Location and architecture of the Caulobacter crescentus chemoreceptor array

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Summary
A new method for recording both fluorescence and cryo-EM images of small bacterial cells was developed and used to identify chemoreceptor arrays in cryotomograms of intact Caulobacter crescentus cells. We show that in wild-type cells preserved in a near-native state, the chemoreceptors are hexagonally packed with a lattice spacing of 12 nm, just a few tens of nanometers away from the flagellar motor that they control. The arrays were always found on the convex side of the cell, further demonstrating that Caulobacter cells maintain dorsal/ventral as well as anterior/posterior asymmetry. Placing the known crystal structure of a trimer of receptor dimers at each vertex of the lattice accounts well for the density and agrees with other constraints. Based on this model for the arrangement of receptors, there are between one and two thousand receptors per array.

Introduction
Bacterial chemotaxis depends on two large macromolecular complexes: a chemosensory system that senses the environment and a motor that propels the cell (Bourret and Stock, 2002). The core of the chemosensory system is an array of transmembrane ‘chemoreceptors’ which binds to attractants or repellents in the surrounding media and regulates the activity of CheA histidine kinases attached to the receptors’ cytoplasmic domains. CheA phosphorylates CheY, which in turn diffuses to and binds the C-ring of flagellar motors. The relative ratio of phosphorylated and unphosphorylated CheY acts as a toggle switch, causing flagellar motors to switch the sense of their rotation (i.e. from clockwise to counter-clockwise or vice-versa), which in turn causes cells to either continue swimming forward (‘run’) or change their direction randomly (‘tumble’) in multilagellated Escherichia coli cells (Berg, 2003). In Caulobacter crescentus, the switch in sense of rotation causes short reversals in swimming direction (Alley et al., 1991). The chemoreceptors are also called methyl-accepting chemotaxis proteins (MCPs) because their activity is regulated by the methyltransferase CheR and the methyl-esterase CheB.

Chemoreceptors are linked at their proximal (cytoplasmic) tips by CheA and an additional protein CheW in such a way that they form tightly coupled arrays at the cell poles (Maddock and Shapiro, 1993; Bray et al., 1998; Lybarger and Maddock, 2001; Boukhvalova et al., 2002; Gestwicki and Kiessling, 2002). While high-resolution structures of all these components are available, including the cytoplasmic portion of MCPs (Kim et al., 1999; Park et al., 2006), CheA (Bilwes et al., 1999), CheW (Griswold et al., 2002), CheR (Djordjevic and Stock, 1997; 1998) and CheB (Djordjevic et al., 1998), the arrangement, size and position of the arrays with respect to the flagellar motors inside cells remains unclear. Understanding this ‘mesoscale’ architectural is critical, however, as it is vital to the arrays’ cooperative signal amplification and regulation (Armes et al., 2002; Gestwicki and Kiessling, 2002; Sourjik, 2004; Sourjik and Berg, 2004; Li and Hazelbauer, 2005; Parkinson et al., 2005). Based on crystal structures, two models of the arrangement of the MCPs, CheA and CheW in the array have been suggested (reviewed in Weis, 2006). In the ‘trimer of dimers’ model, the unit cell of a hexagonal lattice is formed by three receptor dimers (Shimizu et al., 2000). In the ‘hedgerows of dimers’ model, receptor dimers are arranged in rows (Park et al., 2006).

Cryo-EM studies of soluble complexes of a partial receptor with CheA and CheW show an alternative ‘barrel’-like pattern (Francis et al., 2004; Wolanin et al., 2006). In order to elucidate the architecture of the chemotaxis system, we visualized it directly within intact C. crescentus cells by electron cryotomography (ECT; Lucic et al., 2005; Jensen and Briegel, 2007). C. crescentus is a

