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Article

His-Tag-Mediated Dimerization of Chemoreceptors Leads to Assembly of Functional Nanoarrays

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Supporting Information

ABSTRACT: Transmembrane chemotaxis receptors are found in bacteria in extended hexagonal arrays stabilized by the membrane and by cytosolic binding partners, the kinase CheA and coupling protein CheW. Models of array architecture and assembly propose receptors cluster into trimers of dimers that associate with one CheA dimer and two CheW monomers to form the minimal "core unit" necessary for signal transduction. Reconstructing *in vitro* chemoreceptor ternary complexes that are homogeneous and functional and

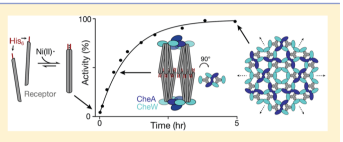


exhibit native architecture remains a challenge. Here we report that His-tag-mediated receptor dimerization with divalent metals is sufficient to drive assembly of nativelike functional arrays of a receptor cytoplasmic fragment. Our results indicate receptor dimerization initiates assembly and precedes formation of ternary complexes with partial kinase activity. Restoration of maximal kinase activity coincides with a shift to larger complexes, suggesting that kinase activity depends on interactions beyond the core unit. We hypothesize that achieving maximal activity requires building core units into hexagons and/or coalescing hexagons into the extended lattice. Overall, the minimally perturbing His-tag-mediated dimerization leads to assembly of chemoreceptor arrays with native architecture and thus serves as a powerful tool for studying the assembly and mechanism of this complex and other multiprotein complexes.

Many fundamental biological processes are performed by multiprotein complexes.^{1,2} To fully understand how these processes occur within the cell, methods are needed both for monitoring such processes *in vivo* and for isolation and/or reassembly of such complexes for *in vitro* structure–function analysis. Isolation and reassembly of complexes often require modifications or deletions of portions of the protein components: for instance, deletion of transmembrane domains of membrane proteins may be needed to prepare homogeneous complexes for structural or biophysical studies. Typically, such truncations also delete stabilizing interactions, and it becomes difficult to reassemble complexes with nativelike architecture and activity. We have developed a novel strategy for the stabilization of assemblies with native structure and activity for *in vitro* mechanistic studies.

Bacterial chemotaxis receptors are an ideal system for understanding the molecular details of signal transduction by membrane proteins and their complexes. Chemotaxis receptors are dimeric transmembrane proteins anchored in the inner membrane in large patches at the poles of the cell, which are responsible for sensing and responding to environmental stimuli. When the ligand binds to the receptor periplasmic domain, a signal is transmitted down the ~300 Å length of the receptor to control the autophosphorylation activity of an associated histidine kinase, CheA, which plays a central role in the signal transduction pathway. In addition to forming a complex with CheA and a coupling protein CheW, receptors form large clusters (a membrane surface area of 53000 nm² for Escherichia coli)³⁻⁵ that enable extraordinary levels of cooperativity and sensitivity.^{6,7} The native system is built of trimers of receptor dimers that each form one vertex of a hexagon within the membrane and are stabilized at the membrane-distal tip of the receptor by hexagonal rings of alternating CheA and CheW.⁸⁻¹⁰ It is challenging to prepare homogeneous samples of functional chemoreceptor complexes in vitro for structure-function analysis. Complexes of the intact membrane-bound receptor with CheA and CheW are not homogeneous, in part because of the inability to control the orientation of the receptor within membrane vesicles^{11,12} or nanodiscs.¹³ In principle, the membrane orientation issue can be addressed by using a cytoplasmic fragment of the receptor, but this truncated construct does not form functional complexes with CheA or CheW in solution. Presumably, this is because the missing transmembrane and periplasmic domains are needed to provide protein-protein interactions that stabilize the receptor dimer and geometrical alignment within the membrane that promotes assembly of the array.

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In an elegant combination of membrane mimetics and nanoscale assembly tools, Weis and co-workers developed templating vesicles that promote the assembly of membraneassociated multiprotein complexes from soluble component proteins. This technology harnesses the high-affinity interaction between histidine imidazole groups and the Ni(II)-NTA functional group originally developed for immobilized metal affinity chromatography protein purification.¹⁴ The N-terminally His-tagged cytoplasmic fragment (CF) of the aspartate chemoreceptor binds to vesicles containing lipids with nickelchelating headgroups (DOGS-NTA), which enables binding of CheA and CheW into ternary complexes with kinase and methylation activity that mimic native signaling states.^{15,16} Molecular crowding agents such as PEG8000 provide an alternate means of driving assembly of functional complexes, with no requirement for membrane vesicles.¹⁷ Both assembly methods result in extended hexagonal arrays with nativelike architecture,¹² and PEG-mediated assembly forms sandwich lattices remarkably similar to cytoplasmic chemoreceptor arrays seen in some bacterial species.^{18,19}

Here we report a novel method for assembling functional arrays *in vitro* and for monitoring the stages of assembly. We show that adding divalent metal salts to ternary mixtures of Histagged CF, CheA, and CheW promotes receptor dimerization and assembly into active complexes with native stoichiometry and architecture. Moreover, kinetic analysis with this new assembly method provides the first experimental evidence that initial binding of CheA and CheW into ternary complexes proposed to be the "core unit" does not yield high activity, and that activity reaches its maximum upon formation of larger arrays. His-tag-mediated dimerization provides a new approach to stabilizing multiprotein assemblies for analysis of protein mechanisms within their native complexes.

MATERIALS AND METHODS

Vesicle Preparation. DOPC (1,2-dioleoyl-sn-glycero-3phosphocholine) and the nickel-chelating lipid DOGS-NTA- Ni^{2+} (1,2-dioleoyl-sn-glycero-3-{[N-(5-amino-1-carboxypentyl)iminodiacetic acid]succinyl}) (Avanti Polar Lipids) were combined in chloroform at a 1.5:1 DOPC:DOGS-NTA-Ni²⁺ molar ratio. Lipids were dried into a thin film with a stream of N₂ gas and left under a vacuum for 1 h before being rehydrated with 1× PKB {phosphate kinase buffer $[50 \text{ mM K}_{r}H_{r}PO_{4} (pH)]$ 7.5), 50 mM KCl, and 5 mM MgCl₂] and vortexed extensively (30 s vortex, 5 min rest, repeated three times) to form multilamellar vesicles, followed by five cycles of 3 min freezing/ thawing. Freeze/thaw cycles were accomplished by dipping the lipids in liquid nitrogen for 20 s to freeze and left on the bench for 3 min and then held for 3 min in a 37 °C water bath. Extrusion was then performed with an extrusion apparatus (Avanti Polar Lipids) using a 100 nm diameter pore size polycarbonate membrane. High-concentration stocks at 3 mM total lipid (1.8 mM DOPC and 1.2 mM DOGS-NTA-Ni²⁺) were prepared, and a final [lipid]_{total} of 725 μ M was used for assembly (see below). The lipid concentrations were optimized in parallel with CheA and CheW concentrations to maximize kinase activity under conditions under which the available vesicle surface area could accommodate all CF as hexagonal arrays.

Plasmid Construction. The pHTCF4Q plasmid encoding the cytoplasmic fragment (CF) of the aspartate chemoreceptor with glutamine at all four primary methylation sites (4Q) was isolated from DH5 α F' *E. coli* cells.²⁰ A TEV-protease (tobacco

etch virus) recognition and cleavage sequence (ENLYFQG) was inserted directly following the N-terminal His6-tag of pHTCF4Q using site-directed mutagenesis and polymerase chain reaction (PCR). Forward (5'-GAAAACCTGTATTTT-CAGGGCGGATCCCCTATGCAACG-3') and reverse (5'-GCCCTGAAAATACAGGTTTTCGTGATGGTGATGGT-GATG-3') overlapping primers were designed to include the TEV-protease recognition sequence and were purchased from Eurofins Genomics. The PCR was performed in a thermocycler (Bio-Rad MJ Mini), and reagents, including Phusion DNA polymerase, dNTPs, and DpnI, were purchased from New England Biolabs. The PCR product (pTEVCF4Q, amp^R) was subjected to DpnI digestion and transformed into DH5 α F' for plasmid purification. Following sequence verification (Genewiz), pTEVCF4Q was co-transformed with pCF430 (encoding lacI^q and tet^R) into BL21(DE3) for protein expression.

