GPCR Genes as Activators of Surface Colonization Pathways in a Model Marine Diatom

HIGHLIGHTS
- The model diatom *Phaeodactylum tricornutum* shifts morphology to form biofilms
- G-protein-coupled receptors (GPCRs) can modulate diatom surface colonization
- GPCR1A expression can induce biofouling morphotype and UV resistance
- Identified genes and pathways can serve as targets for anti-biofouling discoveries
A hallmark of the diatom cell wall is its siliceous structure, called the frustule, which is typically made of silicon dioxide ($\text{SiO}_2$) particles having diameter ranging from 10 to 100 nm (De Tommasi et al., 2017; Vardi et al., 2008). Unlike other known diatoms, the diatom *Phaeodactylum tricornutum* can grow in the absence of silicon and exists in different morphotypes such as fusiform, oval, and triradiate cell forms. Among these forms, only the oval cells make silicified frustules (De Martino et al., 2011; Lewin et al., 1958). The oval cell forms only become dominant in *P. tricornutum* culture population in response to environmental stress and during biofilm formation on solid surfaces in which cells form aggregates (De Martino et al., 2011; Finlay et al., 2013; Garacci et al., 2019; Salta et al., 2013; Varela et al., 2012). Marine biofouling has significant impacts on immersed artificial structures such as ship hulls, aquaculture cage facilities, and seawater handling pipes (Leterme et al., 2016; Salta et al., 2013). Particularly, biofouling on ships increases fuel and maintenance costs significantly (Salta et al., 2013). Conversely, surface colonization and biofilm formation have substantial physiological advantages for microalgae in response to stress conditions and also play important roles in microbial adaptation to fluctuations in marine environments (Dang and Lovell, 2016; Schaum, 2019).

 SUMMARY

Surface colonization allows diatoms, a dominant group of phytoplankton in oceans, to adapt to harsh marine environments while mediating biofouling to human-made underwater facilities. The regulatory pathways underlying diatom surface colonization, which involves morphotype switching in some species, remain mostly unknown. Here, we describe the identification of 61 signaling genes, including G-protein-coupled receptors (GPCRs) and protein kinases, which are differentially regulated during surface colonization in the model diatom species, *Phaeodactylum tricornutum*. We show that the transformation of *P. tricornutum* with constructs expressing individual GPCR genes induces cells to adopt the surface colonization morphology. *P. tricornutum* cells transformed to express GPCR1A display 30% more resistance to UV light exposure than their non-biofouling wild-type counterparts, consistent with increased silicification of cell walls associated with the oval biofouling morphotype. Our results provide a mechanistic definition of morphological shifts during surface colonization and identify candidate target proteins for the screening of eco-friendly, anti-biofouling molecules.

 INTRODUCTION

Diatoms (Bacillariophyta) are the dominant group of microalgae in today’s oceans (Benoist et al., 2017; Mock et al., 2017) and one of the most diverse and ecologically important clades of phytoplankton, contributing up to 20% of the global primary production (Bowler et al., 2008; Levitan et al., 2014; Vardi et al., 2008). Diatoms are recognized as the primary contributor to biofilm formation in marine or underwater environments, a process that leads to biofouling (Bruckner et al., 2011; Callow and Callow, 2011; Dang and Lovell, 2016; Finlay et al., 2013; Garacci et al., 2019; Salta et al., 2013; Varela et al., 2012). Marine biofouling has significant impacts on immersed artificial structures such as ship hulls, aquaculture cage facilities, and seawater handling pipes (Leterme et al., 2016; Salta et al., 2013). Particularly, biofouling on ships increases fuel and maintenance costs significantly (Salta et al., 2013). Conversely, surface colonization and biofilm formation have substantial physiological advantages for microalgae in response to stress conditions and also play important roles in microbial adaptation to fluctuations in marine environments (Dang and Lovell, 2016; Schaum, 2019).

An important aspect of the diatom cell wall is its siliceous structure, known as the frustule, which is typically made up of silicon dioxide ($\text{SiO}_2$) particles having a diameter ranging from 10 to 100 nm (De Tommasi et al., 2017; Vardi et al., 2008). Unlike other known diatoms, the diatom *Phaeodactylum tricornutum* is capable of growing without silicon and exists in different morphotypes such as fusiform, oval, and triradiate cell forms. Among these forms, only the oval cells make silicified frustules (De Martino et al., 2011; Lewin et al., 1958). The oval cell forms only become dominant in *P. tricornutum* culture population in response to environmental stress and during biofilm formation on solid surfaces in which cells form aggregates (De Martino et al., 2011; Francius et al., 2008). Silica plays an essential role in cell metabolism in *P. tricornutum* when it makes morphology shift from fusiform to oval in response to environmental stress. To date, *P. tricornutum* is the only known diatom species that can grow without silicon, and this feature makes it an ideal model to study cell morphology and how related molecular regulation underlies adaptive capability in marine environments. Although different strains/accessions of *P. tricornutum* differ in phenotypes as well as genotypes...
De Martino et al., 2007, 2011, it has been reported that only the oval cell morphotype of P. tricornutum can adhere to surfaces and give rise to colonies to form compact biofilms (Stanley and Callow, 2007).

Diatoms have evolved intricate signaling pathways in response to environmental fluctuations, allowing them to be the dominant clade of microalgae in oceans (Thompson and Coates, 2017). Among the signaling pathways in cell differentiation and morphogenesis (Basson, 2012), G-protein-coupled receptor (GPCR) signaling pathway is highly conserved across eukaryotes and plays an essential role in signal transduction and response to extracellular stimuli (Chan et al., 2015; Cohen et al., 2017; Port et al., 2013). GPCRs, also known as seven-transmembrane domain receptors, are present in most organisms and mediate many vital signaling processes and physiological functions through their interactions with G proteins and downstream effectors (Shaw et al., 2019). Fungal GPCR-mediated signaling and the understanding of receptor-ligand interactions could facilitate the development of receptor-interfering compounds for disease control (Brown et al., 2018). By manipulating the expression levels of the GPCR components, researchers have engineered the yeast Saccharomyces cerevisiae and tuned the receptor and reporter to function in a controllable manner to obtain a platform for biosensor applications (Shaw et al., 2019).