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Gram-negative bacteria characterized by a dimorphic life cycle. Motile ‘swarmer’ cells mature by releasing their single polar flagellum and developing a tube-like stalk and terminal holdfast which attaches the cells to surfaces (Wagner and Brun, 2007). These non-motile, ‘stalked’ cells then elongate and undergo multiple rounds of division, each time producing and releasing a swarmer progeny. In *C. crescentus*, the chemosensory and propulsion systems are coupled in space and time: chemoreceptor and flagellar gene expression is co-ordinated (Jensen et al., 2002), chemoreceptors localize to the flagellated pole in swarmer cells (Alley et al., 1992), and stalked cells contain neither chemoreceptors nor flagella (Shaw et al., 1983; Gomes and Shapiro, 1984; Alley et al., 1992). By recording both fluorescent light microscopical and electron tomographical images of particular *C. crescentus* cells, here we identify the chemoreceptor array in *C. crescentus* tomograms and describe its size, location and structure within intact cells. The results confirm previous electron microscopical studies that identified a similar structure in *E. coli* as the chemoreceptor array (Zhang et al., 2007), and support a modified ‘trimers of dimers’ network model for its architecture.

**Results**

**Tomograms of wild-type cells**

About 50 three-dimensional reconstructions (tomograms) of intact, frozen-hydrated, log-phase *C. crescentus* cells at various stages of the cell cycle were produced which revealed the typical ultrastructure of *C. crescentus*: an S-layer, outer membrane, peptidoglycan and inner membrane enclosing a cytoplasm containing ribosome-like particles, cytoskeletal filaments and occa- sional vesicles (Briegel et al., 2006). Most cells exhibited a stalk, a minority exhibited a flagellum, and still others exhibited no appendages at all, presumably because the cell was in the transitional state between shedding the flagella and forming a stalk. In 12 cells, we observed a plate-like cytoplasmic assembly (Fig.1) that closely resembled the assemblies identified previously as chemoreceptor arrays in *E. coli* (Zhang et al., 2007). The ‘base’-plate was positioned 31 nm below (proximal to) the inner membrane (measured ‘peak-to-peak’), bent to follow the curvature of the cell wall. A series of thinner densities perpendicular to the base plate and membrane spanned the distance between them. These thin connecting densities passed through a second, fainter plate parallel to the cell wall about 10 nm below the inner membrane. In the nine cells where pili and/or flagella were observed, the plate-like assembly colocalized to the convex side of the same pole (see Movie S1). The distance between the plate-like assemblies and the flagellar motors ranged from 17 to 130 nm. The assembly was not found in any of the stalked cells. The area of the inner membrane covered by these assemblies ranged from ~11 000 to 22 800 nm². The assembly’s location, timing within the cell cycle, and resemblance to the chemoreceptor arrays of *E. coli* all suggested that it was the chemoreceptor array.

**McpA mutants**

To confirm whether this assembly was in fact the chemoreceptor array, we imaged a *C. crescentus* strain carrying a fusion of the red-fluorescent protein mCherry to the C-terminus of the mcpA gene on a xylose-inducible plasmid. Because expression of the mcpA–mCherry fusion was uncoupled from the cell cycle, after cells were incubated in the presence of xylose for a few hours every cell exhibited a bright fluorescent signal at one of the poles, regardless of its developmental stage. In the case of dividing cells, bright signals were seen at both poles. Such induced cells were plunge-frozen and tomograms of 13 poles were produced. Plate-like assemblies were now found in every cell, whether stalked or flagellated (Fig. 2), and always at the pole with the appendage, as had been seen previously in *Caulobacter* mutants where the carboxyl-terminal degradation domain of McpA was eliminated (Alley et al., 1993). The assemblies could also now be found on the concave as well as convex side of the cell or at the tip of the pole. Thus the pattern of mislocalization of the fluorescently tagged McpA matched that of the plate-like assemblies.