Protein Purification. BL21(DE3) E. coli cells expressing H_6CF (pHTCF4Q, amp^R) or H_6TEV -CF (pTEVCF4Q, amp^R) co-transformed with pCF430 (tet^R) were grown in Luria-Bertani (LB) broth with ampicillin (100 μ g/mL) and tetracycline (10 μ g/mL) at 37 °C until the optical density at 600 nm reached approximately 0.6. The temperature was decreased to 15 °C for induction with 1 mM IPTG for 16-18 h. Cells were pelleted, resuspended in lysis buffer [75 mM K₂H₂PO₄ (pH 7.5), 500 mM NaCl, 5 mM imidazole, and 1 mM EDTA], and lysed with a microfluidizer at 16K psi. PMSF (1 mM) was added every hour following cell lysis to limit proteolysis. Cell debris was separated by centrifugation and the supernatant applied to a HisTrap FF Ni²⁺-NTA affinity column (GE Healthcare) equilibrated with 10 column volumes of 75 mM K_xH_xPO₄ (pH 7.5), 500 mM NaCl, and 5 mM imidazole. The column was washed with 5 column volumes of 75 mM K_xH_xPO₄ (pH 7.5), 500 mM NaCl, and 50 mM imidazole, before elution with 75 mM K_xH_xPO₄ (pH 7.5), 500 mM NaCl, and 500 mM imidazole. Eluted protein was verified by SDS-PAGE, and fractions were pooled prior to treatment with 5 mM EDTA to chelate any $\mathrm{Ni}^{2\mathrm{+}}$ stripped from the column and dialysis against 75 mM K_xH_xPO₄ (pH 7.5) and 75 mM KCl with 7 kDa molecular weight cutoff SnakeSkin tubing (Thermo Scientific) to remove EDTA and imidazole and lower the NaCl concentration. Half of the purified H₆TEV-CF was subjected to His-tag removal by TEV-protease cleavage (details below).

Plasmids encoding TEV-cleavable His-tagged CheA (pTEVcheA, kan^R), CheW (pTEVcheW, kan^R), and CheY (pTEVcheY, kan^R) (plasmid construction details described elsewhere)²¹ were expressed in BL21(DE3) and grown at 37 °C in LB broth supplemented with 50 μ g/mL kanamycin. At an optical density at 600 nm of ~0.7-0.9, 1 mM IPTG was added to induce expression for 3 h before the cells were harvested and purified with HisTrap affinity chromatography as described for CF above. A different buffer system for CheA, CheW, and CheY purification was used: lysis buffer [75 mM Tris-HCl (pH 7.4), 100 mM KCl, and 1 mM EDTA], equilibration buffer [75 mM Tris-HCl (pH 7.4) and 100 mM KCl], wash buffer [75 mM Tris-HCl (pH 7.4), 100 mM KCl, and 10 mM imidazole], and elution buffer [75 mM Tris-HCl (pH 7.4), 100 mM KCl, and 250 mM imidazole]. Following elution, fractions containing protein were verified by SDS-PAGE, pooled and treated with 5 mM EDTA, and then dialyzed as for CF into 75 mM Tris-HCl (pH 7.4) and 100 mM KCl prior to His-tag removal.

Plasmid pRK793 (amp^R) encoding N-terminally His-tagged TEV-protease (a gift from D. Waugh, Addgene plasmid 8827)²²

was expressed in BL21(DE3)-RIL (amp^R). Cells were grown at 37 °C in LB broth containing 150 μ g/mL ampicillin and 50 μ g/mL chloramphenicol. Protein production was induced at 30 °C with 1 mM IPTG for 4 h when an optical density of 0.6 was reached. The protein purification protocol and buffer system for CheA, CheW, and CheY was also used here (see above).

His-Tag Cleavage with TEV-Protease. Following purification, CheA, CheW, CheW, and H_6 TEV-CF were incubated with TEV-protease (purification details given above) at a 50:1 His-tagged protein:TEV-protease molar ratio, and the mixture was shaken at 4 °C overnight and 25 °C for 3 h. Complete cleavage was confirmed by a gel shift observed by SDS–PAGE. Cleaved proteins were separated from TEV-protease by being passed through the equilibrated HisTrap column, collected, and concentrated with 10 kDa centrifugal concentrators (Amicon); protein concentrations were measured by the BCA assay (Thermo Scientific), and proteins were frozen in liquid nitrogen and stored at -80 °C.

Complex Assembly. Preparation of ternary complexes was performed as previously described for vesicles¹⁵ and PEG¹⁷ with some modifications. Briefly, vesicle samples were prepared by combining the following (in order): autoclaved water, 1 mM PMSF dissolved in 100% ethanol, phosphate kinase buffer (PKB) from a 5× stock (the 1× stock being 50 mM $K_xH_xPO_{4y}$ 50 mM KCl, and 5 mM MgCl₂), 12 µM CheA, 24 µM CheW, 30 μ M CF (H₆CF, H₆TEV-CF, or Δ H₆CF), and 725 μ M vesicles (prepared as described above). PEG-assembled samples were made by combining the following (in order): autoclaved water, 1 mM PMSF, PKB, 12 µM CheA, 20 µM CheW, 50 µM CF (H₆CF, H₆TEV-CF, or Δ H₆CF), 7.5% (w/v) PEG8000 [from a 40% (w/v) stock], and 4% (w/v) D-trehalose [from a 40% (w/v) stock]. Metal-assembled samples were prepared under protein and buffer conditions identical to those optimized for vesicle assembly, except that vesicles were replaced with metal salts (NiCl₂·6H₂O, ZnCl₂ anhydrous, CuCl₂·2H₂O, MnCl₂·4H₂O, and CoCl₂·6H₂O, all obtained from Sigma-Aldrich) prepared at ~1 mM in autoclaved Milli-Q H₂O. Metal stock concentrations were verified before being used (see below).

Once combined, samples were incubated in a 25 $^{\circ}$ C water bath for 4 h or overnight before use. To measure kinetics of assembly, both activity and sedimentation (below) were measured at time points immediately following addition of all complex components.

Activity and Binding Assays. Kinase activity was measured using an enzyme-coupled ATPase assay that couples NADH oxidation to consumption of ATP.^{15,23,24} Briefly. activity measurements were conducted immediately following a 100-fold dilution of assembled complexes into PKB containing 55 µM CheY, 2.2 mM phosphoenolpyruvate, 4 mM ATP, 250 µM NADH, and 20 units of PK/LDH enzyme (Sigma-Aldrich). The background activity of CheY under identical conditions in the absence of the complex was subtracted before calculation of kinase activity. The kinase activity (inverse seconds) was determined from the linear change in absorbance at 340 nm $[d[ATP]/dt = -6220(dA_{340}/$ dt] over 1.5 min. Total activity is based on the full amount of CheA in the sample (12 μ M), while the specific activity considers only the amount of CheA in the complex, based on quantification from the sedimentation assay (below).

