Cloning and Sequencing of G Protein encoding genes of *Balanus improvisus*

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Examinar: Prof. Anders blomberg

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Cover: Picture shows the adult form of the Balanus improvisus (Photo by Sergej Olenin)
I dedicate this little effort of mine to my parents for their never ending moral support, encouragement and prayers. Specially dedicated to my husband for his cooperation, financial support and love that enabled me to achieve this goal.
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ABSTRACT

The barnacle *Balanus improvisus* is a major fouling organism in Swedish waters. Settling of barnacles onto ship hulls leads to biofouling. Antifouling paints are used to prevent the settlement of the barnacles on ship hulls, but some of these have severe impact on marine ecosystems. G-protein-coupled octopamine receptors of *B. improvisus* have already been cloned and sequenced. It was discovered that these octopamine receptors are activated by the novel antifouling agent medetomidine, leading to inhibition of the settling process of *B. improvisus*. Research on *B. improvisus* has been conducted to provide alternatives to toxic substances such as tributyl tin (TBT), copper metal oxides or organic biocides in marine paints.

The knowledge about G proteins of barnacles is very limited. We have cloned and sequenced two G protein subunits; Ga<sub>q</sub> and Ga<sub>s</sub>. Cloning of Ga<sub>q</sub> and Ga<sub>s</sub> was performed by PCR amplification using sequence information based on ESTs from a *B. improvisus* cDNA library. cDNA was used as template. The obtained sequences were aligned with lobster *Homarus americanus* sequence. In Ga<sub>q</sub> different variants of 5′ ends were found whereas in Ga<sub>s</sub> different variants at the 3′ end are present. Interestingly Ga<sub>s</sub> in *B. improvisus* has 165 nucleotides longer open reading frame as compared to the one in *H. americanus*.

This is a first step in the cloning of G-protein subunits of *B. improvisus*. Functional studies of G-protein with GPCRs will be performed in the future. The enhanced knowledge of G-protein and GPCRs in barnacles will contribute to the development of more specific antifouling paints.
LIST OF ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPCRs</td>
<td>G-protein Coupled Receptors.</td>
</tr>
<tr>
<td>GDP</td>
<td>Guanosine Diphosphate</td>
</tr>
<tr>
<td>GTP</td>
<td>Guanosine Triphosphate</td>
</tr>
<tr>
<td>RT-PCR</td>
<td>Reverse Transcription Polymerase Chain Reaction</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl Sulfoxide</td>
</tr>
<tr>
<td>cPCR</td>
<td>Colony-Polymerase Chain Reaction</td>
</tr>
<tr>
<td>RLM-RACE</td>
<td>RNA Ligase Mediated Rapid Amplification of cDNA Ends</td>
</tr>
<tr>
<td>EST</td>
<td>Expressed Sequence Tags</td>
</tr>
<tr>
<td>GSP</td>
<td>Gene Specific Primer</td>
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</table>
AIM

The aim of this project is to clone and sequence different subunits of Gα proteins from the barnacle, *B. improvisus*.

This report focuses on:

Cloning and sequencing of the two G protein subunits Gαs and Gαq.

INTRODUCTION

Barnacles and biofouling

Biofouling is a worldwide problem in marine systems and barnacles are the most problematic biofouling marine organisms. Biofouling results in an increase in drag, which in turn leads to increased fuel consumption, hull cleaning and removal of paint (Callow and Callow, 2002). *B. improvisus* is the barnacle species causing the most severe fouling problems in the Baltic sea (Lind et al., 2010). The life cycle of *B. improvisus* has seven larval stages, and between the sixth and seventh transitions, nauplii larva transforms into cyprid larva. In order to complete the transition to adult life, cyprid larvae must attach to a hard substrate (Fig.1).

![Figure 1: The Life Cycle of B. improvisus](drawing by Susanna Falkbring)
Adhesion of barnacles
After identifying a suitable surface to adhere cyprid larva secretes adhesive proteins by a pair of cement glands. The cement glands are connected by cement ducts, which widens into a muscular sac. The muscular sac is connected to an antenna by another cement duct and pours proteinaceous cement into antenna. Cyprid cement flows around and embeds the attachment organs and the cyprid larva is able to attach itself to the surface and metamorphoses into a calcified adult barnacle (Odling et al., 2006).