Here, we interrogate the molecular mechanism underlying the morphological shift in P. tricornutum (strain Pt18.6F) in relation to surface colonization (Figure 1). The Pt18.6F strain of P. tricornutum was selected as the model in our study as this strain is known to be an optimal reference model for genetic and genomic studies (Fu et al., 2017), with the ability to shift its morphotypes during surface colonization and biofilm formation, in addition to the availability of its complete genome sequence (Bowler et al., 2008). We identify a set of signaling genes that are differentially regulated during morphotype switching. We demonstrate that engineering the strain to overexpress a GPCR gene is sufficient to shift the dominant morphotype from fusiform to oval cells during non-stress growth conditions. The engineered strain also achieves similar photosynthesis performance as its fusiform wild-type (WT) counterpart, and it shows more resistance to UV stress, indicating its potential biotechnological applications.

RESULTS
Genome-wide Transcriptome Analysis of Two Different Morphotype Cultures
To identify transcriptional shift of cells between solid culture (oval > 75%) and liquid culture (fusiform > 75%), we performed RNA sequencing (RNA-seq) on RNA isolated from the WT P. tricornutum strain Pt18.6F grown in liquid and on solid media under low light intensity of 30–50 µmol photons m−2 s−1 (see Methods). The genome sequence of P. tricornutum (ASM15095v2 [2013-07-EBI-Phatr3]) was used as a reference to align the transcriptome reads, and normalized counts were calculated to characterize gene expression in cells (see Methods). Among all the expressed genes detected, differentially expressed genes (DEGs) were identified with a threshold of greater than 2-fold and a false discovery rate (FDR) of <0.05.
A total of 2,468 genes were up-regulated and 1,878 genes were down-regulated in the solid WT (Figure 2 and Table S1) when compared with the liquid WT cells. Gene set enrichment analysis (GSEA) (Bindea et al., 2009; Love et al., 2014) was carried out to identify overrepresented Gene Ontology (GO) terms for DEGs from the WT, highlighting cellular biosynthetic processes and signaling-related GO terms in up-regulated genes (Figure S1) and photosynthesis-related GO terms in down-regulated genes (Figure S2). We then performed KEGG Orthology annotations (Figure S3) on a total of 2,468 up-regulated genes from the WT using the KEGG Automatic Annotation Server (http://www.genome.jp/tools/kaas/) and identified 61 genes involved in 44 potential signaling pathways (Figure 2 and Table S2). GSEA for the 61 identified signaling genes was used for analysis. GO terms are represented as nodes in the graph, each pathway and its related nodes were presented with a unique color (p < 0.05), and node sizes indicate the relative numbers of genes that represent the GO term, whereas the edges represent genes shared between the GO terms.

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(Phatr3 J47068, Phatr3 J49346, and Phatr3 J35620) were up-regulated in the solid culture of *P. tricornutum* when compared with its liquid culture counterpart (Figure 2). A protein-protein interaction network was also predicted using these 61 signaling genes (59 of 61 were identified in the database) based on the STRING database (Figure S4). We note that the network is fragmented due to the lack of experimental evidence in diatoms.

**GPCR1A Transforms Show Distinct Morphological Shift and Strong Resistance to Light Stress**

We queried the KEGG Orthology annotation server with the 2,468 up-regulated genes and identified 61 signaling genes among them. To experimentally interrogate their potential roles in the morphological shift, we synthesized a group of 14 candidate signaling genes with a focus on several GPCR genes because GPCRs are on the top level of signal transduction hierarchy (Figure S5). We expressed these individually in *P. tricornutum* cells (Miyahara et al., 2013) with a controllable nitrate reductase promoter (see Methods) to test their ability to shift the cell morphology from fusiform to oval forms. We observed that cells transformed with either a GPCR1A or a GPCR4 construct shifted the population to be dominated by the oval cells.

We conducted surface colonization experiments with these GPCR transformants (i.e., GPCR1A transformants and GPCR4 transformants) on glass slides and found that both transformants showed much stronger attachment and developed more clumps on glass slides over 72 h when compared with their WT counterpart (Figure S6). The oval cell morphology-dominant transformants were able to colonize surfaces effectively and showed stronger adhesion patterns than their WT counterpart (Figure S6). These results are also in line with those of previous studies on the adhesion strength of different *P. tricornutum* species (De Martino et al., 2011; Stanley and Callow, 2007).

The GPCR1A transformants were selected for further experimental investigation. The typical cell morphotypes of the GPCR1A transforms and WT in liquid culture under non-stress growth conditions were observed by light and scanning electron microscopies (Figures 3A and 3B). The average cell size of the GPCR1A transformants (Table S3) in liquid culture was 6.63 ± 0.12 μm, which is similar to that of WT cells on solid culture (6.35 ± 0.12 μm) but significantly larger (p = 0.0016) than those of the cells in liquid culture (5.13 ± 0.25 μm). The transformed oval cells were morphologically similar to the WT oval cells.

To evaluate the strain capability in response to immediate light stress, we determined the effective quantum yields (QY) of photosystem II (PSII) of the transormants using the WT as controls under light stress of 220 μmol photons m⁻² s⁻¹ (Figure 3C). The effective QYs of the transformants were similar to those of the WT of liquid culture (Figure 3C). Moreover, the effects of UV treatment on different strains were studied (Figure 3D) to assess the role of diatom frustules in UV protection as silica cell walls in diatoms can protect DNA from UV light damage (Aguirre et al., 2018). As indicated by their different survival rates (p = 0.031), we found that the transformant cultures, with >75% oval cells, were more resistant to UV-C than the WT cultures of over 90% fusiform cells (Figure 3D). This result indicates that the oval cell form could provide an additional layer of protection against UV irradiation damage.