The amount of bound protein was determined with a sedimentation assay performed at 25 °C in a benchtop ultracentrifuge (Beckman TLX, TLA 120.2 rotor, 125000g, 30

min). Typically, a 35 μ L aliquot of sample (in either assembled ternary complexes or individual proteins) was centrifuged. Following sedimentation, the supernatant was carefully removed and placed into a clean Eppendorf tube to minimize contamination of free protein in the pellet containing the bound fraction of proteins. The pellets were slowly resuspended to the original volume in autoclaved Milli-Q H₂O and vortexed to homogenize the resuspension. Aliquots of Total (free plus bound protein, before sedimentation), Supernatant (free protein), and Pellet (bound protein) were subjected to SDS-PAGE (12.5% acrylamide) and stained with Gel-code Blue (Pierce Chemical Co.). Gels were imaged by densitometry with a Gel Doc EZ Imager (Bio-Rad), and the integrated intensities were analyzed with ImageJ.²⁵ The bound concentrations were computed as (I_{pellet}/I_{total}) [protein]_{total} for each protein to be quantified. For excess CheA and CheW conditions used in this study, quantification of bound protein by the pellet was more reproducible than $I_{\text{total}} - I_{\text{supernatant}}$. Gels included two additional lanes of the Total sample diluted 6- and 12-fold such that a calibration curve from three intensities for known concentrations could be calculated. The y-intercept was then subtracted from each integrated intensity to correct for the gel background intensity and more accurately estimate the fraction of protein bound in a complex.

Determination of Metal Concentration. Metal concentrations of the ~ 1 mM aqueous metal stocks (NiCl₂, ZnCl₂, $CuCl_2$, $MnCl_2$, and $CoCl_2$) and the amounts of metal bound to H_6CF (alone or in complex) were measured using a PerkinElmer Optima DV4300 inductively coupled plasma optical emission spectroscopy (ICP-OES) instrument. Metal standards (TraceCERT by Sigma-Aldrich) were used to produce a calibration curve for each metal at 0.1, 0.2, 0.75, 1, 2, and 10 ppm. Metal stocks were diluted 100-fold, and protein/metal samples were diluted 10-fold into Milli-Q water that was also used to blank the instrument. Protein samples were prepared by overnight incubation of 30 μ M H₆CF (alone or with 12 μ M CheA and 24 μ M CheW) with the metals that produced active complexes at optimal concentrations (180 μ M NiCl₂, 300 µM ZnCl₂, or 360 µM CoCl₂) in a 25 °C water bath. Aliquots of these samples were then subjected to ultracentrifugation (60K rpm for 30 min at 25 °C) to separate large metal-protein complexes from unbound protein and metal in the supernatant. Bound protein and metals in the pellet were carefully resuspended to the initial volume in Milli-Q water. The entire resuspended pellet was used for ICP-OES analysis, so that any chunks of protein pellet were still part of the sample that was measured.

SEC–MALS. Size exclusion chromatography and multi-angle light scattering (SEC–MALS) were used to monitor metalmediated complex formation. The SEC instrument is coupled to an 18-angle static light scattering detector (DAWN HELEOS-II), a dynamic light scattering detector (Wyatt-QELS), and a differential refractometer (Optilab T-rEX, Wyatt Technology). Proteins were injected onto a TSKgel G2000SW_{XL} column maintained at 4 °C and pre-equilibrated overnight with PKB buffer (pH 7.0) kept on ice. Typical injections were 75 μ L of 0.5–3 mg/mL protein. Data were collected every second for 30 min at a flow rate of 0.5 mL/min. Protein elution was monitored at A_{280} and A_{214} , in addition to static and dynamic light scattering. The accompanying ASTRA VI software was used to determine molar mass, peak polydispersity, and ultraviolet (UV) peak areas for all individual

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peaks and the entire elution to determine the percent loss in a precolumn 0.22 μ m filter.

Electron Cryo-Tomography (ECT). Samples were prepared by mixing protein assemblies (as above) with a bovine serum albumin-treated 10 nm colloidal gold solution (Cell Microscopy Core, Utrecht University, Utrecht, The Netherlands). After being briefly vortexed, 3 μ L mixtures were applied to freshly plasma-cleaned R2/2 copper Quantifoil grids (Quantifoil Micro Tools). Blotting and plunge freezing in ethane were performed for approximately 1 s with a Leica EMGP instrument (Leica microsystems, Wetzlar, Germany), set at 20 °C and 95% humidity. Grids were stored in liquid nitrogen until they were imaged. Data were collected on a Titan Krios transmission electron microscope from Thermo Fisher Scientific (formerly FEI; Hillsboro, OR) with a field emission gun operating at 300 kV. The microscope was equipped with a Gatan (Pleasanton, CA) image filter and a Gatan K2 Summit direct electron detector. Data were acquired using Tomography version 4.0 (Thermo Fisher Scientific) in batch mode. Tilt series for PEG and Ni(II)-assembled H₆CF ternary complexes were collected at nominal magnifications of 42000× and 33000×, respectively. A discontinuous tilt scheme was used for imaging, taking images from 0° to -60° followed by 0° to 60° with a 2° tilting increment. The cumulative dosage for each tilt series was 80 e/A². Defocus was set to $-8 \ \mu m$. The tilt series for Ni(II)-assembled H₆CF (without CheA or CheW) and PEG-assembled ΔH_6CF complexes were collected using the same tilt scheme but at a defocus of -300 nm with a volta phase plate (VPP) (Thermo Fisher Scientific). The VPP was heated to 225 °C and activated for 80 s before each tilt series aiming to generate an approximate phase shift of 90°.26 An extra 5 s of conditioning time was applied between each tilt image. All images were acquired using the low-dose routine integrated in the Tomography software. Drift correction and tilt series alignment were performed with software package IMOD.^{27,2} ⁸ Tomograms were reconstructed using both weighted back-projection (WBP) and nine iterations of simultaneous iterative reconstruction (SIRT). PEET was used for subvolume averaging.²⁹ Visualization and image analysis were performed with ImageJ and Chimera.^{25,30}

RESULTS

Divalent Metals Restore Chemoreceptor Ternary Complex Formation and Function. Under conditions similar to those used to form functional complexes of aspartate receptor CF, CheA, and CheW with templating vesicles, we have discovered that some divalent metals are sufficient to restore function, with no requirement for vesicles. A series of metal titrations reveal the conditions needed to restore both kinase activity of CheA (measured with an enzyme-coupled ATPase assay, using the native phosphoryl acceptor protein CheY) and ternary complex formation (measured with sedimentation assays). Five divalent metal chlorides were tested (NiCl₂, ZnCl₂, CoCl₂, CuCl₂, and MnCl₂) at concentrations ranging from 0 to 540 μ M while protein concentrations were kept constant at levels that promote maximal incorporation of CF into functional complexes [30 μ M CF with excess CheA (12 μ M) and CheW (24 μ M)]. As shown in Figure 1, NiCl₂ promotes active complex formation at the lowest concentrations, with maximum kinase activity reached at 180 μ M NiCl₂. The optimal ZnCl₂ concentration for kinase activity is 300 μ M, and the total kinase activity is 90% of the highest observed for NiCl₂. While CoCl₂ also produced active

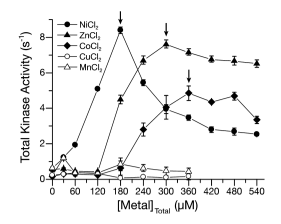


Figure 1. Three divalent metals promote kinase activity at different optimal metal concentrations. All experiments use excess CheA and CheW, under conditions optimized for maximal incorporation of CF into complexes with maximal kinase activity (30 μ M H₆CF, 12 μ M CheA, and 24 μ M CheW). Additions of NiCl₂ (\bullet), ZnCl₂ (\blacktriangle), or $CoCl_2$ (\blacklozenge) each promote kinase activity, with NiCl₂ promoting the highest activity, followed by $ZnCl_2$ and then $CoCl_2$. $CuCl_2$ (O) and $MnCl_2$ (\triangle) do not promote activity: activity comparable to that of the no-metal control is observed at all metal concentrations. As a positive control, PEG-mediated samples were run in parallel, with typical total kinase activities of ~11 s⁻¹. Comparisons of specific kinase activity (per bound CheA) among PEG-, vesicle-, and NiCl2-mediated assemblies are plotted in Figure 2B. Arrows denote maximally active conditions used for further studies, including metal:protein stoichiometry measurements listed in Table 1. Activities are averages of four to eight replicates measured on two or three days; error bars indicate ±one standard deviation. For this plot, measurements on each day were averaged, and the averages for all days were then combined, with propagation of the errors.

