

The Citrate Carrier CitS Probed by Single-Molecule Fluorescence Spectroscopy

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ABSTRACT A prominent region of the Na⁺-dependent citrate carrier (CitS) from *Klebsiella pneumoniae* is the highly conserved loop X-XI, which contains a putative citrate binding site. To monitor potential conformational changes within this region by single-molecule fluorescence spectroscopy, the target cysteines C398 and C414 of the single-Cys mutants (CitS-sC398, CitS-sC414) were selectively labeled with the thiol-reactive fluorophores AlexaFluor 546/568 C₅ maleimide (AF₅₄₆, AF₅₆₈). While both single-cysteine mutants were catalytically active citrate carriers, labeling with the fluorophore was only tolerated at C398. Upon citrate addition to the functional protein fluorophore conjugate CitS-sC398-AF₅₄₆, complete fluorescence quenching of the majority of molecules was observed, indicating a citrate-induced conformational change of the fluorophore-containing domain of CitS. This quenching was specific for the physiological substrate citrate and therefore most likely reflecting a conformational change in the citrate transport mechanism. Single-molecule studies with dual-labeled CitS-sC398-AF_{546/568} and dual-color detection provided strong evidence for a homodimeric association of CitS.

INTRODUCTION

The Na⁺-dependent citrate carrier (CitS) of *Klebsiella pneumoniae* is expressed during anaerobic growth on citrate and enables this enterobacterial species to utilize citrate as sole carbon and energy source via the citrate fermentation pathway (Bott et al., 1995; Dimroth and Thomer, 1986). The *citS* gene encodes a highly hydrophobic integral membrane protein with a postulated topology of 11 transmembrane helices connected by hydrophilic loops (van Geest et al., 1999). CitS catalyzes the highly specific, electroneutral uptake of Hcitrate²⁻ in cotransport with two Na⁺ ions (Lolkema et al., 1994; Pos and Dimroth, 1996). Citrate counterflow experiments revealed that CitS features two binding sites for citrate, an external high-affinity site and an internal low-affinity site. Active citrate uptake is driven by a chemical ion gradient ($\Delta p\text{Na}^+/\Delta\text{pH}$), which is in part generated by the oxaloacetate decarboxylase Na⁺ pump, a key enzyme of citrate fermentation in *K. pneumoniae* (Dimroth, 1997). According to multiple-sequence alignments with closely related members of the 2-hydroxy-carboxylate transporter family (TC 2.A.24) (Saier, 2000) the primary structure of CitS shares 26% and 28% identical residues with CitP from *Leuconostoc mesenteroides* and MleP from *Lactococcus lactis*, respectively, which function as Hcitrate²⁻/lactate⁻ and malate²⁻/lactate⁻ antiporter, respectively (Bandell et al., 1997; Marty-Teyssset et al., 1995). In particular, the C-terminal part including loop X-XI of these three transporters is highly conserved. Evidence

indicates that this region is involved in substrate binding and hence of particular importance for the function of these transporters (Bandell and Lolkema, 2000a; Bandell and Lolkema, 2000b; Pos et al., 1998).

Single-molecule spectroscopy has developed to a powerful tool to investigate biological systems (Weiss, 1999). This method has distinct advantages over conventional ensemble spectrophotometric studies, e.g., much higher sensitivity, the possibility to discriminate between static and dynamic quenching in a single experiment, or direct observation of binding events by colocalization studies (Schutz et al., 1998). Inherently single-molecule experiments are free from unwanted interactions between proteins due to high-concentration artifacts. Recently, fluorescence quenching has been used to study protein folding (Zhuang et al., 2000) and conformational heterogeneities of membrane proteins (Peleg et al., 2001). We now used fluorescence quenching of single molecules to get further insights into the role of loop X-XI in the citrate translocation mechanism of CitS. Our results show that single-molecule fluorescence studies can be used to examine functional conformational changes in membrane proteins, specifically in transporters. In a second part, single-molecule studies with dual-color labeling allowed us to report on the oligomerization state of CitS.

MATERIALS AND METHODS

Bacterial strains and plasmids

Escherichia coli DH5 α (Bethesda Research Laboratories, Frederick, MD) and plasmid pFXa3 (Kästner et al., 2000) were routinely used for cloning procedures and mutagenesis. *E. coli* C43(DE3) (Miroux and Walker, 1996) and plasmids derived from pCitS_{H3} (Pos and Dimroth, 1996) were used for high-level expression of the *citS* mutants. *E. coli* BL21(DE3) (Studier and Moffatt, 1986) was used as host for whole-cell transport assays. To test the ability of cells to grow on citrate as sole carbon and energy source Simmons' citrate agar was used (Simmons, 1926).

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Construction of single-Cys mutants

The CitS single-Cys mutants were constructed by polymerase chain reaction (PCR) mutagenesis using the two-step method described by Weiner et al. (1994). Four of the five native cysteines (C278, C317, C347, C398, C414) were subsequently replaced with serines resulting in the two CitS single-Cys mutants CitS-sC398 and CitS-sC414. For each mutation, two complementary mutagenic oligonucleotide primers were synthesized containing the desired mutation (indicated by boldface type) and a unique restriction site (underlined and specified in parentheses) to facilitate mutant identification (only the forward primers are shown):

C278S: GGTGCTGTCCACTACA**AGCTT**CCTGCTGGCC (*HindIII*)
 C317S: CCTCCGGGCT**GAGCTC**ACCGGAAATC (*SacI*)
 C347S: GGGGTCGGCGT**CAGCTACACAGATCT**ACAGGAAATTA-
 TC (*BglII*)
 C398S: CACTGCAGGCCTGT**CCATGGC**CAACCGC (*NcoI*)
 C414S: CCTGGAAGTTCTCT**CTGCTAGCA**ACCGTATGAACCTTA-
 TCTC (*NheI*)