To further investigate these correlated mislocalizations, we developed a new method to obtain both a fluorescent image and a cryotomogram of the same cell. After being induced with xylose, mutant cells were concentrated, lightly fixed, immobilized with poly-L-lysine on gold TEM ‘finder’ grids, sandwiched between a glass slide and coverslip together with excess media, and imaged in a high-powered oil-immersion fluorescence light microscope (fLM) (Fig. 3A). The grids were then gently but quickly removed from the glass slide, plunge-frozen and transferred to the electron cryomicroscope. The same region observed by fLM was located in the TEM (Fig. 3B) and tilt-series of specific cells were recorded. Altogether, seven data sets from two different grids were matched containing a total of nine cell poles (two poles are shown in Fig. 3C and D). The presence and absence of the putative chemoreceptor arrays in the tomograms correlated perfectly with the presence or absence of fluorescent foci (seven poles exhibited both a fluorescent focus and a plate-like assembly in the same location and two poles showed neither). We conclude that the plate-like assemblies are in fact the chemoreceptor arrays of *C. crescentus.*
Fig. 1. Reconstructions of wild-type *C. crescentus* swarmer cells. Representative ~12 nm thick slices through the centres of three *C. crescentus* swarmer cells are shown. At the convex side of the flagellated pole, a plate-like assembly (white arrows) is visible. The prominent base plate is connected to the inner membrane by thin, wall-like densities. A second, fainter layer of density parallel to the membrane can be discerned between the base plate and inner membrane. In most cases, pili (black arrows in A) and flagella (not shown) were also present at the same cell pole, but not in the same slice where the plate-like assembly was most visible (the case for the cells in B and C). Scale bar: 100 nm.
C. crescentus chemoreceptor arrays are hexagonally packed

In ‘side’ views such as that seen in Fig. 1, the chemoreceptor arrays were composed of the base plate and thin, regularly spaced, wall-like densities connecting the base with the inner membrane (an enlarged view of another example is shown in Fig. 4A). ‘Top’ views of the region right above the base layer (toward the inner membrane) showed a honeycomb-like arrangement of the wall-like densities (Fig. 4B). The power spectrum of this region revealed a hexagonal lattice with a spacing of 12 nm (Fig. 4B, inset). While applying a median filter to the tomogram reduced background noise (Fig. 4C and D), to further improve the signal-to-noise ratio we extracted and averaged seven subvolumes with obvious hexagonal ordering from one (unfiltered) tomogram and applied sixfold rotational symmetry to the resulting average. The side view of this averaged structure is shown in Fig. 5A, as well as several ‘top-views’ at indicated positions between the base plate and inner membrane (Fig. 5B–F). From the base plate (pink in Fig. 5G), a honeycomb-like hexagonal net was seen rising towards the membrane. Closer to the inner membrane the hexagonal order quickly deteriorated until it was no longer discernable. By measuring the surface area on the inner membrane covered by the arrays on several representative cells, the arrays were estimated to contain between 90 and 180 hexagons.

Discussion

Our understanding of many biological systems has progressed from knowledge of their basic functions to identification of the macromolecules involved to atomic models of each of their structures, but before accurate mechanistic computational models can be constructed, the details of how the components are arranged within living cells must also be elucidated. Work on bacterial chemotaxis has followed this pattern, resulting in a need for architectural details to constrain and refine already sophisticated computational models (Bray et al., 1998; Lipkow et al., 2005; Bray et al., 2007). Here we used a combination of fLM and ECT of wild-type and mutant cells to identify and describe the C. crescentus chemoreceptor array in intact cells.

A major technical challenge in ECT has been to unambiguously identify structures seen in tomograms. There are as yet no ‘GFP-like’, genetically encodable, electron-dense labels that would work in this context (ECT of intact bacterial cells). Instead, structures like cytoskeletal filaments have been identified by correlating their size, position or number with changes in their expression levels or stability (Li et al., 2007). Likewise, chemoreceptor arrays had previously been identified in images and tomograms of frozen-hydrated E. coli cells based on their polar localization, increase in number in cells overexpressing chemoreceptors, or dependence on CheA and CheW (Weis et al., 2003; Lefman et al., 2004; Zhang et al., 2004;
**Fig. 3.** Correlated fLM and ECT of McpA mutant cells.

A. fLM image of slightly fixed, xylose-induced McpA–mCherry mutant cells immobilized on an EM finder grid with regularly spaced, 2 μm diameter holes in a thin carbon film. Most cells exhibit a single fluorescent focus at one pole, but elongated pre-divisional cells exhibit foci at both poles.