complexes in this titration series, the highest kinase activity (reached at 360 μ M CoCl₂) is only 50% of that of the optimal NiCl₂ sample. Thus, the relative efficiencies of these metals for promoting assembly of functional complexes decrease as follows: NiCl₂ > ZnCl₂ > CoCl₂. Interestingly, the sedimentation trends for CF, CheA, and CheW with each metal titration track mostly with the respective onset of activity (Figure S1). For example, NiCl₂-assembled complexes show both activity and sedimentation beginning at low NiCl₂ concentrations, and ZnCl₂-assembled complexes show nearly zero sedimentation until 180 μ M ZnCl₂, where activity is first observed. Furthermore, maximal sedimentation of CF, CheA, and CheW is reached for most cases under conditions that give maximal activity, and the CoCl₂ titration reaches activity and sedimentation of only 50% of those of the optimal NiCl₂ sample. Thus, three divalent metals restore the ability of the chemotaxis receptor CF to assemble with CheA and CheW into sedimentable complexes that activate the kinase CheA.

Interestingly, NiCl₂ concentrations of >180 μ M are clearly detrimental to kinase activity, and sedimentation of CF increases an additional 15%; CheA and CheW sedimentations remain constant (Figure S1). In contrast, ZnCl₂ and CoCl₂ cause an only modest decrease in activity at high concentrations (and no significant additional sedimentation).

Neither CuCl₂ nor MnCl₂ restored formation of active complexes. Kinase activity throughout the titration is similar to that of the proteins with no metal (<1 s⁻¹). Furthermore, all samples with CuCl₂ and MnCl₂ also had very low protein sedimentation values of <15%, again similar to levels for the nometal control, with the exception of CuCl₂ and CheA. At high

 $CuCl_2$ concentrations, nearly all of CheA sedimented (data not shown). Because there is no appreciable kinase activity or sedimentation of CheW and H₆CF, we can conclude that $CuCl_2$ is most likely causing aggregation of CheA.

Measurements of bound metals indicate that the active complexes have metal:protein stoichiometries close to 1:1 [Ni(II) and Co(II)] and 3:1 [Zn(II)]. The metal ion concentrations bound to CF were determined using inductively coupled plasma optimal emission spectroscopy (ICP-OES), in samples of active complexes (prepared with CheA and CheW) as well as CF alone. Metals producing active complexes lead to sedimentation of both CF alone and CF with CheA and CheW (with the exception of $CoCl_2$ + CF alone, which does not sediment). Samples were prepared with each metal at the optimal concentration based on the activity data (Figure 1, arrows) and incubated for long assembly times that maximize kinase activity (>4 h). The bound protein (CF alone or in complex with CheA and CheW) and metal were separated from free protein and metal by ultracentrifugation. Resuspended pellets were then analyzed for metal content by ICP-OES and protein content by SDS-PAGE (via comparison to a CF standard measured by the BCA assay). Unfortunately, we were unable to implement sulfur quantification by ICP-OES, which would have enabled measurement of a more accurate metal:protein ratio as the number of sulfur atoms per CF is known. As listed in Table 1, CF alone and CF in complex have

Table 1. Stoichiometries of Metal Binding to CF Measured by ICP-OES^{*a*}

metal added ^b	metal bound to CF $alone^c$	metal bound to CF in complex ^d
Ni(II)	1.2 ± 0.04	1.1 ± 0.05
Zn(II)	3.8 ± 0.16	3.1 ± 0.12
Co(II)	ND ^e	1.1 ± 0.15

^{*a*}Data are averages \pm one standard deviation for the number of metal ions bound per CF monomer from 10 independent ICP-OES measurements of each metal. ^{*b*}Metal added at concentrations found to give maximally active complexes (Figure 1, arrows). ^{*c*}Samples contain 30 μ M CF. ^{*d*}Samples contain 30 μ M CF, 12 μ M CheA, and 24 μ M CheW. ^{*e*}CoCl₂ with CF alone does not sediment, and thus, metal content could not be measured.

nearly identical stoichiometries for Ni(II) and Co(II), which suggests that metals bind similarly to CF in both cases. Moreover, the calculated stoichiometries are both close to one metal ion per CF monomer. In contrast, Zn(II) exhibited both a higher and a more variable stoichiometry of 3.8 Zn(II) per CF alone and 3.1 per CF in complexes. Thus, for all three metals, we can conclude that a large excess (6–12 equiv) is necessary to drive assembly to maximally active CF ternary complexes, but only a fraction of metal is directly involved in the assembly. All further studies were performed on Ni(II)-assembled samples at 180 μ M NiCl₂.

The Receptor His-Tag Is Essential for Metal-Mediated Assembly. We hypothesized that the His-tag could be involved in the metal-mediated assembly of functional chemoreceptor complexes, perhaps via stabilization of CF dimers or oligomers. To test this, we engineered a variant of the CF construct (henceforth called H₆CF for the sake of clarity) by inserting the recognition and cleavage sequence for TEVprotease after the His-tag (H₆TEV-CF). TEV-protease cleavage of H₆TEV-CF yields Δ H₆CF, with the native CF sequence plus a single glycine at the N-terminus. For each version of CF (H₆CF and H₆TEV-CF with His-tags and Δ H₆CF with no Histag, as shown in Figure 2A), we tested the assembly of ternary complexes with CheA and CheW by three *in vitro* methods [vesicles, PEG, and Ni(II)] and measured both the kinase activity and protein stoichiometry of sedimented complexes.

The three proteins together in buffer with no vesicles, PEG, or divalent metal show low kinase activity (Figure 2B, None) and <15% binding (not shown). We had previously observed variable levels of kinase activity for H₆CF, CheA, and CheW assembled in the absence of vesicles, PEG, or added metals and realized H₆CF may contain variable amounts of Ni(II), stripped from the NTA affinity column during purification.³¹ We incorporated a 5 mM EDTA treatment step into all purifications, following elution of protein from the column, and this EDTA is removed in the subsequent dialysis step (see Materials and Methods). Thus, the low level of activity and sedimentation observed in the absence of vesicles, PEG, or metal (Figure 2B, None) is not due to adventitious Ni(II) but presumably reflects the low stability of these complexes formed with the truncated CF.

As shown in Figure 2B, both H_6CF and $H_6TEV-CF$ complexes are functional with all assembly methods, indicating that the TEV insertion does not interfere with assembly of functional CF complexes. As anticipated, removing the His-tag results in no kinase activity and background levels of protein binding for both vesicle and Ni(II) assembly methods, demonstrating that the His-tag is required for metal-mediated assembly.

The native stoichiometry of chemoreceptor arrays is thought to be 6:1:2 receptor:CheA:CheW.^{9,10} As shown in Figure 2C, all active complexes of the different constructs assembled with vesicles, PEG, or Ni(II) display near-native stoichiometries. The variation in CheW (ranging from 2.1 to 3.0) may be due to weak staining of CheW, resulting in less accurate quantification. Overall, the stoichiometries are consistent with the nativelike architecture of PEG arrays, which have high CheA occupancy observed by electron cryotomography.¹² Thus, PEG mediates assembly via molecular crowding that does not require the Histag; both vesicle and Ni(II) assembly require the His-tag to mediate assembly of complexes with comparable kinase activity and protein stoichiometry.

