonged culturing of GC is not feasible because the bacteria undergo autolysis shortly after reaching stationary phase [48–52]. Therefore to ensure significant reduction in the amount of Obg_{GC} , the conditional knockout strain FA1090 P_{lac} :: obg_{GC} was first treated as described above, cultures were collected 3 h after initial inoculation (indicated by an arrow in Fig. 4b), washed, and back diluted into fresh GCBL with or without the inducer. Culture density and bacterial viability, measured as Optical





Density at 600 nm (OD₆₀₀, Fig. 4b) and Colony-Forming Units (CFUs, Fig. 4c), respectively, were monitored every hour. Growth kinetics of FA1090 Plac::obg^{GC} cultured in the presence of IPTG (Fig. 4a) closely followed the pattern observed in parental wild type strain (Fig. 5a). In contrast, under non-permissive conditions, the culture density and bacterial viability were decreased significantly (Fig. 4b and c, respectively), concomitant with the reduction in Obg_{GC} level (Fig. 4d). At 6 h of the experiment, the Obg_{GC}-depleted culture contained on average 3.46×10^5 live bacterial cells, whereas 1.87×10^8 were present in permissive conditions (Fig. 4c). Further, FA1090 Plac::obg_{GC} grown in liquid media in the presence of IPTG was unable to survive upon plating on GCB lacking the inducer (Fig. 4e), confirming our prior observations (Fig. 4a).

Together, these studies demonstrate that Obg plays a pivotal function in GC physiology, as the depletion of Obg_{GC} caused loss of GC viability.

Expression of Obg_{GC}

The expression of Obg protein varies in different examined bacterial species. For instance, in *S. coelicolor* Obg is expressed in a growth-dependent manner with a sharp decline right after the beginning of aerial mycelium development and at the end of vegetative growth [28], whereas constant levels of Obg are maintained throughout the *C. crescentus* life cycle [30].

To examine the expression of Obg_{GC} , wild type FA1090 was maintained under routine aerobic cultivation in GCBL. Bacterial proliferation was monitored by measurements of cell density at OD_{600} within 6 h of the experiment (Fig. 5a). Every hour GC samples were collected and the whole cell lysates were probed with anti- Obg_{GC}

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antisera. The same samples were also examined using antibodies against an unrelated protein, Ng-MIP (Fig. 5b) and by SDS-PAGE coupled with colloidal coomassie staining (Additional file 2: Figure S2B) as loading controls. Immunoblotting and densitometry analyses (Fig. 5b and c, respectively) showed that Obg_{GC} reached maximum expression in the early logarithmic phase of GC growth at $OD_{600} \sim 0.5$ (2 h from the start of the experiment), and remained constant until stationary phase.

We also asked whether the conditions that more closely resemble clinical infection, such as anoxia, iron deprivation and, in the event of disseminated infection, exposure to human serum [2, 53], influence expression of Obg_{GC} . As expected, immunoblotting analysis showed increased levels of TbpB and AniA, which are well-recognized protein markers for iron-limited [54] and anaerobic [55] growth conditions, respectively (Fig. 5d). In contrast, the cellular concentrations of Obg_{GC} remained unaltered during growth of wild type FA1090 on GCB aerobically, in iron-limited conditions, in the presence of 7.5 % normal human sera, and anaerobically in the presence of nitrite as a terminal electron acceptor.

Subcellular localization of Obg_{GC}

The subcellular fractionation experiments showed that Obg was localized in the cytosol [35] and partially associated with the crude cell envelopes in *E. coli* [56], whereas in *S. coelicolor*, immunoelectron microscopy indicated that Obg was associated with the cytoplasmic membrane [28]. In addition, a growing number of reports show that different members of the Obg family cofractionate primarily with the 50S ribosomal subunit [17, 22, 32, 57]; however, in *Mycobacterium tuberculosis*, Obg is present in the 30S, 50S, and 70S ribosomal fractions [37].

