Structural basis for feed-forward transcriptional regulation of membrane lipid homeostasis in Staphylococcus aureus.

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Introduction

The cell membrane is an essential structure to bacteria. It primarily consists of a fluid phospholipid bilayer in which a variety of proteins are embedded. Most steps involved in phospholipid biosynthesis are therefore explored as targets for designing new antibacterial drugs [1]. The central events in the building of phospholipids enclose the biosynthesis of fatty acids, which are the most energetically expensive lipid components, by the type II fatty acid synthase (FASII) on the cytoplasmic side of the membrane, and subsequent delivery of the latter to the membrane-bound glycerol-phosphate acyltransferases (Fig. S1). Due to the vital role of the membrane lipid bilayer, bacteria have evolved sophisticated mechanisms to finely control the expression of the genes responsible for the metabolism of phospholipids [2].

Transcriptional regulation of bacterial lipid biosynthetic genes is poorly understood at the molecular level. Indeed, the only two well-documented examples are probably those of the transcription factors FadR and DesT, which regulate the biosynthesis of unsaturated fatty acids (UFA) in *Escherichia coli* and *Pseudomonas aeruginosa*, respectively. FadR was discovered as a repressor of the β-oxidation regulon [3,4] and subsequently found to also activate the transcription of *fabA* and *fabB*, two essential genes for the biosynthesis of UFA [5–7]. Binding of FadR to its DNA operator is antagonized by different long-chain acyl-CoAs [8–11], in agreement with the proposed structural model [12–14]. DesT is a TetR-like repressor that primarily controls the expression of the...
Author Summary

An opportunistic Gram-positive pathogen, Staphylococcus aureus is a major threat to humans and animals, being responsible for a variety of infections ranging from mild superficial to severe infections such as infective endocarditis, septic arthritis, osteomyelitis and sepsis. The increasing resistance of S. aureus against most current antibiotics emphasizes the need to develop new approaches to control this important pathogen. The lipid biosynthetic pathway is one appealing target actively pursued to develop anti-Staphylococcal agents. Despite its potential biomedical importance, however, little is known about the signaling mechanisms that allow S. aureus to control its phospholipid content. In order to shed light on this fundamental mechanism, we studied S. aureus FapR (SaFapR) a transcription factor that senses the levels of malonyl-CoA, a key intermediate in fatty acid biosynthesis, and modulates the expression of genes involved in fatty acid and phospholipid biosynthesis. Our studies of SaFapR uncovered the mechanistic basis of a complex biological switch that controls membrane lipid homeostasis in S. aureus. We also discovered that disruption of the ability of SaFapR to recognize malonyl-CoA, the ligand that controls SaFapR binding to DNA, compromises S. aureus viability, thus revealing new opportunities for the development of antibiotics against this major human pathogen.

Results

FapR senses malonyl-CoA and regulates lipid biosynthesis in S. aureus

The S. aureus fap regulon is organized in an B. subtilis [18], except for two missing genes (yhcC and fabHB) (Table S1). Electrophoretic mobility shift assays revealed that SaFapR binds to its own promoter (PfapR) and that malonyl-CoA specifically disrupts the repressor-operator complex (Fig. S2). Furthermore, the unlinked genes plsX and fabH from the fap regulon are upregulated in two distinct S. aureus strains lacking the repressor (Fig. S3A) and SaFapR is able to complement a B. subtilis fapR mutant strain (Fig. S3B). These results clearly demonstrate that SaFapR conserves the same regulatory function originally described for the B. subtilis orthologue [20].