**Comparative Transcriptomics and Reconstruction of Signaling Network during Surface Colonization**

The GPCR1A transformants displayed oval cells as the dominant form in the population when compared with their WT counterparts under the same non-stress growth condition (Figure S6). The transcriptome analysis of GPCR1A transformants identified 1,568 up-regulated genes and 1739 down-regulated genes when compared with its WT counterpart in liquid culture (Table S4). The DEGs are fewer in the transformants than in the solid WT. Furthermore, 685 of the 1,568 up-regulated genes in the transformants were shared with the identified up-regulated DEGs in the solid WT during the surface colonization process (Figure 4). Among the 685 up-regulated DEGs, four GPCR genes including GPCR1A, GPCR1B, GPCR3, and GPCR4 were highly expressed in the transformants when compared with their WT counterpart in the liquid culture, showing a similar signaling pattern as the solid WT growing on agar plates (Figure 4 and Table S4). However, the fold changes of the top 100 up-regulated genes from the solid WT, ranging from 21.0 to 279.8 with a median level of 36.2, were much higher than those from transformants, ranging from 8.3 to 143.7 with a median level of
12.1 (Table S5). In addition, 14 of 16 shared genes of the two top-100 lists (Table S5) were at much greater fold changes in the solid WT, indicating that the expression shift was moderate in the transformants compared with those in the condition of harsh environmental stress (e.g., surface colonization). As a consequence of GPCR1A overexpression, downstream effector and regulator genes, including a GTPase-binding protein gene (Phatr3_J26387) and a protein kinase C gene (Phatr3_J14202), were also found to be up-regulated in the transformants. In addition to the GPCR signaling genes and their downstream effectors, we analyzed the transcription factors (TFs) from the two groups of DEGs, i.e., GPCR1A transformants and the solid WT, when compared with the wild type (with oval cells at <10% of the population). Although many TF families showed both up- and down-expression pattern, one TF, PHF5-like protein (PHD finger-like domain-containing protein 5A), was up-regulated both in the solid culture and in the transformants, when compared with the liquid WT. We note that PHF5-like proteins are reported with a role in cell cycle progression in yeast and morphological development in worms (Oltra et al., 2004).

We reconstructed and manually curated the proposed GPCR-mediated signaling pathway based on the integration of up-regulated DEGs in solid culture when compared with liquid culture information using Figure 3. Morphological Shift in the Diatom Population and Its Characteristics
(A) Bright-field microscopic images showing the dominant morphotypes in the GPCR1A transformants when compared with the wild-type (WT).
(B) Scanning electron microscopic images of different morphotypes in transformants and WT.
(C) Determination of the theoretical photosynthetic efficiency ($F_v/F_m$) and effective quantum yield (QY) of photosystem II under a light intensity of 220 $\mu$mol photons m$^{-2}$ s$^{-1}$.
(D) Enhanced cell resistance to UV radiation in transformants (with oval cells at >75% of the population) when compared with the wild type (with oval cells at <10% of the population). Values were averaged from either two or three independent experiments. WT, wild-type; PAR, photosynthetically active radiation. Error bars indicate SEM. * Indicates statistical significance between the groups (p < 0.05).
KAAS (KEGG Automatic Annotation Server, Tables S1, S2, and S6). The proposed pathways included major up-regulated signaling genes as a consequence of growth on solid media (Figure 6 and Table S6). Key effectors for the surface colonization process, including AMPK, cAMP, FOXO, MAPK, and mTOR signaling pathways and their related kinases and activators, are proposed (Figure 6). Among the 32 up-regulated signaling genes in the solid WT when compared with the liquid WT, only 11 signaling genes were found to be up-regulated in the transformants compared with their liquid WT counterpart (Figure 6). The transformants appeared capable of shifting the dominant morphotype from fusiform to oval cells with fewer up-regulated signaling genes than in the solid WT (Figure 6 and Table S6). Moreover, the fold changes of four of the GPCR signaling genes, i.e., GPC1A, GPC1B, GPC3, and GPCR4, were $2.87 \pm 0.07$ ($n = 3$, $\pm$ SEM), $2.41 \pm 0.11$, $2.55 \pm 0.04$, and $2.86 \pm 0.08$ times, respectively, when compared with their liquid WT counterparts; these fold differences were statistically significant (FDR < 0.05). However, the fold change of expression in these GPCR genes in transformants only ranged from 2.41 to 2.87. In contrast, the fold changes for these genes ranged from 4.14 to 49.18 in the solid WT when compared with the liquid WT (Tables S2 and S4), indicating that the expression changes of GPCR signaling genes were more affected by environmental factors.

A schematic wiring diagram of the key signaling components and the downstream targets is illustrated, and the polyamine pathway is highlighted (Figure S7). We note that polyamines play an important role in the promotion of silica precipitation for the formation of siliceous shells of diatoms (Kroger, 2007). Genes encoding well-known cell wall proteins such as silaffins have not been identified in P. tricornutum; however, the up-regulation of major genes involved in the polyamine pathway alludes to the role of polyamines in the construction of the frustule during oval cell formation (Figure S7).
DISCUSSION

The model diatom *P. tricornutum*, a planktonic species that also has a benthic morphotype (i.e., oval cell form), usually appears as fusiform in liquid cultures under non-stress conditions, and it contributes to biofilm formation and biofouling upon morphology shift and surface colonization (Thompson and Coates, 2017). The cell and molecular biology of biofouling are largely unknown, even though environment-friendly antifouling coatings are urgently needed. In this study, we identified a group of 61 signaling genes involved in *P. tricornutum* during the morphological shift of cell populations (Figures 4 and S3). These up-regulated signaling genes include five annotated GPCR genes in diatoms, i.e., GPCR1A, GPCR1B, GPCR2, GPCR3, and GPCR4. We note that these GPCRs might also be involved in responding to sexual cues in diatoms, such as the planktonic species *Pseudo-nitzschia multistriata* (Basu et al., 2017). In this study, we transformed the WT model diatom species, *P. tricornutum*, with GPCR expression constructs, to shift its morphotype and demonstrated that the engineered strain enhanced its resistance to UV light. Moreover, we show that the induced expression of GPCR1A is sufficient to shift morphotypes, pointing to this protein as a potential target protein of screening anti-biofouling compounds. A comparative RNA-seq study on *Phaeodactylum* morphotypes was recently reported on the strain Pt3 (Ovide et al., 2018), in which the major morphotype is oval (60%–75%) under liquid culture conditions. This phenotypic ratio is uncommon and is different in other strains of *P. tricornutum*, in which the major morphotype is fusiform (De Martino et al., 2007).