The mutagenic PCRs for CitS-sC398 (CitS-sC414) were performed in the following order: C278S → C317S → C347S → C414S (C398S). In addition, a Cys-less mutant was constructed by replacing even the last remaining cysteine residue with serine (CitS-C-less). After each PCR the methylated nonmutated parental DNA was digested with *DpnI* and 2 μ l of the reaction were transformed into *E. coli* DH5 α by electroporation (Holo and Nes, 1989). Mutated DNA was identified by the acquisition of the new restriction site and sequenced by using the dideoxy chain termination method (Sanger et al., 1977) to ensure that no other mutations occurred during the amplification process. For expression and purification the mutagenized *citS* genes in pFXa3 were cloned into pCitS_{H3} comprising an N-terminally attached 10x polyhistidine tag. Therefore the pFXa3 plasmids were digested with *Bam*HI to produce 1.36 kb fragments carrying the mutagenized *citS* gene (*citS**). Then the *Bam*HI fragment of pCitS_{H3} was replaced by the mutagenized *citS* fragments resulting in pCitS_{H3}(sC398), pCitS_{H3}(sC414), and pCitS_{H3}(C-less). Again, the DNA sequence of the ligated fragments was confirmed by sequencing.

Expression and purification of the mutant proteins

For expression and purification of the His-tagged fusion proteins, essentially the same protocol was used as described earlier (Kästner et al., 2000). For the final Ni²⁺ affinity chromatography we used a Poros MC20 column (4.6 × 100 mm, PerSeptive Biosystems, Foster City, CA) operated on a BioCAD workstation (PerSeptive Biosystems, Foster City, CA) with ultraviolet (UV) detection at 280 nm. The Triton X-100 extract of the membrane vesicles was applied to the column preequilibrated with 10 column volumes of buffer C (50 mM potassium phosphate, 100 mM potassium chloride, 0.5 mM imidazole, 10% (v/v) glycerol, 0.05% (w/v) n-dodecyl- β -D-maltoside, pH 7.0). The column was washed with a linear gradient of 0.5–180 mM imidazole in 10 column volumes buffer C followed by 10 column volumes of buffer C containing 180 mM imidazole. The fusion proteins were eluted with buffer C containing 500 mM imidazole. The flow rate was 10 ml/min and 1 ml fractions were collected. The fractions containing the purified protein were pooled and concentrated by ultrafiltration using Centricon YM-30 centrifugal filter devices (Amicon, Danvers, MA). To remove imidazole, the samples (2 ml) were dialyzed twice for 6 h at 4°C in 500 ml buffer C using SpectraPor regenerated cellulose membranes (25 kD MWCO, Spectrum Laboratories, Rancho Dominguez, CA).

Reconstitution and transport experiments

The purified proteins were reconstituted using the freeze/thaw/sonication procedure as described previously (Pos and Dimroth, 1996). For transport

experiments the proteoliposomes (2 μ l) prepared in buffer R (50 mM potassium phosphate, pH 7.5) were diluted into 98 μ l of buffer A (50 mM sodium phosphate, pH 6.0) containing 12 μ M of [1,5-¹⁴C]citrate (180–220 cpm/pmol). The transport assays were terminated according to a time scale (0–120 s) by the addition of 0.9 ml of ice-cold 0.1 M LiCl and rapid filtration through 0.22 μ m filters (GSTF, Millipore, Bedford, MA). The filters were washed once with 1 ml of ice-cold 0.1 M LiCl and inserted into wells of a Picoplate (24-well, Canberra Packard). After the addition of 1 ml Microscint-20 (Canberra Packard, Pangbourne, UK), the entrapped [1,5-¹⁴C]citrate was determined by liquid scintillation counting on a Microplate scintillation counter (TopCount, Canberra Packard, Pangbourne, UK). Initial rates were determined by the same procedure, but the reaction was immediately stopped after 5 s incubation time.

Labeling with AF₅₄₆ and AF₅₆₈

To obtain fluorophore-conjugated CitS, 5 μ l of Alexa Fluor 546 C₅ maleimide (AF₅₄₆, Molecular Probes, Leiden, The Netherlands) stock solution (800 μ M) in 50 mM potassium phosphate buffer (pH 7.0) were added to 100 μ l of purified and concentrated CitS-sC398 or CitS-sC414 (40 μ M) and incubated in a thermomixer (Eppendorf, Hamburg, Germany) for 10 min at 25°C in the dark. The reaction was stopped by passage through two centrifuge columns (1 ml, Sephadex G-25 medium) to separate labeled CitS from unreacted fluorophore. The degree of labeling was calculated by measuring the absorption of the fluorophore protein conjugate using a molar extinction coefficient of 92,000 M⁻¹ cm⁻¹ at 575 nm (Molecular Probes, Leiden, The Netherlands) and applying the Beer-Lambert law. For dual-color labeling a stock solution containing equimolar concentrations (800 μ M) of Alexa Fluor 568 C₅ maleimide (AF₅₆₈, Molecular Probes, Leiden, The Netherlands) and Alexa Fluor 546 C₅ maleimide (AF₅₄₆) was used. The labeled protein was stored under liquid nitrogen.

SDS-PAGE and BN-PAGE analysis

The fluorophore attachment of fluorophore-protein conjugates was verified by SDS polyacrylamide gel electrophoresis using a 15% separating gel (Laemmli and Faure, 1973). The SDS-PAGE gel was first illuminated at 366 nm and then stained with Coomassie Blue G250.

