Sensing Layer for Ni Detection in Water Created by Immobilization of Bioengineered Flagellar Nanotubes on Gold Surfaces

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ABSTRACT: The environmental monitoring of Ni is targeted at a threshold limit value of 0.34 μM, as set by the World Health Organization. This sensitivity target can usually only be met by time-consuming and expensive laboratory measurements. There is a need for inexpensive, field-applicable methods, even if they are only used for signaling the necessity of a more accurate laboratory investigation. In this work, bioengineered, protein-based sensing layers were developed for Ni detection in water. Two bacterial Ni-binding flagellin variants were fabricated using genetic engineering, and their applicability as Ni-sensitive biochip coatings was tested. Nanotubes of mutant flagellins were built by in vitro polymerization. A large surface density of the nanotubes on the sensor surface was achieved by covalent immobilization chemistry based on a dithiobis(succimidyl propionate) cross-linking method. The formation and density of the sensing layer was monitored and verified by spectroscopic ellipsometry and atomic force microscopy. Cyclic voltammetry (CV) measurements revealed a Ni sensitivity below 1 μM. It was also shown that, even after two months of storage, the used sensors can be regenerated and reused by rinsing in a 10 mM solution of ethylenediaminetetraacetic acid at room temperature.

KEYWORDS: flagellin, water contamination, electronic sensing, spectroscopic ellipsometry

1. INTRODUCTION

Toxicity and other environmental effects of heavy metal traces in natural waters have attracted increasing attention and concern in recent years. Heavy metals exist in natural waters in colloidal, particulate, and/or dissolved phases. Their toxic effects are different, but all of them (e.g., Hg, Cd, Ni, As, Cr, Ti, or Pb) are dangerous, especially due to their potential bioaccumulation.1–3 Heavy metals can enter into a water supply by industrial or consumer waste or due to the acidic rain that breaks down the soil and releases heavy metals into streams, lakes, rivers, and groundwater.4–9 Environmental monitoring generates a steadily growing need for heavy metal ion detection in the concentration range of ppm or below. Requirements for this analytical task usually can be met by applying methods such as atomic absorption spectrometry (AAS), atomic fluorescence spectroscopy (AFS), and inductively coupled plasma mass spectroscopy (ICP-MS).10 However, these methods need laboratory facilities (expensive hardware and trained staff) and they are time-consuming and costly. There is a well-established need for field-applicable methods and equipment using disposable sensors.11

Efforts for constructing and manufacturing portable heavy metal sensors have been increasing in the past years, applying a wide range of concepts, including optical- (e.g., colorimetric,11 fluorescent,12,13 Raman,14 or plasmon-enhanced15 methods), electrochemical-10,16 and nanomaterial-based realizations.14 Each method has advantages and disadvantages depending on the requirements and target elements. Many elements have been investigated, including Hg,17,18 Cu,19 Ni,20 Pb,16 and Fe,21 with some approaches being sensitive but requiring complex preparation17,18,20 and with others utilizing robust detection schemes and portability.22,23 In this work, using genetic engineering, our aim was to develop a novel type of heavy-metal-sensing layer, having good regeneration capability, stability, and flexibility, due to the modification potential of the
surface of the applied protein nanotube according to the target element. By describing this method, a framework is demonstrated in which there are many opportunities for element-optimized detections.

Electrochemical analytical methods are versatile tools for the measurement of heavy metals, due to the fact that most chemical processes related to their compounds involve electron transfer. Among the available electrochemical techniques, cyclic voltammetry (CV) is prominent, with its advantages of simplicity, sensitivity, speed, and low costs, which result in a wide range of applications.26 Since heavy metal contamination must be detected in the ppm range, in order to enhance sensitivity, the application of nanoparticles or other selective binding agents is strongly demanded on the electrode surface.

In this work, we fabricated Ni-sensitive layers from bioengineered bacterial flagellar filaments. The flagellar filament is a natural protein nanotube, which is formed by self-assembly from thousands of flagellin (FliC) subunits.25 Flagellin is an excellent object for protein engineering. Its terminal regions are highly conserved and required for filament formation, in contrast to the variable middle portion that forms the D3 domain exposed on the surface of the filaments. The D3 domain can be modified or replaced by heterologous segments, enabling the introduction of specific functionalities into the host molecule.26−29 These flagellin variants reengineered in the D3 region typically retain their polymerization ability and can be used to make functionalized protein nanotubes in vitro. They can be readily produced by bacteria because they form filaments on the cell surface, which can be removed and purified easily.