To examine the cellular localization of Obg_{GC}, wild type FA1090 was cultured under standard laboratory conditions in GCBL and harvested at the midexponential phase of growth. The bacterial cells were lysed and the cell envelope proteins were separated from the cytosolic proteome by a sodium carbonate extraction procedure [58]. The crude cell envelopes were treated with sarkosyl to solubilize the inner membrane proteins, and the outer membrane was recovered by ultracentrifugation [59]. The same total amounts of purified subproteomes (15 µg) were separated by SDS-PAGE and probed with anti-Obg_{GC} antisera (Fig. 6a). The outer membrane protein, MtrE, which contains an extended periplasmic tunnel [60], was used as the cell envelope protein marker. As expected, MtrE was absent in the cytosol and enriched in the sarkosyl-insoluble fraction, whereas the vast majority of Obg_{GC} was present in the cytosolic protein fractions (Fig. 6a). A faint band of the same molecular weight was also detected in the cytoplasmic membrane. Many cytoplasmic proteins are repeatedly identified in different cell envelope proteomics studies and are often considered "contaminants" [61]. However, recent thorough sequential biochemical fractionations of E. coli, combined with mass spectrometry, demonstrated that many of these proteins, including Obg, form an actual peripheral inner membrane proteome linked via functional and/or structural oligomeric complexes [62].

Next, we addressed whether Obg_{GC} cofractionates with the ribosomes by ultracentrifugation of GC cell lysates through sucrose gradients and analysis of the polyribosomes profiles. Most of the cellular proteins accumulated at the top of the gradient followed by the peaks for small and large ribosomal subunits, the 70S monosomes, and the polyribosomes (Fig. 6b). Immunoblotting analysis with anti-Obg_{GC} antisera showed that under these conditions, the greatest amounts of Obg_{GC} were in 50S fractions and at the top of the gradient.

Based on these results, we conclude that Obg_{GC} is largely localized to the cytosol and primarily associates with the 50S ribosomal particle and not with the 70S monosomes or with translating ribosomes. Association of a part of the Obg_{GC} cellular pool with the cytoplasmic membrane may have functional implications, as Obg has been shown to be involved in key cellular processes such as ribosome maturation, DNA synthesis, cell division and morphology. Obg could be recruited to the membranebound complexes on demand, depending on the metabolic status of the bacterial cell.

$\operatorname{Obg}_{\operatorname{GC}}$ is expressed by contemporary clinical isolates of GC

Finally, the conservation of the predicted amino acid sequence of Obg_{GC} was assessed using the completed



genome sequences of strains FA1090 (Gen Bank accession number AE004969) and NCCP11945 (Gen Bank accession number CP001050), as well as the draft genome sequences of 14 different GC strains (downloaded from the Broad Institute website http://www.broadinstitute.org/annotation/genome/neisseria_gonorrhoeae/MultiHome.html). These analyses demonstrated that Obg_{GC} is 100 % identical among 14 strains and has a single amino acid change in the GC isolates designated as DGI2 and PID18 (Additional file 3: Figure S3).

Subsequently, to examine expression of Obg_{GC} , a diversified panel of GC isolates was utilized. This panel included common laboratory strains MS11, F62, and 1291, as well as 32 strains isolated from different gonorrhea patients from distinct geographical areas and at different time points. The anti- Obg_{GC} antibodies detected a band of the same size in all examined GC isolates, albeit the level of expression varied between some strains (Fig. 7).

Conclusions

Targeting essential proteins and critical cellular processes that are widely conserved remains an attractive avenue in antibacterial drug discovery programs. Compounds interfering with ribosome function and biogenesis and thus inhibiting different aspects of protein synthesis are among the most clinically useful antibiotics in spite of evolutionary conservation of bacterial and mitochondrial ribosomes [63].

Here, we show for the first time that Obg_{GC} is a GTPase essential for GC viability, mainly associated with the 50S large ribosomal subunit, abundant during different growth phases as well as under environmental conditions relevant to infection, and conserved in GC isolates. Together, these findings underscore the potential of Obg_{GC} as a target for the development of therapeutics against gonorrhea.

Methods

Bacterial strains, plasmids, and growth conditions

Strain of GC FA1090 [64] was primarily used in this study. Additionally, we employed: MS11 [65], 1291 [66], F62 [67], FA19 [68], isolates LGB1, LG14, LG20, and LG26 collected from two public health clinics in Baltimore between the years 1991–1994 [58], 13 strains derived from different patients seen at the Public Health–Seattle & King County STD clinic in 2011–2013 (Wierzbicki, et al., manuscript in preparation), as well as 14 WHO reference strains [69, 70]. Clinical isolates were kindly provided by Olusegun O. Soge, and King K. Holmes (Departments of GC was maintained on GCB Medium Base agar plates with Kellogg's supplements in 5 % CO₂ atm at 37 °C or in GCBL Medium Base Broth containing Kellogg's supplements and sodium bicarbonate (at a final concentration of 0.042 %) at 37 °C [48, 71, 72]. To achieve iron limited conditions, GCB without ferric nitrate in Kellogg's Supplements and with deferoxamine mesylate salt (Desferal, Sigma) at 5 μ M final concentration was utilized [73]. In addition, when stated in the text, GCB supplemented with 7.5 % normal human serum [74] was used to grow GC.

