The Roles of the Dimeric and Tetrameric Structures of the Clock Protein KaiB in the Generation of Circadian Oscillations in Cyanobacteria

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Background: The function of KaiB remains to be solved.

Results: Dimeric KaiB1–94 generated circadian oscillation in vitro, but it did not in cells.

Conclusion: KaiB tetramer-dimer transformation is responsible for the regulation of the SasA-mediated clock output pathway.

Significance: We demonstrated the role of KaiB in the regulation of the SasA–KaiC interaction, involved in the transmission of time-information from KaiABC-machinery to transcription apparatus.

The molecular machinery of the cyanobacterial circadian clock consists of three proteins, KaiA, KaiB, and KaiC. The three Kai proteins interact with each other and generate circadian oscillations in vitro in the presence of ATP (an in vitro KaiABC clock system). KaiB consists of four subunits organized as a dimer of dimers, and its overall shape is that of an elongated hexagonal plate with a positively charged cleft flanked by two negatively charged ridges. We found that a mutant KaiB with a C-terminal deletion (KaiB1–94), which lacks the negatively charged ridges, was a dimer. Despite its dimeric structure, KaiB1–94 interacted with KaiC and generated normal circadian oscillations in the in vitro KaiABC clock system. KaiB1–94 also generated circadian oscillations in cyanobacterial cells, but they were weak, indicating that the C-terminal region and tetrameric structure of KaiB are necessary for the generation of normal gene expression rhythms in vivo. KaiB1–94 showed the highest affinity for KaiC among the KaiC-binding proteins we examined and inhibited KaiC from forming a complex with SasA, which is involved in the main output pathway from the KaiABC clock oscillator in transcription regulation. This defect explains the mechanism underlying the lack of normal gene expression rhythms in cells expressing KaiB1–94.

The circadian clock is an endogenous biological mechanism that generates daily cycles in physiological activity (circadian rhythms) (1, 2). Cyanobacteria are the simplest organisms that exhibit circadian rhythms (2), and the circadian clock gene cluster kaiABC is essential for circadian rhythms in cyanobacteria (3). The rhythms involve circadian oscillations in the phosphorylation level (4) and ATPase activity (5) of KaiC and complex formation among KaiA, KaiB, and KaiC (6, 7). KaiC has a duplicated RecA/DnaB structure. Its N-terminal domain shows ATPase activity (8), whereas its C-terminal domain shows both ATPase activity (5, 8) and intersubunit phosphorylation activity (9, 10). KaiA and KaiB have opposite effects on KaiC ATPase activity (5, 8) and phosphorylation level; KaiA increases them (10–13), and KaiB decreases them (5, 12, 14). KaiB is composed of two asymmetric dimers (15, 16) that form a tetramer. KaiB directly associates with the C-terminal clock oscillator domain (13) of KaiA (17). Its overall shape is that of an elongated hexagonal plate with a positively charged cleft (PC) that is flanked by two negatively charged ridges (NRs) (16). Rhythm analysis of cyanobacterial cells carrying a mutant kaiB gene show that the PC is necessary to KaiB clock function (16). Deletion of the C-terminal residues from amino acids 95–108 of KaiB (yielding mutant KaiB1–94) derived from the cyanobacterial circadian clock gene show that the PC is necessary to KaiB clock function (16). Deletion of the C-terminal residues from amino acids 95–108 of KaiB (yielding mutant KaiB1–94) derived from the cyanobacterial circadian clock gene show that the PC is necessary to KaiB clock function (16).
teria Synechococcus sp. strain PCC 7942 (hereafter Synechococcus) and Thermosynechococcus elongatus results in loss of the NRs and extensively weakens in vivo circadian rhythms (16). Here we examine the structure of KaiB1–94 and the role it plays in the generation circadian oscillations.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification**—KaiA, KaiB, KaiC, and SasA derived from *T. elongatus* (18) were expressed in *Escherichia coli* BL21 cells and purified as previously described (16, 17, 19–21). We generated gene constructs for a truncated mutant KaiB with a C-terminal deletion from residues 95–108 (KaiB1–94) and a mutant KaiC with alanine (KaiC_AAla) or aspartate (KaiC_DDA) substitutions at the two KaiC phosphorylation sites using PCR-mediated site-directed mutagenesis and cloned them in the pGEX-6P-1 vector (GE Healthcare) as described previously (20).

To determine their purity, we subjected the purified proteins to SDS-PAGE on 15% gels (22) and stained the gels with Coomassie Brilliant Blue (CBB). We estimated protein concentrations using the Bio-Rad Protein Assay with BSA as the standard as previously described (20). Unless otherwise stated, KaiA, KaiB, KaiC, and SasA refer to the KaiA dimer, KaiB tetramer, KaiC hexamer, and SasA trimer (21), respectively, and their concentrations are expressed in terms of their oligomeric status.