Metal-Mediated Complexes Form Arrays with Nativelike Architecture. Chemoreceptor complexes with CheA and CheW form an extended lattice of hexagonal arrays located within the bacterial inner membrane that is responsible for ensuring a sensitive and integrated response to chemical gradients. Because ternary complex formation and kinase activity have been observed in the absence of arrays, $^{32-34}\ensuremath{$ we used electron cryotomography (ECT) to assess the structural features and homogeneity of the metal-mediated complexes. ECT images of Ni(II)-mediated H₆CF ternary complexes prepared under conditions leading to maximal activity reveal clear high-contrast hexagonal lattices (Figure 3A). With 12 nm center-to-center spacing between hexagons, these are indistinguishable from in vivo arrays and from our other in vitro assemblies formed with templating vesicles and PEG-mediated crowding.¹² Furthermore, isosurface averages extracted from the Ni(II)- and PEG-mediated tomograms are remarkably similar to the electron density from in vivo intact receptors (Figure S2). Interestingly, Ni(II)-mediated arrays also form the sandwichlike structures seen with PEG (Figure 3B). This architecture is likely necessary to stabilize the membrane-

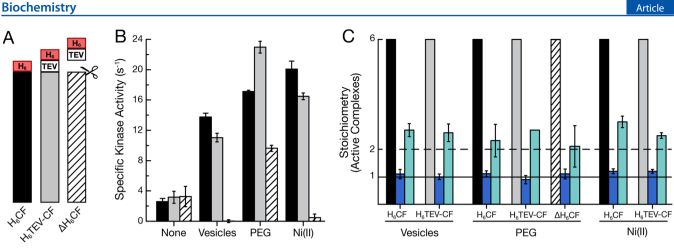


Figure 2. Characterization of in vitro-assembled ternary complexes reveals the receptor His-tag is required for metal-mediated assembly of active complexes with nativelike stoichiometry. (A) Cartoon depicting constructs tested to determine the role of the His-tag (H₆, red box) in metalmediated assembly of active complexes. (B) Specific activity of CheA kinase assembled with CheW and His-tagged receptor (black for H₆CF or gray for H₆TEV-CF) or non-His-tagged receptor (striped for Δ H₆CF). Activities are averages of four to eight replicates measured on two to four days; error bars indicate ±one standard deviation. (C) Stoichiometries for active complexes determined as the ratio of proteins in the sedimented complexes. The $\sim 25-28 \mu$ M sedimented CF was set to 6 for calculations of the relative amounts of bound CheA (blue) and CheW (cyan), based on ratios of band intensities in SDS-PAGE of sedimented complexes. Horizontal solid and dashed lines correspond to native 6:1:2 molar stoichiometries. Stoichiometries are averages of two to four replicates measured on two to four days; error bars indicate ±one standard deviation. Error bars are either large or missing for CheW stoichiometries in PEG complexes, because of gel distortions from PEG in the CheW molecular weight range.

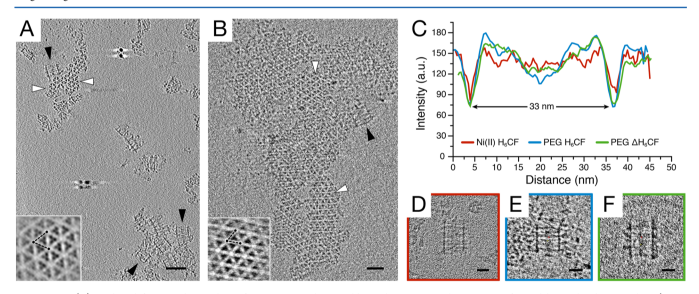


Figure 3. Ni(II)-mediated functional complexes form arrays with nativelike architecture similar to arrays formed by molecular crowding (PEGmediated arrays). Tomographic slices of (A) Ni(II)-mediated H₆CF and (B) PEG-mediated H₆CF show both top-down patches of hexagonal arrays (white arrows) and side view sandwichlike structures (black arrows) made from two layers of CF with overlapping N- and C-termini that are sandwiched between baseplates of CheA and CheW on the outside. Scale bars are 50 nm, and insets show enlarged subvolume averages of the hexagonal array with the canonical 12 nm center-to-center spacing (dotted lines in the insets of panel A and B are 12 nm) that is identical to that of in vivo arrays.^{8,12} (C) Comparison of intensities of the three-dimensional volume reveals that sandwich thicknesses for three sample types are all close to 33 nm. Five separate cubes containing sandwiches of similar proportions were averaged for each sample (Figure S3). (D-F) Example twodimensional tomographic slices of sandwiches from Ni(II)-mediated H₆CF, PEG-mediated H₆CF, and PEG-mediated Δ H₆CF ternary complexes used to determine intensity across the sandwich length. Scale bars are 20 nm.

proximal ends of the long CF in the absence of organization by a membrane.

To further investigate the structural similarities between arrays assembled by these methods, we measured the baseplate distances between CheA/CheW layers for three assemblies that formed sandwiches. Figure 3C, a plot of the intensity of the three-dimensional volume of the side view for Ni(II)-mediated H₆CF, PEG-mediated H₆CF, and PEG-mediated Δ H₆CF sandwiches (examples in Figure 3D-F) reveals identical

distances of 33 nm between baseplates. Given an approximate H_6CF receptor length of ~21.8 nm (145 residues, assuming they are all α -helical) from the N-terminus to the membranedistal tip, this 33 nm sandwich thickness predicts that CFs from each side of the sandwich overlap with each other in the central \sim 10.6 nm region. This may be consistent with the disorder (low electron density) observed in the approximately central one-third of the sandwich, as seen in a side view of the isosurfaces of the electron density calculated by subvolume

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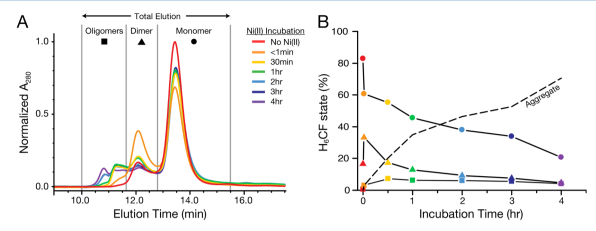


Figure 4. Ni(II) stabilizes the H₆CF dimer, which in turn favors larger oligomers. (A) SEC–MALS chromatograms of 30 μ M H₆CF incubated with 180 μ M NiCl₂, colored by rainbow according to the increasing incubation time. Vertical lines indicate elution time bounds used to distinguish ther monomer (12.8–15.5 min), dimer (11.7–12.8 min), and oligomers (10–11.7 min). Although identical volumes (75 μ L) of the same sample were injected, some H₆CF aggregate was caught in the precolumn filter. Chromatograms are normalized to a value of 1 for the largest peak in the series. These data are further scaled to the same total area to account for aggregated protein (B, dashed line). (B) The peak areas of the monomer (circle), dimer (triangle), and oligomers (square) were used to calculate the amount of H₆CF in each state for all incubation times. The percent H₆CF in aggregates larger than the 0.22 μ m precolumn filter (dashed line) was calculated from the loss of area under the SEC elution curve.

averaging of tomograms for Ni(II)- and PEG-mediated H_6CF (Figure S2). Remarkably, the similarity in baseplate distances indicates that the overlap interactions that stabilize the sandwich assembly do not depend critically on what stabilizes the complexes (metal or PEG) or on the sequence of the N-termini (H_6CF vs ΔH_6CF) that are present in the overlap region. Finally, the similarity in the architecture of the PEG-assembled H_6CF and ΔH_6CF arrays indicates that the ECT resolution is not sufficient to detect whatever differences in structure and dynamics account for the 2-fold difference in kinase activity (Figure 2B) between these samples.