The *E. coli* strain NEB5 α was used for genetic manipulations. Bacteria were streaked from -80 °C on Luria-Bertani (LB) agar supplemented with kanamycin (50 µg/mL) when needed. *E. coli* strains were cultured in LB medium at 37 ° C. All media utilized in this study were purchased from Difco.

Construction of recombinant wild type and mutated versions of Obg_GC

All oligonucleotide primers designed and used in this study were synthesized by IDT DNA Technologies. Relevant restriction sites within primers are underlined. Recombinant N-His-Obg_{GC} and C-His-Obg_{GC} were generated by amplifying the obg gene from genomic DNA using two pairs of primers, respectively: obg-f 5'GAATTC CATATGAAATTCATCGACGAAGCAAAA3' and obg-r 5'GAATTCAAGCTTTTACTCCGGCTTAAACACG3'; as well as obg-c-f 5'GACTCCATGGAATTCATCGACGAA GCAAAAATCG3' and obg-c-r 5'GACTAAGCTTCTCC GGCTTAAACACGCC3'. The corresponding PCR products were digested with NdeI-HindIII or NcoI-HindIII (respectively) and cloned into similarly digested pET28a(+) to create plasmids pET-N-His-Obg_{GC} and pET-C-His-Obg_{GC}. Mutagenesis of T192 and T193 residues into alanine were accomplished using the QuickChange II



Protein purification

The E. coli BL21(DE3) was used as a host for expression of all Obg_{GC} variants. Bacteria harboring N-His-Obg_{GC}, C-His-Obg_{GC}, or T192AT193A were cultured in LB at 37 °C until the cultures reached OD_{600} of ~ 0.5, and the expression of individual Obg_{GC} variants was induced by the addition of IPTG to a final concentration of 1 mM. Cells were pelleted 3 h after induction and resuspended in lysis buffer (20 mM Tris-HCl pH 8.0, 10 mM imidazole, 450 mM NaCl). The cell lysis was carried out by passing the suspension five times through a French pressure cell press at 12,000 psi. Bacterial debris was removed by centrifugation and the clarified crude cell extracts were loaded onto HisPur Cobalt Resin (Thermo) equilibrated with lysis buffer. Columns were subsequently washed with solution containing 20 mM Tris-HCl pH 8.0, 20 mM imidazole, and 450 mM NaCl. Proteins were eluted with the same buffer containing 250 mM imidazole. The eluate was dialyzed two times for an hour and overnight against 20 mM Tris-HCl pH 8.0. The purified proteins were concentrated using Microsep Advance Centrifugal Devices with 10 K molecular cutoff (PALL). Protein concentrations were determined using the Bradford method with a Protein Assay Kit (BioRad). Glycerol was added to purified proteins to a final concentration of 10 % and proteins were stored at -80 ° C until further use.

Polyclonal rabbit anti-Obg_{GC} antisera

The polyclonal anti-Obg_{GC} antibodies were prepared by Pacific Immunology (Ramona, CA, USA) using the purified N-His-Obg_{GC}, two New Zealand White rabbits, and a 13-week antibody production protocol approved by IACUC Animal Protocol #1, in a certified animal facility (USDA 93-R-283) and the NIH Animal Welfare Assurance Program (#A4182-01).

Biochemical assays

The binding and hydrolysis of guanidine nucleotides were executed as described [26]. All fluorescence measurements were performed at 37 °C using a Synergy HT plate reader (BioTek). To determine the concentration of Mg²⁺ required for binding of mant-GTP or mant-GDP (Life Technologies), N-His-Obg_{GC} (2 μ M) was incubated with either mant nucleotide (0.3 μ M) and increasing concentrations of Mg²⁺ (from 0 to 10 mM) in binding buffer B containing

50 mM Tris–HCl pH 8.0, 50 mM KCl, 2 mM dithiothreitol, 10 μ M ATP, and 10 % (wt/vol) glycerol. The experiments were repeated on three separate occasions and the data is presented as the percentage of maximal measured relative fluorescence units.