Previous studies demonstrated that binding proteins, enzymes, or reporter proteins can be inserted into the variable central portion of the flagellin protein, resulting in a plethora of building blocks applicable for functionalized nanotube formation.26−29 Flagellin-based fusion proteins with catalytic and molecular recognition functionalities can be assembled to form mixed or block copolymers in a rationally designed manner, opening new horizons for applications in medical diagnostics, environmental monitoring, or bionanotechnology.

In this work, we aimed at creating Ni-binding flagellin variants that are capable of forming nanotubes displaying thousands of regularly arranged binding sites on their surface. Two approaches were followed. In the first approach, the D3 domain of flagellin was engineered to create a metal-binding site by introducing His residues at appropriate positions. Alternatively, the D3 domain was removed and replaced by an oligopeptide with a known Ni-binding ability. These flagellin constructs were produced by bacteria, easily obtained from the surface of cells by mechanical treatment and centrifugation, and were further purified through depolymerization/polymerization steps. Their metal-binding properties were characterized by isothermal titration calorimetry (ITC), and the thermal stability of flagellar nanotubes built from the mutant flagellins was also assessed. Filaments with Ni-binding ability were deposited by thiol-based surface chemistry as a thin layer on the working electrode of an electrochemical microsensor specially designed for this purpose. The immobilization of the developed metal-binding filaments was followed by a recently developed in situ optical method30,31 and the prepared filament layers were also characterized using atomic force microscopy (AFM). To assess the sensitivity, detection limit, reproducibility, and regenerability, the deposited filament layers were electrochemically tested in a buffer solution (pH 7.4) by CV.

This paper demonstrates that mutant flagellar filaments have the potential to form protein-based Ni-sensing layers on a gold substrate, which can be employed in low-cost portable electrochemical biosensors to fulfill the requirements for sensing Ni in water at concentrations as low as the World Health Organization (WHO) health threshold limit (maximum allowable concentration).

2. EXPERIMENTAL SECTION

2.1. Cloning, Expression, and Purification of Mutant Flagellin Variants. For the construction of Ni-binding flagellin mutants, a pKOT-based plasmid containing the gene of a D3-deleted variant of flagellin served as a cloning vector.32 Within this plasmid, the D3-domain coding region of flagellin was replaced with a short gene cassette, including the recognition sites of XhoI, AgeI, XmaI, and SacI (EC: 3.1.21.4) restriction enzymes.33,34

The 4H_D3 variant was constructed by the mutation of four amino acids within the D3 domain, residues 190−284 of wild-type Salmonella typhimurium flagellin, to histidine. The 4H_D3 domain coding sequence containing the mutations L209H−V235H−K241H−S264H, where the numbers correspond to the position of the amino acid in wild-type flagellin, and bearing the XhoI and SacI restriction enzyme recognition sequences on its 5′ and 3′ end, respectively, was synthesized and cloned into a pUC57 plasmid by GenScript (Piscataway, NJ).

Due to the length of the HG12 motif (HGGMGHHGGGHHG), complementary coding sequences were designed with the appropriate overhangs generated by the digestion of XhoI and SacI restriction sites, attached to both ends (forward, 5′-TCGAG CAC GGT GGT GTT CAC GGT CAT GGC GCC GCC CAT GGC GAGCT-3′; reverse, 5′-GCC ATG GCC GCC GCC ATG ACC GTG ACC ACC ACC GTG C-3′) and hybridized using oligo annealing techniques.

Both fragments (HG12 and 4H_D3) were inserted into the gene of a D3-deleted variant of flagellin between the XhoI and SacI restriction sites using a T4 DNA ligation kit (Thermo Scientific). Escherichia coli TOP10 cells (Invitrogen) were transformed by the resulting plasmid constructs, purified by a plasmid purification kit (Viogene), and checked by digestion and DNA sequencing.