The Blue Native PAGE analysis of wild-type CitS and CitS-sC398 was performed as described by Schägger and Jagow (1991). The purified samples (25 μ g) were applied to a 5–15% gradient polyacrylamide gel with a 4% stacking gel and subjected to electrophoresis at 4°C. Immediately after electrophoresis, the gel was stained with Coomassie Blue G250.

Ni-NTA coating of cover glasses

The glass surface was coated with Ni-NTA using a modified procedure described by Adachi et al. (2000). The cover glasses (24 × 24 mm, Menzel, Braunschweig, Germany) were precleaned by baking at 500°C for 2 h. Successively, the glasses were incubated in sealing solution (2% (v/v) 3-glycidyloxypropyl-trimethoxysilane (Fluka, Buchs, Switzerland), 0.01% (v/v) acetic acid) for 3 h at 90°C; coating solution (2% (w/v) N,N-bis(carboxymethyl)-L-lysine (Fluka, Buchs, Switzerland), 2 mM KHCO₃, pH 10.0) for 16 h at 60°C; and Ni²⁺ solution (10 mM NiSO₄, 5 mM glycine, pH 8.0) for at least 2 h at room temperature. After each coating step the glasses were washed with ultrapure water.

Sample preparation

For each series of measurements aliquots of labeled CitS-sC398 were thawed and diluted to a concentration of 30 nM in buffer F (50 mM potassium phosphate, 100 mM potassium chloride, 0.05% n-dodecyl- β -D-maltoside, pH 7.0). A Ni-NTA coated cover glass was attached to the piezo

scan stage and covered with 60 μl of buffer F. Upon addition of 2 μl His-tagged CitS or fluorescent derivatives (30 nM), the proteins become immobilized. In the case of fluorescence quenching studies, 5 μl of citrate or Na^+ in buffer F in various concentrations was added.

Scanning confocal fluorescence microscopy

The measurements were performed using a home built scanning confocal fluorescence microscope (Prummer et al., 2000). A scheme of the set up is shown in Fig. 1. A pulsed frequency-doubled Nd:YAG laser (Antares; Coherent, Santa Clara, CA) was used to excite the fluorophores. The polarization was controlled by a polarizer, an electrooptical modulator, an analyzer, and a quarter- or a half-wave plate. To achieve a Gaussian beam profile, the light was spatially filtered by an optical fiber. Thereafter, the light was reflected by a dichroic (540DRLP, Omega, Brattleboro, VT) and an ordinary mirror into a high numerical aperture objective lens (Planapo 60 \times , NA 1.4; Nikon, Tokyo, Japan) resulting in a diffraction limited excitation spot (full-width at half maximum \sim 250 nm). A confocal fluorescence image was built up by scanning the sample with the piezo stage pixel-by-pixel through the excitation volume while counting the fluorescence photons at each point.

The fluorescence light emerging from the excitation volume was collected by the objective lens and subsequently filtered by the dichroic mirror and a holographic notch filter (532 nm Notch plus; Kaiser, Ann Arbor, MI). At the polarizing beam splitter the fluorescence light was decomposed in its vertically polarized component, which was detected by a single-photon counting avalanche photo diode (APD 1) (SPCM-AQR-14; Perkin Elmer, Fremont, CA; dark counts less than 100 s^{-1}) and its horizontal component, which was projected onto a dichroic mirror (595DRLP, Omega). The dichroic mirror decomposes the fluorescence light according to its wavelength. Light that had a wavelength shorter than 595 nm was reflected onto APD 2, and light with a longer wavelength than 595 nm passed the dichroic mirror and was detected by APD 3. Artifacts caused by the polarization dependency of the dichroic mirror were avoided by using this polarization resolved detection scheme. The pulses from the three APDs were fed into a time-correlated single-photon counting board (SPC 402, Becker and Hickl, Berlin, Germany), which allowed for the storage of arrival time after pulsed excitation, channel number, and absolute time on a photon-by-photon basis.

From the absolute time when a photon was detected, the intensity was calculated for each channel (I_1 , I_2 , I_3) with variable integration time. The dichroic ratio

$$D = \frac{I_2 - I_3}{I_2 + I_3}$$

depends on the emission spectrum of the fluorophore. It is independent of the emission intensity and was used to identify and distinguish the different fluorophores AF₅₄₆ and AF₅₆₈.

In addition, the polarization

$$P = \frac{I_1 - (I_2 + I_3)}{I_1 + I_2 + I_3}$$

was monitored to observe changes in the orientation of the dye labels.

RESULTS

A prominent region of the Na^+ -dependent citrate carrier CitS and other members of the 2-hydroxycarboxylate (TC 2.A.24) carrier family is the C-terminal part including the hydrophilic loop X-XI and transmembrane segment XI. This region contains numerous conserved residues and has been described to harbor the binding site for the transport substrate. Transporters are generally believed to perform a catalytic cycle, which starts with the binding of the transport substrate(s) from one side of the membrane and succeeds with a conformational change through which the bound substrates gain access to the opposite membrane surface where they dissociate. We reasoned that a conformational change of CitS after citrate binding might be observable by fluorescence quenching of a fluorophore attached to an appropriate site of the molecule.

Single-Cys mutants CitS-sC398 and CitS-sC414

CitS has two endogenous cysteine residues (C398, C414) in the particularly interesting loop X-XI region, which could be used as targets for the binding of the thiol-reactive fluorophores (Fig. 2). However, as five cysteines are present in CitS, single cysteine mutants had first to be created genetically to achieve specific labeling of only a single cysteine residue. With this goal in mind we replaced the five cysteines step-by-step with serine and determined the citrate transport activity of the carrier after each step. Fortunately, all cysteine/serine substitutions were tolerated because the *E. coli* transformants expressing the various *citS* mutants (DH5 α /pFXa3-*citS**) were all able to grow on Simmons' citrate agar as recognized by a color change of the pH indicator bromthymol blue due to alkalization of the medium. All citrate carriers containing cysteine substitutions were confirmed to be functional by [$1,5\text{-}^{14}\text{C}$]citrate uptake into the *E. coli* cells expressing the CitS mutants. For a more quantitative assessment of the citrate transport activities of particularly interesting mutants, the N-terminally His-tagged fusion proteins CitS-sC398, CitS-sC414, and CitS-C-less were expressed, purified, and reconstituted into proteoliposomes.