In addition to the observed cell differentiation in diatoms, other eukaryotic microorganisms, such as the yeast, are able to differentiate into specific subpopulations with different characteristics to form organized microbial communities such as biofilms, colonies, and pseudohyphae on solid surfaces with unique
The budding yeast *Saccharomyces cerevisiae* can differentiate from unicellular to pseudohyphal filamentous form in response to stress under the regulation of cyclic AMP-protein kinase A (cAMP/PKA) and the filamentation mitogen-activated protein kinase (fMAPK), rat sarcoma/protein kinase A (RAS/PKA), sucrose nonfermentable (SNF), and target of rapamycin (TOR) signaling pathways (Kim and Rose, 2015). Notably, GPCRs such as Gz subunit Gpa2 of the RAS pathway were recognized during cell differentiation to the filamentous form (Cullen and Sprague, 2012; Pothoulakis and Ellis, 2018). Furthermore, *S. cerevisiae* was found capable of forming one colony with two major

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**Figure 6. A Reconstructed Putative Signaling Network Involved in Surface Colonization**

The *P. tricornutum* signaling network reconstruction was done using information obtained from KAAS (KEGG Automatic Annotation Server), as well as through manual curation. Annotations were obtained by performing BLAST and assigning orthologs by the BBH (bidirectional best hit) method, against the manually curated KEGG GENES database. Compounds are indicated within circles: AMP, 5'-adenosine monophosphate; Ca²⁺, calcium cation; cAMP, cyclic AMP; DAG, diacylglycerol; PA, phosphatidic acid. Proteins are indicated in square-shaped boxes: 14-3-3, 14-3-3 protein epsilon; AC, adenylate cyclase 1; ACC1, acetyl-CoA carboxylase/biotin carboxylase 1; ARF-EF, ARF guanyl-nucleotide exchange factor; ATM, serine-protein kinase ATM; ATR, serine/threonine-protein kinase ATR; CBP, E1A/CREB-binding protein; CytC, cytochrome c; DAGK, diacylglycerol kinase; GABARAP, GABA(A) receptor-associated protein; GATOR, GATOR complex protein; GPCR, G-protein-coupled receptor; HSP70, heat shock 70-kDa protein; HSP90, heat shock 90-kDa protein; IPO7, importin-7; MOB, MOB kinase activator; mTORC1, serine/threonine-protein kinase mTOR; NDUF7, NADH dehydrogenase (ubiquinone) Fe-S protein 7; PI3K, phosphatidylinositol 3-kinase; PIIRH-2, RING finger and CHY zinc finger domain-containing protein; PKA, protein kinase A; PKC, protein kinase C; PLCD, phosphatidylinositol phospholipase C; PLD, phospholipase D1/2; PMCA, P-type Ca²⁺-transporter type 2B; SMEK, protein phosphatase 4 regulatory subunit 3; SPK, sphingosine kinase; UBCD, ubiquitin-conjugating enzyme E2 D; V-ATPase, V-type H⁺-transporting ATPase. Detailed information on differentially expressed genes is included in Table S6. Solid-line arrows represented direct transduction between two proteins/compounds, whereas dashed-line arrows indicated multiple steps of transduction between two proteins/compounds. The coloring of the lines was done to enhance the diagram. Blue-colored text indicated that the genes encoding the proteins were up-regulated in solid wild-type (WT) culture compared with liquid WT culture, yellow-colored boxes represented the genes encoding the proteins that were up-regulated in liquid transformants culture compared with liquid WT culture; the gene expression information is based on whole-transcriptome analysis carried out in this study.
subpopulations, i.e., U and L cells, occupying the upper and lower colony regions, respectively, and the optimized metabolic properties for U cell long-term existence were activated and controlled by the glutamine-induced TOR pathway amino acid sensor systems (e.g., SPS) together with other signaling pathways under lowered respiration (Cap et al., 2012). Mpk1 (the MAPK) and Bck1 (the MEKK) as well as the receptors that activate and regulate the pathway together with Rlm1 (a TF phosphorylated and activated by Mpk1), all played important roles during yeast cell differentiation into sporulation in colonies (Piccirillo et al., 2015). In this study, PKA and mTOR signaling pathways were also found up-regulated during diatom surface colonization (Figure 6; Tables S1, S4, and S6). Moreover, PKA was up-regulated in the GPCR1A transformants when compared with its WT counterpart (Figure 6). These findings highlighted the common function of these signaling components during cell differentiation across the distant phyla of diatoms and yeast.

P. tricornutum is the only known diatom species that can grow without silicate and has a significant distinction between its two main morphotypes, i.e., fusiform and oval cells, and only oval cells form silicified cell walls (Tesson et al., 2009). With the function of matrix peptides and proteins, silicic acid is converted into amorphous hydrated silicon dioxide with nanostructural properties in diatom cells (Pamirsky and Golokh-vast, 2013). However, there is no clear information on the native slialfins in P. tricornutum. The engineered strain demonstrated enhanced tolerance to UV stress when compared with the WT counterpart, most likely due to the silicified cell walls as diatom SiO2 frustules can protect DNA from UV light (Aguirre et al., 2018; Elleegaard et al., 2016). It has also been reported that genetically engineered diatom biosilica can be used for targeted drug delivery (Delalat et al., 2013). These findings may facilitate developing UV stress-resistant strains in industries as well as designing novel applications in biomedicine (Baio et al., 2014).