In order to express the flagellin variants, flagellin-deficient SJW2536 strain of Salmonella typhimurium was electroporated by the fusion gene constructs. Purification of the fusion protein variants was carried out as described previously,35 except in order to avoid metal-ion-induced protein aggregation, 3 mM ethylenediaminetetraacetic acid (EDTA) was added to the washing buffer after the first polymerization step.

The protein concentration of samples was determined from spectrophotometric measurements at a wavelength of 280 nm using molar extinction coefficients (ε280 = 17880 M−1 cm−1) for wild-type flagellin and 4H_FliC and ε280 = 10430 M−1 cm−1 for HG12_FliC), calculated from the known aromatic amino acid contents of the molecules using the ProtParam program.36 The purity of protein samples was checked by SDS-PAGE using 12.5% polyacrylamide gels, stained with Coomassie blue R-250 (Merck).

2.2. Formation of Flagellar Nanotubes. In vitro polymerization of the flagellin constructs was performed by inducing nucleation and polymerization by ammonium sulfate (AS).37 The protein solutions of 2−3 mg/mL were prepared in phosphate-buffered saline (PBS), and 4 M AS was added to achieve a final concentration of 0.6 M. The filament formation was judged after at least 3 h or after overnight incubation at room temperature (RT). The filaments were observed by dark-field microscopy with an Olympus BX50 microscope. The size distribution of reconstructed filaments was checked by dynamic light scattering (DLS) using a Zetasizer Nano-ZS (Malvern) instrument.

2.3. Preparation of Sensor Structures. 2.3.1. Microfabrication of Sensor Chips. Si wafer-integrated sensor chips were used for
testing the Ni-binding flagellar filaments. The electrochemical transducer was designed with three electrodes (Ag/Ti—reference electrode, Au/Ti—working electrode, and Pt/Ti—auxiliary electrode), as shown in Figure 1. A 4-channel potentiostat was prepared in which the analyzing solutions were placed independently. The electrochemical sensors detected the targeted species (contaminants), whereas one of the sensors was used as a reference.

2.3.2. Preparation of the Sensitive Flagellar Nanotube Layer. The covalent immobilization of flagellar nanotubes on gold electrodes was carried out using thiol-based surface chemistry. The clean chips were dipped into a dithiobis(succinimidy propionate) (DSP, Sigma-Aldrich) cross-linking agent solution prepared in dimethyl sulfoxide (DMSO from VWR), and they were incubated on a rocking shaker at RT. DSP-activated surfaces were rinsed twice with DMSO, soaked in a DSP solution prepared in dimethyl sulfoxide (DMSO from VWR), and soaked in PBS for 15 min sequential washings in PBS. The prepared filament-coated chips were stored in PBS in closed vials at 4 °C.

Both the concentration (0.04–10 mg/mL) of the DSP solution and the incubation time (5–90 min) were varied to find appropriate conditions for efficient nanotube deposition. The quality of the surface coverage was judged by AFM. On the basis of these experiments, chips for CV measurements were prepared by applying a 10 mg/mL DSP concentration with a 90 min incubation time.

2.4. ITC Studies. The experiments were carried out with a VP-ITC titration calorimeter (Malvern). The measurements were performed in 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) solution containing 150 mM NaCl (pH 7.0) at 25 °C. The protein samples were extensively dialyzed against the buffer at 4 °C. All solutions were thoroughly degassed before use by stirring under vacuum. The Ni-binding flagellin variants were loaded into the calorimetric cell, and 10 μL portions of the concentrated NiSO4 solution were injected. To measure the heats of dilution, two blank titrations were performed: one with the injection of NiSO4 solution into the buffer and another with the injection of buffer into the flagellin solution. The averaged heats of dilution were subtracted from the main experiment. The calorimetric data were analyzed using MicroCal Origin software, fitting them to a single binding site model.

2.5. Circular Dichroism Measurements. The temperature-induced unfolding of flagellar nanotubes was monitored by far-UV circular dichroism (CD) in a rectangular quartz cell with a path length of 0.1 cm at 222 nm in 10 mM PBS (pH 7.4) over the temperature range of 20–70 °C with a heating rate of 60 °C/h. Measurements were performed with a Jasco J-1100 spectropolarimeter equipped with a Peltier thermostated cell holder.