After applying ΔpNa^+ and ΔpH the kinetics of [$1,5\text{-}^{14}\text{C}$]citrate uptake were determined (Fig. 3) yielding initial

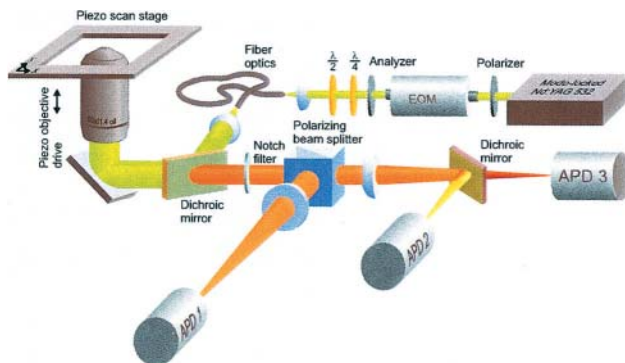


FIGURE 1 Schematic drawing of a scanning confocal fluorescence microscope for imaging and spectroscopy of single molecules in aqueous solution. For the immobilization of His-tagged fusion proteins a Ni-NTA coated cover glass was attached to the piezo scan stage and covered with buffer solution.

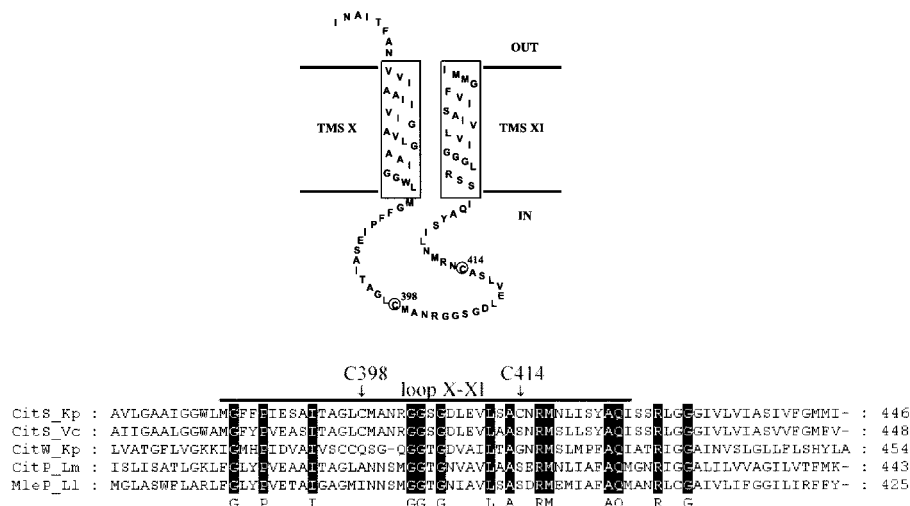


FIGURE 2 Topology model and amino acid sequence alignment of the C-terminal part of CitS from *K. pneumoniae* (*Kp*) with homologs of the 2-hydroxy-carboxylate transporter family: CitS from *Vibrio cholerae* (*Vc*), CitW from *K. pneumoniae* (*Kp*), CitP from *Leuconostoc mesenteroides* (*Lm*), and MleP from *Lactococcus lactis* (*Ll*). Circles and arrows indicate the position of the target cysteines for thiol-reactive labeling of CitS-sC398 or CitS-sC414. The consensus sequence and the loop X-XI region are indicated by black bars. The boxes represent transmembrane segment X and XI.

rates of citrate uptake of $267 \text{ nmol mg}^{-1} \text{ min}^{-1}$, $109 \text{ nmol mg}^{-1} \text{ min}^{-1}$, $123 \text{ nmol mg}^{-1} \text{ min}^{-1}$, or $77 \text{ nmol mg}^{-1} \text{ min}^{-1}$ for wild-type CitS, CitS-sC398, CitS-sC414, or Cys-less CitS, respectively. We conclude therefore that CitS mutants in which four or all five endogenous cysteine residues are replaced by serine are functional citrate carriers with transport activities of $\sim 40\text{--}50\%$ for the single-Cys mutants and $\sim 30\%$ for Cys-less CitS.

Labeling of CitS-sC398 and CitS-sC414 with AF₅₄₆ and AF₅₆₈

The thiol-reactive fluorophores AF₅₄₆ and AF₅₆₈ were the most suitable probes for our spectroscopic studies with the Nd:YAG excitation laser (532 nm) due to their absorption and emission spectra (Haugland, 2000). For site-specific