Marine biofouling has long constituted a major problem for the global maritime and aquaculture industries to alleviate as it has adverse effects and increases operating costs significantly (Callow and Callow, 2011; Salta et al., 2013). The International Maritime Organization banned many traditional biocide-based antifouling coatings, including the very effective and widely used tributyltin (TBT) coatings at the beginning of the twenty-first century due to their high toxicity toward all organisms in the ecosystem and negative impacts on non-target organisms. In this context, an alternative antifouling technology has been in great demand, and much effort has been focused on developing environment-friendly antifouling systems (Magin et al., 2010). Recently, researchers started to explore biomimetic designs for antifouling methods that can combat fouling without toxic side effects (Bixler and Bhushan, 2012; Kang et al., 2016). Understanding the biology of biofouling is important for the development of a successful and robust nontoxic biofouling remedy. As diatoms are the dominant algal group in the ocean and also one of the early colonizers involved particularly in the micro-fouling stage in which biofilm is formed (Dang and Lovell, 2016; Salta et al., 2013), biological understanding of the molecular regulation underlying diatom biofilm formation and dispersal may be beneficial to develop new strategies to control and mitigate biofouling. P. tricornutum is a polymorphic pennate diatom capable of strong adhesion to the substrates, common in pennate diatoms that all mediate biofouling (Dugdale et al., 2006; Herbstova et al., 2017; Stanley and Callow, 2007). Our findings may pave the way for developing effective and environment-friendly antifouling technology with promising biological targets from signaling pathways. As such, the GPCR1A gene identified here is a promising antifouling target. Furthermore, its orthologs in other biofouling organisms can be considered targets to interrogate for developments of a universal, bio-based, antifouling strategy. As morphological shift is a coordinated process during cell development, other regulatory factors such as TFs (Figure 5) may also play important roles and could be of interest for future studies. We expect the novel findings and resources described in this study to serve as a starting point for the discovery of new antifouling targets in the future.

Limitations of the Study

P. tricornutum strain Pt1 8.6v was used as an optimal model diatom species in this study as this strain can make a clear shift of morphotype from fusiform to oval cells during surface colonization. We identified and reconstructed a signaling network underlying surface colonization, highlighting the roles of signaling genes, including genes encoding GPCRs. However, the network analysis in this study relied on the reference genome sequence of P. tricornutum (2013-07-EBI-Phatr3), which is incomplete in functional annotations, as observable in KEGG pathway maps. As a consequence, there might be other key signaling genes and regulatory TFs involved that our study could not identify. In addition, although the analysis of the gene expression of the whole population allowed the identification of DEGs in conjunction with morphological shifts, our RNA-seq analysis looked at asynchronous populations of cells enriched in one or the other morphotype; therefore
our reported gene expression changes are limited to measurements of population averages and may not fully reflect the fold changes between individual morphotypes. Surface colonization is a long process (i.e., weeks), and the shift of dominant morphotype in population is a distinctive feature of *P. tricornutum* during such process and biofilm formation. There might be transient gene expression events that we likely could not cover with our approach as we only looked at gene expression changes at a single time point; however, our main goal was to identify gene products that could shift cell morphology and maintain it, which we were able to achieve. Finally, the identification of involved signaling genes was based on mRNA expression profiles and not protein levels due to the lack of antibodies for the detection of GPCRs in diatoms. Although we validated the role of GPCR1A experimentally, our analysis could have missed other effectors that their activity was modulated, but their expression level at mRNA level was not significantly altered.

**Resource Availability**

**Lead Contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Kourosh Salehi-Ashtiani (ksa3@nyu.edu).

**Materials Availability**

All sequences of synthesized genes can be found in Table S7.

**Data and Code Availability**

RNA-seq data from this article can be found in the GenBank/NCBI data libraries (GenBank: PRJNAS66271). The RNA-seq data have also been deposited in Dryad with a unique identifier (https://doi.org/10.5061/dryad.ns1m8ppx).

**METHODS**

All methods can be found in the accompanying Transparent Methods supplemental file.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101424.

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**AUTHOR CONTRIBUTIONS**

W.F. and K.S.-A. designed the research and wrote the paper. W.F. performed the laboratory experiments, including vector construction, transformation, and strain validation, and analyzed the data. W.F., A.C., B.D., and A.J. performed transcriptome analysis. M.S. performed confocal microscopy experiments. W.F. and K.B. conducted morphotype analyses. W.F. and J.W. conducted scanning electron microscopic analyses. W.F. and M.S. performed the RNA-seq experiments. W.F., D.A.-K., S.D., A.M., and A.A. contributed to strain screening and PCR verification. J.W. carried out scanning electron microscopic imaging of diatom samples. All authors contributed to the editing of the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing financial interests.
REFERENCES

Aguirre, L.E., Ouyang, L.Q., Elving, A., Hedblom, M., Wulff, A., and Inganas, O. (2018). Diatom frustules protect DNA from ultraviolet light. Sci. Rep. 8, 5138.

Baio, J.E., Zane, A., Jaeger, V., Roehrich, A.M., Lutz, H., Pfendtner, J., Drobny, G.P., and Weidner, T. (2014). Diatom mimics: directing the formation of biosilica nanoparticles by controlled folding of tyrosine-leucine peptides. J. Am. Chem. Soc. 136, 15134–15137.

Basson, M.A. (2012). Signaling in cell differentiation and morphogenesis. Cold Spring Harb. Perspect. Biol. 4, a008151.

Basu, S., Patil, S., Mapleson, D., Russo, M.T., Vitale, L., Fevola, C., Maumus, F., Casotti, R., Mock, T., Caccamo, M., et al. (2017). Finding a partner in the ocean: molecular and evolutionary bases of the response to sexual cues in a planktonic diatom. New Phytol. 215, 140–156.

Benoiston, A.S., Ibarbazi, F.M., Bittner, L., Guidi, L., Jahn, O., Dutkiewicz, S., and Bowler, C. (2017). The evolution of diatoms and their biogeochemical functions. Philos. Trans. R. Soc. B 372, 20160397.

Bindea, G., Milecnik, B., Hackl, H., Charoentong, P., Tosolini, M., Kirilovsky, A., Fridman, W.H., Pages, F., Trajanoski, Z., and Galon, J. (2009). ClueGO: a Cytoscape plug-in to decipher multidisciplinary survey. Mar. Genome Biol. 3, 6.

Bowler, C., Allen, A.E., Badger, J.H., Grimwood, J., Babij, K., Kuo, A., Maumus, U., Martens, C., Maumus, F., Ottillar, R.P., et al. (2008). The Phaeodactylum genome reveals the evolutionary history of diatom genomes. Nature 456, 239–244.