**Estimation of the Molecular Mass of KaiB by Gel Filtration Chromatography**—We estimated the molecular mass of KaiB using analytical gel filtration chromatography on a Superdex 75 5/150GL column (GE Healthcare) equilibrated with 20 mM Tris-HCl buffer (pH 7.5) containing 150 mM NaCl. Samples were dialyzed ultracentrifugation using a Beckman Optima XL-A analytical ultracentrifuge with an An60Ti rotor. Samples were dialyzed against 20 mM HEPES-NaOH buffer (pH 7.5) containing 150 mM NaCl (sedimentation equilibrium buffer), which was used as the blank. We performed measurements at 20 °C at 36,000, 38,000, and 40,000 rpm for KaiB1–94 and at 20,000, 22,000, and 24,000 rpm for wild-type KaiB (KaiBWT). We monitored the elution profiles of proteins by absorbance at 280 nm (A280) and performed all analyses in 20 mM Tris-HCl buffer (pH 7.5) containing 150 mM NaCl (gel filtration buffer) at 4 °C using an AKTA explorer (GE Healthcare) and a Gel Filtration Calibration Kit LMW (GE Healthcare) for the molecular mass standards. We estimated the molecular mass of KaiB1–94 and 0.759 ml/g for KaiBWT for the analyses.

**Sedimentation Equilibrium by Analytical Ultracentrifugation**—We performed sedimentation equilibrium analytical ultracentrifugation using a Beckman Optima XL-A analytical ultracentrifuge with an An60Ti rotor. Samples were dialyzed against 20 mM HEPES-NaOH buffer (pH 7.5) containing 150 mM NaCl (sedimentation equilibrium buffer), which was used as the blank. We performed measurements at 20 °C at 36,000, 38,000, and 40,000 rpm for KaiB1–94 and at 20,000, 22,000, and 24,000 rpm for wild-type KaiB (KaiBWT). We estimated the molecular mass of KaiB1–94 and 0.759 ml/g for KaiBWT for the analyses.

**Estimation of the Molecular Mass and Stoichiometry of the KaiB-KaiC Complex by Gel Filtration Chromatography**—Reaction mixtures containing 5 μM KaiB1–94 or 2.5 μM KaiBWT and 1 μM KaiCDD in 20 mM HEPES-NaOH buffer (pH 7.5) containing 1 mM ATP, 5 mM MgCl2, and 150 mM NaCl (HEPES reaction buffer) were incubated at 40 °C for 18 h and analyzed by gel filtration chromatography on a Superdex 200/HR 10/300 column (GE Healthcare) equilibrated with HEPES reaction buffer containing 0.1 mM ATP and 5 mM MgCl2 at 4 °C. We estimated their purity, we subjected the purified proteins to SDS-PAGE on 15% gels (22) and stained the gels with Coomassie Brilliant Blue (CBB). We estimated protein concentrations using the Bio-Rad Protein Assay with BSA as the standard as previously described (20). Unless otherwise stated, KaiA, KaiB, KaiC, and SasA refer to the KaiA dimer, KaiB tetramer, KaiC hexamer, and SasA trimer (21), respectively, and their concentrations are expressed in terms of their oligomeric status.

**Labeling of KaiB with Cy3 and Fluorescence Correlation Spectroscopy (FCS) Measurements**—Cy3-NHS fluorescent dye esters (GE Healthcare) were covalently coupled to the amines of lysine residues and the N-terminal amino acid residue of KaiB. KaiB (8 μM KaiB1–94 or 4 μM KaiBWT) was incubated with 80 μM Cy3-NHS ester in gel filtration buffer at 4 °C for 2 h. After labeling, the reaction mixtures were loaded on a PD MidiTrap™ G-25 column (GE Healthcare) for removal of the remaining unbound dye from the labeled proteins. We calculated the amounts of Cy3 introduced onto KaiB from the absorbance of the labeled proteins at 552 nm (A552). Under these conditions, 0.85 ± 0.10 and 0.64 ± 0.01 molecules of Cy3 (n = 3) per subunit of KaiB1–94 and KaiBWT were introduced into KaiB1–94 and KaiBWT, respectively.