The SaFapR-DNA complex

The crystal structure of SaFapR in complex with a 40-bp oligonucleotide comprising the PfapR promoter was determined at 3.15 Å resolution (Table 1). Two SaFapR homodimers were observed to bind to a single DNA molecule in the crystal (Fig. 1A). This observation is in agreement with isothermal titration calorimetry (ITC) studies of protein-DNA interactions, which showed a complex isotherm with a large exothermic component and two distinct binding reactions (Fig. 1B). Sequence analysis of the promoters of the fap regulon from B. subtilis [18] and S. aureus (Fig. S4) revealed the presence of a conserved inverted repeat. Our previous DNAse I footprinting analyses performed with BsFapR on both strands of the B. subtilis fapR promoter demonstrated that this symmetric element covers half of the DNAse protected region [20]. Interestingly, this region corresponds to the recognition site of one SaFapR homodimer in the crystal, suggesting a sequential mechanism of binding. Indeed, a sequential binding model fits well the observed isotherm (Fig. 1B), indicating that the two SaFapR homodimers bind the operator with nanomolar dissociation constants of 0.5±0.1 nM for the first and 51±8 nM for the second binding reactions. In the crystal structure of the complex, however, the two SaFapR homodimers are related by a local two-fold symmetry axis, and the DNA molecule is bound in a 50:50 orientation, as confirmed by determining the crystal structure of SaFapR in complex with an asymmetrically Br-labelled oligonucleotide using the Br anomalous scattering signal (data not shown).

Each protein homodimer exhibits an elongated asymmetric conformation in the crystal structure, in which the two DNA-bound DBDs are structurally detached from the central dimeric ‘hot-dog’ EBD. Indeed, the positioning of the EBD (stabilized by crystal packing contacts) relative to the DBD-DNA complex requires partial unwinding of the C-terminal end of αL, as
confirmed by comparing the structures of the ligand-free and DNA-bound forms of SaFapR (see below). The amphipatic linker helix \( \alpha_L \) plays a major role in stabilizing the molecular architecture of the SaFapR-DNA complex. Helix \( \alpha_L \) interacts with \( \alpha_L \) from the second protomer mainly through its exposed hydrophobic face (including residues Ile59, Val62 and Ala63; Fig. S5A). This dimerization region is further stabilized by hydrophobic contacts (Phe26) and hydrophilic interactions between \( \alpha_L \) and the \( \alpha_1-\alpha_2 \) connecting loop from both DBDs. In addition, the guanidinium group of Arg59 gets engaged in electrostatic interactions with the main-chain carbonyl groups of Pro25 and Ile27, and the hydroxyl group of Tyr67 is within hydrogen bonding distance to the main-chain of Ser23 at the end of \( \alpha_1 \) on the second protomer (Fig. S5A). Altogether this contact network largely contributes to the correct relative positioning of the two helix-turn-helix motifs on the DNA major groove.

The two DBDs from the asymmetric homodimer interact in a similar manner with DNA. The recognition helix \( \alpha_3 \) from the helix-turn-helix motif penetrates the double-helix major groove and makes extensive interactions with the two DNA chains (Fig. 2A). The SaFapR-DNA interface mainly involves the DNA backbone phosphates (Fig. 2B), except for base-specific interactions of Gln41 (from the recognition helix \( \alpha_3 \)) in the major groove and Arg56 (from the linker helix \( \alpha_L \)) in the minor groove. Importantly, the insertion of the guanidinium groups of this arginine from both protomers promotes the opening of the minor groove, inducing a pronounced local bending of DNA (Fig. 2C). The two phosphate-contacting residues from helix \( \alpha_1 \) (Lys10 and Arg13), as well as residues from the recognition helix \( \alpha_3 \) and the C-terminal half of the linker helix \( \alpha_L \) (which include most other DNA-contacting residues) are highly conserved in the entire FapR protein family (Fig. S6), indicating a conserved mode of DNA-binding. The N-terminus of the protein (preceding helix \( \alpha_1 \)) faces the adjacent minor groove in the crystal structure (Fig. 2A), suggesting that it could be engaged in additional DNA interactions. However, this region is poorly conserved in the FapR protein family and is disordered in the crystal structure.