Brown, N.A., Schrevens, S., van Dijck, P., and Goldman, G.H. (2018). Fusulin G-protein-coupled receptors: mediators of pathogenesis and targets for disease control. Nat. Microbiol. 3, 402–414.

Bruckner, C.G., Rehm, C., Grossart, H.P., and Kroth, P.G. (2011). Growth and release of planktonic diatom. New Phytol. 215, 1052–1063.

Bruckner, C.G., Rehm, C., Grossart, H.P., and Callow, J.A. (2013). A novel biofilm channel for evaluating the adhesion of diatoms to non-biological surfaces. Environ. Microbiol. 15, 117–124.

Budler, G.D., and Bhushan, B. (2012). Biofouling: lessons from nature. Philos. Trans. R. Soc. A 370, 2381–2417.

Callow, J.A., Allen, A.E., Badger, J.H., Grimwood, J., Babij, K., Kuo, A., Maumus, U., Martens, C., Maumus, F., Ottillar, R.P., et al. (2008). The Phaeodactylum genome reveals the evolutionary history of diatom genomes. Nature 456, 239–244.

De Tommasi, E., Giels, J., and Rogato, A. (2017). Diatom frustule morphology and function: a multidisciplinary survey. Mar. Genom. 35, 1–8.

Delalat, B., Sheppard, V.C., Ghaemi, S.R., Su, S., Prestidge, C.A., McPhee, G., Rogers, M.L., Donoghue, J.F., Pillay, V., John, T.G., et al. (2015). Targeted drug delivery using genetically engineered diatom biosilica. Nat. Commun. 6, 679.

Dugdale, T.M., Willis, A., and Wetherbee, R. (2006). Adhesive modular proteins occur in the extracellular mucilage of the motile, penate diatom Phaeodactylum tricornutum. Biophys. J. 90, L58–L60.

Ellegaard, M., Lenau, T., Lundholm, N., Maibohm, C., Friis, S.M.M., Rottkat, K., and Su, Y.Y. (2016). The fascinating diatom frustule-can it play a role for attenuation of UV radiation? J. Appl. Physiol. 28, 3295–3306.

Finlay, J.A., Schultz, M.P., Cone, G., Callow, M.E., and Callow, J.A. (2013). A novel biofilm channel for evaluating the adhesion of diatoms to non-biocidal coatings. Biofouling 29, 401–411.

Francis, G., Tesson, B., Dague, E., Martin-Jezequel, V., and Dufrene, Y.F. (2008). Nanostructure and nanomechanics of live Phaeodactylum tricornutum morphotypes. Environ. Microbiol. 10, 1344–1356.

Fu, W.Q., Chaiboonchoe, A., Khraiwesh, B., Sultana, M., Jaiswal, A., Jijakli, K., Nelson, D.R., Miyahara, M., Aoi, M., Inoue-Kashino, N., Kashino, Y., and Ifuku, K. (2013). Highly efficient transformation of the diatom Phaeodactylum tricornutum by multi-pulse electroporation. Biosci. Biotechnol. Biochem. 77, 874–876.

Mock, T., Ottillar, R.P., Strauss, J., Mulligan, M., Paajanen, P., Schmitz, J., Salamon, A., Sanges, R., Toseland, A., Ward, B.J., et al. (2012). Evolutionary genomics of the cold-adapted diatom Fragilariopsis cylindrus. Nature 491, 536–540.

Oltra, E., Verde, F., Werner, R., and D’Urso, G. (2004). A novel RING-finger-like protein NIN1 is essential for cell cycle progression in fission yeast. J. Cell Sci. 117, 967–974.

Ovide, C., Kiever-Meyer, M.C., Berard, C., Vergne, N., Lecroq, T., Plisson, C., Burel, C., Bernard, S., Drouich, A., Lerouve, P., et al. (2018). Comparative in depth RNA sequencing of P. tricornutum’s morphotypes reveals specific features of the oval morphotype. Sci. Rep. 8, 14340.

Pamirsky, I.E., and Golokhvast, K.S. (2013). Silaffins of diatoms: from applied biotechnology to biomedicine. Mar. Drugs 11, 3155–3167.
Piccirillo, S., Morales, R., White, M.G., Smith, K., Kapros, T., and Honigberg, S.M. (2015). Cell differentiation and spatial organization in yeast colonies: role of cell-wall integrity pathway. Genetics 201, 1427–1438.

Port, J.A., Parker, M.S., Kodner, R.B., Wallace, J.C., Armbrust, E.V., and Faustman, E.M. (2013). Identification of G protein-coupled receptor signaling pathway proteins in marine diatoms using comparative genomics. BMC Genomics 14, 503.

Pothoulakis, G., and Ellis, T. (2018). Synthetic gene regulation for independent external induction of the Saccharomyces cerevisiae pseudohyphal growth phenotype. Commun. Biol. 1, 7.

Salta, M., Wharton, J.A., Blache, Y., Stokes, K.R., and Briand, J.F. (2013). Marine biofilms on artificial surfaces: structure and dynamics. Environ. Microbiol. 15, 2879–2893.

Schaum, C.E. (2019). Enhanced biofilm formation aids adaptation to extreme warming and environmental instability in the diatom Thalassiosira pseudonana and its associated bacteria. Limnol. Oceanogr. 64, 441–460.

Shaw, W.M., Yamauchi, H., Mead, J., Gowers, G.F., Bell, D.J., Oling, D., Larsson, N., Wigglesworth, M., Ladds, G., and Ellis, T. (2019). Engineering a model cell for rational tuning of GPCR signaling. Cell 177, 782–796.e27.

Stanley, M.S., and Callow, J.A. (2007). Whole cell adhesion strength of morphotypes and isolates of Phaeodactylum tricornutum (Bacillariophyceae). Eur. J. Phycol. 42, 191–197.

Tesson, B., Gaillard, C., and Martin-Jezequel, V. (2009). Insights into the polymorphism of the diatom Phaeodactylum tricornutum Bohlin. Bot. Mar. 52, 104–116.

Thompson, S.E.M., and Coates, J.C. (2017). Surface sensing and stress-signalling in Ulva and fouling diatoms - potential targets for antifouling: a review. Biofouling 33, 410–432.