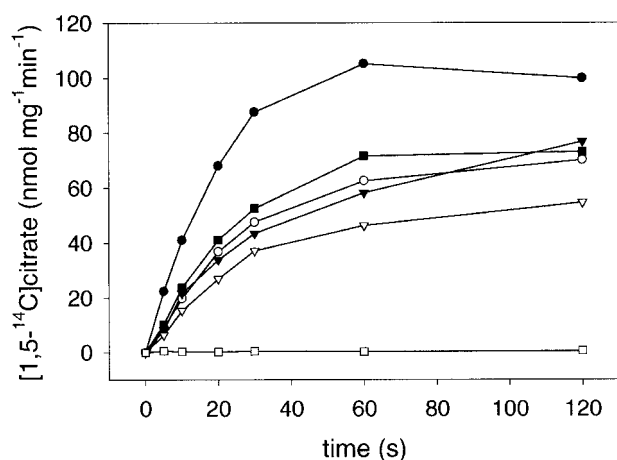


FIGURE 3 Citrate uptake into proteoliposomes containing purified wild-type CitS (●), CitS-sC398 (○), CitS-sC414 (■), CitS-sC398-AF₅₄₆ (▼), CitS-sC414-AF₅₄₆ (□), and Cys-less CitS (▽). The transport assay was initiated by diluting $2 \mu\text{l}$ proteoliposomes into $98 \mu\text{l}$ buffer A containing $12 \mu\text{M}$ of $[1,5\text{-}^{14}\text{C}]$ citrate.

labeling CitS-sC398 and CitS-sC414 were reacted with AF₅₄₆ or an equimolar mixture of AF₅₆₈ and AF₅₄₆ in case of dual-color labeling. To confirm the labeling reaction, the conjugates CitS-sC398-AF₅₄₆ (CitS-sC398¹) and CitS-sC414-AF₅₄₆ (CitS-sC414¹) were subjected to SDS-PAGE and examined by UV illumination and Coomassie staining (Fig. 4).

According to our calculations, the degree of labeling was $\sim 80\%$ for both conjugates. Thus, the region comprising C398 and C414 must be accessible to this rather hydrophilic fluorophore. The effect of the bound fluorescent probe on the activity of the conjugates was again investigated by $[1,5\text{-}^{14}\text{C}]$ citrate uptake studies with CitS-sC398¹ and CitS-sC414¹ reconstituted into proteoliposomes (Fig. 3). The initial transport rate of the conjugate CitS-sC398¹ ($98 \text{ nmol mg}^{-1} \text{ min}^{-1}$) was only slightly reduced compared to the nonmodified protein CitS-sC398 ($109 \text{ nmol mg}^{-1} \text{ min}^{-1}$) and showed almost the same kinetic pattern (Fig. 5). This suggested that the bound fluorophore at position 398 did not affect imperative interactions during citrate translocation or result in partial inactivation. In contrast, with conjugate CitS-sC414¹ a complete loss of transport activity

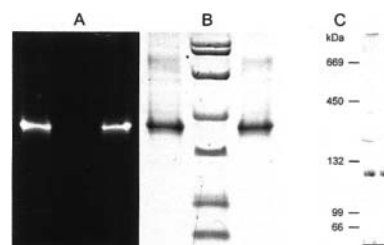


FIGURE 4 (A and B) SDS-PAGE analysis of the protein fluorophore conjugates CitS-sC398¹ (left lanes) and CitS-sC414¹ (right lanes) and a nonlabeled protein marker (center lane). The gel was examined by UV illumination at 366 nm (A) and Coomassie staining (B). (C) BN-PAGE analysis of purified wild-type CitS (left lane) and CitS-sC398 (right lane) as described under Materials and Methods.

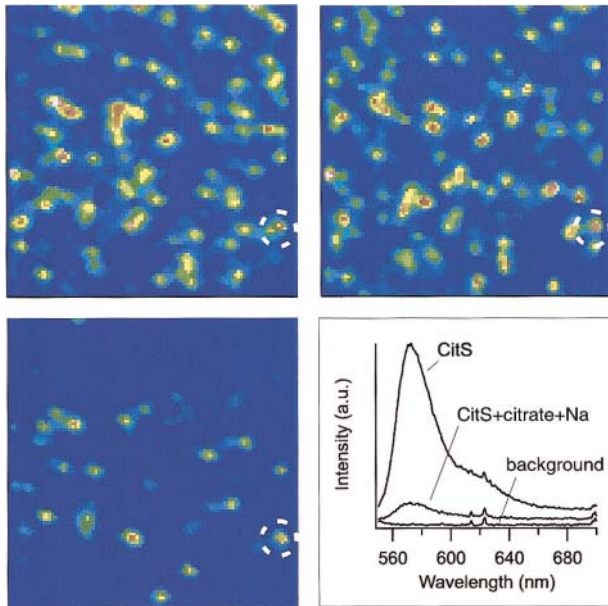


FIGURE 5 Scanning confocal microscopy images of individual CitS-sC398¹ molecules immobilized on Ni-NTA coated cover glass. The first two images (*top row*) show the same frame ($10\ \mu\text{m} \times 10\ \mu\text{m}$) recorded between a time interval of 2 min. The third image (*bottom left*) represents the identical frame upon addition of $250\ \mu\text{M}$ citrate. The dashed circle marks an identical spot in all three images. (*Bottom right*) Ensemble spectra taken under the same conditions. (*From top to bottom*) Pure AF₅₄₆ labeled CitS-sC398¹, after addition of citrate and Na⁺, and background.

was observed. Most likely, the bound probe AF₅₄₆ at position 414 blocks citrate binding or inhibits mandatory interactions during the transport process. Therefore, we decided to concentrate our studies on the functional conjugate CitS-sC398¹.

Single-molecule fluorescence microscopy with CitS-sC398¹

Preliminary experiments with ensemble measurements using the functional conjugate CitS-sC398¹ already indicated that the addition of citrate caused fluorescence quenching of the probe bound to C398 in loop X-XI and a slight spectral blue shift (Fig. 5, *inset*). Because single-molecule fluorescence spectroscopy provides much higher sensitivity and additional spectroscopic information, we decided to use this technique to study the putative conformational changes of the loop X-XI region.