2.6. Plasmon-Enhanced Ellipsometry Measurements. In order to verify the effect of DSP coating on the formation of a dense flagellar nanotube layer on the gold surface, in situ spectroscopic ellipsometry (SE) measurements were carried out. SE is an optical method with outstanding sensitivity to the thickness and dielectric properties of thin layer structures. By utilizing its high sensitivity, SE has been long used as an excellent tool for following biorelated processes.

In this study, a Woollam M2000 rotating compensator spectroscopic ellipsometer was used with a homemade Kretschmann–Raether flow cell (Figure 2) for internal reflection surface plasmon-enhanced ellipsometry (SPR-SE) that utilizes the surface plasmon resonance (SPR) effect. SPR is one of the most sensitive optical methods in the field of biorelated research, and its combination with SE provides further advantages, such as the possibility of constructing a complex optical model and utilizing the phase information.

The substrate was a glass (BK7) hemicylinder that enabled focusing without the distortion of the measured ellipsometric angles. A glass plate covered by a 40 nm gold layer was attached to the bottom of the hemicylinder, with the gold layer facing the liquid. The surface of the gold layer was carefully cleaned using a UV/ozone cleaner (Microvacuum Ltd., Hungary), followed by rinsing with DI water and flowing nitrogen gas. Subsequently, half of the slide was immersed into a DSP solution for 90 min, while the other half stayed intact (reference channel). This sample preparation method made the

Figure 1. Sensor chip and electrochemical setting with three metal electrodes on a Si substrate. Note that the three electrodes of the conventional arrangement are integrated on a silicon chip using planar semiconductor technology.

Figure 2. (A) Rendered visualization of the ellipsometric Kretschmann–Raether configuration for the internal reflection in situ measurements. (B) Location of the cell on the mapping stage of a commercial ellipsometer. The inset in the bottom right shows the structure, layers, and materials in the cell. “BK7” is the glass hemisphere on the top, which can also be seen on the photograph.
investigation of the impact of DSP on the filament layer formation significantly more reliable, since the measuring and reference channels were located on the same substrate (only one sample was required for two measurements); thus, the quality of the surface or any unintended temperature fluctuations in the flow cell were compensated by the reference channel and had no effect on the result.

For the in situ SPR-SE measurement, a focused light beam was used with a spot size of 0.9 mm × 0.3 mm. The location of the spot was repeatedly changed between the two points, with one corresponding to a position covered with the DSP layer. This method is called two-channel spectroscopic ellipsometry (here, “channel” refers to the measurement channels).30 The SPR-SE measurement was carried out in a wide wavelength range of 400–1690 nm and at the optimal angle of incidence that had been carefully chosen prior to the adsorption process. The angle of incidence was held constant during the measurement.

The baseline of the measurement was provided by a 10 mM PBS solution that was delivered to the surface for 10 min by the help of a peristaltic pump. Subsequently, a flagellar filament solution of 0.5 mg/mL was circulated for 120 min, which corresponds to the incubation time of the chips made for the voltammetric measurements.

For the data evaluation, a complex optical model was constructed, since an SE measurement usually only provides the amplitude and phase differences of the p- and s-polarized lights, referred to as Ψ and Δ, respectively. The optical model consists of all dielectric functions and thicknesses of the layers in the structure, including different corrections as well, such as those that correct the effect caused by the focusing lenses or by the inner stress of the BK7 glass prism. The optical properties of the PBS solution were provided by the Sellmeier equation:

\[
n_a(\lambda) = \sqrt{\frac{\varepsilon_\infty + \frac{A_{\varepsilon}}{\lambda^2} - \varepsilon_S}{1 - \frac{B_S}{\lambda^2}}}
\]

(1)

where \(\lambda\) and \(n_a(\lambda)\) are the illumination wavelength and the refractive index spectrum of PBS, respectively. Additionally, \(\varepsilon_\infty, A_{\varepsilon}, B_S,\) and \(\varepsilon_S\) denote the index offset, the amplitude, the center energy, and the position of a pole in the infrared region, respectively. The filament layer is described by using the Cauchy equation of

\[
n_a(\lambda) = A_C + \frac{B_C}{\lambda^2} + \frac{C_C^4}{\lambda^4}
\]

