The convergent xenogeneic silencer MucR predisposes \( \alpha \)-proteobacteria to integrate AT-rich symbiosis genes

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ABSTRACT

Bacterial adaptation is largely shaped by horizontal gene transfer, xenogeneic silencing mediated by lineage-specific DNA bridgers (H-NS, Lsr2, MvaT and Rok), and various anti-silencing mechanisms. No xenogeneic silencing DNA bridger is known for \( \alpha \)-proteobacteria, from which mitochondria evolved. By investigating \( \alpha \)-proteobacterium Sinorhizobium fredii, a facultative legume microsymbiont, here we report the conserved zinc-finger bearing MucR as a novel xenogeneic silencing DNA bridge. Self-association mediated by its N-terminal domain (NTD) is required for DNA–MucR–DNA bridging complex formation, maximizing MucR stability, transcriptional silencing, and efficient symbiosis in legume nodules. Essential roles of NTD, CTD (C-terminal DNA-binding domain), or full-length MucR in symbiosis can be replaced by non-homologous NTD, CTD, or full-length protein of H-NS from \( \gamma \)-proteobacterium Escherichia coli, while NTD rather than CTD of Lsr2 from Gram-positive Mycobacterium tuberculosis can replace the corresponding domain of MucR in symbiosis. Chromatin immunoprecipitation sequencing reveals similar recruitment profiles of H-NS, MucR and various functional chimeric xenogeneic silencers across the multipartite genome of S. fredii, i.e. preferring AT-rich genomic islands and symbiosis plasmid with key symbiosis genes as shared targets. Collectively, the convergently evolved DNA bridge MucR predisposed \( \alpha \)-proteobacteria to integrate AT-rich foreign DNA including symbiosis genes, horizontal transfer of which is strongly selected in nature.

INTRODUCTION

The prevailing symbiogenic theory of eukaryogenesis involves the symbiosis between a facultative \( \alpha \)-proteobacterium and an archaeal host (1,2). This putative \( \alpha \)-proteobacterium evolved into the present-day mitochondria with probably 3000–4000 genes transferred into the nucleus (3). Many mitochondria within a single eukaryote cell produce more ATP per gene than prokaryotes and contribute to the development of multicellularity and genome expansion of eukaryotes (4). On the other hand, the archaeal host provided the complex information processing machinery including histones which compacted the ever-growing eukaryotic genome, the majority of which is mobile or ‘selfish’ DNA (4,5). Despite extensive horizontal gene transfer (HGT) between prokaryotes in microbiota of various niches (6), it has been proposed that prokaryote genome size is limited by the ability to manipulate and control gene expression rather than the cost of synthesizing DNA (5). Without histone proteins, the organization of bacterial DNA depends on nucleoid-associated proteins (NAPs) involved in bending and looping of DNA (7). Known bacterial DNA bending factors are Fis (factor for inversion stimulation), IHF (integration host factor), and HU (histone-like protein from Escherichia coli strain U93) (7,8). DNA looping in bacteria is actively mediated by the conserved SMC (structural maintenance of chromosomes) complexes driven by ATP (9), and passively by lineage-specific DNA bridgers including H-NS (histone-like nucleoid structuring protein) (10), MvaT (11,12), Lsr2 (13,14) and Rok (15,16) reported in Enterobacteriaceae (Gram-), Pseudomonas (Gram-), Actinomycetes (Gram+) and Bacillus (Gram+), respectively. Despite low sequence homology between these lineage-specific DNA bridgers, they share convergent features such as self-association mainly mediated by their N-terminal...
domain (NTD), and xenogeneic silencing function mediated by the DNA binding preference of C-terminal domain (CTD) to AT-rich foreign DNA sequences (7,17–19). However, it remains elusive whether DNA bridger(s) of such convergent features also evolved in α-proteobacteria which are enriched with symbiotic and pathogenic bacteria facultatively associated with various eukaryotes (20,21).

In 1980s, ros and mucR mutants forming rough outer surface colonies were independently identified for the plant pathogen Agrobacterium tumefaciens and legume nodule microsymbiont Sinorhizobium meliloti, respectively (22,23). MucR and Ros, or named as RosR or MI in other species, were later demonstrated as homologs belonging to the Cys2His2 zinc-finger bearing MucR/Ros family (24–27). Evolutionary analysis further suggested a possible ancestral origin of MucR/Ros in α-proteobacteria and δ-proteobacteria, followed by multiple independent HGT and gene loss events in the evolution of the tree of life (20). The MucR/Ros family proteins have received intensive studies in plant and human pathogens (Agrobacterium and Brucella), legume microsymbionts (Sinorhizobium, Rhizobium and Mesorhizobium), and also in the cell cycle model Caulobacter (20). MucR/Ros functions as a pleiotropic transcriptional regulator for cellular processes such as biosynthesis of various exopolysaccharides (28–30), cell cycle (31), c-di-GMP signaling (32), motility and quorum sensing (33–37), secretion systems including T1SS, T3SS and T4SS (38,39), some of which are involved in virulence (40–45) and symbiosis (25,32,33,38,46–48). A combined ChIP-seq (Chromatin immunoprecipitation sequencing) and RNA-seq analysis of Sinorhizobium fredii revealed that MucR mainly acts as a silencer for AT-rich foreign genes with predisposed high transcription potential (38). This feature of MucR is similar to non-homologous xenogeneic silencers H-NS, Lsr2, MvaT and Rok from other bacterial classes (17,20), i.e. banking genes with potential benefits while reducing energy cost and toxicity due to random transcription of AT-rich foreign DNA (32,38,49,50).

Extensive studies of the Cys2His2 zinc-finger bearing CTD of MucR/Ros (such as the region 56–142 of Ros from Agrobacterium tumefaciens) have led to a hypothesis that the Cys2His2 zinc-finger structure of MucR/Ros can be the ancestral form of eukaryotic Cys2His2 zinc-finger (27,51–54). The single CTD of MucR/Ros can bind target DNA in vitro (27,55). Several cases of DNA binding by single zinc-finger domain proteins from eukaryotes also have been reported (56,57), while in most cases, a minimum of two zinc-fingers is required for high-affinity DNA binding and a protein harboring tandemly repeated zinc-fingers (as many as 37) is common in eukaryotes (54,55,58). Recently, it was found that NTD is essential for in vitro self-association of MucR homologs from Mesorhizobium loti and Brucella abortus, resulting in high-order oligomerization of MucR (59,60). These cumulative evidences support a hypothesis that the zinc-finger bearing MucR/Ros family protein is a novel xenogeneic silencer conserved in α-proteobacteria (20,38,59–61), though the significance of NTD mediated self-association in shaping MucR-DNA complex remains unexplored (20,59). Does MucR bridge DNA in α-proteobacteria as other lineage-specific xenogeneic silencers? Could MucR in α-proteobacteria be replaced by non-homologous xenogeneic silencers from other bacterial classes?

In this study, we focused on MucR from the broad-host-range S. fredii CCBAU45436 (SF45436) associated with diverse legumes including important crops such as soybean and pigeonpea (62,63), which is essential for effective symbiosis of SF45436 (32,38,46,47). We systematically investigated the key residues and conserved sequence signatures essential for function of NTD, and the regulatory role of self-association mediated by NTD during MucR-DNA interaction. It was revealed that the Cys2His2 zinc-finger bearing MucR is a novel DNA bridger conserved in α-proteobacteria, with its NTD as the key modulation domain. The ability of NTD, CTD, or full-length protein of xenogeneic silencers from E. coli (Gram-) and Mycobacterium tuberculosis (Gram+) to replace corresponding non-homologous regions or full-length protein of MucR was investigated by testing symbiotic efficiency and genome-wide ChIP-seq profiles of related SF45436 derivatives. The significance of the convergently evolved zinc-finger DNA bridger MucR was further discussed in the context of bacterial pangenome evolution and adaptation.

MATERIALS AND METHODS

Strains, plasmids and growth conditions

The strains and plasmids used in this study are listed in Supplementary Table S1. S. fredii strains were grown at 28°C in TY medium (5 g tryptone, 3 g yeast extract, 0.6 g CaCl2 per liter). E. coli strains were grown at 37°C in Luria-Bertani (LB) medium. Antibiotics were supplemented as required at the following concentrations (μg/ml): nalidixic acid (Na), 30; gentamicin (Gen), 30; ampicillin (Amp), 100; kanamycin (Kan), 50. To test the colony phenotype, rhizobial cells pre-cultured for 48 h in TY medium were collected and resuspended in physiological saline to OD600 = 0.2 after twice of wash. 10 μl of each rhizobial suspension was spotted onto the agar plates of MOPS-buffered minimal medium (64) supplemented with or without Congo Red. Plates were incubated at 28°C for 3 days before taking pictures. For E. coli strains, cells were cultured in LB medium for 12 h to stationary phase. The bacterial suspension was collected, washed twice with physiological saline, and resuspended to OD600 = 0.2. The serially diluted bacterial suspension (10 μl) was spotted on LB agar plates. Colony morphology was photographed after culturing at 37°C for 24 h.