The ability to form GTP-and GDP-Obg complexes was studied using 0.3 μ M mant-GTP or mant-GDP analogs in binding buffer B with or without 5 mM Mg²⁺, respectively, and 2 μ M purified Obg_{GC} protein (N-His-Obg_{GC}, T192AT193A, or C-His-Obg_{GC}). These studies were carried out with at least eight biological replicates, and means with corresponding standard error of the mean (SEM) are reported.

To assess the GTP hydrolysis rate of Obg_{GC} , N-His- Obg_{GC} or C-His- Obg_{GC} (16 μ M) was prebound to 0.3 μ M mant-GTP in binding buffer B supplemented with 5 mM MgCl₂. The decrease in fluorescence associated with conversion of mant-GTP-Obg complexes into mant-GDP-Obg was recorded in 1 min intervals for 3 h. Data were fitted to a single exponential decay equation using Graph-Pad Prism 6.0f (Graph Pad Software). The single turnover rate constant and the half-life of hydrolysis were obtained from at least four independent experiments.

Construction of conditional obg_{GC} mutant strain

The conditional obg_{GC} mutant strain, FA1090 P_{lac}:: obg_{GC} , was constructed using a similar approach as we previously described [58]. Briefly, a gene encoding the Obg homolog in GC, Obg_{GC} (NGO1990), including an upstream region containing its indigenous ribosome-binding site (16 bp) was amplified with primers 5'CAAACAAGAGCATTT AATG3' and 5'AAGTTGGGCCGGCCTTACTCCGGCT TAAAC3'. To place the obg_{GC} under the control of the Plac promoter, the resulting 1189 bp PCR product was digested with FseI and sub-cloned into ScaI-FseI digested pGGC4. This vector contains an IPTG-inducible promoter, which enables the controlled expression of a cloned gene [75]. The upstream region of obg_{GC} was amplified with primers 5'ACTAGTGAATTCGCCTTGCTGTCGC TTTG3' and 5'ATCGATGGTACCTTGGTTTTAAATAG GGTTTCAGGC3'. The obtained 531 bp product was digested with EcoRI and KpnI, and cloned into pUC18K [76], yielding the pUC18K-obg_{GC}-up. Next, the DNA fragment containing lacl repressor gene, Plac promoter and obg_{GC} gene carried on the pGCC4 was amplified with primers 5'ACTCAATAGGATCCTCACTGCCCGCTTTC CAG3' and 5'CATAAGCAGTCGACTTCAGACGGCG-GAGACGGCGGTAATCAGG3'. The PCR product was purified, digested with BamHI-SalI and cloned into pUC18K [76], generating pUC18K-P_{lac}::obg_{GC}. This final construct encompassing nonpolar kanamycin resistance cassette apha-3 [76] flanked by homologous regions for recombination and allelic exchange was used to introduce the mutation onto the FA1090 chromosome. The plasmid was linearized by digestion with NdeI and piliated bacteria were transformed with 0.1 µg of plasmid DNA in liquid media as described [77] with the exception that the growth media were supplemented with 100 µM IPTG. The resulting GC transformants were selected on GCB agar containing 40 µg/mL kanamycin and 100 µM IPTG. The FA1090 P_{lac} :: obg_{GC} clones were verified by PCR with primers 5'GCCTTGCTGTCGCTTTG3' and 5' GGAGACGGCGG TAATCAGG3' and immunoblotting analysis with anti-Obg_{GC} antibodies.

Obg_{GC} depletion studies

To assess the consequence of Obg_{GC} depletion on GC viability, the GC FA1090 Plac::obgGC strain was plated from freezer stocks onto GCB supplemented with 100 µM IPTG. The following day, single nonpiliated colonies were passaged onto fresh GCB plates containing 100 µM IPTG and incubated approximately 18 h in 5 % CO₂ atm at 37 °C. The bacteria were swabbed from plates, suspended in GCBL to OD_{600} of 0.1, and washed two times with GCBL that was pre-warmed to 37 °C to ensure removal of IPTG. Equal amounts of bacterial suspensions were divided into two flasks and cultured aerobically with and without 100 µM IPTG for 3 h at 37 °C. Subsequently, the cultures were diluted to OD_{600} of 0.1 into fresh GCBL with and without 100 µM IPTG. For measurements of OD₆₀₀, bacterial viability (CFUs), and immunoblotting analysis, samples were withdrawn every hour and processed as described [58]. Experiments were performed in biological triplicates and mean values with corresponding SEM are presented.

