Bacterial Energy Sensor Aer Modulates the Activity of the Chemotaxis Kinase CheA Based on the Redox State of the Flavin Cofactor*

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Flagellated bacteria modulate their swimming behavior in response to environmental cues through the CheA/CheY signaling pathway. In addition to responding to external chemicals, bacteria also monitor internal conditions that reflect the availability of oxygen, light, and reducing equivalents, in a process termed “energy taxis.” In Escherichia coli, the transmembrane receptor Aer is the primary energy sensor for motility. Genetic and physiological data suggest that Aer monitors the electron transport chain through the redox state of its FAD cofactor. However, direct biochemical data correlating FAD redox chemistry with CheA kinase activity have been lacking. Here, we test this hypothesis via functional reconstitution of Aer into nanodiscs. As purified, Aer contains fully oxidized FAD, which can be chemically reduced to the anionic semiquinone (ASQ). Oxidized Aer activates CheA, whereas ASQ Aer reversibly inhibits CheA. Under these conditions, Aer cannot be further reduced to the hydroquinone, in contrast to the proposed Aer signaling model. Pulse ESR spectroscopy of the ASQ corroborates a potential mechanism for signaling in that the resulting distance between the two flavin-binding PAS (Per-Arnt-Sim) domains implies that they tightly sandwich the signal-transducing HAMP domain in the kinase-off state. Aer appears to follow oligomerization patterns observed for related chemoreceptors, as higher loading of Aer dimers into nanodiscs increases kinase activity. These results provide a new methodological platform to study Aer function along with new mechanistic details into its signal transduction process.

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The abbreviations used are: MCP, methyl-accepting chemotaxis protein; ETC, electron transport chain; MSP, membrane scaffold protein; ASQ, anionic semiquinone; HQ, hydroquinone; DT, dithionite; cw-ESR, continuous wave electron spin resonance; PDS, pulsed-dipolar ESR spectroscopy; DEER, double electron-electron resonance; DQC, double quantum coherence; DDM, n-dodecyl-β-D-maltopyranoside.

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reduction to the anionic semiquinone (ASQ). Aer assembles into higher-order complexes with CheA and CheW, where it regulates CheA activity based on the redox state of FAD. No evidence is found for a stable hydroquinone form of the protein.

Results

Detergent-solubilized Aer Undergoes Reversible Chemical Reduction—N-terminally His$_6$-tagged *E. coli* Aer was expressed in BL21(DE3) cells and purified in 0.1% n-dodecyl-$\beta$-D-maltopyranoside (DDM) using Co$_2$ affinity chromatography (Fig. 1B). The purified Aer shows signature UV-visible spectra of protein-bound FAD in the fully oxidized state with three absorption peaks at 416, 441, and 465 nm and a relatively broad shoulder centered at 365 nm (Fig. 1C). Aer binds approximately one FAD cofactor per subunit (Fig. 1D). Upon treatment with 10 mM dithionite in anaerobic conditions, the UV-visible spectrum of the Aer FAD converts to that of the single-electron reduced ASQ state as reflected by the concomitant increase at 372 and 500 nm, and disappearance of the 441 nm peak. A shoulder at 395 nm, which is a defining characteristic of the FAD ASQ state, could also be detected. The ASQ reverts back to the fully oxidized cofactor on exposure to air (Fig. 1C) (17).

The continuous wave electron spin resonance (cw-ESR) spectrum confirms the radical nature of the dithionite-reduced FAD, which becomes ESR-silent upon reoxidation (Fig. 1E). The magnetic dipolar coupling between the flavin radicals can be used to measure their distance of separation by pulsed-dipolar ESR spectroscopy (PDS). PDS of the FAD ASQ state reveals one distance peak centered at 41.3 Å, suggesting purely dimeric association of Aer in DDM (Figs. 1B and 2A). These data also define a structural constraint on the disposition of the PAS domain in Aer (Fig. 2B). Interestingly, prolonged anaerobic incubation of Aer with 16 mM dithionite (DT) does not yield the two-electron reduced HQ state. DDM is detrimental to CheA activity, and thus nanodisc incorporation was applied to stabilize Aer in the absence of detergent (18–20). Nanodisc incorporation is extremely effective at reconstituting transmembrane MCPs in active forms (14, 21, 22), and the same holds true for Aer.