Yeast-two-hybrid assay

The coding sequences of MucR1, MucR2, different fragments of MucR1 and MucR1NTD carrying various mutations were amplified by high fidelity PCR and cloned into the pGADT7 (AD) and/or pGBKTK7 (BD). The desired point mutations were firstly generated on the pTOPO (T-vector, GenStar) derivative carrying the wild-type MucR1 coding sequence by site-directed mutagenesis using high fidelity circular PCR. The derived plasmids were further used as PCR templates for amplification. After validation by PCR and Sanger sequencing, the various pGADT7
and pGBK7 plasmids were co-transformed into Saccharomyces cerevisiae AH109 competent cells. The yeast-two-hybrid assay was performed according to the manual of Matchmaker GAL4 two-hybrid system 3 (TAKARA). Briefly, the co-transformed yeast colony was screened on synthetically defined medium lacking Trp/Leu (SD/-Trp/-Leu). The obtained yeast single colony was resuspended in physiological saline to OD600 = 0.2. The serially diluted yeast suspension was spotted on the SD/-Trp/-Leu agar plate and SD/-Trp/-Leu/-His/X-α-Gal agar plate (containing 6.7 g/l yeast nitrogen base without amino acids, 20 g/l glucose, 1.17 g/l Dropout Supplement-Trp-Leu-His, 40 mg/l/X-α-Gal, 15 g/l agar, pH5.8), and cultured at 28°C for 72 h before recording. Reagents used in SD medium were purchased from Coolaber (Beijing). The colony growth and blue color on the SD/-Trp/-Leu/-His/X-α-Gal agar plate indicate protein interaction. pGADT7-T (SV40) and pGBK7-T53 (p53) were used as positive control pair, and pGADT7-T (SV40) and pGBK7-Lam (Lam) were used as negative control pair in the yeast-two-hybrid assay.

Genetic procedures

The primers used are listed in Supplementary Table S2. All plasmid constructs were transformed into E. coli DH5α and verified by Sanger sequencing before conjugation into rhizobia via triparental mating with pRK2013 as the helper plasmid. To construct the ΔmucR1&2 double mutant, upstream and downstream ∼500-bp flanking regions of mucR2 gene were amplified and assembled with the linearized allelic exchange vector pJQ200SK (65). The derived plasmid was introduced into the ΔmucR1 mutant without gentamicin resistance constructed previously (32). Single-crossover clones were screened on TY-agar plates with gentamicin and subject to passage cultivation in TY medium without any antibiotics. The double deletion mutant clones were further selected on TY-agar plates with 5% sucrose and verified by PCR and Sanger sequencing. Construction of ΔmucR1&2 mutant derivatives carrying MucR1-FLAG, Lsr2(1–50)-MucR1<sub>CTD</sub>-FLAG, Lsr2(1–65)<sub>α</sub>-MucR1<sub>CTD</sub>-FLAG, MucR1<sub>NTD</sub>-Lsr2<sub>(34–112)</sub>-FLAG, H-NS<sub>(1–89)</sub>-MucR1<sub>CTD</sub>-FLAG, MucR1<sub>NTD</sub>-H-NS<sub>(77–137)</sub>-FLAG and H-NS-FLAG were also achieved by using allelic exchange strategy. The corresponding coding sequences of these FLAG-tagged proteins were amplified using SF45436 genomic DNA, pTOPO derivatives and chemically synthesized DNA as templates. These fragments, together with the upstream and downstream regions of mucR1 were tandemly cloned into pJQ200SK. The pJQ200SK derivatives were conjugated into the ΔmucR1&2 double mutant for screening single-crossover and double-crossover clones. Finally, pJQ200SK mediated allelic exchange strategy introduced a single copy gene into the genome. For the plasmids used for complementing the Δhns mutant of E. coli (66), the promoter of hns (300 bp), the coding sequences of H-NS-FLAG, Lsr2-FLAG, MucR1-FLAG, Lsr2(1–65)<sub>α</sub>-MucR1<sub>CTD</sub>-FLAG, MucR1<sub>NTD</sub>-Lsr2<sub>(51–112)</sub>-FLAG and MucR1<sub>NTD</sub>-Lsr2<sub>(34–112)</sub>-FLAG were assembled into Smal I-linearized pBBR1MCS2 and then transformed into Δhns.

Protein 3D structure modeling

All 3-dimensional models of MucR1, H-NS and chimeric proteins explored in this study were predicted by AlphaFold2_mmmseqs2 server using default setting parameters and oligomer states (67,68). Alignment and visualization of models were carried out by using Open-Source PyMOL 2.1.0. Interface residues between two chains of MucR1<sub>NTD</sub> dimer were identified by running InterfaceResidues.py script from PyMOLWiki (https://pymolwiki.org).

Plant assay

Symbiotic performance of S. fredii strains was tested on Glycine soja WSD (62). Seeds were surface-sterilized in 3% NaClO (wt/vol) solution after treatment with concentrated sulfuric acid for 10 min and germinated for 36-48 h under dark condition. Seedlings were then sown into vermiculite moistened with low-N nutrient solution in Leonard jars and inoculated with 1 mL physiological saline suspension of rhizobia with OD<sub>600</sub> = 0.2. Plants were grown at 24°C with a 16-h illumination period for 28 days. Leaf chlorophyll content was measured by a SPAD-502 Meter (Konica Minolta) as described previously (63). Two independent experiments were carried out.

Western blotting

To monitor the protein levels of MucR1 and its derivatives at different growth phases, rhizobial cells cultured in TY medium were sampled every 3 h in a time-course manner that started from OD<sub>600</sub> ≈ 0.3. The cell pellets were resuspended in corresponding volume of SDS-loading buffer to normalize the cell density and then lysed by boiling for 10 min. The lysates were separated on 12% SDS-PAGE and transferred to nitrocellulose membranes. For immunodetection of individual proteins, mouse monoclonal anti-FLAG M2 antibody (Sigma), HRP (horseradish peroxidase)-conjugated Goat Anti-Mouse IgG (Abcam), and eECL Western Blot Kit (CW BIO) were used and chemiluminescence signals were captured and quantified using a Fusion FX6 (Vilber) and the Evolution-Capt Edge software.

Protein stability assay

To detect the protein stability of MucR1 and its derivatives under in vivo conditions, the promoter of mucR1 (486 bp) and coding sequences of these two proteins and C-terminal fused FLAG-tag were tandemly cloned into pBBR1MCS-2. The ΔmucR1 mutant derivatives carrying pBR-MucR1-FLAG and pBR-MucR1<sub>Y24G</sub>-FLAG were cultured in TY medium until OD<sub>600</sub> = 0.8. Cells were pelleted and resuspended in PBS supplemented with 50 μg/ml chloramphenicol. After treatment with indicated time, samples were stopped by adding 5× SDS-loading buffer and boiling for 10 min. The lysates were separated in 12% SDS-PAGE and further analyzed by western blotting.
RNA extraction and RT-qPCR

Bacterial cell cultures were collected in a time-course manner as indicated, and the cell pellets were stored at −80°C before use. The bacterial total RNA kit (ZOMANBIO) was used for total RNA extraction from frozen cell pellet samples. The integrity of the resulting RNA samples was validated by agarose gel electrophoresis. RNA sample concentration and purity were measured by NanoPhotometer™ (Implen). Only qualified RNA samples were used for subsequent experiments. cDNA was synthesized from 500 ng RNA samples by using FastKing-RT SuperMix (TIANGEN). The resulting cDNA was diluted 500 times and used as the template for subsequent quantitative PCR. Reverse transcription quantitative PCR was performed with gene-specific primers using RealStar Green Fast Mixture (Genstar) and a QuantStudio™ 6 Flex Real-Time PCR System (Thermal Fisher Scientific; pre-denature stage, 95°C, 10 min; PCR stage, denature, 95°C, 15 s; anneal and elongate, 60°C, 1 min; 40 cycles). The 16S rRNA gene was used as internal reference to normalize the expression level of the wild-type and mutated mucR1 genes. The specificity and reliability of primers were assessed by melting curves and Sanger sequencing of RT-qPCR products. Algorithm 2–ΔΔCq was used to quantify the target gene’s transcription level relative to the 16S rRNA gene. Three independent experiments were performed.

