Dynamic Stabilization of Expressed Proteins in Engineered Diatom Biosilica Matrices

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ABSTRACT: Self-assembly of recombinant proteins within the biosilica of living diatoms represents a means to construct functional materials in a reproducible and scalable manner that will enable applications that harness the inherent specificities of proteins to sense and respond to environmental cues. Here we describe the use of a silafln-derived lysine-rich 39-amino-acid targeting sequence (Sil3T8) that directs a single chain fragment variable (scFv) antibody or an enhanced green fluorescent protein (EGFP) to assemble within the biosilica frustule, resulting in abundance of >200 000 proteins per frustule. Using either a fluorescent ligand bound to the scFv or the intrinsic fluorescence of EGFP, we monitored protein conformational dynamics, accessibility to external quenchers, binding affinity, and conformational stability. Like proteins in solution, proteins within isolated frustules undergo isotropic rotational motion, but with 2-fold increases in rotational correlation times that are indicative of weak macromolecular associations within the biosilica. Solvent accessibilities and high-affinity (pM) binding are comparable to those in solution. In contrast to solution conditions, scFv antibodies within the biosilica matrix retain their binding affinity in the presence of chaotropic agents (i.e., 8 M urea). Together, these results argue that dramatic increases in protein conformational stability within the biosilica matrices arise through molecular crowding, acting to retain native protein folds and associated functionality with the potential to allow the utility of engineered proteins under a range of harsh environmental conditions associated with environmental sensing and industrial catalytic transformations.

Synthetic biology offers a means to develop cellular factories that allow the low-cost synthesis of chemicals and designer materials (see "A New Biology for the Twenty-first Century"). Examples of these approaches currently involve well-studied organisms, including E. coli and S. cerevisiae, whose genetics facilitate the introduction of genes encoding proteins within entire synthetic pathways that permit the generation of industrial quantities of precursor chemicals (e.g., 1,3-propanediol) or low-cost pharmaceuticals (e.g., artemisin). Additional advances are needed to create materials that permit "cell-free" bioprocessing that might include, for example, the cell walls (frustules) of diatoms, as biosilica materials functionally stabilize many bioorganic macromolecules. Along these lines, prior measurements have demonstrated an ability to target specific proteins within living diatoms to organelles linked to biosilica formation, resulting in the creation of highly porous biosilica structures with 18 ± 3 nm spherical nanopores containing embedded proteins capable of mediating analyte binding and catalytic transformation. Such functionalized materials are mechanically strong and potentially useful for a range of existing industrial applications, including those that already use diatomite, in filtration, toothpaste, insecticides, as an absorbent in liquids, and as an activator of blood clotting. Further, genetically engineered diatom biosilica expressing the highly stable immunoglobulin G binding domain of protein G (GB1) permits the selective binding of cell-targeting antibodies. However, current applications are limited to the targeting of highly stable proteins to the biosilica matrix, and have not considered the mechanisms whereby the biosilica matrix can alter analyte binding affinity, substrate accessibility, or catalytic activity that may limit the usefulness of these materials. Furthermore, current approaches of protein immobilization in biosilica result in large reductions in (i) rates of analyte binding and (ii) binding affinities. Such losses in protein function upon immobilization in biosilica may arise from mechanisms similar to those previously observed following protein adsorption onto synthetic silicate surfaces, which can restrict catalytically important domain motions. In this respect, new bioassembly approaches are needed, like those developed in hydrogels, which tether and stabilize proteins within porous materials while maintaining critical domain movements linked to function.

Received: March 25, 2016
Revised: May 2, 2016
Published: May 3, 2016

DOI: 10.1021/acs.bioconjchem.6b00165
Bioconjugate Chem. 2016, 27, 1205−1209
To address the possible utility of biosilica-immobilized proteins, we have used the silaffin-derived lysine rich 39-amino-acid sequence (Sil3T8), previously described by Kröger and colleagues, to target enhanced green fluorescent protein (EGFP) or a single chain fragment variable (scFv) antibody to the biosilica matrix of the diatom Thalassiosira pseudonana. The stability and natural fluorescence of EGFP aid in the quantification of the protein microenvironment. The scFv antibody was chosen as a prototype of a class of reagents routinely used for detection and diagnostics, which commonly involve engineered protein scaffolds that include domain elements of immunoglobulins and engineered single domain proteins. Furthermore, current clinical applications of engineered protein scaffolds use antibody fragments, such as single chain antibodies (scFv), which represent a dominant new approach in the development of antibody-directed therapies. Advantages of the scFv scaffold include rapid selection against target antigens and the ability to obtain large amounts of antibodies using cost-efficient expression systems. However, widespread applications suffer from significant challenges associated with a need to engineer protein stability, while retaining sufficient flexibility within the antigen binding domain to maintain high-affinity binding. Further, engineered protein scaffolds have a tendency to exhibit poor solubility and commonly aggregate in solution.