Vanelslander, B., Paul, C., Grueneberg, J., Prince, E.K., Gillard, J., Sabbe, K., Pohnert, G., and Vyverman, W. (2012). Daily bursts of biogenic cyanogen bromide (BrCN) control biofilm formation around a marine benthic diatom. Proc. Natl. Acad. Sci. U S A 109, 2412–2417.

Vardi, A., Thamatrakoln, K., Bidle, K.D., and Falkowski, P.G. (2008). Diatom genomes come of age. Genome Biol. 9, 245.
Supplemental Information

GPCR Genes as Activators of Surface Colonization

Pathways in a Model Marine Diatom

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Supplemental Information
Supplemental Information includes Transparent Methods, eight figures, and seven tables (shown as datasets in Excel files).

Transparent Methods
Algal culture and growth conditions
The strain, *P. tricornutum* Pt1 8.6F (CCAP 1055/1, Culture Collection of Algae and Protozoa, Scottish Marine Institute, Scotland, UK), was used as a model diatom in this study. The wild type *P. tricornutum* was grown using f/2 +Si medium on either solid agar plates (for surface colonization) or in liquid culture (in flasks) under continuous cool white fluorescent lights of 50 μmol photons m\(^{-2}\) s\(^{-1}\) at 22° ± 2°C (Fu et al., 2017). Cell quantification was performed by determining the absorbance (at 600 nm) using a spectrophotometer regularly.

Chlorophyll fluorescence measurements for photosynthetic parameters
Chlorophyll fluorescence was measured using a PHYTO-PAM-II (Heinz Walz GmbH). The maximum photosynthetic efficiency of PSII was determined as \(F_v/F_m = (F_m - F_o)/F_m\), where \(F_m\) is the maximal fluorescence, and \(F_o\) is the ground fluorescence in the dark- or low-light adapted cells. The effective quantum yield of PSII in cells was determined as \(F_v'/F_m'\) when exposed to the white light of 220 μmol photons m\(^{-2}\) s\(^{-1}\) over 10 minutes, where \(F_m'\) is the maximum fluorescence emission level in the light-acclimated cells. The relative electron transport rate (Cohen et al.) was also obtained for samples during the same experiments.

Generation of genetically modified *P. tricornutum* cells
A group of 14 cDNAs encoding candidate signaling genes (Gene IDs: Phatr3_J22677, Phatr3_J41807, Phatr3_Jdraft1756, Phatr3_J10677, Phatr3_J54411, Phatr3_EG02512, Phatr3_J51511, Phatr3_J12877, Phatr3_J2097, Phatr3_Jdraft1000, Phatr3_J54505, Phatr3_J55230, Phatr3_Jdraft1740, and Phatr3_J44133), including GPCR genes (as shown in Table S7), were synthesized and cloned in the shuttle vector pPha-NR (Fu et
al., 2017) with a nitrate reductase promoter by Twist Bioscience (San Francisco, CA, U.S.A.). The plasmids were then extracted from the bacterial cell cultures using the Qiagen Plasmid Mini kit (Hilden, Germany) after growth and selection in the presence of ampicillin (100 μg/ml). For the transformation of signaling genes in the diatom *P. tricornutum* (Miyahara et al., 2013), the multipulse electroporation protocol was followed as previously described (Fu et al., 2017). PCR verification of the transformants was conducted using genomic DNA and specific primers (Fig. S8).

**Sample preparation, RNA extraction, and transcriptome sequencing (RNA-seq)**

For solid culture experiments of wild type *P. tricornutum*, cells were grown on solid agar (1.2% agar plus the growth medium) for approximately two weeks, while control liquid cultures were grown in flasks for approximately one week (OD$_{600}$=~0.3). Both culture types were generated using f/2 +Si medium under continuous cool white fluorescent lights of 50 μmol photons m$^{-2}$ s$^{-1}$ at 22° ± 2°C. Cells were collected during the exponential phase for RNA extraction. For each group, three independent experiments were conducted, and triplicates of samples were collected for RNA sequencing studies.

For comparison between the transformed (engineered) *P. tricornutum* and its wild type counterpart, both cultures were grown under the same conditions, in liquid, using f/2 +Si medium in flasks for approximately one week under continuous cool white fluorescent lights of 50 μmol photons m$^{-2}$ s$^{-1}$ at 22° ± 2°C. Cultures were added with 5.0 mM KNO$_3$ for one day, and cells were collected during the exponential phase for RNA extraction. For each group, three independent experiments were carried out, and triplicate samples were collected for RNA extraction.

Total RNA extraction was performed using the MagMAX-96 Total RNA Isolation kit AM1830 with all necessary reagents (Thermo Fisher Scientific Inc.) according to the manufacturer’s instruction. The complementary DNA (cDNA) libraries were prepared using the TruSeq RNA Library Preparation kit v2 (Illumina) according to the manufacturer’s instruction, as previously described (Fu et al., 2017). After quality inspection, the library was quantified by quantitative PCR for cluster generation on the Bot system and then
sequenced using paired-end sequencing of 2 × 100–base pair read length on an Illumina HiSeq 2500 system (Illumina).

Quantitative comparison of gene expression, and gene set enrichment analysis (GSEA)
The genome sequence of *P. tricornutum* (2013-07-EBI-Phatr3) was used as a reference to align the transcriptome reads (http://protists.ensembl.org/Phaeodactylum_tricornutum/Info/Annotation/). To characterize gene expression in different samples, DESeq2 (Love et al., 2014) was used to identify DEGs between two groups of samples (triplicates within each group). Following the removal of the raw read count <10 in each sample, the total read count was normalized using DESeq2, which scales the read counts by a reference sample based on the geometric mean of the read counts across all samples. RPKM (Reads Per Kilobase of transcript, per Million mapped reads) was used as a normalized unit of transcript expression. Functional analysis and GSEA of DEGs were performed using ClueGO plugin (v2.5.0) in Cytoscape (v3.6.0; (Bindea et al., 2009). ClueGO determined the statistical overrepresentation of GO terms associated with the identified DEGs. The P values were calculated using a two-sided hypergeometric test, and Benjamini-Hochberg false discovery rate (FDR) correction was used to identify the statistical significance of GO terms with corrected P < 0.05 for multiple testing. The network was automatically laid out based on the ClueGO visual style, where nodes represented the GO terms, the node size referred to the number of genes in a particular GO, and the color of the nodes reflected the enrichment significance of the GO terms.