Protein purification

_E. coli_ BL21(DE3) carrying pET30a-SUMO derivatives harboring His6-SUMO-MucR1, His6-SUMO-MucR1Y24G, His6-SUMO-MucR1-FLAG or His6-SUMO-MucR1Y24G-FLAG was cultured in 500 ml LB medium until OD$_{600}$ = 0.8. Then IPTG was supplied to a final concentration of 0.6 mM in these cultures which were then cultured at 18°C, 150 rpm for 12 h. Cells were harvested by centrifugation (5000 g, 5 min, 4°C) and resuspended in 30 ml lysis buffer (25 mM Tris, pH 8.0, 250 mM NaCl, 10 mM imidazole) supplemented with 0.1 mg/ml DNase I, 0.4 mg/ml lysozyme and protease-inhibitor cocktail (Roche). After 30 min of incubation and 120 cycles of sonication (300 W, 10 s on, 10 s off) on ice, the lysate was cleared by centrifugation (18,000 g, 30 min) and filtration through a 0.22 μm membrane. The supernatant was loaded onto lysis buffer pre-washed Ni-Agarose Resin (CWBCO), and washed three times with wash buffer (lysis buffer with 20/50 mM imidazole) and then eluted with imidazole gradient elution (lysis buffer with 100, 200, 300 mM imidazole). The purified proteins were concentrated by ultrafiltration and redissolved in storage buffer (25 mM Tris, pH 8.0, 250 mM NaCl, 10% glycerol) before use or storage at −80°C. Due to the strong hydrophobicity of MucR1_NTD, it is difficult to obtain high concentration of soluble His$_6$-MucR1 protein. To avoid potential influence of His$_6$-SUMO tag on the subsequent experiments, HRV-3C protease was used to cleave His$_6$-SUMO tag before in vitro protein cross-linking assay where 0.021 μM protein was sufficient. By contrast, for electrophoretic mobility shift assay (EMSA) and DNA-bridging assay which need high concentration of MucR1 and MucR1Y24G proteins in the gradient (from 0.5 μM to 64 μM), the N-terminal His6-SUMO tag was cleaved by HRV-3C protease after incubation of DNA with His$_6$-SUMO-MucR1 or His$_6$-SUMO-MucR1Y24G to avoid protein precipitation as described previously (32).

_In vivo and in vitro protein cross-linking assay

For _in vivo_ protein cross-linking assay, 50 ml TY culture (OD$_{600}$ = 1.2) of each _S. fredii_ strains was pelleted by centrifugation (5000 g, 5 min, 4°C) with twice wash in pre-cooled PBS. Each cell pellet was resuspended in 12 ml PBS supplemented with 10 μl EDTA-free protease-inhibitor cocktail (Roche), 0.1 mg/ml DNase I and 0.4 mg/ml lysozyme, and incubated on ice for 30 min. 50 μl lysates were transferred to new tubes as the control sample. Glutaraldehyde was added to the lysates at a final concentration of 0.02% and incubated on ice for 15 min. 50 μl of cross-linking sample was pipetted and quenched by adding 10 μl 1 M Tris (pH 8.0). For _in vitro_ protein cross-linking assay, the purified His$_6$-SUMO-MucR1-FLAG and His$_6$-SUMO-MucR1Y24G-FLAG were diluted to 10 μg/ml in PBS containing 1 mM dithiothreitol and were then digested with HRV-3C protease to cleave the N-terminal His$_6$-SUMO tag. The cleaved proteins were further dialyzed in PBS (pH 7.4) and incubated with Ni-Agarose Resin (CWBCO) to remove dithiothreitol, His$_6$-SUMO and noncleaved proteins. The MucR1-FLAG and MucR1Y24G-FLAG samples were normalized to 0.021 μM (0.4 μg/ml). Cross-linking reactions were conducted on ice for 8 min or 16 min in the presence of 0.02% glutaraldehyde under conditions with or without various concentrations of DNA probes: the 228-bp Puwx1, the 1069-bp probe harboring this 228-bp region, and the pTOPO-1069 carrying this 1069-bp region. The samples were separated in 12% SDS-PAGE and further analyzed by western blotting.

Electrophoretic mobility shift assay (EMSA)

The electrophoretic mobility shift assay was performed as previously described (32), with modifications as follows. DNA probes were prepared by PCR amplification with the pTOPO plasmid carrying corresponding intergenic regions as templates or annealing complementary synthesized DNA strands using 5’-Cy5-labeled and unlabeled primers as listed in Supplementary Table S2. The binding reaction mixture (10 μl) consisted of 1 mg/ml bovine serum albumin, 1 mg/ml sonicated salmon sperm DNA, 1, 10 or 12.3 mM Cy5-labeled DNA probe, and test proteins in the Tris–KCl buffer (25 mM Tris (pH 8.0), 100 mM KCl, 5% glycerol, 0.05% dodecyl-β-D-maltopyranoside) or the PBS buffer (pH 7.4). The samples were incubated at 25°C for 30 min. Next, 1 μl HRV-3C protease (200 ng/μl; 10 mM dithiothreitol) was added and incubated for 30 min to cleave the N-terminal His$_6$-SUMO tag. To test whether MucR1 can bind to the minor or major groove of DNA, the reagents netropsin and methyl green that specifically bind to the minor and major groove of DNA, respectively, were added as indicated (69,70). The samples were separated in a 6% TB polyacrylamide gel (no EDTA), and the gel was scanned with an Azure Biosystems.
DNA-bridging assay

DNA-bridging assay was performed by using the method described previously (71), with modifications as follows. The DNA probes were prepared by PCR amplification using 5'-Cy5 or 5'-biotin labeled primers as listed in Supplementary Table S2. For each bridging assay, 100 μl hydrophilic streptavidin magnetic beads (NEB) were washed twice with 500 μl PBS and then resuspended in 500 μl coupling buffer (20 mM Tris–HCl, pH7.4, 1 mM EDTA, 500 mM NaCl). Next, 10 pmol biotin-labeled DNA was added to the suspension and incubated with the beads for 30 min at room temperature with gentle rotation. After incubation, the beads were washed twice with 500 μl incubation buffer (20 mM Tris, pH 7.4, 150 mM NaCl, 1 mM dithiothreitol, 5% glycerol (vol/vol), 0.05% Tween 20) and resuspended in incubation buffer supplemented with 10 pmol Cy5-labeled DNA and 10 μl HRV-3C protease to a final volume of 500 μl. The beads suspension was then divided into 50 μl aliquots. Two-fold serially diluted protein samples were added into each of the 50 μl aliquots and supplemented with incubation buffer to a final volume of 60 μl. After a 30 min incubation at room temperature with gentle rotation, the mixture was placed on a magnetic stand for 5 min. The supernatant was transferred into a new tube and marked as sample A. 60 μl elution buffer (incubation buffer supplemented with 0.1% SDS and 20 μg/ml biotin) was added to the beads and incubated in boiling water for 10 min. The eluted sample was transferred into a new tube and marked as sample B. The Cy5 fluorescence signals of sample A and sample B were detected by a Microscale Thermophoresis Monolith NT.115 system (NanoTemper). The Cy5 fluorescence signal of sample A from the reaction without adding protein was defined as 100% input signal. The sample A was also subject to EMSA to check the relative amount of free DNA probe and that bound by MucR1 or MucR1y24G.