We performed FCS measurements using a multiphoton FCS/fluorescence cross-correlation spectroscopy system using Fluor-Twin Dual Emission Uni-laser eXcitation (Fluc DEUTX™) (MBL, Nagoya, Japan). Cy3 was excited by the 445-nm laser, and its emission was detected at 615–690 nm. Cy3-KaiB1–94 and Cy3-KaiBWT (0.1 μM) were separately incubated with 0, 0.03, 0.05, 0.10, and 0.30 μM KaiCDD in the presence of 0, 0.05, 0.10, or 0.30 μM SasA in HEPES reaction buffer at 25 °C for 18 h, and the reaction mixtures were subjected to FCS measurements at 25 °C for 5 s (5 times consecutively). The diffusion time defined as the average time required for the diffusion of fluorescent particles across the detection area reflects the size of the particles. If the effect of the molecular shape on its diffusion is negligible, diffusion time is proportional to the cubic root of the molecular mass, as described in Equation 1 (23, 24),

\[
\frac{DTa}{DTb} = \left(\frac{MMa}{MMb}\right)^{3/2}
\]

where \(DTa\) and \(DTb\) are the diffusion times of molecules a (Cy3-KaiB1–94 or Cy3-KaiBWT) and b (Cy3-KaiB1–94 or Cy3-KaiCDD complex or Cy3-KaiBWT-KaiCDD), respectively, and MMa and MMb are the molecular masses of molecules a and b, respectively. Accordingly, we calculated the size of the Cy3-KaiB1–94-KaiCDD and Cy3-KaiBWT-KaiCDD complexes to be 320 ± 70 and 370 ± 90 kDa, respectively, using the molecular masses of Cy3-KaiBWT (~50 kDa) and Cy3-KaiB1–94 (~25 kDa). These
values are consistent with the molecular masses determined by gel filtration chromatography (see Fig. 2A).

**Surface Plasmon Resonance Analysis of Interaction between KaiB and KaiC**—We analyzed the interaction between KaiB<sub>1–94</sub> and KaiC<sub>DD</sub> and between KaiB<sub>WT</sub> and KaiC<sub>DD</sub> at 25 °C by surface plasmon resonance (25) using a Biacore and its associated software (GE Healthcare). KaiC<sub>DD</sub> (10 μM) was immobilized on a CM5 sensor chip using an amine coupling kit (GE Healthcare) in HEPES reaction buffer containing 0.1 mM ATP, 0.1 mM dithiothreitol (DTT), and 50 mM NaCl. We monitored the association to, and dissociation from, the immobilized KaiC<sub>DD</sub> of KaiB<sub>1–94</sub>, KaiB<sub>WT</sub>, KaiA, and SasA continuously at stepwise concentrations of 0.1, 0.25, 0.5, 1.0, and 1.5 μM in HEPES reaction buffer. We computed the dissociation (k<sub>d</sub>) and association (k<sub>a</sub>) rates by fitting a 1:1 binding model to the experimental data. We calculated the value for apparent equilibrium dissociation constant (K<sub>D</sub>) as k<sub>d</sub>/k<sub>a</sub>.

**Measurement of the Phosphorylation Levels of KaiC**—Reaction mixtures of KaiA and KaiC (each 0.5 μM) were incubated in the presence or absence of 1.0 μM KaiB<sub>1–94</sub> or 0.5 μM KaiB<sub>WT</sub> in gel filtration buffer containing 1 mM ATP and 5 mM MgCl<sub>2</sub> at 25 or 40 °C, and 10-μl aliquots of the reaction mixtures were removed to stop the reaction at specific time intervals. The aliquots were subjected to SDS-PAGE on 12.5% gels (acrylamide:bisacrylamide = 144:1), and the gels were stained with CBB. The intensities of bands were measured by densitometry using a CS Analyzer. KaiC showed a triplet band on SDS-PAGE (4, 9); the two upper bands correspond to phosphorylated KaiC (p-KaiC), and the lowest band corresponds to nonphosphorylated KaiC (np-KaiC). We calculated the relative amount of p-KaiC to total KaiC (the level of p-KaiC) from band intensities and analyzed the circadian oscillations of the level of p-KaiC using the program RAP (26).

**In Vivo Bioluminescence Rhythm Assay**—We measured the bioluminescence rhythms of the *T. elongatus* strains carrying a P<sub>paAB</sub>: lux<sub>AB</sub> reporter gene at 41 °C using a newly developed bioluminescence monitoring apparatus with a robotic plate conveyor. The apparatus was 1.3 times as sensitive to luminescence as our previous apparatuses (27). We analyzed bioluminescence data by a modified cosinor method of RAP as previously described (26, 28). We examined the wild-type strain, a kaiB-null strain carrying a mutant kaiB gene with a nonsense mutation downstream of its initiation codon, and a strain carrying an additional mutant kaiB gene encoding KaiB<sub>1–94</sub>, KaiB<sub>1–94</sub>, in the kaiB-null genetic background (16, 29).