Medetomidine and barnacles
Traditional antifouling substances like tin and copper are highly toxic to non-target organisms in the marine environment. Organotin compounds such as TBT are now banned worldwide. Medetomidine is now being used as effective alternative in marine paints (Krang and Dahlstrom, 2006). Since the 1980’s, medetomidine has been used as a sedative agent in veterinary medicine. Medetomidine functions as α2–adrenoceptor agonist and binds to G protein coupled octopamine receptors in invertebrates and α2–adrenoceptor in vertebrates. Medetomidine strongly inhibits the settling process of the barnacles *B. improvisus* at very low (nanomolar) concentrations (Lennquist et al., 2010).

G protein coupled receptors (GPCRs)
G protein coupled receptors represent one of the largest and most diverse groups of proteins present in most eukaryotes. All GPCRs have seven transmembrane α helical regions, an extracellular N-terminus and an intracellular C terminus. GPCRs bind with a wide range of ligands that modulate vital physiological events like neurotransmission, sensory perception, chemotaxis and cell communication (Tuteja, 2009). The ligands of GPCRs can be various types of substrates like amino acids, proteins, biogenic amines, nucleotides, lipids and Ca^{+2} ions. When activated by ligands, G protein coupled receptors undergo a conformational change that leads to G protein activation by promoting the exchange of GDP/GTP associated with the Ga subunit (Oldham and Hamm, 2008).

Heterotrimeric G protein
Heterotrimeric G proteins are molecular switches that turn on signaling processes for hormones, neurotransmitters, and photons. The G proteins are composed of three subunits, α, β and γ. G proteins exist as trimers with GDP bound to the α subunit in its inactive form. Receptor activation causes G proteins to dissociate into Ga and Gβγ because GDP is replaced by GTP (Xu et al., 1998). The switching function of the G protein depends upon the identity of the α subunit; four Ga subunit classes (Gaαs, Gaαi, Gaαq, and Gaα12) have been proposed based on amino acid sequence analysis, each performing specific functions (Han and Watson, 2005). The Gaαs activates an enzyme called adenylyl cyclase which in turn catalyses the synthesis of the second messenger cAMP. In contrast, Gaαi inhibits the activity of adenylyl cyclase. The Gaαq activates phospholipase C that catalyses the synthesis of two second messengers (DAG and IP3). The Gaα12 proteins regulate cytoskeletal assembly (Lodish, 2008). Within the adrenergic receptor family for
example, the receptors couple to different G-proteins. β-adrenergic receptors couples to GaS, the α1-adrenergic receptor couples to Gaq and the α2-adrenergic receptor is coupled to Gain.

Figure 2: Interaction of GPCR and G protein
G protein and barnacles

Five different G protein coupled octopamine receptors, one α and four β-like receptors, have been recently cloned and characterized by Lind and Blomberg at the University of Gothenburg (Lind et al., 2010). Before this only two GPCRs of the barnacle had been cloned and sequenced (Isoai et al., 1996) (Kawahara et al., 1997). The interaction of the novel antifouling agent medetomidine with the cloned octopamine receptors was studied. All five receptors increase cAMP signaling in the cell, however the main signaling pathway for the α like receptor was via increasing Ca^{2+} levels in the cell. In order to produce more specific antifouling paints it is important to know more about the GPCRs and G-proteins of the barnacles. More knowledge about G-proteins of barnacles and their interactions with GPCRs will help to design non toxic substances. This work focuses on the cloning and sequencing of G-protein encoding genes of B. improvisus. Interaction between G-proteins and GPCRs of the barnacles will be studied in the future.
MATERIAL AND METHODS

cDNA Synthesis

cDNA was synthesized from 1 µg of total RNA from cyprid by following the protocol of Superscripts™ III First-strand synthesis system for RT-PCR (Invitrogen life technologies). RNA together with 1 µl of oligo dT and 1 µl of 10 mM dNTP was heated to 65 °C and then incubated on ice for 1 min. Then cDNA mix was added and incubated in 50 °C for 50 min.

cDNA synthesis mix

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>10XRT Buffer</td>
<td>2 µl</td>
</tr>
<tr>
<td>25 mM MgCl₂</td>
<td>4 µl</td>
</tr>
<tr>
<td>0.1 M DDT</td>
<td>2 µl</td>
</tr>
<tr>
<td>RNase OUT™</td>
<td>1 µl</td>
</tr>
<tr>
<td>Super Script™ III RT</td>
<td>1 µl</td>
</tr>
</tbody>
</table>

The cDNA synthesis was terminated by incubating at 85°C for 5 min. The cDNA was then treated with 1 µl RNase H and incubated at 37°C for the removal of RNA.