B. Low magnification EM image of the same region after the grid had been removed from the light microscope, plunge-frozen and inserted into the EM. Full tilt-series were recorded of the five circled cells, which maintained their positions well during the transfer.

C and D. 15 nm thick slices through the cell poles of cell #3 (C) and #4 (D). The presence, absence and positions of the plate-like assemblies (white arrow) correlated perfectly with fluorescence signals from McpA. Scale bars: 50 nm.
Immunolabelling has also been used to help identify the chemoreceptor array (Weis et al., 2003; Lefman et al., 2004; Zhang et al., 2007). Our work here confirmed that identification through correlated fLM and ECT, a powerful combination of complementary microscopies.

To facilitate correlated light and electron cryomicroscopy, two groups recently developed cryostages that allow frozen-hydrated samples to be observed in light microscopes (Sartori et al., 2007; Schwartz et al., 2007). These stages allow near-native preservation of the sample while both fLM and cryo-EM images are recorded, without any changes in the sample except the unavoidable radiation damage. Unfortunately, because the cold temperatures prohibit the use of immersion oil and all but long-working-distance objective lenses, the resolution of these cryo-light microscopes (including the one we tested) is inadequate to discern localization patterns within small bacterial cells.

In order to obtain both high-resolution fLM and ECT images of the small C. crescentus cells of interest here, we immobilized them on EM finder grids, investigated their fluorescence patterns at room temperature and then plunge-froze them in preparation for electron cryomicroscopy. Minor changes in the sample did occur during the transfer from the light to the electron microscope, including positional shifts of some bacterial cells or even their complete detachment from the grid, and the structures of individual cells were slightly perturbed by the light fixation. We were nevertheless able to unambiguously identify the plate-like assemblies as chemoreceptor arrays by showing that they consistently and precisely colocalized with fluorescently tagged chemoreceptor subunits.

Once the arrays were identified, their architectural features were deduced from cryotomograms of unfixed, wild-type cells. Chemoreceptor arrays were seen to comprise a plate-like base parallel to and ~31 nm below the inner membrane; a series of wall-like densities that extended from the base to the inner membrane; and a second and weaker plate-like density ~10 nm below the inner membrane. The wall-like densities formed a hexagonal net with 12 nm spacing whose order deteriorated rapidly with distance from the base. Given that (i) CheA and CheW are known to bind each other and the cytoplasmic tips of MCPs (Falke and Hazelbauer, 2001; Boukhvalova et al., 2002); (ii) that at the other end of the receptors there is a hydrophobic region that must traverse the membrane to put the periplasmic domain in contact with extra-
cellular stimuli (Maddock and Shapiro, 1993; Sourzik and Berg, 2004); and (iii) that the distance between the base layer and the inner membrane (31 nm) is consistent with the estimated length of MCPs (Kim et al., 1999; Park et al., 2006), we interpret the thin walls as the receptors and the ‘base plate’ their complex with CheA and CheW.

While the resolution in our tomograms was not high enough to reveal the exact packing of MCPs, trimers of receptor dimers, as seen in the crystal structure (Kim et al., 1999), fit compellingly well into the vertices of the hexagonal lattice to form linked rings of receptors (Fig. 6). Cross-linking assays had previously pointed to a ‘trimer-of-dimers’ organization of the receptors in vivo (Studdert and Parkinson, 2004), and a model derived from these studies predicted that the trimers would form the vertices of a hexagonal lattice (Studdert and Parkinson, 2005). Our model also agreed with genetic studies that found an important signalling role for trimers (Ames et al., 2002).

Finally, recent studies using small particles of lipid bilayer containing one or a few chemoreceptors, called nano-discs, further implicated ‘trimers-of-dimers’ as the core structure of kinase activation and control (Boldog et al., 2006). In contrast, other models based on ‘hedgerows’ (Park et al., 2006) or larger numbers of receptors per hexagon were incompatible with our reconstructions. Our data were also unreconcilable with models where trimers of receptor dimers stood alone in the centres of CheA/CheW hexagons (Shimizu et al., 2000), as the receptors formed apparently continuous walls which emerged from the thickest regions of the CheA/CheW base plate.