Insight into Array Assembly and Properties. The results presented above suggest that metal binds to the His-tag of H₆CF and somehow stabilizes formation of the functional, nativelike array with CheA and CheW. Within the native architecture, what interactions could be stabilized by metal bridging two His-tags? The N-terminal His-tags are adjacent within the H₆CF dimer. In contrast, His-tags on two different dimers are not adjacent in the trimer of dimers (they are >65 Å apart in the ECT-derived structural model, Protein Data Bank entry 3JA6),¹⁰ and His-tags on the two H₆CF layers are not likely to be adjacent in the sandwich (they are on opposite sides of the ~ 10 nm overlap region). Thus, it seems likely that metal binding bridges two His-tags within a H₆CF dimer, and stabilization of the dimer drives assembly of the functional array. To investigate this, we monitored the formation of metalmediated complexes of H₆CF using SEC-MALS. The three component proteins and their complexes can be partially resolved via SEC, with the largest complexes and proteins eluting first, but SEC does not yield accurate molecular weights for variably shaped proteins like the long rod-shaped chemoreceptor. The MALS data enable the determination of hydrodynamic properties for better estimation of MW. However, MW estimation is not accurate for overlapping peaks or high-MW complexes that elute near the void volume and are highly polydisperse. SEC-MALS was used to monitor formation of complexes of H_6CF alone (Figure 4) and of functional complexes of H₆CF with CheA and CheW (Figure 5).

Complex formation of H₆CF alone provided the best means of assessing the effect of metal on the H₆CF monomer-dimer equilibrium, because H₆CF dimer and CheA have overlapping peaks (see Figure 5B). We anticipate that binding of Ni(II) to the H₆CF His-tag is similar in the presence and absence of CheA and CheW, because ICP-OES indicates 1:1 binding in both cases. As discussed above, incubation of H₆CF alone with NiCl₂ results in sedimentable aggregates. Apparently, these aggregates can become too large to pass the 0.22 μ m precolumn filter, as the total eluted protein was observed to decrease over the time course of the experiment. Figure 4A shows the overlaid chromatograms from the time course series, all scaled to the same total integrated peak area. The amount of aggregate lost in the precolumn filter (dashed line in Figure 4B) was calculated from the decrease in area with each injection. To quantify the fraction of monomer, dimer, and oligomer, elution time ranges were set at the points of minimum A_{280} between peaks (Figure 4A, gray vertical lines at 10, 11.7, 12.8, and 15.5 min), and these peak areas were used to calculate the percentage of H₆CF in monomer, dimer, and oligomer forms (Figure 4B).

SEC-MALS of H_6CF incubated with 180 μ M NiCl₂ (Figure 4) demonstrates that Ni(II) stabilizes the H_6CF dimer. Prior to the addition of NiCl₂, the monomer:dimer ratio is approximately 10:1 (Figure 4, red) based on the integrated UV peak area ratio of 5.1:1 (area of the dimer peak divided by 2 gives the dimer concentration). Strikingly, within the minimal time to add metal to the protein and inject it onto the column (<1 min), there is a significant decrease in the monomer:dimer ratio to 3.7:1 (Figure 4, orange), which corresponds to a 2.7-fold decrease in this ratio. At remaining time points, both monomer and dimer concentrations continue to decrease in parallel, because of a significant increase in the aggregate fraction (Figure 4B). The system is not at equilibrium, but the monomer:dimer ratio remains in the range of 6–8:1, consistent with Ni(II) stabilization of H_6CF dimers.

To understand the mechanism of metal-mediated formation of functional complexes, we investigated the assembly kinetics by measuring the onset of kinase activity and sedimentation immediately after combining the H_6CF , CheA, CheW, and 180

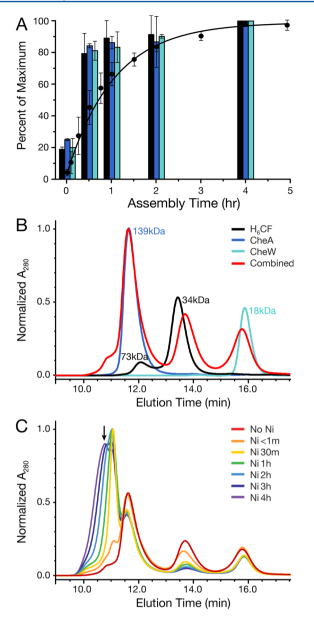


Figure 5. Kinetics of Ni(II)-mediated assembly of H₆CF ternary complexes. (A) Time course of kinase activity (\bullet) and sedimentation of H₆CF, CheA, and CheW (black, dark blue, and cyan bars, respectively) combined with 180 µM NiCl₂. The zero time point is before NiCl₂ addition. Activity and sedimentation data are normalized to their respective maxima, achieved at 4 h (maximum total activity of 7.6 s⁻¹, specific activity per bound CheA of 16 s⁻¹, and maximum sedimentations of 27.0 µM H₆CF, 5.8 µM CheA, and 13.6 µM CheW). Activities and sedimentation percentages are averages of two (sedimentation) and four (activity) replicates measured on 2 days; error bars indicate ±one standard deviation. (B) SEC chromatograms of separately injected H₆CF (black), CheA (blue), and CheW (cyan) at concentrations used for assembly (Figure 1, legend), as well as a single injection of all three in solution without NiCl₂ (red). Molecular masses estimated from multi-angle light scattering (MALS) are indicated and are reasonably consistent with the molecular weights of dimeric CheA (142.7 kDa), dimers and monomers of H₆CF (32.7 kDa), and CheW (18.1 kDa). (C) SEC chromatograms of H₆CF, CheA, and CheW incubated with NiCl₂, colored by rainbow according to increasing incubation times. A high-MW complex of H₆CF, CheA, and CheW (11 min elution time) that forms in low yield in the absence of NiCl₂ (0 min sedimentation in panel A and red in panels B and C) and in high yield at early incubation times (30 and 60 min in panel A and yellow and green in panel C) sediments but does not have

Figure 5. continued

high kinase activity. Maximal kinase activity appears to correlate with formation of a higher-MW complex indicated by the arrow (near void volume at a 10.5 min elution time). All chromatograms in each plot are normalized to a value of 1 for the largest peak in the series.

 μ M NiCl₂. Assembly with either vesicles²⁴ or PEG (unpublished observations) requires incubation at 25 °C for at least 4 h before maximal activity is reached. Although the NiCl₂ assembly activity similarly achieves maximal activity after 4 h (Figure 5A), the sedimentation profile shows nearly complete binding (80–90%) of all three proteins within the first 30 min. Note that complex formation may continue during the 30 min centrifugation, whereas the kinase activity measurements take <2 min; therefore, complete binding occurs at 30–60 min. Thus, the kinase activity is only 40–60% when the CheA and CheW incorporation is maximal, which suggests that initially the three proteins bind quickly but require conformational changes or rearrangement to reach maximal activity.

We returned to SEC-MALS to monitor complex formation with NiCl₂. First, separate injections of the individual proteins demonstrate they are well-resolved and that MALS estimates of the molecular masses are reasonably close to the actual values (Figure 5B and Figure S4): the 32.7 kDa H₆CF elutes at 13.5 min with an apparent MW of 34 kDa (black), the 142.7 kDa CheA dimer elutes at 11.5 min with an apparent MW of 139 kDa (blue), and the 18.1 kDa CheW monomer elutes at 16 min with an apparent MW of 18 kDa (cyan). These proteins were injected at the concentrations used for metal-mediated assembly (30 μ M H₆CF, 12 μ M CheA, and 24 μ M CheW). The H₆CF peak observed at 12 min with an apparent MW of 73 kDa (black) corresponds to the 65.4 kDa dimer but is hidden under the CheA peak once all proteins are combined. With all three proteins combined and injected at the same concentration as the individual injections (Figure 5B, red), the two peaks of H₆CF and CheW move together slightly, which suggests a possible interaction between the two proteins in solution or on the column. There is also a small shoulder at 11 min, which is likely to be a small amount of ternary complex formation, consistent with the <15% sedimentation observed under these conditions for an identical sample [0 time in Figure 5A, which has no Ni(II)].