**Assay for the Complex Formation of SasA with KaiC by Gel Filtration Chromatography**—Reaction mixtures containing 1.5 μM SasA and 0.6 μM KaiC<sub>DD</sub> were incubated in the presence or absence of 3.0 μM KaiB<sub>WT</sub> or 6.0 μM KaiB<sub>1–94</sub> in HEPES reaction buffer containing 0.1 mM DTT at 25 °C for 18 h and analyzed by gel filtration chromatography on a Superdex200/HR 10/300 column (GE Healthcare) equilibrated with HEPES reaction buffer containing 0.1 mM ATP, 5 mM MgCl<sub>2</sub>, and 0.1 mM DTT at 4 °C. Aliquots of the chromatography fractions were subjected to SDS-PAGE on 15% gels, and the gels were stained with CBB. We determined the amount of SasA in each fraction by densitometry.

**RESULTS**

**Oligomeric Structures of KaiB<sub>1–94</sub> and KaiB<sub>WT</sub> in Solution**—To determine the oligomeric structure of KaiB<sub>1–94</sub> in solution, we estimated the molecular mass of KaiB<sub>1–94</sub> by gel filtration chromatography in gel filtration buffer. KaiB<sub>1–94</sub> (subunit molecular mass, 10.9 kDa) eluted as a single peak (Fig. 1A) corresponding to a 23.4 ± 0.4-kDa protein (Fig. 1B and Table 1) that corresponds to the approximate molecular mass of a dimer, whereas KaiB<sub>WT</sub> (subunit molecular mass, 12.4 kDa)
eluted as a single peak (Fig. 1A) corresponding to a 55.4 ± 0.6-kDa protein (Fig. 1B and Table 1) that corresponds to the approximate molecular mass of a tetramer. We obtained similar results under varying buffer conditions (Table 1). In sedimentation equilibrium analysis, KaiB1–94 sedimented as a dimer with a molecular mass of 23 kDa (Fig. 1C and Table 1), and KaiBWT sedimented as a tetramer with a molecular mass of 48 kDa (Fig. 1D and Table 1), confirming the gel filtration chromatography results. Hereafter, when their concentrations are expressed, KaiBWT is considered a tetramer and KaiB1–94 a dimer.

Complex Formation of KaiB1–94 and KaiBWT with KaiC and the Stoichiometry of the KaiB1–94–KaiCDD and KaiBWT–KaiCDD Complexes—First, we examined the complex formation of KaiB1–94 with KaiCWT, KaiCDD, and KaiCAA by FCS analysis (supplemental Fig. S1). The diffusion time of Cy3-KaiB1–94 incubated alone at 25 °C for 18 h was 0.13 ± 0.04 ms, and it increased in the presence of KaiCDD (0.37 ± 0.09 ms). On the other hand, the diffusion time scarcely increased in the presence of KaiCAA (0.16 ± 0.03 ms). These results indicate that KaiB1–94 formed a complex with KaiCDD whereas it scarcely formed a complex with KaiCAA. Furthermore, the diffusion time of Cy3-KaiB1–94 increased in the presence of KaiCWT when the phosphorylation level of KaiCWT had been elevated to about 70% by KaiA (0.26 ± 0.03 ms), whereas it increased only slightly in the presence of KaiCWT when the phosphorylation level of KaiCWT (about 30%) had not been elevated by KaiA (0.18 ± 0.03 ms). Therefore, the complex formation of KaiB1–94 with KaiC depends on the phosphorylation level of KaiC; the higher the phosphorylation level, the more efficient complex formation occurred. Probably, KaiB1–94 associates with p-KaiC more strongly than with np-KaiC. These results are consistent with a previous observation that KaiBWT associates with p-KaiC more strongly than with np-KaiC (30, 31). Hereafter, we used KaiCDD to examine the complex formation of KaiB with KaiC because the phosphorylation level of KaiCWT changes during incubation, especially in the presence of KaiB and/or KaiA (Ref. 4 and see Fig. 3A).

Next we examined the KaiB1–94–KaiCDD complex by gel filtration chromatography followed by SDS-PAGE. KaiB1–94 and KaiCDD each eluted as a single peak, KaiB1–94 with an apparent molecular mass of 25 ± 2 kDa and KaiCDD with an apparent molecular mass of 319 ± 7 kDa (n = 3) (Fig. 2A). When KaiCDD (1 μM) was incubated with an excess of KaiB1–94 (5 μM) at 40 °C for 18 h (n = 3), the reaction products eluted as two peaks, one with a molecular mass of 26 ± 1 kDa (KaiB1–94) and the other with a molecular mass of 350 ± 20 kDa (Fig. 2A), suggesting the formation of a KaiB1–94–KaiCDD complex. We also confirmed the formation of a KaiBWT–KaiCDD complex (366 ± 20 kDa) by gel filtration chromatography (Fig. 2A). The apparent molecular masses of the KaiB1–94–KaiCDD and KaiBWT–KaiCDD complexes were slightly higher than that of the KaiCDD hexamer (Fig. 2A).