RLM- Race

RNA ligase mediated rapid amplification of 5´ and 3´ cDNA ends was done by following Gene Racer™ kit (invitrogen life technologies). In the first step total RNA was treated with calf intestinal phosphatase (CIP) to dephosphorylate non-mRNA or truncated mRNA. Dephosphorylated mRNA was then precipitated and the mRNA cap structure was removed. An RNA Oligo was ligated to decapped mRNA following the kit protocol. mRNA was reverse transcribed into cDNA using Gene Racer Oligo dT primers. For the amplification of the 5´ end of cDNA Gene Racer 5´ primer and Nested 5´ primer together with gene specific primers were used. The amplification of 3´ end was done by using Gene Racer 3´ primer and Nested 3´ primer with gene specific primers.

PCR amplification

For the cloning of Ga₆ and Ga₉ genes in B. improvisus, cDNA was used as template in PCR amplification by using different primer combinations based on cDNA library of B. improvisus made by M. Alm Rosenblad and A. Blomberg.

The PCR reactions were performed by preparing 50 µl of Pfu Ultra enzyme reaction mix in the following concentrations; 1 µl of cDNA was added to 49 µl of reaction mix containing 1x Buffer, dNTP (0.2 mM), DMSO (5%) , primers (fw and rev) 0.5 µM, and Pfu Ultra enzyme 1 µl.
Master mix for cPCR

For colony PCR 19 µl of master mix was prepared for each reaction containing the following. M13 Fw, and M13 Rev (0.5 µM each), 1x Buffer true start, dNTP (0.2 mM), DMSO 5%, MgCl2 (1.5 mM), and 0.20 µl of True start Taq Polymerase. 1 µl cell suspension was then added to obtain final volume of 20 µl for each reaction.

Gel Electrophoresis of DNA

Agarose gel electrophoresis was used to analyze the sizes of DNA fragments. In order to prepare 1% agarose gel to separate bigger fragments of more than 1000 bp, 1g of gel was weighed and melted in 100 ml 1X TAE buffer. For smaller fragments 1.5% of gel was formed by adding 1.5g of gel into 100 ml of buffer. Ethidium bromide was added to the gel to enable the visualization of DNA using UV light. DNA samples were prepared by mixing DNA solution with gel loading dye (Fermentas). Gel electrophoresis of DNA was performed with the constant voltage of 100 volts.

Purification and digestion of PCR product

After separating PCR products on gel, they were purified with a gel extraction kit (Qiagen) following the manufacturer’s instructions. Shortly, the gel was solubilized by adding 3 volumes of buffer QG and incubated at 50°C for 50 min. The sample was added to the QIA quick column and the DNA was washed by buffer PE and finally eluted in 50 µl elution buffer. In some cases purified PCR product was cleaved with the restriction enzyme Eco RI. 2 µl of 10x buffer was mixed with 1µl of DNA then 16 µl of H2O and 1µl of Eco RI was added and incubated at 37°C for 1 hour.

TOPO Cloning of Gaq and Gαs PCR products

The PCR products were cloned into plasmids using the Zero Blunt® TOPO® PCR Cloning Kit for Sequencing (Invitrogen). 4 µl of PCR product and 1µl of salt solution were added to 1ul TOPO vector, mixed and incubated for 30 minutes. Plasmids containing the ligated PCR product were transformed to E. coli by following the One Shot Chemical Transformation Protocol (Invitrogen life technologies). Shortly, cells were heat shocked for 30 s at 42°C, incubated at 37°C in SOC medium and then spread on ampicillin containing plates and incubated at 37°C overnight. Colonies were picked to perform colony PCR (cPCR). Positives colonies from the cPCR were cultured in 3 ml LB overnight and the plasmid was purified by following the QIA miniprep Kit Protocol (Qiagen). Plasmid concentration was determined using a Nano drop method, and purified DNA was sent to MWG biotech Germany for Sanger sequencing.

Expressed sequence tags (ESTs) Library

RNA was prepared from approximately 1000 cyprids. cDNA was prepared from RNA and then sequenced.