Seven major MCP classes have been identified by large-scale comparative genomic analyses based on the...
length and sequence conservation of the domains. \textit{C. crescentus} contains 18 different MCPs. Sixteen of these are of type 36H, as are all five MCPs found in \textit{E. coli} (Alexander and Zhulin, 2007). Of these 16 type 36H MCPs in \textit{C. crescentus}, six contain the C-terminal pentapeptide tether to which CheR and CheB bind in \textit{E. coli} (Alexander and Zhulin, 2007). The other two MCPs in \textit{C. crescentus} are of type 38H and an un-aligned, ‘cytoplasmic’ type. Interestingly, despite these sequence similarities, the lattice spacing we observed in wild-type \textit{C. crescentus} cells (12 nm) differed significantly from the lattice spacing reported in EM studies of \textit{E. coli} overexpressing the Tsr receptor (7.5 nm) (Weis et al., 2003; Lefman et al., 2004). It is unclear whether the receptor packing in the \textit{E. coli} study was altered owing to the elevated Tsr receptor concentrations, if the arrays were in different states, or if the architectures of \textit{E. coli} and \textit{C. crescentus} chemoreceptor arrays are simply different.

In any case, based on the surface area of the arrays observed here (11 000–22 800 nm$^2$), typical \textit{C. crescentus} chemoreceptor arrays contain $\sim$90–180 hexagons. If in fact each intersection in the hexagonal lattice is composed of a trimer of receptor dimers, as we propose, the threefold axes of nearest-neighbour trimers of dimers would be 6.9 nm apart, and there would be $\sim$1100–2200 individual receptors per array. When grown in rich media, the number of chemoreceptors per \textit{E. coli} cell has been estimated biochemically to range from 3600 to 15 000, depending on the bacterial strain, the growth phase and the media (Li and Hazelbauer, 2004). Given that \textit{E. coli} cells are roughly four times as voluminous as \textit{C. crescentus} cells, the concentration of receptors appears to be similar.

The cytoplasmic base layer then must correspond to the complex formed by CheA, CheW and the cytoplasmic tips of the MCPs. Early models of the CheA-CheW layer were based on atomic models of CheA, CheW and the cytoplasmic portion of a receptor (Shimizu et al., 2000). Our results fail to support this model, as the predicted lattice spacing (20 nm) is almost twice as large as we observed. More recently, a structure of a CheA-CheW dimer was determined by crystallography and pulsed ESR of \textit{Thermotoga maritima} proteins, from which a different model of the CheA-CheW-MCP$^{1143C}$ complex was proposed (Park et al., 2006). In this model, the CheA and CheW proteins are arranged in a lattice with a spacing of 13.7 nm, which is just slightly larger than the lattice spacing we observed here in \textit{C. crescentus}, but the CheW dimer cleft in the \textit{T. maritima} model is too small to accommodate the trimers of receptor dimers that appear to be present here (Park et al., 2006).