To determine the stoichiometry of the KaiB1–94–KaiCDD complex, we collected it by gel filtration chromatography (Fig. 2A, 9.0–9.5 ml) and estimated the amount of KaiB1–94 and KaiCDD it contained by SDS-PAGE (Fig. 2B) followed by densitometry (Fig. 2, C and D). When KaiCDD (1 μM) was incubated with an excess of KaiB1–94 (5 μM) at 40 °C for 18 h, the fraction containing the complex contained 3.8 ± 0.3 subunits of KaiB1–94 per molecule of hexameric KaiCDD (Fig. 2E). When we applied the same procedure to the KaiBWT–KaiCDD complex, the stoichiometry showed that it contained 4.2 ± 0.4 subunits of KaiBWT per molecule of hexameric KaiCDD (Fig. 2E), indicating that the KaiB1–94–KaiCDD complex consisted of two molecules of KaiB1–94 (a dimer) and one molecule of hexameric KaiCDD whereas the KaiBWT–KaiCDD complex consisted of one molecule of KaiBWT (a tetramer) and one molecule of hexameric KaiCDD.

We confirmed these results using FCS with Cy3-NHS fluorescent dye ester (Cy3)-labeled KaiB proteins (Cy3-KaiB1–94 and Cy3-KaiBWT). When 0.2 μM Cy3-KaiB1–94 and 0.1 μM Cy3-KaiBWT were incubated in the presence of 0.0, 0.025, 0.05, 0.15, or 0.3 μM KaiCDD, the diffusion time varied directly with the KaiCDD concentration (Fig. 2F), indicating complex formation between KaiCDD and Cy3-KaiB1–94 or Cy3-KaiBWT.

The counts per particle of Cy3-KaiB1–94 (0.2 μM) also varied with KaiCDD concentration (Fig. 2G). The counts per particle values of Cy3-KaiB1–94 in the presence of KaiCDD at molar ratios greater than 0.5 were approximately twice those in its absence, which suggested that one molecule of the Cy3-KaiB1–94–KaiCDD complex contained two molecules of the Cy3-KaiB1–94–KaiCDD dimer (Fig. 2G). In contrast, counts per particle did not significantly change when Cy3-KaiBWT (0.1 μM) was incubated with an excess of KaiCDD (0.3 μM), which suggests that one molecule of the KaiBWT–KaiCDD complex contained one molecule of the KaiBWT tetramer (Fig. 2G).

Interaction of KaiB1–94 Versus KaiBWT with KaiCDD—Surface plasmon resonance analysis showed that the association rate (kA) of KaiCDD was 8 times as high with KaiB1–94 as with KaiBWT, and its dissociation rate (kD) was 0.6 times as high (Table 2). Thus, the binding affinity of KaiB1–94 for KaiCDD was 15 times the binding affinity of KaiBWT. We also examined the binding affinities of KaiA and SasA, other known KaiC-binding proteins, and determined that KaiB1–94 had the highest binding affinity in the group (Table 2).

Circadian Oscillations in the Level of p-KaiC in an In Vitro KaiABC Clock System and Circadian Gene Expression Rhythms in T. elongatus Cells—When 0.5 μM KaiA, KaiB (KaiB1–94 or KaiBWT), KaiC, and ATP were incubated together at 25 or 40 °C
The Function and Oligomer Structure of KaiB

in an in vitro KaiABC clock system (4), the level of p-KaiC oscillated in the presence of KaiB1–94 and KaiBWT (Fig. 3A). At 40 °C, the period length was 21.7 ± 0.9 h for the oscillations driven by KaiB1–94 and 22.6 ± 0.2 h for those driven by KaiBWT (Fig. 3A), which was consistent with the circadian oscillations observed in T. elongatus cells (24.4 ± 0.3 h) (Ref. 16 and Fig. 3B). Thus, KaiB1–94, like KaiBWT, can generate circadian oscillations in the in vitro KaiABC clock system. At 25 °C, the period length was 21.7 ± 1.3 h for the oscillations driven by KaiB1–94 and 23.3 ± 0.5 h for those driven by KaiBWT (Fig. 3A), indicating that the period lengths were temperature-compensated in our system.

Circadian rhythms were normal in the kaiB-null host cells carrying kaiBWT but showed a greatly reduced amplitude in those carrying kaiB1–94 (Fig. 3B).