Designing Gene Specific Primers (GSPs)

Several GSPs were designed based on the B. improvisus ESTs library. Primer optimization was done by using different parameters of primer design like length, Tm and GC content (%).
<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence</th>
<th>Number of Base Pairs</th>
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</tr>
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<td>60</td>
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<tr>
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<td>Gaq1_rev_fl</td>
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<tr>
<td>Gaq_fl_2fw</td>
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<tr>
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<td>Gas_fw2</td>
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<tr>
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<td>74</td>
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</tr>
<tr>
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<td>AAAAAACGAGCACCATTCCGACAGGA</td>
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<td>74</td>
</tr>
<tr>
<td>race_3´_Gas_2</td>
<td>CTAATCCACCACAGAAACAGATATTC</td>
<td>26</td>
<td>76</td>
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<tr>
<td>race_3´_Gas_3</td>
<td>AGTCCCTACGTCGCGCATATCTCA</td>
<td>24</td>
<td>74</td>
</tr>
</tbody>
</table>
RESULTS

To clone and sequence G protein encoding genes of the *B. improvisus* cDNA was used as template. PCR amplification was done by using different primer combinations based on ESTs (expressed sequence tags) from a *B. improvisus* cDNA library constructed by M. Alm Rosenblad and A. Blomberg (unpublished data).

There were two ESTs for Gαq in the EST cDNA library with a gap in between. To cover this gap, two primers were designed on either side of the gap, called gap fw and gap rev. PCR was performed and a band of expected size, about 300 bp was obtained (Fig. 3).

![Fig. 3. Gαq fragment obtained from PCR. Sample (1) 100bp DNA ladder. An approximately 300bp band was obtained (well 3) covering a sequence gap of aligned ESTs. The following PCR program was used: 3 min denaturing step at 98˚C followed by 35 cycles with a 98˚C denaturing step for 30 s, 52˚C annealing for 30 s, 72˚C extension for 2 min and final extension at 72˚C for 10 min.](image)

After performing gel extraction of the obtained DNA band it was ligated into the plasmid pCR® 4Blunt TOPO. The ligated fragment was transformed into *E. coli* and DNA was purified and sent for sequencing. In an attempt to amplify the Gαq full length sequence, different primers were designed located before the start ATG and after the stop codon (Fig. 4A).
PCR was performed by using different full length primers for Gαq. A. Location of the primers. B. Bands were obtained only with gap fw, rev 1 (well 9) and with gap fw, gap rev (well 10). The following PCR program was used: 2 min denaturing step at 98˚C followed by 35 cycles with a 98˚C denaturing step for 30 s, 49˚C annealing for 30 s, 72˚C extension for 2 min and final extension at 72˚C for 10 min.

The full length sequence of Gαq was not amplified. Only a smaller fragment of Gαq from the gap to the 3’ end was amplified which was approximately 1000 bp in length (Fig. 4B).

In a second attempt to amplify the full length sequence of Gαq new primers were designed with a higher annealing temperature (Fig. 5A). Eight different primer combinations covering different parts of the coding region were used, but only four bands were obtained. The full length sequence of Gαq was not amplified, but all the Gαq coding region from gap to the 3’ end was amplified (Fig. 5B).
5′ Race for Gαq

Our primers are based on ESTs from a cDNA library. In order to confirm that these EST sequences are correct, Race 5′ Gαq primers were designed to amplify the complete 5′ end of Gαq down to just below the start ATG. Touchdown and nested PCR were performed by using these primers together with Gene race primer (RLM-RACE kit, Invitrogen). RLM-RACE cDNA was used as template instead of ordinary cDNA. Touchdown PCR is a variant of PCR that aims to reduce non specific primer annealing by gradually lowering the annealing temperature. Nested PCR increases the specificity of PCR reaction. Two sets of primers are used in two successive PCR reactions.

The results of the touchdown PCR showed several faint bands, but after nested PCR a major band product of about 500 bp was obtained (Fig. 6). This PCR product was purified, cloned and sent for sequencing.

An EST from the cDNA library was aligned with the 5′ Race sequences, both were similar. Thus sequence errors in ESTs could not explain the failure to obtain the full length Gαq, when using 5′ primers based on the ESTs.