### Structure of the repressor-effector complex

The crystal structure of full-length SaFapR in complex with malonyl-CoA, determined at 1.9 Å resolution (Table 1), revealed a different, more compact conformation of the repressor. Most notably, the two amphipatic linker helices \( \alpha_L \), instead of interacting with each other as in the DNA-bound repressor, now bind to either side of the central dimeric EBD (Fig. 3A). This results in a protein conformation with the two

| Table 1. Data collection, phasing and refinement statistics. |
|---|---|---|---|---|
| **SaFapR-DNA** | **SaFapR-malonyl-CoA** | **SaFapR (form 1)** | **SaFapR (form 2)** |
| **Data collection** |
| Space group | C222, | P2, | H32 | P3,21 |
| Cell dimensions (Å) |
| \( a, b, c \) | 114.69, 249.94, 179.21 | 53.38, 68.5, 61.16 | 116.58, 116.58, 123.23 | 110.52, 110.52, 115.11 |
| \( \alpha, \beta, \gamma \) | 90, 90, 90 | 90, 111.3, 90 | 90, 90, 120 | 90, 90, 120 |
| Resolution (Å) | 35-3.15 (3.32-3.15) | 44-1.9 (2-1.9) | 47-2.4 (2.53-2.4) | 48-2.6 (2.74-2.6) |
| \( R_{	ext{sym}} \) or \( R_{	ext{merge}} \) | 0.099 (0.572) | 0.043 (0.349) | 0.060 (0.634) | 0.061 (0.426) |
| \( I/\sigma \) | 24.3 (2.4) | 16.4 (3.6) | 15.6 (2.5) | 16.7 (3.4) |
| Completeness (%) | 98.2 (93.7) | 99.3 (100) | 99.9 (100) | 99.7 (98.5) |
| Redundancy | 16.0 (4.9) | 3.7 (3.7) | 5.5 (5.6) | 3.7 (3.6) |
| **Refinement** |
| Resolution (Å) | 3.15 | 1.9 | 2.4 | 2.6 |
| No. reflections | 41820 | 30557 | 12145 | 24146 |
| \( R_{	ext{work}}/R_{	ext{free}} \) | 0.186/0.201 | 0.198/0.235 | 0.196/0.246 | 0.185/0.211 |
| No. atoms |
| Protein | 6040 | 2538 | 1460 | 2431 |
| Ligand/Ion | 1634 | 98 | 25 | 9 |
| Water | - | 150 | 20 | 52 |
| \( B \)-factors |
| Protein | 104 | 42 | 63 | 47 |
| Ligand/Ion | 149 | 41 | 67 | 57 |
| Water | - | 28 | 34 | 18 |
| Ramachandran outliers \( ^1 \) (%) | 0.5 | 0 | 0 | 0 |
| Ramachandran favored \( ^1 \) (%) | 95.8 | 98.1 | 97.8 | 97.0 |
| R.m.s. deviations |
| Bond lengths (Å) | 0.007 | 0.012 | 0.012 | 0.017 |
| Bond angles (°) | 1.14 | 1.39 | 1.64 | 1.61 |

Values in parentheses are for highest-resolution shell.

\(^1\)According to the MolProbity server [53].

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helix-turn-helix domains far apart from each other, corresponding to an incompetent DNA-binding state. Interactions of helix $\alpha_L$ with the lateral face of the EBD play a major role in stabilizing the observed quaternary organization of the protein, mainly through extended hydrophobic contacts between aliphatic and aromatic side chains (Fig. 3B). In addition, several hydrogen bonding and salt bridge interactions (Fig. S5B) lock a well-defined structure for the loop connecting helix $\alpha_L$ to the first $\beta$-strand of the EBD (residues 70–82), which further restrict the mobility of $\alpha_L$ and contrasts with the loose asymmetric conformation observed for this loop in the DNA-bound SaFapR homodimer (Fig. 1A).