Subcellular fractionation

Non-piliated and translucent GC colonies of wild type FA1090 were swabbed from solid media, suspended in 500 mL of GCBL to OD₆₀₀ of 0.1 and cultured at 37 °C with aeration (220 rpm) to OD_{600} of 0.6 - 0.8. Cells were harvested by centrifugation (20 min, $6000 \times g$) and the crude cell envelope fraction was separated from the cytosolic proteins using a sodium carbonate extraction procedure and subsequent ultracentrifugation steps [58]. The inner membrane proteins were solubilized using 2 % Sarkosyl in 20 mM Tris-HCl pH 7.5 according to the method described by Leuzzi et al. [59]. The Sarkosylinsoluble outer membrane fractions were recovered by ultracentrifugation for 1 h at $100,000 \times g$ and 4 °C. The pellet was suspended in PBS containing 1 % SDS. The total protein amount in each isolated subproteome fraction was assessed using a Protein Assay Kit (Bio Rad).

Preparation of GC cell lysates for ribosome profiles

Wild type FA1090 bacteria were incubated in GCBL medium until the mid logarithmic phase of growth $(OD_{600} \sim 0.5 \text{ to } 0.7)$. Chloramphenicol was added to a final

concentration of 100 µg/mL one minute before harvesting. Cells were harvested at 4,000 × g for 10 min at 4 °C and immediately frozen at -80 °C. After thawing on ice, bacteria were resuspended in 1 mL of lysis buffer comprised of 10 mM Tris–HCl pH 7.5, 10 mM MgCl₂, 30 mM NH₄Cl, 100 µg/mL chloramphenicol [78]. Subsequently, an equal volume of glass beads (100 µm; Electron Microscopy Sciences) was added to the solution and bacterial cells were lysed by vortexing every minute for 10 min with a 1 min cooling interval on ice. Lysates were clarified by centrifugation for 15 min at 21,000 × g and 4 °C.

Polyribosome fractionation

The isolated GC cell lysates in amounts corresponding to 15 OD_{260} units were overlaid on top of a 10 to 47 % step sucrose gradient as described previously [79]. Ribosomal subunits were separated by centrifugation in a Beckman SW41 rotor at 174,000 × g for 4 h and 4 ° C. Separated ribosomal subunits were fractionated using an Econo Pump and Econo UV Monitor (Bio-Rad). UV traces were recordered using a Model 1325 Econo Recorder (Bio-Rad) and 500 µL fractions were collected. Protein samples were precipitated with 15 % trichloroacetic acid. The resulting precipitates were solubilized in SDS loading buffer, and after separation in 10-20 % Criterion Tris-Tricine TGX (BioRad) acrylamide gel, all collected fractions were subjected to immunoblotting with anti-Obg_{GC} antisera as described below.

SDS-PAGE and immunoblotting

Whole cell lysates were obtained from GC grown in GCBL with aeration and on GCB plates maintained under growth conditions as stated in the text. When bacterial colonies reached approximately the same size, all strains were harvested, suspended in pre-warmed GCBL, and the cell density was examined by OD₆₀₀ measurement. Fractions containing either cytoplasmic, inner- or outer- membrane proteins (15 µg of proteins loaded per lane), ribosomal particles, or whole cell lysates matched by equivalent OD₆₀₀ units, were prepared in SDS sample buffer in the presence of 50 mM dithiotreitol and separated in either 10-20 % Criterion Tris-Tricine TGX (BioRad) or 4-20 % Mini-PROTEAN TGX precast gels (Bio-Rad). The proteins were transferred onto 0.2 µm nitrocellulose membrane (Bio-Rad) using a Trans-blot Turbo (Bio-Rad). A solution of 5 % milk in phosphate buffered saline pH 7.0 (PBS, Li-Core) supplemented with 0.1 % Tween 20 (PBST) was used for blocking. Following 1 h of incubation, polyclonal rabbit antisera against Obg_{GC} (1:5,000), polyclonal anti-AniA antibodies (1:10,000; [58]), monoclonal mouse anti-MtrE antisera (1:10,000; a gift of Ann Jerse, Uniformed Services University, Bethesda), monoclonal mouse anti-Ng-MIP antibodies (1:10,000; a gift of Mariagrazia Pizza, Novartis Vaccines, Italy), or polyclonal rabbit antiTbpB antisera (1:1,000; a gift of Cynthia Cornelissen, Virginia Commonwealth University, Richmond) diluted in PBST as indicated in parenthesis were added to the membranes. The horseradish peroxidase conjugate of goat anti-rabbit IgG antisera (BioRad) or goat antimouse IgG antibody (ThermoFisher Scientific), correspondingly, were utilized as secondary antibodies at 1:10,000 dilution. The reactions were developed using Clarity Western ECL-Substrate (BioRad) and a Chemi-DocTM MP System (BioRad) was used for western blot imaging.