Facilitating analysis of the function of the scFv, we have coupled Alexa fluorophores to a derivative of the trinitrotoluene ligand (TNT), permitting a determination of how protein targeting affects ligand affinity and accessibility. Dynamic structural measurements of highly fluorescent EGFP were used to identify how the biosilica environment may act to restrict protein motions to enable a mechanistic understanding of how the biosilica environment stabilizes protein function. Finally, possible alterations in protein stability were used to quantify whether protein stabilization requires associated reductions in protein dynamics, as previously reported for adsorbed proteins using mesoporous silicate and hydrogel materials.

To understand the utility of engineered frustules as a platform to detect small molecule ligands, we transformed diatoms with a well characterized scFv antibody against TNT linked to the Sil3T8 biosilica targeting peptide. The scFv antibody is tethered within the biosilica matrix, as Sil3-directed fusion proteins are (i) retained within the biosilica despite repeated washing and (ii) capable of interacting with large amounts of frustules/mL = 2 nM scFv) (Figure 1). The observed binding of nanomolar concentrations of scFv (6 × 10^6 frustules/mL) resulted in >500 000 EGFP in each transformed frustule,5 resulting in >500 000 EGFP in each transformed frustule (Figure S4). These observations indicate that there is minimal nonspecific binding between TNB-Alexa555 and the biosilica matrix. Maximal TNB-Alexa555 binding stoichiometries indicate that there are >200 000 scFv antibodies present in isolated frustules. This was accomplished through measurements of the fluorescence intensity of bound TNB-Alexa555, which binds with high affinity to this scFv (K_d = 90 pM; Figure S2). In the presence of a large excess TNB-Alexa555, less than 0.2% binds to transformed frustules expressing scFv antibodies (Figure S3). These results indicate that there is minimal nonspecific binding between TNB-Alexa555 and the biosilica matrix. Maximal TNB-Alexa555 binding stoichiometries indicate that there are >200 000 scFv antibodies within each transformed frustule. Similar abundances of enhanced green fluorescent protein (EGFP) are apparent when the same Sil3T8 targeting peptide is used to express and target EGFP to the biosilica matrix within the diatom frustule, resulting in >500 000 EGFP in each transformed frustule (Figure S4). These observations indicate that the transformed frustules contain nanomolar concentrations of scFv (6 × 10^6 frustules/mL = 2 nM scFv) (Figure 1). The observed binding of nanomolar amounts of TNB-Alexa488 (2 nM) upon incubation with transformed frustules expressing nanomolar amounts of scFv antibodies provides strong evidence that the high-affinity binding between TNB-Alexa555 and scFv antibodies observed in solution is retained in transformed biosilica.

To assess whether the scFv antibodies in the transformed diatoms retain a similar high-affinity ligand binding capacity to that observed in solution, we measured the abundance of scFv antibodies present in isolated frustules. This was accomplished through measurements of the fluorescence intensity of bound TNB-Alexa555, which binds with high affinity to this scFv (K_d = 90 pM; Figure S2). In the presence of a large excess TNB-Alexa555, less than 0.2% binds to transformed frustules expressing scFv antibodies (Figure S3). These results indicate that there is minimal nonspecific binding between TNB-Alexa555 and the biosilica matrix. Maximal TNB-Alexa555 binding stoichiometries indicate that there are >200 000 scFv antibodies within each transformed frustule. Similar abundances of enhanced green fluorescent protein (EGFP) are apparent when the same Sil3T8 targeting peptide is used to express and target EGFP to the biosilica matrix within the diatom frustule, resulting in >500 000 EGFP in each transformed frustule (Figure S4). These observations indicate that the transformed frustules contain nanomolar concentrations of scFv (6 × 10^6 frustules/mL = 2 nM scFv) (Figure 1). The observed binding of nanomolar amounts of TNB-Alexa488 (2 nM) upon incubation with transformed frustules expressing nanomolar amounts of scFv antibodies provides strong evidence that the high-affinity binding between TNB-Alexa555 and scFv antibodies observed in solution is retained in transformed biosilica.

The utility of proteins tethered within isolated frustules for environmental monitoring requires that mass transfer (translational diffusion) is not restricted by the biosilica matrix. Using the fluorescence quencher trypan blue (mass = 873 Da), we find that the solvent accessibility is very similar for TNB-Alexa555 bound to scFv (K_d = (20 ± 1) × 10^4 M^-1) and EGFP (K_d = (35 ± 1) × 10^4 M^-1) in transformed frustules, which is expected given the similar fluorescence lifetimes (3.3 ns vs 2.4 ns) and localization patterns (Figure 2). In comparison, quenching efficiencies are lower in solution for...
comparison to EGFP in solution, there is a reduction in the times associated with protein rotational dynamics described, two rotational correlation times are required to reduce in rates of rotational motion. As previously expressed in transformed frustules that is indicative of a lifetime of EGFP, suggesting that observed differences in brightness of EGFP within the biosilica matrix to measure the physical interactions between proteins and the biosilica matrix of proteins for a range of industrial applications. To understand modulated light, there is an increase in the differential phase and modulated anisotropy for EGFP within the biosilica matrix indicates an increase in the environment where binding and catalysis for embedded proteins will not be diffusion limited.