The phosphopantetheine and malonyl groups of malonyl-CoA are well defined in the electron density map (Fig. 3C). The phosphopantetheine group binds within a tunnel at the interface between the two protomers in the SaFapR homodimer, and adopts the same conformation as observed in many acyl-CoA-binding proteins displaying the hot-dog fold [20,24]. This binding mode results in the complete occlusion of the ligand malonate from the bulk solvent. The charged carboxylate group of malonate is

Figure 1. Overall structure of the SaFapR-operator complex. (A) Surface representation of the DNA operator (in red) with two bound FapR homodimers looking down the non-crystallographic two-fold symmetry axis. For one homodimeric repressor (in yellow and orange) the DNA-binding domains (DBDs), the linker helix $\alpha_L$ and the dimeric effector-binding domain (DBD) are indicated. (B) ITC study of SaFapR binding to the PfapR operator at 25°C. The top panel shows the raw heat signal for 6 µl injections of a 68 µM solution of SaFapR dimer into a 4 µM solution of the 40 bp DNA oligonucleotide (curve 1 obtained by subtraction of the SaFapR dilution energy curve 3 from the raw titration curve 2). The bottom panel shows the integrated injection heats after normalization fitted with a sequential binding model. Two SaFapR dimers bind the operator, with parameters ($K_d = 0.5\pm0.1$ nM, $\Delta H_i = -22.5\pm0.2$ kcal/mol, $T i S_i = -9.8\pm0.2$, kcal/mol) and ($K_d = 51\pm8$ nM, $\Delta H_{ii} = -6.95\pm0.2$ kcal/mol, $T_{i S_{ii}} = 3.0\pm0.3$ kcal/mol). doi:10.1371/journal.ppat.1003108.g001
neutralized through a salt bridge with the guanidinium side-chain of Arg110, and makes additional hydrogen bonding interactions with the main-chain nitrogen from Gly111 and the side-chain of Asn119 from the second protomer (Fig. 3C). The engagement of Arg110 in malonyl-CoA-binding triggers a local reorganization of hydrogen-bonding interactions (Fig. S5B) and surface reshaping that further stabilize the loop connecting aL to the first b-strand of the hot-dog fold in the non-productive conformation (Fig. 3A), thus preventing DNA binding. On the other hand, the adenosine 39-phosphate moiety of malonyl-CoA is largely exposed to the solvent (it is partially disordered in one of the two protomers) and makes no specific contacts with the protein. This implies that SoFapR specifically recognizes the malonyl-phosphopantetheine moiety of the ligand, in agreement with the observation that either malonyl-CoA or malonyl-acyl carrier protein (malonyl-ACP) can indistinctly function as effector molecules [28].

A detailed comparison of the malonyl-CoA complexes between full-length SoFapR and the truncated form of BoFapR (lacking the DBDs) revealed a conserved structural arrangement of the EBD core, and ligand binding promoted the same conformation of the connecting loop aL-B1 (Fig. S7). On the other hand, helix aL displays a different organization, due in part to the absence of the DBDs. This helix protrudes away from the EBD core in BoFapR to get involved in crystal packing contacts [20]. Interestingly, hydrophobic residues engaged in these inter-domain interactions are largely invariant in the whole FapR family (Fig. S6), as are also key residues involved in electrostatic interactions (Fig. S5). Altogether, the structural alignment indicates not only an identical mode of malonyl-CoA binding but also the conservation of the DBD – aL – EBD interactions required to stabilize the FapR-malonyl-CoA complex as visualized in the SoFapR model.

The overall structure of SoFapR

The crystal structure of full-length SoFapR in the absence of ligands has been obtained in two different crystal forms (Table 1). Interestingly, in three out of four crystallographically independent protomers the repressor has the same non-productive quaternary arrangement as observed in the structure of malonyl-CoA-bound SoFapR, with helix aL bound to the lateral face of the EBD (Fig. 4A), strongly suggesting that this compact structure likely represents a conformation of the protein in solution. However, the helix-turn-helix motifs display high temperature factors or even partial disorder, and are engaged in extensive crystal contacts, suggesting the coexistence of alternative conformational states in solution characterized by flexible DBDs. In that sense, in one protomer of the crystal form 2 (Table 1) both helix aL and its associated DBD are highly flexible and could not be modeled, suggesting a marginal stability of the observed quaternary arrangement. Moreover, the first visible residues of this same monomer (positions 72–77, connecting helix aL with the first b-strand of the EBD) adopt a conformation that differs from that observed in the other ligand-free or malonyl-CoA-bound proto-
mers, but resembles that found for one subunit of the asymmetric DNA-bound form of the repressor (Fig. 4B).