(2)

with initial parameters of \(A_C = 1.45, B_C = 0.01,\) and \(C_C = 0.00.\) The parameter \(A_C\) was fitted during the data evaluation, and only the thickness of the layer was fixed at 40 nm. The amount of adsorbed molecules was then calculated by de Feijter’s equation,42 which provides the mass density of the adsorbed molecules:

\[
\Gamma = \frac{d_i(n_{a,C} - n_s)}{dn/dc}
\]

(3)

where \(d_i\) is the thickness of the adsorbed protein layer and \(n_{a,C}\) and \(n_s\) are the refractive indices of the adsorbed protein layer and the ambient solution at a wavelength of 632.8 nm, respectively. Additionally, \(dn/dc\) is the refractive index increment of the protein for which a value of 0.18 mL/g was used.35

2.7. AFM Measurements. The AFM measurements were performed with a SmartSPM 1000 instrument manufactured by AIST-NT (Novato, US). The instrument was used in the tapping mode, scanning an area of 5 μm by 5 μm. The AFM images were processed using data leveling, background subtraction, and false color mapping with Gwyddion software.35

2.8. CV Measurements. The CV characterization was made using a Gamry Instruments ESA410 electrochemical signal analyzer. The measurements were performed in a pH 7.0 HEPES solution using the basic three electrode arrangement (working + Pt counter + saturated calomel reference electrodes). Si-based sensor chips were used as working electrodes. The typical measurement range was –0.6 V to 0.6 V. The conductivity of the buffer solution was enhanced by adding 100 mM NaCl. After testing the CV characteristics in the Ni-free solution, variable amounts of a NiSO₄ solution were added into the measurement cell in order to study the effect of the Ni(II) concentration on the CV curves.

2.9. Sample Regeneration and Storage. In order to examine the stability and potential reuse of the sensing structure, the following sequence was applied: (i) After acquiring the CV curve in the 5 μM Ni²⁺-containing solution, the flagellin-covered chips were rinsed in DI water and were stored at 4 °C under PBS solution. (ii) After 2 months of storage, the chips were rinsed first in a 10 mM EDTA solution and then in DI water. (iii) The CV measurement procedure was repeated on the samples using 0, 1, 2, and 5 μM Ni²⁺ solutions in HEPES buffer.

3. RESULTS AND DISCUSSION

3.1. Development of Ni/Metal-Binding Flagellin Variants. We aimed at developing flagellin variants possessing Ni-binding ability for biosensing applications. Natural Ni-binding proteins typically contain several properly positioned His residues to coordinate the binding of metal ions.44,45 We aimed at creating a Ni-binding site within the D3 domain using molecular modeling and genetic engineering techniques. On the basis of the known atomic structure of Salmonella flagellin (PDB: 1UCU), computer modeling with Chimera46 helped us to identify amino acid residues L209, V235, K241, and S264, which were near to each other in an appropriate spatial arrangement and orientation to promote the creation of a Ni-binding site by replacing their original side chains for the imidazole side chain of histidine (Figure 3A). The gene segment coding for this mutant D3 domain (4H_D3) was synthesized by GenScript and was ligated into a pKOT-based plasmid containing the gene of a D3-deleted flagellin.

![Figure 3. Construction of Ni-binding flagellin variants. (A) The hypervariable D3 domain of Salmonella flagellin was replaced by the HG12 (HGGGHGHHGGHG) Ni-binding motif. (B) The Cα backbone trace of the D3 domain of flagellin (PDB code: 1UCU) is displayed, showing the residues mutated for histidine. Molecular modeling was performed with Chimera.](https://doi.org/10.1021/acsbiomaterials.0c00280)

In addition to engineering D3 to furnish it with Ni-binding ability, we tried another approach in which the whole D3 domain was replaced with an oligopeptide segment of known Ni-binding affinity (Figure 3B). The histidine-rich HG12 dodecapeptide (HGGGHGHHGGHG) was reported to bind copper and nickel with a high affinity.47 The DNA segment coding for the HG12 oligopeptide was synthesized (IDT), annealed, and ligated into the pKOT-based plasmid containing the D3-deleted mutant FliC gene between XhoI and SacI restriction sites.