Inhibition of SasA–KaiCDD Complex Formation by KaiB1–94—To determine why cyanobacterial cells expressing KaiB1–94 showed weakened circadian gene expression rhythms even though KaiB1–94 is able to generate normal cir-

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**TABLE 2**

Kinetic and steady state parameters for the interaction of KaiBWT, KaiB1–94, KaiA, and SasA with KaiCDD obtained by surface plasmon resonance analysis.

| Protein       | $k_a$ ($\times 10^4$) | $k_d$ ($\times 10^{-2}$) | $K_D$ ($\times 10^{-4}$) |
|---------------|-----------------------|--------------------------|--------------------------|
| KaiBWT        | 1.3 ± 0.057           | 0.46 ± 0.058             | 3.5 ± 0.38               |
| KaiB1–94      | 10.4 ± 0.61           | 0.26 ± 0.040             | 0.24 ± 0.029             |
| KaiA          | 3.9 ± 1.0             | 2.5 ± 0.10               | 6.8 ± 1.4                |
| SasA          | 7.7 ± 1.1             | 1.2 ± 0.066              | 1.5 ± 0.24               |

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circadian oscillations in vitro, we examined the effects of KaiB on the SasA-mediated clock output pathway, that is, on the formation of the SasA-KaiC complex.

Gel filtration chromatography yielded only a single peak for both KaiCDD and SasA corresponding to a 340 ± 20-kDa protein and a 140 ± 10-kDa protein, respectively. When reaction mixtures containing 1.5 μM SasA and 0.6 μM KaiCDD were incubated in the absence or presence of 3 μM KaiBWT or 6 μM KaiB1–94 at 25 °C for 18 h, the elution profile of the sample without KaiB showed 3 peaks (Fig. 4A) that we analyzed by SDS-PAGE (Fig. 4B). Peak 1 (>670 kDa, the molecular mass of the largest standard used) corresponded to the SasA–KaiCDD complex, peak 2 (440 ± 50 kDa) corresponded to the other form of the SasA–KaiCDD complex, and peak 3 (130 ± 5 kDa) corresponded to SasA. The amount of SasA contained in each fraction estimated by SDS-PAGE followed by densitometry showed that 34 ± 10% (n = 3) of the total SasA was unbound (Fig. 4C).

The elution profile of the sample with 6 μM KaiB1–94 showed three peaks (Fig. 4A); peak 1 (360 ± 10 kDa) corresponded to KaiCDD and the KaiB1–94–KaiCDD complex, peak 2 (140 ± 10 kDa) corresponded to SasA, and peak 3 (20 ± 0 kDa) corresponded to KaiB1–94. The presence of KaiB1–94 all but extin-
The Function and Oligomer Structure of KaiB

guished the SasA-KaiC\textsubscript{DD} complex and greatly increased free SasA (Fig. 4A). The free SasA fractions contained 92 ± 10% (n = 3) of the total SasA added (Fig. 4, B and C), which indicated nearly complete inhibition of SasA-KaiC\textsubscript{DD} complex formation by KaiB\textsubscript{1–94}. In the presence of 3 μM KaiB\textsubscript{WT}, the elution profile of the sample showed four peaks (Fig. 4A) corresponding to the SasA-KaiC\textsubscript{DD} complex (>670 kDa), the other form of the SasA-KaiC\textsubscript{DD} complex, KaiB\textsubscript{DD} and KaiB\textsubscript{WT}-KaiC\textsubscript{DD} complex (420 ± 40 kDa), SasA (140 ± 10 kDa), and KaiB\textsubscript{WT} (50 ± 0 kDa). The presence of KaiB\textsubscript{WT} slightly reduced the SasA-KaiC\textsubscript{DD} complex peak and slightly increased the free SasA peak; the free SasA fractions contained 43 ± 5% (n = 3) of the total SasA added (control 34 ± 10%) (Fig. 4C), which suggests that KaiB\textsubscript{WT} inhibited formation of the SasA-KaiC\textsubscript{DD} complex, albeit very slightly. Thus, KaiB\textsubscript{1–94} was a stronger inhibitor of SasA-KaiC\textsubscript{DD} complex formation than KaiB\textsubscript{WT}.

We also examined KaiB inhibition of SasA-KaiC\textsubscript{WT} complex formation by FCS analysis with Cy3-SasA (supplemental Fig. S2). The diffusion time of Cy3-SasA (0.2 μM) incubated alone at 25 °C for 18 h was 0.14 ± 0.00 ms. On the other hand, when Cy3-SasA was incubated with 0.2 μM KaiC\textsubscript{WT}, its diffusion time increased to 0.30 ± 0.06 ms, indicating formation of a Cy3-SasA-KaiC\textsubscript{WT} complex. The addition of 2.0 μM KaiB\textsubscript{1–94} almost completely canceled this diffusion time increase (0.16 ± 0.03 ms), indicating that KaiB\textsubscript{1–94} inhibited SasA-KaiC\textsubscript{WT} complex formation as well as SasA-KaiC\textsubscript{DD} complex formation (see above). However, the addition of KaiB\textsubscript{WT} did not cancel the diffusion time increase under the conditions examined (diffusion time: 0.35 ± 0.03 and 0.33 ± 0.01 ms in the presence of 1.0 μM KaiB\textsubscript{WT} and 2.0 μM KaiB\textsubscript{WT}, respectively) (supplemental Fig. S2).