Ligand-induced structural transition
The structure of SaFapR in complex with the DNA operator revealed a strikingly different overall conformation of the protein, compared to those of the ligand-free and the effector-bound repressor. As highlighted by the crystal structures described above, the transitional switch between the relaxed (DNA-bound) and tense (malonyl-CoA-bound) forms of the repressor involves a large-scale structural rearrangement (Fig. 5). The amphipathic helix $\alpha_L$, whose hydrophobic face binds to the protein core in the tense state (Fig. 5A), structurally dissociates and moves $\sim 30$ Å to interact with the same helix from the other protomer and with DNA in the relaxed state (Fig. 5B). This DNA-driven process requires partial unwinding of the linker helix $\alpha_L$ and the solvent-exposure of hydrophobic side-chains from the EBD (Leu82, Ile83, Val85, Ile94, Val123) and from the loop immediately following helix $\alpha_L$ (Ile70, Phe78) in both protomers. Such a substantial conformational rearrangement between the tense and relaxed states (Fig. 5C) contrasts with the more subtle structural changes observed for other well-studied bacterial classes of allosteric transcriptional regulators such as the tetracycline [29] or lactose [30] repressors, illustrating that specific physiological responses can be achieved by a variety of mechanisms.

Disruption of membrane lipid homeostasis is lethal for S. aureus
Bacterial FASII has been identified as a promising target for antibacterial drug discovery. Nevertheless, Brinster et al [31] have questioned the feasibility of this approach in Gram-positive pathogens based on the finding that FASII is not essential in Streptococcus agalactiae if the bacterium is supplemented with fatty acids or human serum. This controversy was recently clarified by Parsons et al [32], who showed that externally added fatty acids downregulate the activity of the acetyl-CoA carboxylase in some Gram-positive pathogens, such as Streptococcus pneumoniae. This biochemical mechanism suppresses the malonyl-CoA levels allowing fatty acid supplements to replace endogenous fatty acids completely, thus rescuing bacterial FASII inhibition. In S. aureus, this feed-back regulatory mechanism is not present [32] and thus external fatty acids do not circumvent the treatment of this bacterium with FASII inhibitors. These results and the regulatory properties of the $acc$ genes led the authors to propose that pathogens containing FapR would likely be sensitive to FASII inhibitors, regardless of the addition of external fatty acids [32]. These considerations prompted us to test if structure-based mutations predicted to disrupt the SaFapR-malonyl-CoA interaction are lethal for S. aureus even in the presence of extracellular fatty acids. To this end, based on the structural information and on our previous work [20], we substituted Arg110 by alanine to disrupt the key interaction with the malonyl carboxylate, and introduced a double substitution (Gly111Val, Leu132Trp) to
block the ligand-binding tunnel that accommodates the phosphopantetheine moiety in the repressor-effector complex (Fig. 3C). These mutants were expected to retain their DNA-binding capacity and to permanently repress the expression of the \( fap \) regulon, independently of the metabolic conditions. To test this prediction we engineered \( S. aureus \) \( fapR \) null mutants to produce the protein variants in response to an inducer (IPTG). IPTG-induced expression of \( \text{Sa} FapRR_{110A} \) caused a small drop in cell viability (data not shown), while cells expressing \( \text{Sa} FapRG_{111V,L132W} \) failed to grow (Fig. 6). A variety of fatty acids, including oleic acid, a common mammalian fatty acid, and anteiso saturated fatty acids, the most abundant acyl chains found in \( S. aureus \) phospholipids were unable to overcome the growth inhibition caused by the expression of the superrepressor \( \text{Sa} FapRG_{111V,L132W} \) in \( S. aureus \) (Fig. 6). These results clearly show that disruption of the membrane lipid homeostasis mimics the effects of FASII inhibitors in \( S. aureus \) strongly supporting the notion that exogenous fatty acids cannot replace the endogenously produced acyl chain in this bacterium [32].