Chromatin immunoprecipitation sequencing (ChIP-seq) and ChIP-qPCR

S. fredii strains expressing MucR1, H-NS, or chimeric proteins with C-terminal FLAG were cultured in TY medium until OD600 = 1.2. Formaldehyde was added to the culture to a final concentration of 1%. Cross-linking reactions were sustained for 15 min at 28°C with 180 rpm shaking, and quenched for 15 min after adding glycine to a final concentration of 100 mM. The cross-linked cells were harvested (5000 g, 5 min, 4°C) and washed twice with pre-cooled PBS. The cell pellets were ground into fine powder in liquid nitrogen and resuspended in ChIP buffer (50 mM Tris, pH8.0, 150 mM NaCl, 5 mM EDTA, 0.1% deoxycholate sodium, 1% Triton X-100). Cell lysates were sonicated to shear DNA fragments to an average length of 300–400 bp in a bath sonicator (Q800R3, QSonica) and cleared by centrifugation (13 000 g, 20 min, 4°C). For each ChIP reaction, the supernatant containing 4 mg protein was pre-cleared by 10 μl Dynabeads M-280 (Invitrogen) and then mixed with 5 μl anti-FLAG M2 antibody (Sigma) overnight at 4°C with gentle rotation. 50 μl of beads were added to the samples and incubated at 4°C for 4 h with gentle rotation. The beads were washed twice with ChIP buffer, and once with high salt buffer (50 mM Tris, pH 8.0, 500 mM NaCl, 5 mM EDTA, 0.1% deoxycholate sodium, 1% Triton X-100), LiCl wash buffer (10 mM Tris, pH8.0, 250 mM LiCl, 0.05% deoxycholate sodium, 1 mM EDTA) and TE buffer (10 mM Tris, pH 8.0, 1 mM EDTA). The protein-DNA complex was eluted in 500 μl elution buffer (1% SDS, 0.1 M NaHCO3). To reverse the cross-linking, the eluted samples were supplemented with NaCl to a final concentration of 300 mM, incubated overnight at 65°C and further digested by 0.5 mg proteinase K for 2 h at 45°C. Chip DNA was extracted using phenol:chloroform:isoamylalcohol (25:24:1), ethanol precipitated using Dr GenTLE Precipitation Carrier (TAKARA). 100 μl of the pre-cleared supernatant was used for total chromatin input DNA preparation. The recruitment level of FLAG-tagged proteins in PmucR1 and Pnuxx1 was detected by ChIP-qPCR to assess the quality of Chip DNA. The total chromatin input DNA was diluted to 1 ng/μl, and the Chip DNA was diluted 250 times. These diluted DNA samples were used as the template for subsequent qPCR. An 80-bp fragment inside S456436_h52410 with no protein recruitment signal was amplified by the primers ngtv-QF/ngtv-QR and used as an internal reference to normalize protein recruitment levels in PmucR1 and Pnuxx1. Subsequent qPCR was performed as described in RT-qPCR. Algorithm 2 ΔΔCq was used to quantify protein recruitment levels. The primers used are listed in Supplementary Table S2. Library construction and deep sequencing (paired-end 150 using HiSeq 3000: Illumina) were performed by Novogene-Beijing, with the total chromatin input DNA as control. Three biological replicates were tested. ChIP-seq reads mapping, peak calling, peakID unification between samples and target gene association were performed as previously described (38).

Hemolytic activity assay

The hemolytic activity was detected as previously described with minor modifications (13). In brief, two independent colonies per strain were stabbed into the blood agar plates containing 3% tryptic soy powder, 1.5% agar and 5% defibrinated sheep blood (Solarbio) and incubated at 37°C for 24 h before recording. A clear zone around the E. coli colony indicated hemolytic activity caused by the lysis of red blood cells.

RESULTS AND DISCUSSION

The N-terminal domain is essential for self-association and stability of MucR1

The number of MucR copies within a genome ranges from 1 to 45 in α-proteobacteria and 1 to 6 in δ-proteobacteria, showing multiple independent duplication and horizontal transfer events (20). S. fredii CCBAU45436 (SF45436) has two MucR copies, with MucR1 as a functional xenogeneic silencer of pleiotropic regulatory roles while MucR2 being unable to bind DNA due to a frame shift mutation in its C-terminal DNA binding domain (32,38,46) (Figure 1A). The CTD of MucR/Ros alone can bind DNA in vitro (27) and its NMR structure (PDB: 2jsp for Ros87 from A. tumefaciens) showed RMSD (root-mean-square deviation) values of 1.263 Å and 1.099 Å, respectively, com-
Figure 1. MucR1 self-association and its interaction with MucR2 mediated by the N-terminal domain. (A) Schematic representation of MucR1 and MucR2. MucR2 is not a functional DNA binding regulator due to a frameshift mutation at 114th aa. (B) Predicted protein structure of homo-dimer of the full MucR1 protein by AlphaFold2. LDDT, local-distance difference test measuring the accuracy of prediction. Structure for the top 1 model is shown. (C) The yeast-two-hybrid assay identifies the N-terminal domain (NTD) involved in MucR1-MucR1 and MucR1-MucR2 interactions. MucR2 is indicated in red border. (D) The smallest fragment of NTD involved in self-association is MucR1\textsubscript{17–47}. The same fragments were cloned in AD (pGADT7) and BD (pGBK7). (C, D) AD, pGADT7 derived constructs; BD, pGBK7 derived constructs. The colony growth and blue color on the medium lacking Trp/Leu/His but supplemented with X-gal indicate protein interaction. Results of positive (SV40-p53; blue colonies) and negative (SV40-Lam; no colonies formed) controls in the yeast-two-hybrid assay are shown for comparison. The minimal NTD fragment enabling self-association was MucR1\textsubscript{17–47} (Figure 1D).

In order to characterize conserved sequence features allowing self-association of MucR1\textsubscript{NTD}, 2201 MucR homologs in Pfam database were analyzed and a distinct sequence signature of NTD was identified (Figure 2A), i.e. periodic nonpolar branched-chain amino acids Ile/Val/Leu (20, 21, 25, 30, 35, 39, 42 and 46). Substitutions of these hydrophobic residues with the side-chain free Gly (8/8) or hydrophilic Asn (7/8) led to loss of self-association ability of MucR1\textsubscript{1–56} (except the V30N substitution) in the yeast-two-hybrid assay (Figure 2A and B). By contrast, substitutions with another branched-chain amino acid (the second abundant one among 2201 MucR homologs) had no negative effect on self-association (Figure 2A and B). Notably, there are six spacers of 2–4 residues between these conserved adjacent branched-chain amino acids Ile/Val/Leu (Figure 2A). Substitutions of representative residues in these spacers by Gly or Asn did not abolish self-association (Figure 2A). To investigate the flexibility of spacer length, MucR1\textsubscript{1–56} derivatives with either duplication (+1) or dele-
Figure 2. Identification of key residues and their spacers in NTD involved in self-association. (A) The yeast-two-hybrid assay of MucR1 (1–56) derivatives (the same fragments were cloned in AD and BD). The HMM logo is based on 2201 MucR/Ros homologs in the Pfam database (871 species). The key residues (1st) were substituted by G, N or the secondly conserved residue (2nd). 'Dup', duplication; 'Del', deletion. Six spacers between conserved branched-chain Ile/Val/Leu are indicated by blue and purple numbers. (B) Summary for proteins with substitutions at conserved residues. Red and blue arrows represent loss and maintenance of self-association ability, respectively. (C) Summary for proteins with duplication (+1) or deletion (−1) of the same residue 'X' in spacers (B, C) The numbers in brackets (x/y) indicate x out of y test proteins maintained self-association ability (A–C) Hydrophobic, amphipathic, and hydrophilic residues are indicated in different colors. (D) MucR1NTD dimer predicted by AlphaFold2. Orange, conserved residues. Side chains of conserved residues and/or interface residues (ΔASA > 1.0) are shown in stick mode. ΔASA, changes in accessible surface area.

tion (−1) of representative spacer residues were constructed and tested for the self-association ability in the yeast-two-hybrid assay (Figure 2A and C). It turned out that the length of spacers within the L35–I39–V42–L46 fragment was under strong natural selection (spacer ID: 4, 5, and 6; Figure 2C) while those within the V21–V25–V30–L35 region showed flexibility to certain extent (spacer ID: 1, 2 and 3; Figure 2C). Within the V21–V25 spacer (ID: 1), hydrophobic A23 can be substituted by Gly or Asn without defects in self-association whereas substitution by the second abundant S23 had a notable defect (Figure 2A and B), implying potential context effect of surrounding residues. In addition to these conserved hydrophobic residues, the 24th residue of MucR homologs is distinct for its most abundant amphipathic Tyr carrying a –N–H group on its aromatic ring or the second abundant His harboring two titratable –
N–H groups (Figure 2A). The Y24 of MucR1(1–56) can be substituted by the second abundant H24 but not by non-polar Gly or neutral polar Asn (Figure 2A and B). It has been shown that the number of His-containing networks on the subunit surface of a protein together with the strength of surrounding hydrophobic interactions allow pH-driven conformational changes of helical bundles, and this process is mediated by the pH responsive His protonation levels and hydrogen-bond networks (75). Moreover, π–π stacking and hydrogen-bonding interactions mediated by aromatic moieties of Tyr pairs facilitate the interaction between neighboring α helix (76). Therefore, despite a considerable divergence in NTD among 2201 MucR homologs (Figure 2A), key residues and their spacer length have been under strong natural selection to ensure self-association as also indicated in the MucR1(1–56) dimer predicted by Alphafold2 (Figure 2D).