For the single-molecule studies His-tagged CitS-sC398¹ was immobilized on a Ni-NTA coated cover glass and subsequently fluorescence images were recorded with an integration time of 1 ms per pixel. The same frame ($10\ \mu\text{m} \times 10\ \mu\text{m}$) was recorded twice to show the effect of bleaching and additional binding/unbinding of CitS-sC398¹ to the glass surface, leading to vanishing and emerging spots, respectively (Fig. 5). Some of the spots moved laterally and

indicated partial mobility of conjugates but the majority of bright spots did not change significantly. However, the addition of $250\ \mu\text{M}$ citrate caused strong static quenching of AF₅₄₆ at position 398, whereas the addition of $250\ \mu\text{M}$ DL-isocitrate, that served as a control, did not alter the fluorescence emission of the bound probe (Fig. 6).

Because CitS is known to be the most specific transporter among well-characterized members of the 2-hydroxycarboxylate transport protein family, transporting only citrate in uptake mode as well as in exchange mode (Bandell et al., 1997; Pos and Dimroth, 1996), this finding strongly suggests that the interaction of substrate with CitS induced the quenching, not the mere presence of citrate or other 2-hydroxycarboxylates. Also, a change in the local pH can be excluded as the reason for quenching because the fluorescence properties of the Alexa dyes are only very weakly pH-sensitive as stated by the supplier (Haugland, 2000) and

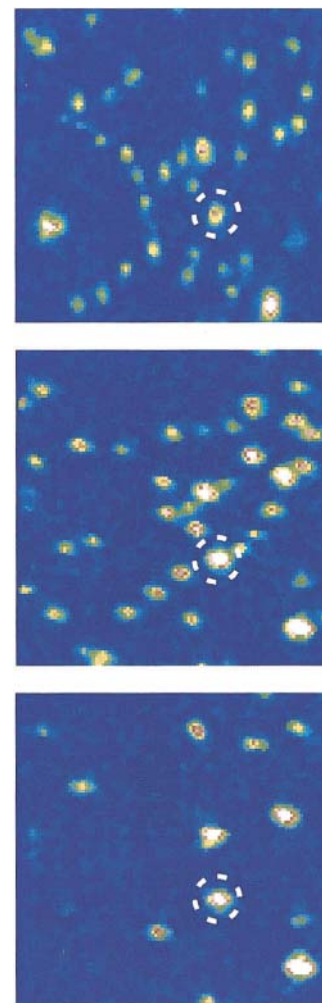


FIGURE 6 Scanning confocal microscopy images of CitS-sC398¹ in the absence of substrate (*top*), in the presence of $250\ \mu\text{M}$ DL-isocitrate (*middle*), and upon addition of $250\ \mu\text{M}$ citrate (*bottom*). All images were recorded consecutively.

confirmed in this study as shown in the top panel of Fig. 8. To exclude direct interactions between citrate and the bound fluorophore, we conducted an additional control experiment by adding citrate in various concentrations to pure AF₅₄₆ at pH 7.2, but this time no fluorescence quenching was observed (Fig. 8).

The rather high intensity of the unquenched spots suggests that these spots correspond to inactive protein aggregates or nonspecifically bound protein that might be partly unfolded. We observed an increase of large aggregates when CitS-sC398¹ was subjected to room temperature for more than 30 min. Single-molecule experiments offer the unique opportunity to disregard these aggregates from further analysis. Only isolated, diffraction limited spots are taken into account. Within these spots the number of fluorescent molecules can be counted during photobleaching. To test if the presence of the physiological coupling ion Na⁺ is

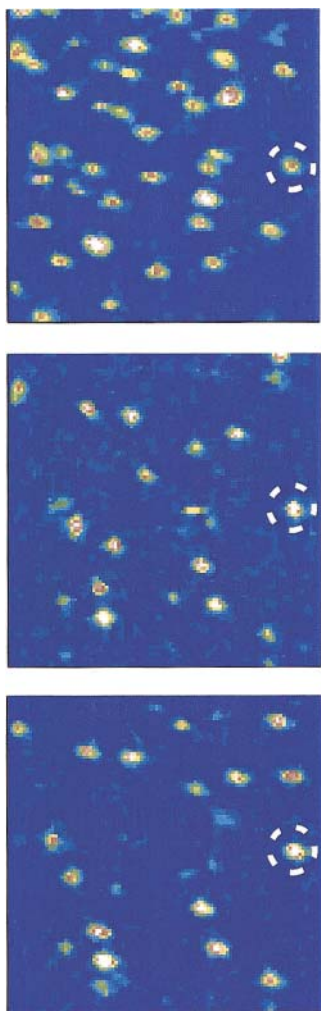


FIGURE 7 Scanning confocal microscopy images of CitS-sC398¹ in the absence of substrate (*top*), in the presence of 250 μ M citrate (*center*), and in the additional presence of 50 mM Na⁺ (*bottom*). All images were recorded consecutively.

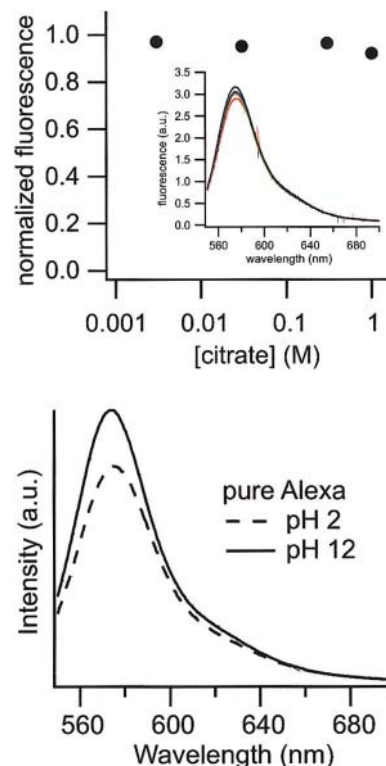


FIGURE 8 Control experiments. (*Bottom*) Two emission spectra taken at extreme pH values to prove that a change in pH does not have the observed quenching effect on the fluorophores. (*Top*) Emission maxima of AF₅₄₆ taken at increasing citrate concentrations (pH 7.0) to prove that a high level of citrate does not have the observed quenching effect. The values are normalized to the fluorescence intensity without citrate. The respective emission spectra are shown in the inset.