**Discussion**

The structures of \( \text{Sa} \)FapR, in three relevant states of its regulation cycle, uncover a complex biological switch, characterized by completely different protein-protein interactions involving in all cases the linker helix \( \alpha_L \) and so leading to a distinct quaternary arrangement for the tense and relaxed states of the repressor (Fig. 5). Similarly to other homodimeric proteins studied by single-molecule or NMR relaxation approaches [33,34], our crystallographic studies suggest that the two conformational states of \( \text{Sa} \)FapR (i.e. with EBD-bound or EBD-detached DBDs) can be populated in the ligand-free repressor species. Thus, a higher cellular concentration of malonyl-CoA would not only trigger the conformational changes that disrupt the \( \text{Sa} \)FapR-operator complex [20], but would also promote the dynamic shift of the ligand-free repressor population towards the tense state.

Our results highlight the ability of FapR to monitor the levels of malonyl-CoA and appropriately tune gene expression to control lipid metabolism, ensuring that the phospholipid biosynthetic
pathway will be supplied with appropriate levels of fatty acids either synthesized endogenously or incorporated from the environment. Bacterial FASII is a target actively pursued by several research groups to control bacterial pathogens [35]. A controversy surrounding FASII as a suitable antibiotic target for *S. aureus* was based on the ability of this pathogen to incorporate extracellular fatty acids [36]. This apparent discrepancy was recently clarified by showing that although exogenous fatty acids indeed are incorporated by *S. aureus*, following its conversion to acyl-ACP, they cannot deplete the generation of malonyl-CoA and malonyl-ACP [32]. Thus, these fatty acid intermediates release FapR from its binding sites [28] and are used by FASII to initiate new acyl chains. Thus, when a FASII inhibitor is deployed against *S. aureus*, the initiation of new acyl chains continues leading to depletion of ACP, which is correlated with diminished exogenous fatty acids incorporation into phospholipids [32]. Strikingly, SaFapR, not only control the expression of the FASII pathway, but also regulate the expression of key enzymes required for phospholipid synthesis such as PlsX and PlsC [18,37]. Most of the Gram-positive pathogens rely on the PlsX/PlsY system to initiate phospholipid synthesis by converting acyl-ACP to acyl-P0₄ by PlsX followed by the transfer of the fatty acid to the 1 position of glycerol-P0₄ by the PlsY acyltransferase [37,38]. A strength of targeting these steps in lipid synthesis is that acyltransferases inhibition cannot be circumvented by supplementation with extracellular fatty acids. Thus, targeting of lipid synthesis with compounds that block the expression of PlsX in *S. aureus* cannot be ignored, specially taking into account that it has been reported that extracellular fatty acids increase the MIC for FASII inhibitors in this important pathogen. [32,39]. In this sense, the unique mode of action of FapR and our encouraging *in vivo* results validate lipid homeostasis as a promising target for new antibacterial drug discovery in Gram-positive bacteria.

Conserved in many Gram-positive bacteria, FapR is a paradigm of a feed-forward-controlled lipid transcriptional regulator. In other characterized bacterial lipid regulators it is the long acyl-chain end products of the FASII pathway that act as feedback ligands of lipid transcriptional regulators [40].

Figure 5. The transitional switch between the relaxed and tense states of the repressor involves a significant rearrangement of the DBDs. (A) SaFapR in complex with malonyl-CoA (shown in stick representation), tense state. (B) SaFapR in complex with DNA, relaxed state in which the amphipathic helix αL from each protomer associates with each other. (C) Superposition of the two conformational states of the repressor illustrating the structural transition. Solvent accessible surfaces are shown in transparent to highlight the DNA-induced dissociation of the invariant effector-binding domain from the DBDs. The molecules are shown in light (relaxed) and dark (tense) grey, except for the helices from one DBD (colored).