To test if MucR1 can self-associate in vivo in SF45436, the cross-linking assay was performed for the ΔmucR1Δ2 double mutant carrying MucR1-FLAG or its derivatives with representative substitutions (Y24G, C82G, Y24G&C82G, V21G) or deletions (V29-, and Δ2–56) (Figure 3A). The wild-type MucR1 can form dimer and putative higher-order oligomers in vivo. The MucR1C82G has a substitution in the Cys2His2 zinc-finger motif (X2-Cys79-X2-Cys82-X9-His92-X2-His96) that is essential for DNA binding (52), and exhibited a stronger signal for monomer, dimer and putative higher-order oligomers (Figure 3A). This is in line with the evidence that MucR1 directly represses its own transcription (38). These cross-linking results were supported by docking analysis of either NTD (Figure 3B) or full-length MucR1 (Figure 3C and Supplementary Figure S1) using Alphafold2 (68). Briefly, the NTD homo-dimer was characterized by its two sets of longitudinally arranged α helices corresponding to the region covering three spacers (ID: 4–6) under strong selection (magenta in Figure 3B), and that covering spacers showing relatively higher flexibility (ID: 2–3), respectively. The NTD homo-tetramer and homo-octamer are organized in a circle-like manner (Figure 3B). These self-association patterns mediated by α helices and the adjacent low complex linker region connecting CTD (as exemplified by tetramer in Figure 3C) may allow the MucR1 oligomers to bind multiple sites of DNA. Therefore, both in vitro (59–61) and in vivo evidences support self-association of MucR1.

Moreover, it is noteworthy that MucR1Δ2–56 with a truncated NTD (lacking the 2–56 fragment) showed a low protein level under test conditions (Figure 3A). Similar results were observed for those MucR1 derivatives with substitutions at a conserved branched-chain amino acid (V21G) or at the amphipathic Y24 (Y24G and Y24G&C82G), and MucR1 with a shortened spacer in the V25–V30 region of NTD (V29-) (Figure 3A). The low protein levels of these MucR1 derivatives in vivo, due to either low expression levels or impaired protein stability, made it impossible to draw any conclusions on the role of NTD in self-association in vivo. To explain why protein levels of these MucR1 derivatives are low in vivo, protein and transcriptional levels of representative MucR1 derivatives (Y24G, Y24G&C82G, C82G) were further monitored at exponential phase and stationary phase (Figure 4A). Throughout the 12 sampling points spanning more than 30 h (Figure 4A), the protein level of MucR1C82G was higher than MucR1 which was in turn higher than MucR1Y24G and MucR1Y24G&C82G (Figure 4B), whereas the transcriptional levels of MucR1Y24G and MucR1Y24G&C82G were indistinguishable from MucR1C82G and higher than MucR1 (Figure 4C). The high transcriptional and protein levels of MucR1C82G are consistent with the essential role of zinc-finger (X2-Cys79-X2-Cys82-X9-His92-X2-His96) bearing CTD of MucR1 in transcriptional repression of mucRI (38,52). We further wondered if the higher transcription level of MucR1Y24G compared to MucR1 was due to derepression mediated by potentially impaired DNA binding ability or/and reduced protein stability of MucR1Y24G. EMSA (electrophoretic mobility shift assay) with an equal quantity of purified MucR1Y24G and MucR1 (Supplementary Figure S2) showed that no discrete bandshift was observed at low protein:DNA ratios implying unstable and disintegrating protein-DNA complex (Figure 4D). Moreover, upon adding increasing amounts of protein for both MucR1 and MucR1Y24G (Figure 4D), the protein-DNA complex shifted further (Figure 4D). The bandshift of the PmucRI probe was more significant in the presence of saturated MucR1 than with saturated MucR1Y24G (Figure 4D), implying more MucR1 than MucR1Y24G bound to DNA probe. These results are in line with a working model that MucR1Y24G is unable to self-associate but still individually binds to multiple binding sites in the PmucRI probe. However, this interaction between MucR1Y24G and PmucRI failed to repress the transcription of mucRI as observed in Figure 4C. On the other hand, when the in vivo translation process at the mid-log phase was inhibited by adding chloramphenicol (Figure 4E), MucR1Y24G exhibited a fast decline in protein stability within 4 h while MucR1 was fairly stable 24 h post translation inhibition. Therefore, the NTD is essential for maintaining the maximum stability of MucR1 in vivo.

As revealed above, it is not feasible to investigate the role of NTD in self-association of MucR1 in vivo (Figure 3A), largely due to the impaired stability of MucR1 derivatives with key mutations in NTD (Figure 4E). Moreover, the potential interactions of MucR1 with other proteins might lead to a false positive signal of high-order multimers observed in vivo cross-linking assay (Figure 3A). Consequently, purified MucR1-FLAG and MucR1Y24G-FLAG were tested in the in vitro cross-linking assay (Figure 5A). It is clear that MucR1-FLAG was able to self-associate in dimer and high-order oligomers while MucR1Y24G-FLAG exhibited significant defects in self-association. To test if the presence of target DNA can affect the self-association process of MucR1, the promoter region of uxxs1 targeted by MucR1, revealed earlier (28,38) and confirmed herein (Supplementary Figure S3), was included in the in vitro cross-linking assay (Figure 5B). The amount of MucR1 dimer increased with the concentration of the Puxxs1 probe (from 0 to 5 nM; 228 bp) (Figure 5B). Similar results were obtained when 5 nM of a longer DNA probe (1069 bp) harboring the 228-bp Puxxs1 region or a plasmid-based probe carrying this 1069-bp region (pTOPO-1069; 2934 bp) (Supplementary Figure S4). In short, MucR1 self-associates mainly in the form of dimer and also in various high-order multimer forms under test conditions, which depends on its NTD and can be enhanced by target DNA.
Figure 3. In vivo cross-linking assay of MucR1 and its derivatives, and predicted homo-multimers of MucR1. (A) Late log phase cultures (OD$_{600}$ = 1.2) were subject to cross-linking by glutaraldehyde for 15 min and resolved by SDS-PAGE. The anti-FLAG antibody was used to detect MucR1 and its derivatives (substitutions and deletions). Samples were normalized by bacterial OD$_{600}$. *, red asterisk indicates dimer. (B, C) Predicted protein structure of homo-multimers of the NTD (B) or homo-tetramer of full MucR1 (C) by Alphafold2. Structure for the top 1 model is shown (based on local-distance difference test; Supplementary Figure S1 for details). The color scheme in (B). Blue, Y24; yellow, conserved branched-chain residues; cyan, spacers ID1-3; magenta, spacers ID4-6. Low complex linker region is indicated in (C).

The N-terminal domain is essential for DNA-bridging ability of MucR1

It has been proposed that xenogeneic silencer H-NS from *E. coli* represses gene transcription by bridging DNA fragments (7,18). The DNA-bridging ability has also been demonstrated for the other documented lineage-specific xenogeneic silencers Lsr2 (77), MvaT (78) and Rok (79). Here we tested whether MucR1 could have this DNA-bridging ability. The notable supershift of protein-DNA complex in EMSA for the 228-bp *puxs1* probe was observed when the protein:DNA ratio was increased in the treatment of MucR1 or MucR1Y24G (Supplementary Figure S3). This is similar to canonical xenogeneic silencers which extensively bind DNA in a way that additional dimers bind next to and associate with dimers or multimers already bound to the DNA (18).

Our previous ChIP-seq analysis of MucR1 uncovered the 10–11 bp periodic repeats of T or TT among MucR1 binding peaks (38), which are in line with the ‘class A flexible patterns’ TTxxxGxxxTxxxxxxxTT (80) and an EMSA verified Ros binding motif TxxxxxGxxxxxT (81). Sequence analysis identified three TTxxxGxxxTxxxxxxxTT motifs (named as motif_a) and ten TxxxxxGxxxxxT motifs (named as motif_b) in this *puxs1* probe (Figure 6A; motifs were sequentially numbered from right to left to reflect their relative position to the *puxs1* start codon). The ability of MucR1 to bind 12 fragments harboring these individual motifs was tested by EMSA (Supplementary Figure S5 and S6). All test fragments except those carrying motif_b_10 (AT% = 31%) or motif_b_9 (AT% = 15%) can be effectively bound by MucR1 at relatively low MucR1:DNA ratios. Within the *puxs1* region, four fragments (20–29 bp) covering the other regions without motif_a or motif_b can also be effectively bound by MucR1 (Supplementary Figure S7), and their AT content range from 55% to 83% (Supplementary Figure S7). These results raised a question of to what extent either motif_a or motif_b can modulate MucR1 recruitment efficiency. To answer this question, MucR1 recruitment efficiency by two representative fragments carrying motif_a_1 and motif_b_8, respectively, was compared to their corresponding random probes (without motif_b of the same nucleotide composition) and motif-substituted probes (the T/A at both ends of these putative high-affinity motifs substituted by G/C). Random probes exhibited smaller difference in MucR1 recruitment efficiency compared to wild-type probes than those motif-substituted probes did (Supplementary Figure S5). For example, in the presence of 9.28 µM MucR1, random probes and motif-substituted probes showed 2.99 ± 1.56% and
Figure 4. Post translation stability of MucR1 depends on its NTD. (A) Growth curves of test strains. (B) Dynamic protein levels of MucR1 and its derivatives. The 1st sampling point is indicated in (A). (C) Dynamic transcription levels of mucR1 and its derivatives (relative to 16S rRNA gene). Error bars represent SD of three biological replicates. The 1stsamplingis indicated in (A). (D) EMSA showing impaired bandshift of the PmucR1 (the promoter of mucR1) in the treatment of MucR1Y24G compared to MucR1. The PmucR1 probe, 12.3 nM. Protein concentration: 0.5, 1.5, 4.5, 13.5, 27 and 54 μM, 0 μM. (E) Protein stability assay of MucR1 and MucR1Y24G in the ∆mucR1 mutant carrying pBR-MucR1-FLAG or pBR-MucR1Y24G-FLAG post translation inhibition by chloramphenicol treatment of mid-log phase culture (OD600 = 0.8).