To monitor the formation of a complex with time, a large volume was prepared to enable multiple injections of a single sample at different time points. While this experiment imparts a dilution (~13-fold) from injection to elution, which favors protein dissociation, dissociation caused by a 100-fold dilution of vesicle-assembled arrays is much slower than the time scale of this experiment.²⁴

The first changes observed by SEC during complex assembly involve primarily H_6CF . The difference between the no NiCl₂ and <1 min with NiCl₂ injections (Figure 5C, red and orange) is a decrease in H_6CF peak height and a corresponding increase in the intensity of a high-MW species that elutes at 11.1 min. The latter may consist of only H_6CF oligomers, because there is no significant change in the intensities of the CheA and CheW peaks. Next, incorporation of CheA and CheW into complexes with H_6CF appears to be nearly complete in 30 min, based on both the sedimentation data (Figure 5A) showing nearly complete binding at 30–60 min and the decrease in the intensities of the CheA and CheW SEC peaks (Figure 5C, yellow) with no further changes at longer time points. CheA

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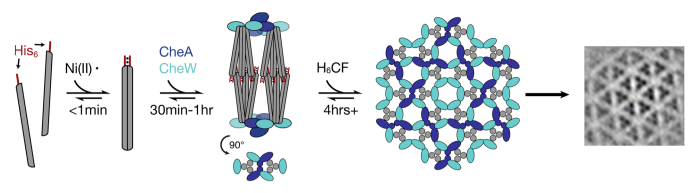


Figure 6. Model of array assembly by Ni(II)-mediated His-tag dimerization. Cartoon representation showing progression of assembly starting with CF monomers (side view, gray bars) rapidly forming dimers upon Ni(II) (black dots) binding to CF His-tags (red line). CheA dimers (blue) and CheW monomers (cyan) bind to stabilized H₆CF dimers to capacity within 30 min to 1 h. Binding of a CheA dimer and four CheWs drives assembly of a pair of trimers-of-receptor dimers into a core unit, which is likely further stabilized as a sandwich. At longer time points (\geq 4 h), the assemblies form hexagons and larger arrays with alternating CheA/CheW and CheW-only rings.

and CheW are presumably incorporated into a complex with H₆CF that elutes as a large SEC peak at 11.1 min. While the elution time is the same as that of the small shoulder seen immediately after the addition of NiCl₂ (Figure 5C, orange), this peak is highly polydisperse and thus likely contains a mixture of species with different molecular weights. At longer time points (1-4 h, green to blue to violet, Figure 5C), a second peak emerges at an even higher MW (arrow, 10.8 min) and the H₆CF peak continues to decrease to nearly zero intensity. These changes coincide with the assembly reaching maximum kinase activity, as shown in Figure 5A. The H₆CF, CheA, and CheW peaks remaining at 4 h are consistent with the assembly conditions: large amounts of free CheA and CheW because these are added in excess and only $\sim 20\%$ free H₆CF, consistent with sedimentation analysis that typically shows ~10-15% free H_6CF after assembly.

DISCUSSION

Here we report a novel method for assembly of the aspartate chemoreceptor cytoplasmic fragment (CF) into nanoarrays through metal-mediated His-tag dimerization and compare these to other in vitro nativelike arrays. Functional complexes of CF, CheA, and CheW form upon addition of specific divalent metals and assemble into an extended hexagonal lattice that exhibits the widely conserved, ^{5,35} native 12 nm center-to-center spacing (Figure 3). Intact chemoreceptor arrays are normally stabilized by a combination of anchoring in the membrane and cytosolic binding of CheA and CheW in rings at the membrane-distal cytoplasmic tip of the receptor. In current structural models of the complex, the long (~200 Å) cytoplasmic domain has protein-protein interactions with itself, CheA, and CheW only at its membrane-distal end, so the constraints imposed by its transmembrane and periplasmic domains are likely critical to the stability of the array. The necessary membrane-proximal stability to form CF arrays can be achieved with a membrane mimetic (templating vesicles)¹⁵ or with a sandwich architecture produced with PEG¹⁷ or divalent metals (this work). Importantly, all three in vitro preparations with CF yield high CheA occupancy and native stoichiometry (Figure 2C), although PEG arrays have previously been shown to lack the alternating hexagons of CheA observed for in vivo arrays.¹² Measured CheW stoichiometries of 2 monomers per 6 CF monomers in these assemblies provide further evidence of CheW-only rings that are proposed to provide structural integrity to the array.⁹

Chemoreceptors have been observed to form alternate nonnative structures in the absence or with insufficient quantities of their cytoplasmic binding partners, CheA and CheW. In particular, so-called "zippers" with interdigitation of the receptor cytoplasmic tips have been reported for overexpressed receptors in vivo,^{4,36} reconstituted intact receptors,³⁷ and CF assembled on templating vesicles.²⁴ A more detailed analysis of the overexpressed intact receptors revealed hexagonal packing with 9 nm center-to-center spacing that is distinct from that of the wild-type 12 nm arrays observed in both membrane-bound and sandwich architectures.¹² In this study, we have identified yet another structure formed by CF in the absence of CheA and CheW, double-stranded fibrils. ECT images (Figure S5A) demonstrate that H₆CF and NiCl₂ in the absence of CheA and CheW produce a mixture of nonspecific aggregates and fibrils with uniform 7 nm widths but varying lengths (40-105 nm). Interestingly, these fibrils and aggregates are at least partially reversible: addition of CheA and CheW after preincubation of H₆CF with NiCl₂ can produce complexes with kinase activity that is 60-80% of that of normal complexes assembled by adding NiCl₂ last (Figure S5B).

In contrast to the non-native "zippers" that form with cytoplasmic tips interdigitated in the center in the absence of sufficient CheA and CheW, "sandwiches" have receptors oriented in the opposite direction: cytoplasmic tips interact with CheA/CheW rings on both surfaces of the sandwich, forming hexagonal arrays with the canonical 12 nm center-to-center packing. Although sandwiches are not the native architecture for the *E. coli* aspartate receptor used in this study, similar cytoplasmic arrays exhibiting both 12 nm hexagonal lattices and sandwich architectures are observed for cytoplasmic chemoreceptors (lacking transmembrane domains) from a variety of bacteria and archaea, including *Vibrio cholerae*,¹⁹ *Rhodobacter sphaeroides*,¹⁸ and *Methanobacterium formicicum*.³⁸

The observed metal dependence of assembly is consistent with the proposed His-tag dimerization mechanism. The relative efficiency of the divalent metals in promoting active complex formation [Ni(II) > Zn(II) > Co(II)] follows the relative affinities of these metals for peptides containing multiple His residues.³⁹ This is consistent with preliminary X-ray absorption spectroscopy (XAS) data (H.-T. Huang and M. J. Maroney, unpublished observations), suggesting that the Ni(II) bound to H₆CF is coordinated by six ligands, including multiple histidines. Importantly, knowing Ni(II) is six-

coordinate and therefore paramagnetic suggests it may be possible to obtain distance constraints with paramagnetic relaxation enhancement NMR experiments involving comparison to the diamagnetic Zn(II) assembly that has an identical array architecture as determined by ECT (Figure S6). This could provide critical structural information in the overlap region of the sandwich where electron density is lost in ECT (Figure S2)

Characterization of the metal-mediated assembly of CF arrays by activity assays, sedimentation assays, and SEC-MALS allows us to propose a model for the mechanism of this assembly (Figure 6) and yields new insights into the assembly and properties of native chemoreceptor arrays. The fast (<1 min) initial increase in the dimer fraction (Figure 4) indicates that assembly begins with the stabilization of CF dimers, most likely by metals bridging two His tags. Subsequently, CheA and CheW are incorporated with metal-stabilized CF dimers into sedimentable complexes. However, binding of CheA and CheW is complete within 30-60 min, forming sedimentable complexes smaller than the fully active complexes that form in 4 h (Figure 5A,C). While our data cannot distinguish the exact size of these initial smaller complexes, Figure 6 shows one possibility that includes full binding of CheA and CheW into a sandwich of two core units, each containing 12 CFs (two trimers of dimers), bridged by one CheA dimer, and also containing four CheW monomers. It is unclear what changes occur between 1 and 4 h to yield larger complexes with higher activity that ultimately have the canonical hexagonal architecture. In one SEC-MALS series (Figure 5C), there was a small increase in the level of CF binding, while in another series (not shown), there was a small increase in the level of binding of all three proteins. We propose that building to larger complexes with maximal activity likely involves assembly of core units into hexagons that coalesce to form larger arrays and likely also involves additional binding of some CF, CheA, and/ or CheW to unoccupied sites in the coalesced array, as well as rearrangement of CheA and CheW into the native pattern that maximizes CheA binding in alternating hexagons of CheA/W¹² and only CheW. In any case, it appears that protein-protein interactions beyond those in the core unit are required for full activity.