mandatory for citrate binding interactions with CitS or has any influence on the citrate-induced fluorescence quenching, we first added 250 μ M citrate to a Na⁺-free buffer and subsequently 50 mM Na⁺ to immobilized CitS-sC398¹ and vice versa. To ensure saturating conditions, the used concentration of Na⁺ was \sim 20 times higher than the apparent affinity of wild-type CitS for sodium ions ($K_{0.5} = 2.7$ mM) (Kästner et al., 2000). The images recorded at each step are shown in Fig. 7. Clearly, the addition of citrate already quenched all of the fluorophores bound to functional proteins, whereas the absence or presence of sodium ions had no effect on the observed static quenching of AF₅₄₆. When first Na⁺ and then citrate was added, quenching again occurred only at the presence of citrate (data not shown).

Dual-color labeling and detection studies with CitS-sC398²

The fluorescence intensity of a number of spots showed strong and irregular variations, usually not seen with single molecules. They could be explained by the existence of CitS oligomers with two or more fluorophores at a distance of 4 nm or less. Fluorescence homotransfer together with some

rotational degree of freedom and intersystem crossing could be responsible for the fluctuating intensity. Indeed, there was preliminary evidence for a dimeric arrangement of CitS in detergent micelles and lipid membranes. Such evidences derived from affinity chromatography, where the CitS fusion protein eluted as heterodimer because of a mixed population of a tagged and a nontagged form (purification tag cleaved off) (Pos et al., 1994). Recent blue native PAGE experiments in our lab correspond with these findings because purified wild-type CitS as well as CitS-sC398 showed an electrophoretic mobility of the homodimeric form (~ 100 kDa) (Fig. 4 C).

Because single-molecule experiments, in contrast to ensemble measurements by UV/VIS spectrophotometry, allow us to identify heterogeneities or different classes in a population, they can be used to study the state of protein-protein interaction. For this purpose, we performed a dual-color labeling and detection experiment with immobilized CitS-sC398-AF_{546/568} (CitS-sC398²). Due to our equimolar labeling procedure with the two different Alexa Fluor derivatives AF₅₄₆ and AF₅₆₈ and $\sim 80\%$ degree of labeling, the following dual-labeled dimeric species of CitS would be expected: (CitS-AF₅₄₆)², (CitS-sC398-AF₅₄₆)(CitS-sC398-AF₅₆₈), and (CitS-sC398-AF₅₆₈)². At low surface density of fluorescent spots two-step bleaching and the colocalization of two fluorophores (AF₅₄₆ and/or AF₅₆₈) at one spot would prove the existence of the dimeric form. In the case of dual-color labeled CitS-sC398² a simultaneous change of the dichroic ratio is expected upon stepwise bleaching of the two fluorophores that could serve as additional proof (Schutz et al. 1998).

For each experiment we selected a specific single-molecule spot, collected the fluorescence photons, and computed the intensity and the dichroic ratio as a function of time. By simultaneously monitoring the polarization we could exclude the possibility that the decrease of emission intensity was caused by a reorientation of the fluorophores. Fig. 9 shows a typical time trace of dual-color labeled CitS-sC398², i.e., (CitS-sC398-AF₅₄₆)(CitS-sC398-AF₅₆₈), including consecutive two-step bleaching of AF₅₆₈ and

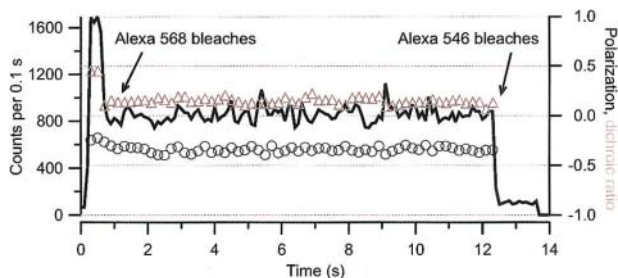


FIGURE 9 Single-molecule fluorescence emission time trace (—) including dichroic ratio (Δ) and polarization (\circ) of dual-color labeled CitS-sC398². The black arrows indicate the consecutive two-step bleaching of AF₅₆₈ and AF₅₄₆ and the associated change of the dichroic ratio upon bleaching of AF₅₆₈.

AF₅₄₆ (1700 \rightarrow 800 \rightarrow ~ 100 counts per 0.1 s) and the associated change of the dichroic ratio (D) upon bleaching of the first fluorophore AF₅₆₈. In summary, 48 spots were selected for evaluation. We observed two-step bleaching for 33 spots of which 17 showed an associated change of the dichroic ratio. Assuming a labeling efficiency of 80%, these numbers indicate a strong evidence of dimer formation of CitS. Intensively fluorescing spots, which exhibited a continuous instead of stepwise bleaching, were also observed. We assigned these spots to nonfunctional aggregates of a large number of molecules and discarded them from further analysis.

DISCUSSION

Single-molecule fluorescence microscopy with immobilized CitS-sC398¹ revealed an almost complete fluorescence quenching of the majority of molecules upon citrate addition (Fig. 5). The fluorescence quenching response was highly specific for citrate and was not observed with DL-isocitrate, which is not a substrate of CitS (Fig. 6). Moreover, the addition of citrate to pure AF₅₄₆ did not cause any significant quenching of the fluorophore, nor did changes of the pH to pH 2 or pH 12 (Fig. 8).