Figure 5. In vitro self-association of MucR1 depends on NTD and enhanced by target DNA. (A) In vitro cross-linking assay of MucR1-FLAG and MucR1Y24G-FLAG. (B) In vitro cross-linking assay of MucR1-FLAG with various concentrations of DNA probe Puxs1. The numbers below the picture refer to the quantified band intensity values of dimers or multimers in each lane. Multimers of different oligomerization levels were collectively quantified.
Figure 6. DNA bridging mediated by MucR1 requires a functional NTD. (A) Distribution of two putative high-affinity binding motifs (with AT content indicated in brackets) and −10/−35 boxes inside the Puxs1. The fragments I (108 bp) and II (120 bp) are indicated. EMSA for 16 fragments fully covering the 228-bp Puxs1 is shown in Supplementary Figures S5-S7. The fragments corresponding to motif b10 and motif b9 show low binding affinity by MucR1 (Supplementary Figure S5-S7). (B) Schematic view of protein-DNA bridging assay. Black ball, streptavidin magnetic bead; gray-ball-labeled green solid line, biotin-labeled DNA probe; yellow-star-labeled green dashed line, Cy5-labeled DNA probe. (C) MucR1 can bridge the fragment I anchored on beads (Bead-I) and Cy5-labeled I in the supernatant (Cy5-I) (108 bp), or ‘Bead-II’ and ‘Cy5-II’. (D) MucR1 can bridge the 228-bp Puxs1 (filled black circles), or ‘Bead-II + Cy5-I’ (open grey circles) while MucR1Y24G almost loses the bridging function on the 228-bp Puxs1 (filled red circles). Error bars represent SD of four measurements for two biological replicates from one out of two independent experiments. (E) EMSA of the supernatant sample in bridge reaction system showing the dosage dependent interactions between MucR1 or MucR1Y24G and Cy5-labeled DNA probes. (F) Working model for DNA bridging mediated by MucR1 NTD (green). MucR1Y24G is not able to efficiently mediate DNA bridging.

13.27 ± 0.02% (mean ± SD) difference, respectively, in their free probe proportion compared to their corresponding wild-type probes. A similar trend can be found when lower concentrations of MucR1 were used (Supplementary Figure S5). Although motif-substituted probes exhibited significantly lower MucR1 recruitment efficiency, these A/T to G/C substitutions did not abolish the interaction (Supplementary Figure S5). This finding was further supported by motif-substituted probes derived from the other 10 fragments carrying either motif a or motif b (Supplementary Figure S6). Therefore, the extensive binding of MucR1 to DNA does not rely heavily on these motifs.

The observed higher MucR1 recruitment efficiency by wild-type probes carrying the 10–11 bp periodic T or TT motifs than the mutated probes with A/T substituted by G/C is more likely due to the decreased AT content. This is in line with the preference for AT-rich sequences by MucR1 and other known xenogeneic silencers (17,38). Indeed, mo-
tif_b.10 (AT% = 31%) and motif_b.9 (AT% = 15%) are of lower AT content compared to the other motifs (54–85%) (Figure 6A) and the genome average (36.9%). Among test fragments harboring motif_b of the same length (21 bp), the highest binding affinity by MucR1 was observed for the one carrying motif_b.8 (AT% = 85%), followed by the other fragments carrying motif_b.5, motif_b.6, motif_b.7, motif_b.1(2), motif_b.3, or motif_b.4 (AT% = 54–77%); Supplementary Figure S3 and Supplementary Figure S6). Overall, MucR1 recruitment efficiency decreased with the increase of DNA GC content. These results provide strong evidence for the potential of this Puxs1 probe to extensively interact with MucR1.

This information allowed us to perform further DNA-bridging assay (Figure 6B), in which biotin-labeled DNA probe is associated with the streptavidin magnetic beads and Cy5-labeled DNA probe can be bridged to bead-associated DNA by DNA-bridging proteins. The whole Puxs1 fragment was roughly split in half: the 108-bp fragment I and 120-bp fragment II (Figure 6A and C). MucR1 can bridge the fragment I (or II) anchored on beads and the Cy5-labeled fragment I (or II) in the reaction buffer. MucR1 can also bridge the bead-anchored fragment II and the Cy5-labeled fragment I (Figure 6D). The Cy5-Puxs1 probe not engaged in DNA-MucR1-DNA bridging was not all free as revealed in EMSA of supernatant samples (Figure 6E). There are two phased A-tracts (5′−AAAAAA−3′ and 5′−ATTT−3′) and four TpA steps between motif_a.3 and motif_b.5 (Figure 6A). A-tract is defined as DNA sequences with three or more consecutive adenines and/or thymines without the TpA step (82). It has been demonstrated that phased short A-tracts (with individual tracts <10 bp) induce a static intrinsic bending in the DNA double-helix (83,84). Moreover, the TpA step is the most flexible dinucleotide step contributing to DNA helix bending (85).

Here we further tested to what extent the two phased A-tracts (5′−AAAAAA−3′ and 5′−ATTT−3′) may be involved in the observed DNA bridging mediated by MucR1 dimers or multimers. Two phased A-tracts were mutated from 5′−AAAAAA−3′ and 5′−ATTT−3′ to 5′−AACAC−3′ and 5′−ATCT−3′, generating a mutated 228-bp Puxs1 which was as efficient as the wild-type Puxs1 in DNA-bridging assay using MucR1 (Supplementary Figure S8). Therefore, MucR1 does not rely on A-tracts to bridge DNA.

In contrast to MucR1, MucR1Y24G exhibited significant defects in DNA bridging (Figure 6D), despite its ability of binding multiple sites of the Cy5-labeled Puxs1 probe in the supernatant (Figure 6E, central panel). Since a portion of the MucR1 pool was trapped in DNA-MucR1-DNA complex on beads while MucR1Y24G was not (Figure 6D), there were more MucR1Y24G than MucR1 available for EMSA of supernatant samples (Figure 6E). Therefore, the NTD-dependent self-association of MucR1 is essential for DNA-MucR1-DNA bridging complex formation (Figure 6F).

As described above, the predicted MucR1 multimers have a quasi-circular structure with their NTDs associating inside and CTDs arranged along an outside ring (Figure 3 and Supplementary Figure S1), which seems to be different from those multimers of H-NS, Lsr2, MvaT and Rok with their CTDs arranged on two sides of the associating NTD backbone (Supplementary Figure S1) (19). It is notable however that the low complex linker region between zinc-finger and N-terminal associating α helices may allow certain flexibility of MucR1 dimer and multimer during the interaction with DNA (Figure 6F).

Moreover, ~10 element lies in the high-affinity region containing motif_b.8 and the region covering ~35 element can also be effectively bound by MucR1 (Figure 6A, Supplementary Figures S5–S7), which together with downstream DNA-bridging events may prevent transcription initiation and/or elongation by RNA polymerase complex. This is supported by earlier evidence that the uxs1 gene is significantly upregulated in the mucR mutants of both SF45436 and Sinorhizobium meliloti (28,46,86).