The insights from this novel in vitro assembly mechanism may also shed light on how these remarkable arrays form in the cell. Briegel and others¹² proposed the following model for in vivo array assembly: (1) receptors dimerize, (2) three receptor dimers form a trimer of dimers, (3) pairs of receptor trimers of dimers are bridged by a CheA dimer and bind two CheWs to form the core unit, (4) three core units coalesce to form hexagons containing CheA and CheW, and (5) the array grows through binding of additional core units or of additional CheA/ W-filled hexagons. Our data support the importance of the initial receptor dimerization in the membrane, because metalmediated stabilization of the CF dimer is sufficient to drive array assembly. We see no evidence of formation of discrete trimers of CF dimers in the absence of CheA and CheW and instead observe that CF dimers form various oligomers and large aggregates, consistent with prior observations in vivo.⁴ We propose that in the absence of a stabilizing membrane or membrane mimetic, CheA and CheW are required to mediate assembly of receptor dimers into trimers of dimers. These are then bridged by CheA dimers into core units that also bind four CheWs. This would satisfy all of the receptor binding sites (two receptor-CheA interfaces, four receptor-CheW interfaces, and

six receptor-receptor interfaces) and thus prevent non-native interactions that lead to aggregation both in vitro and in vivo. Our data support incorporation of four CheWs into the core unit because there is little additional CheW binding upon assembly of larger arrays. Two of the four CheWs in the core unit are weakly bound (lacking the CheA-CheW proteinprotein interaction) and thus would likely be lost during purification of core complexes with full-length receptors in nanodiscs⁴⁰ (which contained two CheWs per core unit). These weakly bound CheWs are more likely to be retained under excess CheW conditions as in our CF assembly conditions. It seems likely that the additional two weakly bound CheWs in the core unit would increase the rate of assembly and, as previously suggested,⁹ increase the stability of the array. Finally, we propose that assembly of core units into hexagons is required to achieve maximal kinase activity.

Several important differences are expected between the Histag-mediated assembly of arrays of receptor fragments and in vivo assembly of membrane-bound chemoreceptor arrays. Studies of assembly of CF arrays cannot provide insights into the effects of the missing periplasmic, transmembrane, and HAMP domains, or the effects of ligand binding.⁴¹ Assembly of core units, hexagons, and arrays likely occurs faster in vivo because of alignment of receptors in the membrane facilitating encounters by two-dimensional diffusion within the plane of the membrane rather than by three-dimensional diffusion. Interestingly, although the cellular concentrations of the E. coli chemotaxis array proteins have been shown to vary significantly with strain and growth conditions, the protein ratios in the cell remain consistent at 2.9 receptors to 1 CheA to 1.2 CheWs.⁴² This excess CheA and CheW is similar to the conditions of our His-tag-mediated assembly of CF arrays, containing a ratio of 2.5 CF to 1 CheA to 2 CheWs. Although in vitro assembly does not duplicate the crowded conditions in the cell that alter protein diffusion rates and affinities, it may be possible to investigate whether excess CheA and CheW promote assembly as previously suggested.¹²

Assembly of large multiprotein complexes in vitro that retain native structure and function is a challenging but essential step in understanding how protein machines operate in the cell. We have demonstrated the value of His-tags to mediate dimerization and stabilize homogeneous, functional assemblies of chemotaxis signaling proteins. This dimerization strategy may prove to be valuable for assembly of other complex systems. Indeed, Cornelissen, van Hest, and co-workers discovered that metal-ion-induced stabilization can be used to stabilize a capsid assembly that is otherwise unstable in the absence of a high negative charge from either encapsulated RNA or low pH. Via the simple addition of divalent metals to the His-tagged cowpea chloric mottle virus capsid protein, capsid structures with native particle size form at neutral pH and can be used to encapsulate other proteins or drugs.43,44 Their model suggests the metal promotes His-tag-to-His-tag interactions that stabilize the large oligomeric capsid structure. His-tag-mediated dimerization is a means of stabilizing protein dimers that is much less perturbing than fusion to a protein dimerization motif. For example, fusion to a leucine zipper (LZ) introduces 30–60 residues, which is more perturbing than a six-residue His-tag and can interfere with assembly of native complexes. Interestingly, fusion of a 45-residue LZ to the chemotaxis receptor CF used in our study restored kinase activation but perturbed the assembly of native complexes with CheA and CheW. The LZ-CF assembled into sandwichlike

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particles in the opposite orientation (two CF layers on the outside of a middle layer of CheA and CheW) that prevented formation of the native array.^{45–47} This comparison of LZ- and His-tag-mediated dimerization of the same system demonstrates that using the least perturbing means of stabilization protein interactions can be critical to assembling native complexes. This approach has significant potential for application to other multiprotein systems, for the stabilization of functional complexes with native architecture. His-tags are widely used, and they are easily introduced at many locations within a recombinantly expressed protein. This should make it possible to drive dimerization at a known dimer interface with minimal perturbation to structure or function, for straightforward and accurate *in vitro* studies of the mechanisms of key processes in the cell.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.bio-chem.7b00698.

Protein sedimentation data for NiCl₂, ZnCl₂, and CoCl₂ assembly titration series; isosurface comparison of Ni(II)- and PEG-mediated sandwich arrays with intact receptor arrays; data sets of five two-dimensional tomographic slices used to estimate baseplate distances; multi-angle light scattering analysis and molar mass estimations for individually injected H₆CF, CheA, and CheW; partially reversible aggregation and fibril formation of H₆CF with NiCl₂; and structural analysis of Zn(II)-assembled complexes (PDF)

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E.R.H. and L.K.T. conceived ideas for the project. E.R.H., W.Y., and A.B. conducted the experiments. All authors analyzed the data. E.R.H. and L.K.T. wrote the manuscript, and all authors have given approval to the final version of the manuscript.

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Notes

The authors declare no competing financial interest.

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ABBREVIATIONS

BCA, bicinchonic acid; CF, cytoplasmic fragment of the *E. coli* aspartate chemotaxis receptor; DOGS-NTA, 1,2-dioleoyl-*sn*-glycero-3-{[*N*-(5-amino-1-carboxypentyl)iminodiacetic acid]-succinyl}; DOPC, 1,2-dioleoyl-*sn*-glycero-3-phosphocholine; ECT, electron cryotomography; ICP-OES, inductively coupled plasma-optical emission spectroscopy; IPTG, isopropyl β -D-1-thiogalactopyranoside; LZ, leucine zipper; PEG, polyethylene glycol 8000; PKB, phosphate kinase buffer; PMSF, phenyl-methanesulfonyl fluoride; SDS–PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis; SEC–MALS, size exclusion chromatography–multi-angle light scattering; TEV, tobacco etch virus; XAS, X-ray absorption spectroscopy.

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