Nontrivial quenching mainly arises from the formation of nonfluorescent products (static quenching), collision with other molecules (dynamic or collisional quenching), and excitation energy transfer (Lakowicz, 1999). In the present study we observed static quenching because only the number of fluorescent molecules decreased, whereas the remaining fluorescent spots showed unchanged photophysical properties. Static quenching can result from the formation of nonfluorescent species, electron or proton transfer, and π -stacking with aromatic systems. Most likely interactions between the fluorophore and proximate aromatic amino acid residues of the protein contribute to the observed effect. Which interaction accounts most for the dramatic quenching of the bound fluorophore AF₅₄₆ remains to be investigated.

We suppose, therefore, that binding of citrate to CitS elicits a conformational change of the protein concomitant with an alteration of the molecular environment of AF₅₄₆, and thus with the observed fluorescence quenching. The fluorescent probe is bound to C398 (loop X-XI) within the highly conserved C-terminal region, which probably includes the substrate binding site (Bandell and Lolkema, 2000b) as proposed for the three closely related 2-hydroxycarboxylate transporters CitS (*K. pneumoniae*), CitP (*L. mesenteroides*), and MleP (*L. lactis*). Hence, the citrate-induced conformational change either involves the loop X-XI region itself or other parts of the molecule interacting with this region. Similar ligand-induced fluorescent changes that were traced back to structural changes associated with substrate binding have been extensively studied for two other secondary active transport proteins using ensemble

measurements, the H⁺-coupled lactose permease (Venkatesan and Kaback, 1998; Wang et al., 1997; Weitzman et al., 1995; Wu et al., 1995) and the H⁺/Na⁺-coupled melibiose permease (Cordat et al., 2000; Mus-Veteau and Leblanc, 1996; Mus-Veteau et al., 1995) from *E. coli*.

CitS has a high-affinity citrate binding site ($K_s \sim 13 \mu\text{M}$) proposed to be responsible in vivo for citrate binding from the periplasm and a low-affinity site ($K_s \sim 14 \text{mM}$) from where citrate is assumed to dissociate into the cytoplasm (Pos and Dimroth, 1996). The concentration of citrate used for the quenching studies (250 μM) was ~ 20 times higher than the binding constant for citrate of the high-affinity site to ensure saturating conditions, and 60 times lower than that of the low-affinity binding site. Hence, the observed conformational change is apparently due to the occupation of the high-affinity site with citrate. Sodium ions, which are obligatory required for citrate uptake, had no noticeable effect on the citrate-induced fluorescence quenching, which probably indicates that citrate binding is Na⁺-independent, but a Na⁺-dependent step occurs later in the transport cycle.

Interestingly, preliminary data derived from citrate binding studies by isothermal titration calorimetry (Kästner et al., unpublished; Wiseman et al., 1989) indicated that the overall enthalpy change at the high-affinity site upon citrate binding was slightly endothermic ($\Delta H_{\text{high}} = 6 \text{kJ/mol}$), whereas the observed enthalpy at the low-affinity site was exothermic ($\Delta H_{\text{low}} = -55 \text{kJ/mol}$). Similar observations have been made for other proteins with ligand-coupled conformational changes like the cAMP receptor protein (Gorshkova et al., 1995) or the inosine monophosphate dehydrogenase (Bruzzeze and Connelly, 1997). There, the exothermic enthalpy of binding (ΔH_{bind}) is overwhelmed by the large endothermic enthalpy of the conformational transition (ΔH_{conf}) that takes place upon binding of the ligand.

For an increasing number of prokaryotic transporters it has been demonstrated that they form dimers. Examples are the tetracycline cation/proton antiporter TetA from *Escherichia coli* (McMurry and Levy, 1995), the H⁺-coupled multidrug transporter EmrE from *E. coli* (Tate et al., 2001), the lactose transporter LacS from *Streptococcus thermophilus* (Friesen et al., 2000), and the Na⁺/H⁺ antiporter NhaA from *E. coli* (Gerchman et al., 2001). For NhaA even two-dimensional crystals showed tightly packed dimers with well-defined interdimer contacts (Williams et al., 1999). Interestingly, for the phosphate carrier (PIC) from *Saccharomyces cerevisiae* it was proven that only the dimeric form is completely active, which enables functional cross talk between the two monomers (Schroers et al., 1998). In contrast, the monomeric as well as the dimeric form of lactose permease from *E. coli* were shown to be functional (Sahin-Toth et al., 1994). Thus, the dimeric arrangement is not necessarily a functional requirement but might be due to protein/lipid interactions and stability reasons in a membrane environment. Moreover, there are theoretical arguments that oligomeric structures are

particularly well-suited to integral membrane proteins with transport function (Klingenberg, 1981). Our single-molecule studies with dual-labeled CitS-sC398² and dual-color detection provided strong evidence for a dimeric association of CitS. The frequent occurrence of two-step bleaching and the simultaneous change of the dichroic ratio upon bleaching of the first fluorophore suggest that functional, detergent-solubilized CitS forms a rather stable homodimer. Together with the biochemical and kinetic indications (Pos et al., 1994; Pos and Dimroth, 1996) these results clearly support our hypothesis that the active form of CitS within the membrane is a homodimer.

In summary, we could demonstrate that fluorescence quenching of single molecules is a powerful technique to follow conformational changes such as that elicited by the binding of citrate within the region of loop X-XI of CitS. Furthermore, with the dual-color labeling and detection technique strong evidence for a dimeric arrangement of CitS was obtained. This technique is generally very useful to determine binding events of single molecules in an unambiguous fashion.

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