When the 4H_FliC and HG12_FliC plasmid constructs were introduced into the flagellin-deficient nonmotile SJW2536 Salmonella strain, they were expressed at a high

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level. The restored swimming ability of the host bacteria (verified by dark-field microscopy) suggested that the fusion proteins preserved their filament-forming ability.

For both HG12_FliC and 4H_FliC variants, the amounts of subunits built into filaments in vivo were studied by gel electrophoresis. The mutant cells were collected by centrifugation, suspended in 20 mM Tris buffer (pH 7.8) containing 150 mM NaCl, and heated to 70 °C to disassemble flagellar filaments, and the depolymerized fractions were analyzed by SDS-PAGE. We observed strong bands at the expected positions for the depolymerized HG12_FliC and 4H_FliC (not shown), but a comparison of their band intensities to the secreted fractions suggested that a significant amount of the flagellin-based fusion proteins was transported into the culture medium in a monomeric form. Transmission electron microscopy studies demonstrated that HG12_FliC filaments preferentially exhibited a straight polymorphic form and were 4–6 μm long, while 4H_FliC filaments had a normal helical waveform and were typically 5–8 μm long (Figure 4).

3.2. Characterization of Metal/Ni-Binding Properties of the Flagellin Variants. The Ni-binding properties of the 4H_FliC and HG12_FliC constructs in their monomeric state were characterized and compared by ITC measurements. Experiments were done done in HEPES buffer because this buffer does not form complexes with Ni ions; therefore, it is ideal for Ni-binding studies. The Ni-binding flagellin variants were loaded into the calorimetric cell, and 10 or 20 μL portions of concentrated NiSO₄ were injected. The measured data were analyzed by a one-binding-site model. As shown in Figure 5, both variants bound Ni with a high affinity. In control experiments, nickel binding to wild-type flagellin was also investigated, but no interaction was observed. The measured data could be fitted well with the applied model, with the binding parameters summarized in Table 1. The stoichiometry of the interaction was typically lower than 1, indicating a slight aggregation of the samples or indicating uncertainties in sample concentrations. While HG12_FliC bound Ni²⁺ with a dissociation constant of 15.3 μM, the 4H_FliC variant exhibited a significantly higher binding affinity, characterized by a K₆ of 1.4 μM. The values obtained for ΔH and ΔS are apparent and include contributions not only from Ni²⁺ binding but also from associated events such as the deprotonation of the histidines and the consequent change in the buffer ionization state. Nevertheless, HG12_FliC showed a substantially smaller heat effect and entropy change than 4H_FliC.

Preliminary experiments were also performed to check the binding affinity of the variants for other heavy metals like Co(II), Cd(II), and Cu(II). Both the 4H_FliC and HG12_FliC variants interacted strongly with Cu²⁺, with a dissociation constant in the submicromolar range. 4H_FliC exhibited an interaction with Co²⁺ and Cd²⁺, with a 2–3 order of magnitude weaker K₆. HG12_FliC did not show a significant interaction with Co²⁺, while it bound Cd²⁺ with a 0.3 mM dissociation constant.

In summary, our results demonstrate that 4H_FliC shows a significantly higher affinity for Ni²⁺ than HG12_FliC. In subsequent experiments, 4H_FliC was used to fabricate capture layers for nickel detection.

3.3. In Vitro Nanotube Formation. To check the in vitro polymerization ability of the purified 4H_FliC fusion protein, filament formation was induced by ammonium sulfate (AS) precipitation. 4H_FliC readily polymerized into flagellar nanotubes upon the addition of 4 M AS to achieve a 0.6 M final concentration. The average length of reconstructed filaments was about 3–500 nm, as judged by DLS experiments. Filaments were stable in PBS (pH 7.4) after the removal of the AS precipitant. The structural stability of reconstructed filaments was investigated via CD spectroscopy by monitoring their temperature-induced disassembly process (Figure 6). 4H_FliC filaments exhibited a similar thermal stability to filaments formed from wild-type flagellin, indicating that this variant fully preserved its filament-forming ability. The depolymerization and unfolding of subunits occurred around 49.4 °C, as shown by the loss of the α-helical CD signal followed at 222 nm.