3′ Race for Gαq

In order to investigate if it was possible to amplify the sequence from just below the ATG to 3′ end of Gαq, 3′ Race was performed. Fw4b and the 3′ Gene racer primer was used. Touchdown PCR and a subsequent nested PCR was performed with fw 6 or fw 5 combined with the nested primer from the RLM-RACE kit. Two fragments were obtained from 3′ race which covers almost all the coding region and un-translated 3′ end of the Gαq (Fig. 7).
Fig. 7. Touchdown and nested PCR to amplify the 3´ end of Gα₉. Two bands were obtained from nested PCR by primer combination fw 6 with 3´ nested Gene racer primer (sample no.4) and fw 5 with 3´ Gene racer primer from kit (sample no.5). PCR was run according to the following: 2 min denaturing step at 98° C followed by 35 cycles with a 98° C denaturing step for 30 s, 58° C annealing for 30 s, 72° C extension for 2 min and final extension at 72° C for 10 min).

Amplification of Gα₉ full length sequence

To amplify the complete sequence of Gα₉ 5´, Gene racer primer and nested 5´ primer were used with gene specific primers (GSPs) rev 1b and rev 3b, respectively. A smear of unspecific bands was obtained, but a band with expected size of about 1400 to 1500 bp in length was also obtained (Fig. 8). This band covers the complete 5´ end sequence and down to below the stop codon. The full length sequence of Gα₉ was thus amplified. By comparing the extreme 5´ end with the previous 5´ race short fragment and the ESTs sequences, it was found that they were not similar (Fig. 9).

Fig. 8. Gα₉ 5´ RACE PCR product. Running a 5´ race with primers at the extreme 5´ end and just below the stop codon, a band obtained which is about 1400 to 1500 bp (arrow indicates right band). The following PCR program was used: 2 min denaturing step at 98° C followed by 35 cycles with a 98° C denaturing step for 30 s, 58° C annealing for 30 s, 72° C extension for 2 min and final extension at 72° C for 10 min.
Sequence alignment of *B. improvisus* Ga\(_q\) 5′ race short fragment with Ga\(_q\) full length Sequence

![Sequence alignment diagram]

**Fig. 9.** Comparison of the 5′ ends of the Ga\(_q\) 5′ RACE short fragment with the Ga\(_q\) full length sequence obtained from 5′ RACE. Nucleotide sequence alignment of *B. improvisus* Ga\(_q\) was performed by using ClustalW program. It was found that the 5′ end of the Ga\(_q\) short fragment is different from the Ga\(_q\) full length 5′ end. Grey color indicates similar sequences while white indicates sequences are not similar.

**Sequence alignment of *B. improvisus* Ga\(_q\) and Homarus Ga\(_q\)**

To compare our Ga\(_q\) obtained sequences with other crustaceans, the sequence alignment of Ga\(_q\) full length with *H. americanus* was done. The nucleotide sequence of the *B. improvisus* Ga\(_q\) 5′ end was aligned with *H. americanus*, the sequences were relatively similar (Fig. 10 a). The nucleotide sequences of Ga\(_q\) full length were translated and were aligned with amino acids sequences of *H. americanus*. Sequence alignment of Ga\(_q\) full length and *H. americanus* shows very high sequence similarity (Fig. 10 b).
Fig. 10. Comparison between *B. improvisus* Gαq and *H. americanus* Gαq sequences. A, Comparison between 5´nucleotide sequences of Gαq is shown. B, comparison between Gαq amino acid sequences.

**Amplification of Gαs**

To amplify the Gαs sequence, primers were designed based on ESTs from the *B. improvisus* cDNA library. In the Gαs EST sequences, parts of the 3` coding region is missing. The sequence
from the cDNA library has a longer open reading frame, than that in *H. americanus* Gαs with one ATG before the start ATG of *H. americanus* and the second ATG are positioned similar as in the *H. americanus* Gαs. Two primers were designed, fw 1 before the first ATG and fw 2 before the second ATG. One rev primer located in the middle of the coding region, was designed and used in combination with two forward primers (Fig. 11).

**Fig. 11.** PCR fragments of Gαs. PCR was done by using two primer combinations fw 1, fw2 with rev primer. Two nice bands were obtained. PCR was run in the following way: 2 min denaturing step at 98°C followed by 35 cycles with a 98°C denaturing step for 30 s, 62°C annealing for 30 s, 72°C extension for 1 min and final extension at 72°C for 10 min.

To amplify the 3′ end of Gαs, Gene racer 3′ primer with nested 3′ primer and GSPs (gene specific primers) race3′ Gas1 and race3′ Gas2 were used. After nested PCR three bands of about 1000 to 1500 bp were obtained (Fig. 12). The obtained products were purified from the gel, cloned and sent for sequencing. The 3′ end of Gαs was amplified. Based on the sequence obtained from the middle band, reverse full length primer was designed to amplify the complete sequence of Gαs. PCR was performed by using primers fw 1 and fw 2 with the rev full length primer, and the complete sequence of Gαs was amplified (Fig. 13). Three different variants of gas were found differing only in the 3′ end of the coding region (Fig.14).