Aer Reversibly Regulates CheA Activity Based on FAD Redox State—Aer was incorporated into nanodiscs assembled with *E. coli* polar lipids and the membrane scaffold protein (MSP), MSP1E3D1 (Fig. 3A, inset). As in detergent, Aer in nanodiscs can be reversibly reduced to the ASQ state upon dithionite treatment as revealed by UV-visible (Fig. 3A) and cw-ESR (Fig. 3B) spectroscopies. Nanodisc-incorporated Aer with fully oxidized FAD activates CheA to produce phosphorylated CheY,
whereas upon chemical reduction to the ASQ, CheA activity is inhibited (Fig. 3C). Aer reoxidation by ambient oxygen recovers a majority of the activity. The reason for incomplete recovery is unknown, although it is not uncommon for DT to adversely affect protein activity. DT has little effect on CheA activity alone. Thus, Aer with oxidized FAD activates CheA and would therefore cause cell tumbling. Reduction of Aer to the ASQ reversibly inhibits kinase activity and would cause smooth swimming.

In nanodiscs, we still could not generate a stable HQ state of Aer FAD by prolonged incubation with dithionite in anaerobic conditions even in the presence of redox mediators, which facilitate outer sphere electron and proton transfer reactions to FAD. Another reductant, Cr-EDTA, also failed to reduce the cofactor to the HQ state. As free FAD is reduced to the HQ state by dithionite, chemical reduction of Aer was tested in the presence of additional FAD with the expectation that the exchange of the Aer FAD with free HQ is not favored.

**FIGURE 2. PAS domain separations in Aer.** A, PDS-derived distance distributions between ASQ states of FAD in Aer solubilized in DDM and incorporated into nanodiscs (Aer:MSPE3D1 = 1.43). Both $P(r)$s are plotted normalized to the area of major peaks. The insets show the fits to the time domain DEER data for both cases. DQC data for detergent are very similar to respective DEER data (not shown). B, model of Aer based on data in Ref. 28. The 42 Å flavin separation agrees well with a tight interaction between the PAS β-sheet and HAMP helix AS2. Dotted lines represent an ~40-residue unmodeled linker between PAS and the first transmembrane helix. An F1 linker helix between PAS and HAMP is also not shown.

**FIGURE 3. Aer activity in nanodiscs depends on redox state.** A and B, UV-visible spectra (A) and cw-ESR spectra (B) at different times during dithionite reduction and aerobic reoxidation of FAD in Aer incorporated in nanodiscs. Abs., absorbance; AU, absorbance units. The inset of A displays the bands for Aer and MSPE3D1 on an SDS-PAGE gel after nanodisc reconstitution and purification. C, reversible regulation of CheA phospho-transfer activity based on the FAD redox state. Phospho-transfer to CheY is shown relative to that for fully oxidized FAD (designated as Aer), which has been normalized to 100. The error bars display the S.E. with $n=6$. The inset displays a representative autoradiograph, and the individual data points are shown as open black circles. D, the effect of additional free FAD on the chemical reduction of Aer. The UV-visible absorbance spectrum of Aer in nanodiscs with additional free FAD (0.1 mg/ml) is shown after subtracting the spectrum for free FAD alone. Likewise, the spectrum for dithionite-treated Aer in nanodiscs with additional FAD is shown after subtracting the spectrum for the reduced, free FAD. The light-colored spectra are for Aer without additional free FAD added, whereas the dark-colored spectra are for the Aer contribution in the presence of oxidized (blue) or fully reduced (red) free FAD.
Activity Depends on Oligomerization of Aer Dimers—Ternary signaling complexes of the major \textit{E. coli} MCPs Tar (aspartate receptor) and Tsr (serine receptor) with CheA and CheW require a trimer of receptor dimers to activate CheA (14, 23–26). To investigate whether association of dimers is important for Aer activity, we varied the molar ratio of Aer to MSP1E3D1 for the reconstitutions to alter the average number of Aer dimers in each nanodisc, as was demonstrated previously for Tar (14). Notably, two MSP proteins are required for nanodisc formation, and thus the preparation with Aer to the MSP1E3D1 ratio of 0.16 will mainly produce single Aer dimers in the discs. Size exclusion chromatograms for two preparations (Aer:MSP1E3D1 = 0.16 and 0.77) reveal a shift of the nanodisc-incorporated Aer to a smaller elution volume in the 0.77 preparation, implying substantially larger particles with higher Aer:MSP subunit ratios (Fig. 4A). Increasing the molar ratio of the receptor in the preparations (14, 27) raises the probability of three dimers aligned on the same side of the nanodisc. Interestingly, distance distribution measurement between FAD ASQs in the highest ratio preparation (1.43) gives two peaks centered at 41.5 and 52.4 Å, respectively (Fig. 2A). The peak at smaller distance arises from the intersubunit separation (which matches well to the distance in detergent). The narrow breadth of the shorter distance distribution indicates a relatively stable juxtaposition of the PAS domains, whereas the ~42 Å separation is too far for a PAS-PAS dimer. However, a close association between the two PAS and HAMP domains (Fig. 2B), in accordance with identified PAS-HAMP interface residues (28), agrees well with the PDS data. Although the PDS peak at longer separation is rather weak and cannot be confirmed with full certainty, the longer distance (absent in detergent) may arise from different dimers on the same side of the nanodiscs. Accordingly, the ability of CheA to catalyze phospho-transfer to CheY is modulated based on the preparation ratio for the Aer incorporation into nanodiscs (Fig. 4B). The kinase activity shows a bell-shaped dependence on the molar ratio, which is very similar to what has been observed for Tar (14), suggesting that Aer requires the similar trimeric assembly of other MCPs to achieve maximum kinase activation. However, due to the low cellular abundance of Aer (~2% of all MCPs (29, 30), it may associate as mixed trimers with the related high abundance chemoreceptors Tar and Tsr (31, 32).