Reconstituted flagellar filaments are known to be highly resistant against proteolytic degradation and can be stored at RT for a long period of time. Filaments built from 4H_FliC subunits were used to create sensing layers of biochips for voltammetric measurements.

3.4. Flagellar Nanotube Deposition on Gold Electrodes. To fabricate capture layers, 4H_FliC filaments were cross-linked by DSP to the surface of gold electrodes. There are several methods suitable for immobilizing flagellar nanotubes on various types of surfaces.48,49 We chose DSP-based covalent surface chemistry for use as a simple but effective method for producing stable surface-filament attachments that can potentially sustain complex media when the sensor is applied in future measurements. The filament layers were characterized using SPR-SE and AFM techniques. The effect of the applied DSP concentration and incubation time was investigated to find appropriate conditions for the efficient nanotube deposition. Some representative AFM images are shown in Figure 7. Our experiments revealed that applying a DSP solution at a concentration higher than 4 mg/mL for at least 30 min results in a sufficiently covered surface layer (Figure 7C). According to Figure 7A, the surface density of filaments was significantly lower in the case of noncovalent adsorption, even compared to the lower-concentration DSP experiment shown in Figure 7B. The filament layer formation was monitored in real-time by in situ SPR-SE techniques, and the obtained data were evaluated using the analysis method described in section 2.6. In Figure 8, the refractive index change is shown for a wavelength of 633 nm as a function of time, induced by the formed filament layer. Our observations are in good agreement with the AFM results and we can conclude that the effect of DSP functionalization is clearly significant. The amount of adsorbed molecules is more than double on the functionalized surface than that on the bare gold site (391 and 184 ng/cm²).
respectively). Note that the adsorptions plotted in Figure 8 were performed in the same process simultaneously by moving the focused spot of the ellipsometer back and forth between the functionalized and nonfunctionalized surfaces. This result demonstrates that the DSP functionalization step has a vital role in the preparation of a dense sensing layer required for detecting Ni molecules in water at low concentrations.

3.5. Electronic Measurements Using Flagellar Nanotube Sensing Layers for Ni Detection. The nickel-detecting capability of the CV limit of detection setting was

![Figure 5](https://dx.doi.org/10.1021/acsbiomaterials.0c00280) & ![Figure 6](https://dx.doi.org/10.1021/acsbiomaterials.0c00280)
tested by the measurement of buffer solutions containing NiSO₄ at different concentrations. As a negative control, a golden electrode covered by “wild-type” flagellar nanotubes (i.e., without Ni-binding sites) was used in the 0−5 μM range. The set of measurements in Figure 9A clearly demonstrates that the addition of the metal ion has a small effect on the CV curves.

Figure 9B shows CV curves taken on a freshly prepared sensor covered by flagellar filaments built from 4H_FliC flagellin. By comparison of the curve obtained from the measurement of the Ni-free solution to other curves belonging to concentrations of 1, 2, 5, 20, 50, and 100 μM Ni²⁺, it is clearly visible that a cathodic peak appears in the potential range around −250 mV, and the peak intensity monotonously increases with the Ni²⁺ concentration, up to saturation. This set of peaks can be associated with the binding and subsequent cathodic reduction of Ni²⁺ from the solution. A comparison of Figure 9A and 9B proves that the effect is due to the presence of Ni(II) on (A) wild-type flagellar nanotubes ([Ni²⁺] = 0−5 μM), (B) Ni-binding flagellar nanotubes ([Ni²⁺] = 0−100 μM), and (C) Ni-binding nanotubes after storage and regeneration ([Ni²⁺] = 0−100 μM). The numbers in the legends show concentrations in μM.
of the Ni-binding site created within the 4H_FliC flagellin. Moreover, a significant and reproducible change of the CV profile for the 1 μM Ni²⁺ solution is in good agreement with the micromolar dissociation constant measured for the 4H_FliC−Ni²⁺ interaction by ITC (section 3.2).