**Fig. 12.** 3′ race for Gαs. Three bands were obtained between 1000 to 1500 bp. The following PCR program was used: 2 min denaturing step at 98°C followed by 35 cycles with a 98°C denaturing step for 30 s, 63°C annealing for 30 s, 72°C extension for 2 min and final extension at 72°C for 10 min.
**Fig. 13.** $\Gamma_{\alpha s}$ PCR product obtained by using full length primers. Band was obtained between 1200 to 1500 bp. PCR program used was: 2 min denaturing step at 98°C followed by 35 cycles with a 98°C denaturing step for 30 s, 58°C annealing for 30 s, 72°C extension for 2 min and final extension at 72°C for 10 min.

**Fig. 14.** An amino acid sequence analysis of 3 different clones of $\Gamma_{\alpha s}$ full length. The nucleotide sequences of three different clones of $\Gamma_{\alpha s}$ were translated and aligned together. After sequence alignment of $\Gamma_{\alpha s}$ different variants of $\Gamma_{\alpha s}$ were found. All of the three clones of $\Gamma_{\alpha s}$ show very high sequence similarity in the 5′ end but in the 3′ end upto 40 amino acid, they are different.
**Sequence alignment of Gα<sub>s</sub> full length and *H. americanus***

Sequence alignment of Gα<sub>s</sub> full length and Gα<sub>s</sub> from *H. americanus* was done. It is found that both are relatively similar (Fig. 15).

![Sequence alignment of Gα<sub>s</sub> full length and *H. americanus*](image)

**Fig. 15.** Comparison between Gα<sub>s</sub> full length and *H. americanus* sequences.
Sequence identity of *B. improvisus* $\text{Ga}_q$ and $\text{Ga}_s$ with G-proteins of related organisms

The amino acid sequence of $\text{Ga}_q$ of *B. improvisus* shows high sequence similarity (83%) with the amino acid sequence of $\text{Ga}_q$ from *H. americanus*. The amino acid sequence of different variants of $\text{Ga}_q$ in Drosophila shows 65 to 83 % sequence similarity with $\text{Ga}_q$ of *B. improvisus*. The amino acid sequence of *B. improvisus* $\text{Ga}_q$ shows 68% sequence similarity with the amino acid sequence of Daphnia. The nucleotide sequence of one of the variants of $\text{Ga}_s$ ($\text{Ga}s_2(6)$) shows high sequence identity 80 to 86% with lobster, Drosophila and Daphnia. The other two variants of $\text{Ga}_s$ in *B. improvisus* show somewhat less sequence identity (Table 2).

**Table 2. Identity table of $\text{Ga}_q$ and $\text{Ga}_s$**

<table>
<thead>
<tr>
<th></th>
<th>$\text{Ga}s_2(6)_\text{Bimp}$</th>
<th>$\text{Ga}s_3(2)_\text{Bimp}$</th>
<th>$\text{Ga}s_3(4)_\text{Bimp}$</th>
<th>$\text{Ga}Q_{\text{Bimp}}$</th>
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<tr>
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<td>98</td>
<td>100</td>
<td>73</td>
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<tr>
<td>$\text{Ga}Q_{\text{Bimp}}$</td>
<td>43</td>
<td>42</td>
<td>39</td>
<td>68</td>
</tr>
</tbody>
</table>

Table 2. Amino acid identity table for $\text{Ga}_q$ and $\text{Ga}_s$ of *B. improvisus*, Drosophila, Daphnia and lobster. Bimp (*B. improvisus*), Dp (Daphnia), DROME (Drosophila) HOMAM (Homarus) and GNAS (Ga$_s$).
Phylogenetic analysis of *B. improvisus* $\alpha_q$ and $\alpha_s$

Comparison of $\alpha_q$ and $\alpha_s$ of *B. improvisus* was done by constructing a phylogenetic tree by using neighbor joining method. In order to construct a phylogenetic tree, nucleotide sequences of $\alpha_q$ and $\alpha_s$ obtained from other invertebrates like drosophila, daphnia and lobster were aligned with the barnacles nucleotide sequences. It is obvious from the tree that $\alpha_q$ of *B. improvisus* has relationship with the $\alpha_q$ of other organisms like daphnia, drosophila and lobster. The $\alpha_s$ variants of *B. improvisus* also show close relationship with the $\alpha_s$ of daphnia, drosophila and lobster.