Discussion

Here, we show that Aer modulates CheA activity based on FAD redox states, and like other chemoreceptors, likely functions as a trimer-of-dimers. The prevailing model (5, 15) conjectured a third HQ state to explain the cellular tumbling in the presence of substrate and absence of terminal electron acceptors. However, in our \textit{in vitro} experimental conditions, even the low-potential reductant DT was unable to produce a fully reduced HQ state. Notably, the related flavin-binding protein \textit{Azotobacter vinelandii} NifL reduces to the HQ upon DT treatment (33, 34). Further reduction of Aer may be limited by its preference to form the anionic instead of the neutral semiquinone. Reduction to the hydroquinone requires proton transfer, which may be kinetically disfavored under these conditions. If oxygen were to react directly with Aer, it would not be expected to produce the observed smooth swimming response because this behavior requires reduced FAD, which \textit{O}_2 destablizes. However, oxygen can favor FAD reduction by rather acting as an acceptor to the ETC.

Interface studies indicate that the HAMP AS2 helix and the PAS β-sheet closely associate in the kinase-off state (28). Such interactions are fully consistent with the well defined ~42 Å separation between the reduced flavin cofactors that we observe. Thus, tight sandwiching of the HAMP domain by the reduced PAS domains appears to send an “off” signal through Aer to CheA.

Experimental Procedures

\textit{E. coli} Aer Expression and Purification—BL21(DE3) cells were transformed with pET28 vector containing N-terminally His\textsubscript{6}-tagged \textit{E. coli} Aer, plated on a kanamycin-added LB-agar plate, and incubated at 37 °C overnight. A single colony was added to an autoclaved and kanamycin-mixed LB medium for overnight growth under 200 rpm of shaking at 37 °C. 1 liter of the autoclaved and antibiotic-added Terrific Broth medium was inoculated with 20 ml of the overnight culture. The cells were then grown at 37 °C until \textit{A}\textsubscript{600} reached ~0.3. The temperature was turned down to 16 °C, whereas cell growth continued until \textit{A}\textsubscript{600} reached 0.6–0.8. At this point, the expression was induced by adding 0.5 mM isopropyl-1-thio-β-D-galactopyranoside. Cells were harvested after overnight expression at 16 °C and stored at ~80 °C.
The cells were resuspended in KCl extraction buffer (50 mM Tris, pH 8.0; 500 mM KCl; 10% glycerol), sonicated for 5–6 min, and centrifuged at 10,000 rpm for 30 min to obtain the membrane fractions in the low-spin pellet, which was again resuspended in KCl extraction buffer added with 1 mM PMSF and 1% DDM. The resuspended low-spin pellet was first sonicated at low power with the 4.5 power setting on the instrument to ensure fine resuspension and then at high power (7.5 setting) for 2 min before rocking at 4 °C for 5 h and centrifuging at 22,000 rpm for 1 h. The supernatant was incubated with Co(II) resin overnight after 0.1 mg/ml FAD was added to it. The next morning, the resin was washed once with wash buffer I (50 mM Tris, pH 8.0; 300 mM NaCl; 10% glycerol) containing 0.1% DDM and 0.1 mg/ml FAD. The resin was then washed several times with wash buffer II (25 mM Tris, pH 8.0; 150 mM NaCl; 10% glycerol) containing 0.1% DDM and 0.1 mg/ml FAD, and then with the buffer without FAD, and finally, Aer protein was eluted from the resin with elution buffer (wash buffer II; 250 mM imidazole, pH 8.0). UV-visible spectrum was acquired to ensure that Aer contained FAD. Normally, 1 liter of Terrific Broth culture typically gave ~12 mg of Aer (1 ml of 100 μM dimer).