Surface colonization on glass slides
Glass microscope slides were first rinsed with deionized water then soaked in 1M HCl for 24 h (Stanley and Callow, 2007). The acid was removed by rinsing the slides using deionized water. The slides were rinsed and dried before adding the prepared diatom culture. The wild type, GPCR1A transformants, and GPCR4 transformants were cultivated under the same condition, then collected during exponential growth. Cells numbers were determined using a hemocytometer under a microscope for all strains. The same amount of cells
(500,000 cells/ml) were used in each experiment. Slides were placed in individual compartments of culture dishes, and 15 ml of cultures at the correct dilution were added. Three replicates were used for each experiment. The cells were allowed to settle for 72 h at 22±2°C in growth chambers, after which the slides were gently shaken/rinsed to remove unattached cells on a shaker at 150 rpm. Cells were then counted from each of the three replicate slides after recording microscope images from 9 randomly sampled views to provide cell adhesion and colonization data.

**Statistical analysis**

A student’s t-test was performed to evaluate the significance of the difference between the two groups of samples for physiology data (p-value < 0.05).
Supplementary figures and figure legends

Fig. S1. Gene set enrichment analysis (GSEA) of up-regulated DEGs, related to Figure 2. GSEA was carried out to identify the overrepresented gene ontology (GO) terms for biological process, and molecular function of 2468 identified differentially expressed genes (DEGs) between wild type (WT) strain grown on solid media compared to liquid media growth. Node sizes indicate the relative numbers of genes that represent a GO term; GO terms are represented as nodes in the graph, the color gradient (red to dark red) represents the statistical term enrichment significance (P < 0.05). The blue circle marks the GO terms related to signaling pathways.
**Fig. S2. GSEA of down-regulated DEGs, related to Figure 2.** The 1878 DEGs were used to identify overrepresented gene ontology (GO) terms for biological process and molecular function between the growth of WT on solid media in comparison with liquid growth. Node sizes indicate the relative numbers of genes representing a GO term; GO terms are represented as nodes in the graph, the color gradient (red to dark red) represents the statistical term enrichment significance (P < 0.05). Blue circle highlights the GO terms identified in relation to photosynthetic processes.
Identification of signaling pathways and genes using BLAST functional annotation and pathway mapping, related to Figure 2. The sequences of the 2468 up-regulated genes in the WT were searched by BLAST against the manually curated KEGG GENES database (https://www.genome.jp/kegg/kaas/) with a complete genome of *P. tricornutum* using BBH (bi-directional best hit of nucleotide by its default setup) to assign orthologs and provide functional annotation and pathway mapping through KEGG Automatic Annotation Server. The signaling pathways are marked with red boxes. Detailed gene IDs of the 61 identified signaling genes are described in Supplemental Table 2.
Fig. S4. Gene expression profile of 61 identified signaling genes, related to Figure 3.  (A) A heatmap of the identified up-regulated signaling genes in the wild type. The color key represents the value of log$_2$ (Normalized counts) of RPKM values. (B) Reconstruction of gene association network using the up-regulated signaling genes using the STRING database (https://string-db.org). The network is grouped using k-means clustering (with k=3). Nodes represent genes, and node colors represent the k-mean clustering; edge (link) colors indicate the type of evidence for the interactions. Known interactions, predicted interactions, and other interactions are indicated in the lower panel of the figure.
Figure S5. A workflow for screening and selection of candidate genes for overexpression in *P. tricornutum*, related to Figure 3. The principal selection criteria are based on the integration of gene expression information with computational enrichment analyses and manual curation of the results, as shown.
Fig. S6. Surface colonization of the wild type and the two transformants on glass slides over 72 hours, related to Figures 3 and 4. (A) Brightfield microscopy image of the cells on glass slides. Oval cells were observed as the dominant form in the population of the two transformants with the 100X objective lens using oil immersion. (B) More cells of transformants were attached to the glass slides in comparison with the wild type. Cells were counted based on a total of 9 fields of view for each glass slide. Detailed information on surface colonization experiments is present in the Methods section. The symbol (*) indicates a significant difference between the two groups (p=0.032 for comparison between the wild type and GPCR1A transformants, and p=0.039 for comparison between the wild type and GPCR4 transformants, respectively); however, there is no significant difference between GPCR1A transformants and GPCR4 transformants (p=0.103).
Figure S7. An overview of key putative signaling and metabolic pathways affected in GPCR1A transformants, related to Figure 6. The illustration of major signaling network elements was derived from Figure 6 based on network reconstruction and whole transcriptome analysis. The environmental signals (putative ligands) can bind to GPCR1A, and the associated G-proteins induce the downstream effector-activated pathways (e.g., MAPK/ERK). The G-protein (consisting of α, β, and γ subunits) is separated in the illustration from GPCR1A for the visualization of its activation by the receptor moiety (the drawing does not reflect the actual localization of the G proteins, which are otherwise closely associated with the receptor). The figure also highlights the GPCR-mediated activation of the polyamine synthesis pathway as polyamines play an important role in silica precipitation for the formation of siliceous shells of diatoms. MAPK/ERK: mitogen-activated protein kinase or extracellular signal-regulated kinase.
Fig. S8. PCR verification of GPCR transformants based on genomic DNA, related to Figures 3 and 4. (A) Primers (Forward, 5’-CACAAACCGAACAGCCCTAC-3’; Reverse, 5’-TCGAGCTTCACAACCTGTCC-3’) were used to amplify DNA fragment across
the nitrate reductase promoter and GPCR1A gene to verify the positive transformation of the synthetic DNA construct into *P. tricornutum* cells; (B) Primers (Forward, 5'-CACAAACCGAACAGCCCTAC-3'; Reverse, 5'-CAGTGACGTTGCGACAATCC-3') were used to amplify DNA fragment across the nitrate reductase promoter and GPCR4 gene to verify the positive transformation of the synthetic DNA construct into *P. tricornutum* cells.