The essential role of MucR1 and/or its NTD in rhizobium-legume symbiosis can be replaced by non-homologous convergent silencers and/or their NTDs

The divergence of various rhizobial genera belonging to α-proteobacteria predates the emergence of legume symbiotic nitrogen fixation (87–89). We have demonstrated that MucR1, conserved in α-proteobacteria, is essential for maintaining nitrogen fixation efficiency of rhizobia in legume nodules (20,46). Since the observed ever-increasing diversity of rhizobia (more than two hundred species) is largely attributable to the horizontal transfer of AT-rich symbiosis genes (90), we wondered if the convergently evolved xenogeneic silencing function of MucR (17,20) pre-disposed α-proteobacteria to successfully integrate horizontally acquired AT-rich symbiosis genes. To test this hypothesis, H-NS from E. coli (Gram-), Lsr2 from M. tuberculosis (Gram+), and various MucR1 derivatives with either NTD or CTD replaced by corresponding fragments from H-NS or Lsr2 were introduced into the chromosome of the ΔmucR1Δ2 mutant of SF45436 as a single copy, which is under the control of the wild-type PmucR1 promoter (Figure 7A). The ΔmucR1Δ2 mutant carrying Lsr2 was not obtained due to the potentially lethal effect. Lsr2(1–50) and Lsr2(1–65) when fused with CTD of H-NS can complement phenotypes of the hns mutant of E. coli to a different extent (14), and therefore these two NTD fragments of Lsr2 were explored herein to generate the ΔmucR1Δ2 mutant carrying Lsr2NTD-MucR1CTD.

Successful expression of H-NS and various chimeric MucR1 derivatives in the ΔmucR1Δ2 mutant was verified by the in vivo cross-linking assay (Figure 7B), and self-association in the dimer form can be observed for H-NS, H-NS(1–89)−MucR1CTD, Lsr2(1–65)−MucR1CTD, MucR1NTD−H-NS(77–137), and MucR1NTD−Lsr2(74–112), but not for Lsr2(1–50)−MucR1CTD. It has been revealed that MucR1 is required for exopolysaccharide production and nitrogen fixation by directly modulating the transcription of exopolysaccharide biosynthesis genes and functional genes essential for maintaining nitrogen fixation process (32,38,46,91,92). The ΔmucR1Δ2 mutant of SF45436 is characterized by forming non-mucoid colonies, and small nodules of low nitrogenase activity which led to a significant reduction in leaf chlorophyll content of soybean plants (38,46) (Figure 7C, D). H-NS, H-NS(1–89)−MucR1CTD,
**Figure 7.** Convergent xenogeneic silencers can rescue symbiotic defects of the ΔmucR1&2 mutant. (A) Schematic diagram of domain swap among convergent xenogeneic silencers Lsr2, H-NS and MucR1. (B) In vivo cross-linking experiment of the ΔmucR1&2 mutant derivatives carrying MucR1-FLAG, Lsr2\(_{2(1-50)}\)-MucR1\(_{CTD}\)-FLAG, Lsr2\(_{2(1-65)}\)-MucR1\(_{CTD}\)-FLAG, MucR1\(_{NTD}\)\(-\)Lsr2\(_{2(74-112)}\)-FLAG, H-NS\(_{1-89}\)\(-\)MucR1\(_{CTD}\)-FLAG, MucR1\(_{NTD}\)\(-\)H-NS\(_{77-137}\)-FLAG or H-NS-FLAG. (C) Symbiotic performance of the ΔmucR1&2 mutant derivatives on wild soybean plants. Leaf chlorophyll content and vertical section of nodules are displayed. Different letters indicate significant differences between means (mean ± SEM; ANOVA followed by Duncan’s test, alpha = 0.05) of 7–12 plants. (D) Colony phenotypes of test strains on MOPS-buffered MM medium with or without Congo Red. Lsr2\(_{2(1-65)}\)-MucR1\(_{CTD}\), MucR1\(_{NTD}\)\(-\)H-NS\(_{77-137}\) rather than Lsr2\(_{2(1-50)}\)-MucR1\(_{CTD}\) and MucR1\(_{NTD}\)\(-\)Lsr2\(_{2(74-112)}\) can effectively rescue the symbiotic performance of the ΔmucR1&2 mutant on soybean plants (Figure 7C). The red non-mucoid colony phenotype of the ΔmucR1&2 mutant on the minimum medium containing Congo Red can be largely rescued by H-NS\(_{1-89}\)\(-\)MucR1\(_{CTD}\), Lsr2\(_{2(1-65)}\)-MucR1\(_{CTD}\) and to a less extent by MucR1\(_{NTD}\)\(-\)H-NS\(_{77-137}\) and H-NS (Figure 7D). MucR1\(_{C82G}\) and MucR1\(_{Y24G}\) were not able to restore the symbiotic and colony phenotypes of the ΔmucR1&2 mutant (Figure 7C and D). These results demonstrated that NTD of MucR1 can be replaced by Lsr2\(_{2(1-65)}\) and H-NS\(_{1-89}\) while H-NS\(_{77-137}\) but not Lsr2\(_{2(74-112)}\) can functionally complement CTD of MucR1 in vivo. Therefore, convergently evolved xenogeneic silencers H-NS and Lsr2 can fully or partially complement the essential role of MucR1 in rhizobium-legume symbiosis.

**MucR1, H-NS, and functional chimeric silencers share similar recruitment profiles in rhizobial genome**

To further explore mechanisms underlying phenotype differences among these ΔmucR1&2 derivatives carrying foreign or chimeric silencers, ChIP-seq analyses were performed (Figure 8A). H-NS, H-NS\(_{1-89}\)\(-\)MucR1\(_{CTD}\), Lsr2\(_{2(1-65)}\)-MucR1\(_{CTD}\) and MucR1\(_{NTD}\)\(-\)H-NS\(_{77-137}\) had similar recruitment profiles across the multipartite genome of SF45436 while Lsr2\(_{2(1-50)}\)-MucR1\(_{CTD}\), MucR1\(_{NTD}\)\(-\)Lsr2\(_{2(74-112)}\), MucR1\(_{Y24G}\) and MucR1\(_{C82G}\) didn’t show significant recruitment signals. These results were confirmed by ChIP-qPCR analyses of P\(_{uxs1}\) and P\(_{mucR1}\) (Supplementary Figure S9). Recruitment profiles were highly reproducible in three independent biological replicates (Supplementary Figure S10). Targets specific to individual wild-type or chimeric silencers had significantly less recruitment levels than those shared by multiple silencers (Figure 8B, C; Supplementary Figure S11). The
Figure 8. ChIP-seq of the ΔmueR1&2 mutant derivatives harboring convergent xenogeneic silencers. (A) ChIP-seq analysis showing recruitment levels of MucR1-FLAG, H-NS-FLAG, Lsr2(1–65)-MucR1CTD-FLAG, H-NS(1–89)-MucR1CTD-FLAG, MucR1NTD-H-NS(77–137)-FLAG, MucR1Y24G-FLAG, MucR1C82G-FLAG, Lsr2(1–50)-MucR1CTD-FLAG, or MucR1NTD-Lsr2(74–112)-FLAG in the ΔmueR1&2 mutant derivatives. The pooled ChIP-seq data from three independent biological replicates (Supplementary Figure S10) are shown. GC% below (black) and above (gray) the genome average is indicated (window size 5000 bp). (B) Venn diagram showing the number of ChIP-seq peaks specific to each silencer or shared by different silencers. Predicted protein structures of homo-dimer by Alphafold2 are shown (the top 1 model based on local-distance difference test). (C) Silencer recruitment levels of ChIP-seq peaks specific to one strain (1) or shared by different strains (2–5). The number of peaks is shown in brackets. Different letters indicate significant differences between medians (Dunn’s test, $\alpha = 0.05$). Error bars represent SD of mean. Results for individual strains are shown in Supplementary Figure S11. (D) Enrichment analysis of 286 common peak-associated genes regarding replicons. Orange and black bars represent proportions of common peak-associated genes and genome size, respectively. Significant enrichment/depletion is indicated (** $P$ value < 0.01; chi-square test).

Targets of high recruitment levels were generally located in AT-rich genomic regions (Figure 8A) and enriched in the symbiosis plasmid but depleted in the chromosome (Figure 8D). This is consistent with that DNA binding preferences of xenogeneic silencers Lsr2, H-NS, MvaT, Rok and MucR1 are positively correlated to the AT% of DNA sequences (17,38). Despite that NTDSs from MucR1, H-NS and Lsr2 show no sequence homology (20), all of them can mediate self-association as demonstrated in cross-linking assay (Figure 7B) and structure prediction by Alphafold2 (Figure 8B). This allowed the DNA-binding domain either from MucR1 or H-NS to effectively target similar AT-rich genomic regions of SF45436 (Figure 8A, C and D). Among the genes associated with 286 shared ChIP-seq peaks (Figure 8D), 144 genes have COG annotation (Supplementary Figure S12). This list includes MucR1 (chromosome), MucR2 (pSFa), and those involved in exopolysaccharide biosynthesis (uxs1 and exoY; chromid), quorum sensing (TraI, TraR; pSymA), chemotaxis (VisN; chromosome), nodulation signal biosynthesis and regulation (NodU, NoI, Noel and NodD2; pSFa), type III secretion system (T3SS) components, effector and transcriptional
The ability of MucR1, H-NS, H-NS (1–89)-MucR1CTD, Lsr2(1–65)-MucR1CTD and MucR1NTD-H-NS(77–137) to target these genes and allow SF45436 to establish efficient symbiosis with soybean plants provides strong evidence for that the convergently evolved xenogeneic silencer predisposed α-proteobacteria to integrate horizontally transferable symbiosis genes.