**Aer Incorporation into Nanodiscs**—Detergent-purified Aer was incorporated into lipidic nanodiscs following the protocols described in various articles (35–37). Briefly, chloroform was evaporated from *E. coli* polar lipid stock by sonication to form vesicles and then by keeping under vacuum overnight. The lipid was then dissolved in sodium cholate by sonication to form vesicles and then by keeping under vacuum overnight. The mixture was rocked at 4 °C for 60 min before pre-hydrated Bio-Beads (SM2) of two-thirds of the volume of the mixture were added and then incubated at 4 °C overnight. The mixture was then centrifuged at 13,000 rpm for 30 min, and the supernatant was further purified either through size exclusion chromatography or in a desalting column to separate the nanodiscs from free FAD and excess lipids. The size exclusion chromatography experiments were done with the Superdex 200 10/300 column from GE Healthcare.

**Sodium Dithionite Reduction of Aer and Aerobic Reoxidation**—10 mM DT was added to Aer from a stock of 100 mM prepared in 1 M Tris, pH 8.0, in an anaerobic glove box, and UV-visible spectra were recorded with a cuvette of 0.2-cm path length that was sealed with Parafilm. To follow aerobic reoxidation, the Parafilm seal was removed, the cuvette was stored in a 4 °C refrigerator, and at different times, the spectra were recorded.

**cw-ESR Spectra Collection**—cw-ESR spectra were collected at 4 °C on Bruker Exelixys E500 EPR instrument at 9.4 GHz with 100-kHz modulation frequency and 1.5- or 2-G modulation amplitude. DT-reduced spectra of Aer contained narrow features because of DT radicals, which were subtracted in MATLAB (MathWorks Inc.).

**Pulsed-dipolar ESR**—Distance distributions between ASQ FADs in Aer were calculated from PDS data as described elsewhere (26, 38, 39). Briefly, ~60 μM DT-treated Aer solutions containing 35% (v/v) glycerol were prepared in ESR tubes. The dipolar evolution was recorded for ~20 h (DDM sample) and for ~40 h (nanodisc sample) at 17.35 GHz on a home-built 2D Fourier transform-ESR instrument using four-pulse double electron-electron resonance (DEER) and double quantum coherence (DQC) ESR. Pump and detection π-pulses were 40 ns, and a 60-MHz frequency separation was used. Data averaging was significantly faster with DQC (4 h for DDM) and is further reduced by the use of a new wavelet-based data denoising method (40). The baseline of the time domain data was corrected with a log-linear polynomial function. The data were then converted to distance distributions between ASQ FAD centers with Tikhonov regularization (41) followed by maximum entropy refinement (42) where the regularization parameter λ of 0.05 was used to achieve acceptable fits of time-domain data (Fig. 2A).

**CheY Phospho-transfer Assay**—As described elsewhere (26) samples containing 2 μM CheA subunit, 3 μM CheW, 15 μM Aer subunit, and 50 μM CheY were produced and treated with no DT (for the fully oxidized state), or 16 μM DT (for the ASQ state). The 20-μL assay mixtures contained ~10% glycerol (v/v) and 2 μL of buffer composed of 62.5 mM Tris (pH 7.5), 625 mM KCl, 6.25 mM EDTA, and 125 mM MgCl2. 16 mM DT was utilized to ensure that the Aer stayed reduced even after the addition of 5 μL of radioactive ATP solution (which was prepared aerobically) during the assays. For the reoxidation experiment, Aer alone was first reduced with 10 mM DT and aerobically reoxidized, and then this reoxidized Aer was mixed with the other proteins.

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