We examined SasA inhibition of KaiB-KaiC\textsubscript{DD} complex formation by FCS analysis with Cy3-KaiB\textsubscript{1–94} and Cy3-KaiB\textsubscript{WT} (Fig. 4D). In the absence of KaiC\textsubscript{DD}, the diffusion time was 0.13 ± 0.02 ms for 0.2 μM Cy3-KaiB\textsubscript{1–94} and 0.11 ± 0.03 ms for 0.1 μM Cy3-KaiB\textsubscript{WT}. In the presence of 0.1 μM KaiC\textsubscript{DD}, it increased to 0.28 ± 0.03 ms for Cy3-KaiB\textsubscript{1–94} and 0.25 ± 0.05 ms for Cy3-KaiB\textsubscript{WT} (Fig. 4D), indicating formation of the Cy3-KaiB\textsubscript{1–94}-KaiC\textsubscript{DD} and Cy3-KaiB\textsubscript{WT}-KaiC\textsubscript{DD} complexes. The diffusion time of 0.2 μM Cy3-KaiB\textsubscript{1–94} was slightly shorter in the presence of 0.3 μM SasA (0.23 ± 0.02 ms) than in its absence (0.28 ± 0.03 ms) (Fig. 4D), which indicates that SasA weakly inhibited formation of the KaiB\textsubscript{1–94}-KaiC\textsubscript{DD} complex. In the presence of KaiC\textsubscript{DD}, on the other hand, SasA reduced the diffusion time of 0.1 μM Cy3-KaiB\textsubscript{WT} from 0.25 ± 0.05 to 0.13 ± 0.01 ms, the level it was in the absence of KaiC\textsubscript{DD} (0.11 ± 0.03 ms) (Fig. 4D), which indicates that SasA almost completely inhibited formation of the KaiB\textsubscript{WT}-KaiC\textsubscript{DD} complex. Thus, SasA inhibited KaiB\textsubscript{WT}-KaiC\textsubscript{DD} complex formation more strongly than KaiB\textsubscript{1–94}-KaiC\textsubscript{DD} complex formation.

**DISCUSSION**

That KaiB\textsubscript{1–94} is a dimer (Fig. 1 and Table 1) but interacts with KaiC\textsubscript{DD} in much the same way as tetrameric KaiB\textsubscript{WT} (Fig. 2) and can generate normal circadian oscillations in vitro (Fig. 3A) indicates that KaiB tetramericity is not required for the generation of circadian oscillations per se. The tetrameric structure of KaiB, however, and its negatively charged C-termin

![Diagram](image-url)
slightly, KaiB1–94 inhibits it strongly (Fig. 4 and supplemental Fig. S2), probably because of the high affinity of KaiB1–94 for KaiCDD (Table 2).

SasA is a KaiC binding histidine kinase in the SasA-RpaA two-component regulatory system (32, 33), and is involved in the main clock output pathway. Ultimately, the time information of the KaiABC clock oscillator is transmitted to the transcription apparatus and generates genome-wide circadian transcription rhythms in cyanobacteria. Disruption of the sasA gene greatly lowers the level of kaiB expression and the amplitude of kaiA and kaiB expression rhythms (32). KaiB1–94 competitively inhibits the SasA–KaiC interaction more strongly than KaiBWT does due to its higher affinity for KaiC (Table 2); it is likely, therefore, that KaiB1–94 hinders the SasA–KaiC interaction that is necessary for the transmission of the time information from the KaiABC clock oscillator to the transcription apparatus through the SasA-mediated clock output pathway (32, 33). We hypothesize that the defective rhythm phenotypes of cyanobacterial cells expressing KaiB1–94 in a ΔkaiB genetic background follow from a KaiB1–94-induced defect in the SasA–KaiC interaction involved in the SasA-mediated clock output pathway.

KaiB (KaiB1–94 and KaiBWT) inhibited formation of the SasA–KaiC (KaiCDD and KaiCWT) complex (Fig. 4, A–C, and supplemental Fig. S2), and SasA inhibited formation of the KaiB (KaiB1–94 and KaiBWT)-KaiCDD complex (Fig. 4D). It is unknown whether the two proteins compete for an identical binding site on KaiC because the binding sites of KaiB and SasA on KaiC are not determined. Previously, the N-terminal domain of SasA was suggested to be homologous to KaiB (32). However, comparison between the NMR structure of SasA N-terminal domain and the x-ray crystal structure of KaiB demonstrated that KaiB and SasA N-terminal domain have different folds (34). Therefore, it is not likely that SasA and KaiB bind to the same site on KaiC. We have demonstrated by surface plasmon resonance analysis and gel filtration chromatography that the SasA binding site of KaiC is located on its N-terminal domain (21), as suggested by NMR analysis (35). The SasA inhibition of KaiBWT–KaiCWT complex formation has been demonstrated by Native-PAGE using proteins derived from Synechococcus (36).