Prior measurements have indicated a functional stabilization of a range of proteins upon their expression within the biosilica frustule. Understanding the mechanisms associated with protein stabilization is necessary for the routine deployment of proteins for a range of industrial applications. To understand the physical interactions between proteins and the biosilica matrix, we have taken advantage of the superior expression and brightness of EGFP within the biosilica matrix to measure the protein rotational dynamics using frequency-domain fluorescence spectroscopy. Upon increasing the frequency of modulated light, there is an increase in the differential phase and modulated anisotropy, which can be fit using the method of nonlinear least-squares to measure the rotational correlation times associated with protein rotational diffusion (Figure 3). In comparison to EGFP in solution, there is a reduction in the differential phase and modulated anisotropy for EGFP expressed in transformed frustules that is indicative of a reduction in rates of rotational motion. As previously described, two rotational correlation times are required to describe the rotational dynamics of EGFP in solution that correspond to a fast (\( \phi_1 = 1.1 \text{ ns} \)) and a slower rotational correlation time (\( \phi_2 = 18.3 \text{ ns} \)); the measured rotational correlation time is inversely related to rates of rotational diffusion. The longer rotational correlation time is in close agreement with the calculated rotational correlation time of 15.2 ns based on the crystal structure 4eul.pdb using the program Hydropro and prior measurements for the rotational motion of GFP in aqueous buffer of 20 ± 1 ns. In comparison, the rotational dynamics of EGFP within the biosilica matrix can be described with a single rotational correlation time (i.e. \( \phi_{rot} = 42 ± 2 \text{ ns} \)) that is very similar to that previously reported for GFP within either the endoplasmic reticulum (\( \phi_{rot} = 39 ± 5 \text{ ns} \)) or cytoplasm of cells grown in culture (\( \phi_{rot} = 36 ± 2 \text{ ns} \)). The large increase in the rotational correlation time for EGFP tethered in biosilica is significant, as there is no overlap in the error surfaces for EGFP in solution and in the biosilica matrix (Figure 3C). The absence of any residual anisotropy in the measurements of the rotational dynamics for EGFP following tethering within the biosilica matrix indicates that EGFP within the biosilica matrix is not adsorbed onto the surface. The approximately 2.5-fold increase in rotational correlation time due to tethering EGFP within the biosilica matrix indicates that EGFP within the biosilica matrix is not adsorbed onto the surface. The approximately 2.5-fold increase in rotational correlation time due to tethering EGFP within the biosilica matrix indicates that EGFP within the biosilica matrix is not adsorbed onto the surface. The approximately 2.5-fold increase in rotational correlation time due to tethering EGFP within the biosilica matrix indicates that EGFP within the biosilica matrix is not adsorbed onto the surface.

Reductions in the rotational dynamics of proteins within the biosilica matrix through biomolecular interactions suggest a crowded environment, which may resist protein denaturation. To assess the possible stability of the scFvTNT antibody within the biosilica matrix, we took advantage of the environmental sensitivity of the fluorescence lifetime of TNB-Alexa555. Using frequency-domain fluorescence spectroscopy to measure the fluorescence lifetime, we observe that in comparison to TNB-Alexa555 nonspecifically associated with the biosilica matrix, there is a large shift toward lower frequencies in the frequency response upon binding to the scFvTNT antibody that is
The two-state model suggests that approximately one-half of the scFv binding is retained between TNB-Alexa555 and scFvTNT in solution (Figure S5), which is consistent with the denaturation of scFvTNT. Under these same denaturing conditions (i.e., 8 M urea), binding is retained between TNB-Alexa555 and scFvTNT within the biosilica matrix, which is apparent from the retention of a significant shift in the frequency response in comparison to unbound TNB-Alexa555 (Figure 4). Analysis of binding using a two-state model suggests that approximately one-half of the scFvTNT retains an ability to bind TNB-Alexa555. A similar stabilization of EGFP is observed following immobilization in biosilica (Figure S6). These results suggest that proteins tethered within the biosilica matrix are stabilized through molecular crowding, and that transformation of diatoms with antibodies represents a path forward toward antibody stabilization that will increase their utility in assays under a range of harsh assay conditions that may, for example, denature antigens to permit their recognition by scFv antibodies within the frustule matrix.

In summary, we have demonstrated that silica-tethered recombinant proteins in diatom frustules are stabilized within a biosilica matrix against chaotropic agents (i.e., 8 M urea) (Figure 4). Silica-tethered recombinant proteins retain the isotropic rotational mobility that is necessary for the retention of high-affinity binding (Figures 1 and 3). Mass transfer is unrestricted through the highly porous biosilica matrix (Figure 2), thereby enabling the use of transformed biosilica for a range of applications involving the detection or chemical transformation of a range of organic molecules, including chemical or biological agents.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.bioconjchem.6b00165.