An even larger complex of CheA, CheW, and the cytoplasmic domain of an MCP has also been generated, first by fusing a leucine zipper dimerization domain to the aspartate receptor Tar (Francis et al., 2002; 2004), and later using the serine receptor Tsr without the dimerization domain (Wolanin et al., 2006). These complexes formed elongated ‘head-to-head’ barrel-like structures with two hollow cavities. The authors interpreted each side of the structure (one barrel) to be two sets of three receptor dimers, standing parallel in asymmetric rings to create each hollow cavity in the centre. The heads of the two barrels were linked by a double CheA/CheW plate in the middle. While a similar number of receptor monomers (24) would be present in each ring of our proposed hexagonal lattice, and the diameter of the barrel and our rings were roughly consistent, the two models differ substantially in other ways. First, the barrel-like structures seemed to be formed by two non-symmetric sides containing three uniquely arranged receptor dimers per side, whereas in our reconstruction the receptors appear to be hexagonally packed, at least near the CheA/CheW base. Second, in the barrel-like structures, two sets of receptors were joined ‘head-to-head’ by a doubly thick CheA/CheW layer. Because a similar arrangement had been seen in \textit{E. coli} cells overproducing the chemotaxis receptor Tsr, where membrane invaginations allowed the signalling domains of two receptor arrays to lie flat against each other (as if a single array had folded over on itself; Lefman et al., 2004), it was proposed that this configuration might represent the activated conformation of the receptors (Wolanin et al., 2006). As in our tomograms of wild-type \textit{C. crescentus} cells, neither membrane invaginations nor doubly thick
receptor arrays were seen, and because CheA, CheW and the receptors are all highly conserved across species, it is unlikely that the signalling domain/signalling domain juxtaposition represents the natural configuration of activated receptors (Zhang et al., 2007), but instead occurs only under artificial conditions.

Thus, none of the existing specific models of the CheA/CheW/MCP array agree well with all our experimental findings, although other patterns such as the one proposed in Studdert and Parkinson (2005) could. Our data also suggest that the receptors are well organized near the CheA/CheW base, but lose that order as they rise towards the inner membrane. The region between the CheA/CheW base and the inner membrane has been called the ‘adaptation compartment’, where it is thought that CheR and CheB regulate the methylation of the receptors (Shimizu et al., 2000). In our tomograms there was a second, faint layer of density where the methylation sites are expected (Ames and Parkinson, 1994; Ames, 1996), ~10 nm below the inner membrane. This layer corresponds to the interaction sites of CheR and CheB with the MCPs, and is consistent with the proposed ‘brachiating mechanism’ of CheR (Levin et al., 2002). In this model, one of CheR’s two binding domains (either the one for the receptor’s flexible tether or the one for its methylation site) stays connected to a first receptor while the other domain binds to a second, neighbouring receptor, leading to a ‘hand-over-hand’ movement of the enzyme through the array. The small number of CheR and CheB molecules compared with receptors in an array (~400–15 000 in E. coli; Li and Hazelbauer, 2004), might not, however, be sufficient alone to explain the observed layer of density in that region.

Finally, the polar localization of the chemoreceptor array in swarmer cells had been shown previously (Alley et al., 1992), but our studies refined its location to the convex side of the flagellated cell pole, 17–130 nm away from the flagellar motor. While it is already known that the cytoskeletal protein Crescentin localizes to the concave side of C. crescentus cells, where it endows cells with their characteristic crescent shape (Ausmees et al., 2003), the presence of the chemotaxis array on only the convex side further highlights how C. crescentus cells maintain dorsal/ventral as well as anterior/posterior asymmetry. Whether this second dimension in intracellular differentiation is unique to C. crescentus or is widespread among bacteria has yet to be determined.

Experimental procedures

Specimen preparation

CB15N cells were grown at 30°C in peptone yeast extract (PYE) medium (Ely, 1991) to an optical density at 600 nm between 0.2 and 0.4. Two millilitres of the cell suspension was pelleted for 5 min at 1500 rcf. The pellet was rinsed in M2 Salt solution (Ely, 1991), then re-suspended in 30–50 μl of fresh M2 solution. The EM R2/2 copper/rhodium grids were glow-discharged and coated with a 3×-concentrated, 10 nm colloidal gold solution (Ted Pella, Redlands, CA). A 5×-concentrated solution of 10 nm colloidal gold was added to the cells immediately before plunge freezing. A 4 μl droplet of the sample solution was applied to the EM grid, then automatically blotted and plunge-frozen in liquid ethane using a Vitrobot (FEI Company, Hillsboro, OR) (Iancu et al., 2007). The grids were stored under liquid nitrogen until data collection.

Synchronized cells

CB15N cells were synchronized according to published protocols (Tsay and Alley, 2001). One millilitre of synchronized cells was pelleted for 5 min at 1500 rcf, re-suspended in 30–50 μl M2 salt solution and plunge-frozen as described above.