Our demonstration that two molecules of KaiB1–94 bind to one molecule of KaiCDD (Fig. 2) suggests that two molecules of KaiB1–94 dimer bind to one molecule of a KaiC hexamer. Because one molecule of KaiBWT binds to one molecule of KaiCDD (Fig. 2), it is likely that upon interaction with KaiC, the KaiB molecule changes from a tetramer to a tetramer to a dimer. This is consistent with the cryo-electron microscopy analysis of the KaiBWT–KaiC WT complex that revealed the possibility of KaiB binding to KaiC as two dimers (37). Based on the crystal structure of KaiB, we have proposed that the interdimer interface of the KaiB tetrameric molecule could be destabilized by changes in pH or ionic strength and dissociate into two dimers (16).

On one surface of the KaiB tetramer, a PC, which is the active site, is covered with two NRs, and the PC may be exposed for ligand binding by NR movement (Fig. 5A) (16). In agreement with this model, KaiB1–94, which lacks NRs and has a continuously exposed PC, is a more stable binding partner for KaiCDD than KaiBWT (Table 2). It is likely that PC exposure by NR movement may play an important role in the interaction of KaiB with KaiC (Fig. 5, A and B). An alternative model is that KaiB associates with KaiC using a surface, which is exposed by the deletion of NRs on KaiB or the dimerization of KaiB. There are two areas on KaiC that are highly negatively charged, one around the pore opening and inside the pore of KaiC N-terminal domain and the other around the intersubunit interface of one of two adjacent KaiC N-terminal domains (Ref. 38 and supplemental Fig. S3). Possible electrostatic interaction between the PC on KaiB and the negatively charged surfaces of KaiC might occur.

Our demonstration that the stable interaction of KaiB1–94 with KaiC inhibited the interaction of SasA with KaiC (Fig. 4) indicated that the C-terminal region of KaiB is necessary for driving normal circadian transcription rhythms through regulation of the KaiC-SasA interaction (Fig. 5, C and D). Furthermore, our recent observations are consistent with our model as follows. Electron spin resonance analysis demonstrated that the mobility of the spin labels introduced into the amino acid residues located on the PC greatly increased on incubation at 40 °C (39), which indicates that the local structure surrounding the spin labels on the PC is relaxed during incubation (39); the 64th
residue located inside the KaiB molecule near the PC interacts with KaiA (17).

In the KaiABC clock oscillator, KaiA associates preferentially with np-KaiC (40) (KaiC_{AA} mimics np-KaiC) to elevate the phosphorylation level of KaiC (11–13). Then, KaiA dissociates from the p-KaiC (KaiC_{PD} mimics completely phosphorylated KaiC) (40). In contrast, KaiB associates preferentially with the p-KaiC (Ref. 6 and supplemental Fig. S1) and then reduces the phosphorylation level of KaiC (12, 14). A series of these reactions and interactions among Kai proteins generate circadian oscillations such as oscillations in the phosphorylation level (4) and ATPase activity (5) of KaiC and complex formation among Kai proteins (6, 7). SasA also associates preferentially with p-KaiC, which enhances the autophosphorylation of SasA (21). KaiB and SasA compete each other for KaiC (Fig. 4 and supplemental Fig. S2). Therefore, it is likely that KaiB regulates the normal SasA-KaiC interaction required for the transmission of time information from KaiC to the transcription apparatus, resulting in genome-wide transcription cycles in cyanobacterial cells (21, 32, 33). SasA trimer and KaiC hexamer associate each other at a 1:1 molar ratio at their N-terminal domains, and the phosphorylation states of their C-terminal domains affect the affinity of the interaction (21). Interestingly, the phosphorylation state of KaiC also affects the rigidity of the C-terminal domains of KaiC hexamer and the interaction between the N-terminal domains and C-terminal domains of the KaiC, and KaiC associates with KaiB when the structure of KaiC C-terminal domains is rigid (35).

KaiB and KaiC occur in Archaea and Proteobacteria as well as in cyanobacteria (41). The C-terminal cluster of negatively charged residues is highly conserved in cyanobacterial clock KaiBs (Ref. 16 and Fig. 6) but not in cyanobacterial non-clock KaiBs (such as Synechocystis KaiB2 (41)) and not in KaiB homologues in Archaea and Proteobacteria (Fig. 6). The negatively charged C terminus of KaiB likely plays an important role in the SasA-mediated clock output pathway through KaiB–KaiC interactions affecting the SasA–KaiC interaction. SasA is present only in cyanobacteria, which suggests that the negatively charged C-terminal region of cyanobacterial KaiB may have co-evolved with SasA as a component of the cyanobacterial circadian system.

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