![Phylogenetic Tree](image)

**Fig. 15.** A phylogenetic tree showing the relationship of *B. improvisus* $\alpha_q$ and $\alpha_s$ with the $\alpha_q$ and $\alpha_s$ of drosophila, daphnia and lobster. GBB2_DROME was used as an out group.
DISCUSSION

Barnacles, like many marine invertebrates cause serious biofouling problems. In this study the barnacle *B. improvisus* G-protein encoding genes Ga<sub>q</sub> and Ga<sub>s</sub> were cloned and sequenced. In a *B. improvisus* EST-library parts of the G-protein sequences were available. In order to amplify the full length proteins, PCR and RACE techniques were chosen to be the method of choice and cDNA was used as template to avoid introns. During the amplification of Ga<sub>q</sub> many difficulties were found, requiring design and optimization of several different primers and amplification conditions. The EST sequences covered the whole coding region except for a small gap in the middle of the gene, as judged by comparison with Ga<sub>q</sub> from *H. americanus*. Running PCR with primers based on ETS sequences from just upstream of the predicted start codon and downstream of the stop codon, did not result in any product despite several optimization efforts. A possible explanation for this was obtained after performing 5´ Race where the whole coding region and the complete 5´ un-translated region was obtained. Comparing the 5´ un-translated region of the EST sequences with the sequences from the RACE product showed that they were not similar. There is thus a possibility that there are two slightly different genes, or splice products of the Ga<sub>q</sub> in *B. improvisus*.

The two different 5´ end sequences (5´ race short and 5´ race full length) of the Ga<sub>q</sub> of *B. improvisus* were aligned with *H. americanus*. The sequence obtained from the extreme 5´ end of Ga<sub>q</sub> 5´ race short was aligned with *H. americanus* sequence, but they were not very similar (about 50% identity). The nucleotides sequence alignment of the coding region of Ga<sub>q</sub> 5´ race full length fragment with *Homarus. americanus* sequences, show that both sequences are relatively similar. It is concluded that Ga<sub>q</sub> has different variants of the 5´ end, which is also seen in other Ga subunits like Ga<sub>o</sub> (Hsu et al., 1990). One explanation could be that they are the products of different genes. The Ga<sub>s</sub> encoding gene of the fruit fly *Drosophila melanogaster* also has two variants of the N-terminus. In drosophila three genes encode Ga<sub>s</sub> and they also differ in N-terminus (www.flybase.org). The other explanation could be that these differences are due to variation found in the sequences from individuals of same species.

In the EST sequences of Ga<sub>s</sub> from *B. improvisus* the start ATG is present upstream as compared with the ATG of *H. americanus*, drosophila and daphnia. Whether the translation will start at the first ATG of the open reading frame in *B. improvisus* is not clear.

In case of Ga<sub>s</sub>, three different variants were found which are very similar at the 5´ end but differ largely in the coding region of the 3´ end. The amino acid sequence of one of the variants is highly conserved and very similar with *H. americanus* sequences, while the two other variants are different in the C-terminal.

The occurrence of different variants of Ga<sub>s</sub> in *B. improvisus* at the 3´ end might be due to alternative splicing at the 3´ end of the mRNA. Another possible reason could be that these variants are the products of different genes. Splice variants of other G proteins also have been shown to have different amino acid sequences at the C-terminus. Three novel splice variants of Ga<sub>s</sub> were found in acute leukemia patients (Ye et al., 1999). The C-terminus of the G protein is known to interact with GPCRs in order to transmit signals (Yoo et al., 2002). We consider that change in the amino acid sequence at the C-terminus of Ga<sub>s</sub> would alter or abolish the interaction with the receptor, since these residues play a critical role in receptor interactions. It has been
shown that a Gαi protein C-terminal splice variant effect intracellular localization of dopamine receptor (Lopez-Aranda et al., 2007).

In the future, the other Gα subunits of the G-protein like Gαi and Gα12 will be cloned and sequenced. The interaction of different Gα subunits with GPCRs will be studied by using the yeast two hybrid interaction system or other systems. The mRNA expression level of different G proteins in different tissues or during different developmental stages will be investigated by QPCR.
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