Caulobacter McpA mutant

A fusion of the C-terminus of the Caulobacter McpA protein to the red-fluorescent protein, mCherry, was expressed in Caulobacter cells from a xylose-inducible promoter. Cells were grown overnight in PYE media containing 5 μg ml⁻¹ kanamycin. One hundred microlitres of the cell suspension was added to 5 ml fresh PYE/kanamycin medium and induced with 0.03% xylose for at least 2 h prior to imaging. For plunge-freezing, 2 ml of the cell suspension was pelleted for 5 min at 1500 rcf and re-suspended in ~100 μl supernatant. Concentrated 10 nm colloidal gold solution treated with BSA was added to the concentrated cells directly before freezing. A 4 μl droplet of the sample solution was manually applied to the grid, automatically blotted, then plunged into a mixture of ethane (37%) and propane (63%) (W.F. Tivol, A. Briegel and G.J. Jensen, unpublished), using a Vitrobot (FEI Company, Hillsboro, OR).

Correlated fLM-ECT of McpA mutants

A 2 ml aliquot of cell culture was pelleted for 5 min at 1500 rcf. The pellet was re-suspended in 1 ml fixing solution containing 0.1% glutaraldehyde, 3% paraformaldehyde and 20 mM sodium phosphate (pH 7) and incubated for 30 min on ice. The cells were then pelleted, washed three times in washing buffer (140 mM NaCl, 3 mM potassium KCl, 8 mM Na₂HPO₄, 1.5 mM KH₂PO₄ and 0.05% Tween) and re-suspended in the imaging buffer containing 50 mM glucose, 10 mM EDTA and 20 mM Tris-HCl (pH 7.6). A 4 μl droplet of this solution was applied to an H2 gold finder TEM grid covered with R2/2 Quantifoil and treated with 4 μl of 0.05% poly-L-lysine in ddH₂O. After blotting the grid from the side, the grid was gently rinsed with buffer. The grid was transferred onto a droplet of imaging buffer on a glass slide and covered with a coverslip. Imaging was carried out in a Nikon 90i fluorescence microscope, then the grid was carefully taken from the slide, 3 μl droplet of BSA-treated, 5×-concentrated, 10 nm gold solution was added to it, and

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then it was plunge-frozen into liquid ethane as described above. Tilt-series were recorded of the same cell clusters imaged in the fLM.

**EM data collection**

EM images were collected using a FEI Polara™ (FEI Company, Hillsboro, OR, USA) 300 kV FEG transmission electron microscope equipped with a Gatan energy filter (slit width 20 eV) on a 2 x 2 k Gatan Ultrascan CCD camera or later a lens-coupled 4 x 4 k Ultracam (Gatan, Pleasanton, CA). Pixels on the CCD represented between 0.67 and 1.2 nm on the specimen. Tilt-series from −60° to 60° with an increment between 0.5° and 1° were recorded semi-automatically around one or two axis (Iancu et al., 2005) at 10 and 12 μm underfocus using the predictive UCSF-Tomo package (Zheng et al., 2004). A cumulative dose of 200 e/nm² or less was used for the directly plunge frozen samples and 100 e/nm² or less for the correlated fLM-ECT tomograms.

**Image processing**

Three-dimensional reconstructions were calculated using the IMOD software package (Mastronarde, 1997). In order to obtain a higher signal-to-noise ratio map of the chemoreceptor array, we manually segmented an array with Amira (Mercury Computer Systems). Using a similar approach as described in Wright et al. (2007), we selected a layer just above the CheA/ChemW base plate (where the hexagonal lattice was most visible) and projected that layer onto the surface of the array. Using a semi-automatic peak detection tool, we picked seven hexagonal centres from the best-ordered regions of the surface projection. Noting their positions, the normal vectors to the surface, and the directions to the closest neighbouring hexagonal centres, we extracted subvolumes surrounding each chosen centre, superimposed them by rotation and translation, averaged the set and finally applied sixfold rotational symmetry.

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Supplementary material
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