The experimentally found lower limit of detection is in good agreement with the WHO threshold limit value for drinking water. This can be considered advantageous since this allows to preselect subsets of samples taken from natural waters. Only these selected ones need to be sent for further expensive and time-consuming analysis.

As noted in section 3.2, the applied flagellin variant, 4H_FliC, also has a strong binding affinity for Cu²⁺, while it shows a very weak interaction with Co²⁺ and Cd²⁺. These results suggest that the developed sensing layer may be applicable for the detection of both copper and nickel contamination. Further studies are needed to characterize in detail the selectivity of the 4H_FliC flagellin-based sensing layer for heavy metal monitoring in the relevant concentration range specified by the health threshold limit value set by the WHO.

In order to determine the stability and reproducibility of the nickel detection ability of the 4H_FliC filament-functionalized electrodes, the CV measurements were repeated on the samples. The storage time between the measurements was two months, during which the chips were stored at 4 °C in PBS buffer. Prior to the second measurement, the chips were regenerated (i.e., they were rinsed in a 10 mM EDTA solution and thoroughly rinsed in DI water). EDTA treatment was intended to completely remove possible Ni contamination on the chips accumulated throughout the first set of measurements.

Figure 9C displays the CV curves measured after storage and regeneration. The measurements clearly demonstrate that the cathodic peak in the potential range at around −250 mV is still apparent and increasing with the applied Ni²⁺ concentration. The saturation tendency also remains observable despite the fact that the Ni-binding sites of the filament layers were previously saturated during the first test. Therefore, this set of experiments confirms that the flagellin layers (i) can be regenerated by rinsing the functionalized electrode in a 10 mM EDTA solution at RT, (ii) can retain their Ni²⁺-binding ability for at least 2 months at 4 °C, and (iii) can be reused as Ni-monitoring sensor layers in the 0–100 μM Ni²⁺ concentration range.

The evaluation of measurements taken on fresh and regenerated samples using the 4H_FliC filament-covered electrode is presented in Figure 10. A linear baseline was determined in the potential range of −150 to −350 mV by linear interpolation of the currents measured at −150 and −350 mV for every single CV curve, and the peak area under this baseline was calculated. Peak areas were expressed in terms of the percentage of the value of the peak belonging to 100 μM and were plotted against the Ni²⁺ concentration (Figure 10).

Fitted exponential trendlines show that increasing the Ni²⁺ concentration resulted in the saturation of the curves at around 50 μM, both in the case of fresh and regenerated chips. This saturation provides further proof of the protein-based mechanism of the sensing process, since the filament layer has a limited number of binding sites. The presence of the concentration-sensitive range (0–50 μM) suggests that this method has the potential to not only detect Ni²⁺ in natural waters but, at the same time, determine its concentration in a broader range around the maximum allowable concentration. A comparison of the fitted trendlines for the fresh and regenerated samples (red and blue lines in Figure 10, respectively) shows that the difference between the two curves does not exceed 10% of their actual values at any point within the relevant concentration range.

4. CONCLUSIONS

In this study, we developed a Ni-sensitive biosensor coating made of protein-engineered flagellar nanotubes. Utilizing the polymerization ability of flagellin variants, we have provided a method for the inexpensive and rapid production of large amounts of protein. The flagellar nanotubes were built from mutant flagellins by in vitro polymerization, and their applicability to form a stable sensing layer for Ni detection was demonstrated by CV measurements. The chips were covered with filaments using a covalent immobilization process that applied thiol-based surface chemistry. The sensing capabilities were tested in a buffer solution (pH 7.0) by CV measurements, showing that the sensitivity allows the detection of Ni²⁺ ions at concentrations as low as 1 μM, which is close to the WHO health threshold limit value of 20 μg/L (0.34 μM) specified for water contamination. On the basis of the CV results obtained after EDTA treatment, we proved that the developed platform has distinguished regeneration properties without showing any detectable loss in its Ni-sensing ability.

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Figure 10. Peak integral (area) values for first-use (blue circles) and after-regeneration (red triangles) samples, calculated using the method described in the text, as a function of the Ni²⁺ concentration. The solid lines show exponential fits to the measured data points.
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