Densitometry

The immunoblot probed with anti-Obg_{GC} antisera was scanned using the ChemiDocTM system (BioRad) and subjected to densitometric analysis using Image LabTM 5.0 software (BioRad). To quantify the intensity of the Obg_{GC} protein bands, the volume tool (rectangle), local background subtraction, and linear regression were used.

Statistical analyses

Statistical analyses were conducted using GraphPad Prism 6.0f (Graph Pad Software) and an unpaired Student's *t*-test was used to analyze the data.

Availability of supporting data

The data supporting the results of this article are included within its Additional files 1, 2 and 3: Figures S1-S3.

Additional files

Additional file 1: Figure S1. Sequence alignment of Obg from *N. gonorrhoeae* and *N. meningitidis* with characterized members of Obg subfamily from various bacterial species. Identical residues are indicated by a star (*); G motifs (from G1 to G5) are underlined. Accession numbers of compared Obg homologs are as follows: *M. tuberculosis* (Mt) (CCE37910), *S. coelicolor* (Sc) (BAA13498), *D. radiodurans* (Dr) (NP_293810), *T. thermophilus* (Tt) (YP_145047), *B. subtilis* (Bs) (AAA22505), *C. crescentus* (Cc) (NP_419134), *N. gonorrhoeae* (Ng) (YP_209010), *N. meningitidis* (Nm) (NP_275074), *E. coli* (Ec) (NP_417650), *V. cholerae* (Vc) (NP_230091).

Additional file 2: Figure S2. Loading controls for immunoblotting experiments. Samples of whole-cell lysates were prepared for SDS-PAGE as described in the text, separated in precast gradient gels and the protein profiles were visualized using colloidal coomassie. Loaded OD₆₀₀ units in individual experiments matched the corresponding samples used in immunoblotting analyses and are indicated below each gel. (A) Loading controls for immunoblotting experiment presented in Figure 2 A. (B) Loading controls for immunoblotting analysis shown in Figure 2 B. (C) Loading controls for experiment shown in Figure 5 B. (D) Loading controls for experiment in Figure 5 D for whole-cell lysates probed with anti-Obg and anti-TbpB antisera. (D) Loading controls for immunoblotting experiment with anti-AniA antisera presented in Figure 5 D.

Additional file 3: Figure S3. Comparison of predicted amino acid sequences of Obg_{GC} between different GC isolates.

Abbreviations

A: Absorbance; GC: *Neisseria gonorrhoeae*; GCB: Gonococcal base agar; GCBL: Gonococcal base liquid medium; OD: Optical density; IPTG: Isopropyl thio-β-D-galactopyranoside; PBS: Phosphate buffered saline; PBST: Phosphate buffered saline supplemented with 0.1 % Tween; SDS-PAGE: Sodium dodecyl sulfate polyacrylamide gel electrophoresis; SEM: Standard error of the mean.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

AES designed the studies. RAZ engineered the recombinant wild type- and mutated- versions of Obg_{GC} performed bioinformatics analyses, polysome profiles, immunoblotting, and densitometry. IHW constructed the conditional FA1090 P_{Iac} : obg_{GC} knockout strain and conducted Obg_{GC} depletion studies, as well as subcellular fractionations. RAZ and BIB purified recombinant proteins. BIB and IHW performed SDS-PAGE coupled with colloidal coomassie staining and immunoblottings. AES, RAZ, and BIB conducted the biochemical analyses of recombinant versions of Obg_{GC} . All authors interpreted the data. AES wrote the manuscript. All authors read and agreed to the content of the manuscript.

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