It remains elusive why chimeric proteins with CTD from H-NS rather than that from Lsr2 work in vivo in the ΔmucR1&2 mutant of SF45436. Indeed, we failed to obtain MucR1NTD-Lsr2(51–112) and full-length Lsr2 complementary strains in the ΔmucR1&2 mutant background. ChIP-seq data (Figure 8) indicated that MucR1NTD-Lsr2(74–112) had no apparent binding peaks in the SF45436 genome. EMSA was further performed to test whether Lsr2, MucR1NTD-Lsr2(51–112) and MucR1NTD-Lsr2(74–112) could be recruited by Puxsi in vitro (Supplementary Figure S13). The data showed that Lsr2 and MucR1NTD-Lsr2(51–112) had stronger DNA binding capacity compared to MucR1 throughout the test gradient of protein:DNA ratios (Supplementary Figure S13). In the treatment of Lsr2 and MucR1NTD-Lsr2(51–112), the corresponding protein-DNA complexes were even stuck in the gel wells. It is likely that such strong protein-DNA interactions may lead to the lethal effect of MucR1NTD-Lsr2(51–112) and Lsr2 in SF45436. By contrast, MucR1NTD-Lsr2(74–112) exhibited a reduced bandshift than MucR1 at high protein:DNA ratios (Supplementary Figure S13), suggesting fewer proteins bound to the DNA probe. This is in line with its weak dimer signals in vivo (Figure 7B). Similarly, Lsr2(1–50)-MucR1CTD and MucR1CTD could show dimer-forming defects in vivo (Figure 7B) but could not be effectively recruited to DNA in ChIP-seq (Figure 8). Therefore, MucR1 derivatives with defects in dimer assembly exhibited impaired DNA binding ability in vitro (Figure 4D, Supplementary Figure S3, and Supplementary Figure S13) and their interactions with DNA in vivo might be too fragile to be detected by the standard ChIP assay (Figure 8A and Supplementary Figure S9). Anyway, the in vitro evidence for CTD-DNA interactions should be carefully addressed when their physiological relevance is discussed. By contrast, the stability and self-association of MucR1 can be ensured by its own NTD, Lsr2(1–65), or H-NS(1–89) (Figure 7B) and H-NS(1–89)-MucR1CTD and Lsr2(1–65)-MucR1CTD can largely restore symbiotic defects and colony phenotypes of the ΔmucR1&2 mutant (Figure C and D).

MucR1 complements characteristic phenotypes of the Δhns mutant of E. coli

To further test if MucR from α-proteobacteria can convergently function in γ-proteobacteria as Lsr2 from Gram+ Mycobacterium, mucR1 from S. fredii was intro-

duced into the Δhns mutant of E. coli (66). The Δhns mutant of E. coli has a characteristic mucoid colony phenotype and generates a clear zone around its colony on the blood agar plate due to hemolysis activity, which can be complemented by Lsr2 from Gram+ Mycobacterium (13). In this work, these characteristic phenotypes of the Δhns mutant of E. coli can be fully complemented by H-NS, Lsr2 and MucR1, and partially by Lsr2(1–65)-MucR1CTD, MucR1NTD-Lsr2(51–112) and MucR1NTD-Lsr2(74–112) (Supplementary Figure S14). The protein expression was validated by Western blot using anti-FLAG antibody (Supplementary Figure S14). Therefore, MucR1 from α-proteobacteria and its various chimeric proteins with Lsr2 from Gram+ Mycobacterium can functionally replace H-NS in E. coli. These results strongly support that MucR1 is a convergently evolved xenogeneic silencer.

MucR1 contacts both minor and major grooves of target DNA

All known prokaryote xenogeneic silencers H-NS, Lsr2, MvaT and Rok bind the minor groove of DNA (17,18,20), while MucR1 and its close homolog Ros belong to the zinc-finger proteins which are well-known for their binding in the major groove in eukaryotes (54). The classic eukaryotic zinc-finger consists of a β-sheet and an α-helix (ββα fold) while the DNA binding domain of prokaryote Ros (Ros87) has a ββαβ topology and two additional C-terminal basic regions (27). Similar to the classic zinc-finger, α1 of Ros87 contacts the major groove of DNA (81), while notably the basic region in α2 and two additional C-terminal basic regions make additional contacts with DNA and stabilize Ros87-DNA interaction (81). These three basic regions are essential for DNA binding ability of Ros87 in vitro (55). In this work (Figure 9), both minor-groove binding drug netropsin (Figure 9A) and major-groove binding drug
methyl green (Figure 9B) can impair the binding of MucR1 to a test fragment of Puxs1. Therefore, MucR1 has the ability to bind both major and minor grooves of DNA. Taken together, the following working model can be proposed: the first α-helix α1 in the zinc-finger structure of Ros/MucR is inserted into the major groove of DNA, and its C-terminal basic regions stabilize protein-DNA interaction by allowing Ros/MucR to bind DNA more extensively from the major groove to the minor groove of DNA.

CONCLUSION

This work reveals that the xenogeneic silencer MucR conserved in α-proteobacteria (20,38) is a novel DNA bridge with its NTD acting as a major modulator. The NTDs from more than 2000 MucR homologs have evolved conserved key residues and their spacer length to ensure self-association (Figure 1 and Figure 2). The NTD-dependent self-association can be enhanced by target DNA (in vivo and in vitro; Figure 3 and Figure 5), and is crucial for maintaining MucR stability (in vivo; Figure 4), transcriptional silencing (in vivo; Figure 4), and DNA–MucR–DNA bridging complex formation (in vitro; Figure 6). Like Lsr2 from M. tuberculosis, MucR can functionally complement H-NS in E. coli (Figure S14). On the other hand, NTDS from H-NS and Lsr2 can functionally replace the NTD of MucR regarding in vivo self-association (Figure 7) and symbiotic interaction of S. fredii with soybean plants (Figure 7). Moreover, the full-length H-NS can also rescue the symbiotic defect of the mucR mutant of S. fredii (Figure 7). H-NS and functional chimeric MucR derivatives have similar recruitment profiles across the multipartite genome of S. fredii, preferring AT-rich genomic islands and symbiosis plasmid, with key symbiosis genes as shared targets (Figure 8 and Supplementary Figure S12). Therefore, the convergently xenogeneic silencer MucR conserved in α-proteobacteria is crucial for the successful integration of later acquired symbiosis genes, horizontal transfer of which is strongly selected in nature (96–98). Notably, functional MucR homologs can be found in symbiosis plasmids of diverse rhizobial species and bacterial phages (20,46), both of which can be horizontally transferred in microbiota (99,100). Since the AT-rich symbiosis genes in symbiosis plasmid are specifically transcribed during symbiotic interactions with legumes rather than under free-living conditions (101,102), it is intriguing to hypothesize that co-transfer of MucR with AT-rich sequences can be an adaptive mechanism by banking foreign genetic materials to fuel the process of exploring novel niches while maintaining regulatory integration. This may at least partially explain the successful spreading of symbiosis genes among hundreds of bacterial species in α-proteobacteria (90).

Since α-proteobacterium is enriched with both symbiotic and pathogenic bacteria associated with eukaryotes and had a putative ancestor for mitochondria (21), the working mechanisms of zinc-finger bearing DNA bridge MucR are of significant importance. The interaction between H-NS-like proteins and DNA can respond to various physico-chemical environmental signals, and diverse anti-silencing mechanisms have been reported (18,103–105). Among hundreds of direct targets of MucR in the multipartite genome of facultative microsymbiont Sinorhizobium fredii (38), many functional genes are involved in stress adaptation or symbiotic interactions (32,48,88,91,106), in which silencing and anti-silencing mechanisms remain unknown. Our current understanding of the biology of MucR in α-proteobacteria is just the tip of the iceberg.

DATA AVAILABILITY

Raw sequence data from ChiP-seq analyses can be accessed via NCBI Sequence Read Archive (PRJNA775862).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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