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Figure 6. Expression of the *SaFapR<sub>G11V, L132W</sub>* superrepressor is lethal for *S. aureus* and fatty acid supplementation cannot overcome growth inhibition. The figure shows the growth of strain RN4220ΔfapR expressing *SaFapR<sub>G11V, L132W</sub>* (REH117) or *SaFapR<sub>WT</sub>* (REH118) under the tight inducible P<sub>spacOid</sub> promoter on THA plates in the absence or presence of 10 mM IPTG. Identical results were obtained for strain REH117 growing in 10 mM IPTG when supplemented with either Tween80 (0.1%) or 500 μM of the following fatty acids: palmitic acid (16:0), oleic acid (18:1), 16:0+18:1, anteiso 17:0 (a17:0), a15:0+a17:0, or 18:0+a17:0.
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binding domains of these proteins, such as E. coli FadR [12] or the TetR-like P. aeruginosa DesT [17], frequently display an α-helical structure with a loose specificity for long-chain acyl-CoA molecules, possibly because the permissive nature of helix-helix interactions provide a suitable platform to evolve a binding site for fatty acid molecules of varying lengths. In contrast, a high effector-binding specificity is required for the feed-forward regulation mechanism of the FapR repressor family [19], which entails the recognition of an upstream biosynthetic intermediate, namely the product of the first committed step in fatty acid biosynthesis. This high specificity is achieved in SfapR by caging the charged malonyl group inside a relatively stiff internal binding pocket, and may be the reason why the hot-dog fold was recruited for this function. Nevertheless, it could be expected that, in organisms using the FapR pathway, a complementary feed-back regulatory loop should also operate at a biochemical level, for instance by controlling the synthesis of malonyl-CoA [37]. This would imply that lipid homeostasis in FapR-containing bacteria would be exquisitely regulated by feed-back and feed-forward mechanisms, as it is indeed the case in higher organisms ranging from Caenorhabditis elegans to humans [41].

Materials and Methods

In vivo studies

In frame fabR deletion mutants of S. aureus strains RN4220 and HG001 [42], RN4220 Δ fabR and HG001 Δ fabR respectively, were constructed and the expression level of the plicX and fabH genes, belonging to the fab regulon, was analyzed by real time PCR as described in Text S1. To evaluate the activity of the S. aureus repressor in B. subtilis, we used the ΔfapR mutant strain GS416 that contains a P6factsbaclacZ reporter fusion and expressed S. aureus fapR from the xylose inducible PysL promoter (for further details see Text S1).

Mutants SfapR R110A and SfapR Q111V.L132W, impaired in malonyl-CoA binding were obtained by site-directed mutagenesis using S. aureus RN4220 genomic DNA as template, mutagenic oligonucleotides and overlap-extension PCR. The replicative vector pOS1 [43] and the tight IPTG-regulated PspacOid promoter from pMUTIN4 were used for ectopic expression of fabR alleles in RN4220 Δ fabR (see Text S1 for details). S. aureus transformants were selected in THA plates supplemented with chloramphenicol (5 μg/ml) and expression of fabR alleles was achieved by addition of 10 mM IPTG. To evaluate the effect of an external source of fatty acids THA plates were supplemented with 500 μM of the stated fatty acids or 0.1% Tween80 (Fig. 6).

Protein expression and purification

The S. aureus fabR gene was cloned into the pET15b vector (Novagen) as described in Text S1 and expressed in E. coli BL21/pLysS. Bacterial cultures were grown in LB supplemented with ampicillin 100 μg/ml and chloramphenicol 10 μg/ml at 37 °C until OD600 0.6. Expression was induced with 0.5 mM IPTG at 20 °C for 17 hours. Cells were harvested by centrifugation at 4 °C, and protein-containing fractions from a Ni2+-affinity chromatography on a HiTrap column (GE Healthcare) were dialyzed at room temperature against 20 mM Tris-HCl pH 7.6 and 50 mM NaCl, concentrated to 15 mg/ml and stored in aliquots at −80°C for further use. Selenomethionine (SeMet)-labeled SfapR was obtained using the E. coli strain B834 (Novagen) grown in M9 medium containing 0.2 g/l selenomethionine and purified as above.

Crystallographic studies

Crystals of the repressor and its complexes with the effector and operator molecules were obtained as described in Text S1. All diffraction datasets were collected from single crystals at 100 K using synchrotron radiation at beamlines ID14.4 and ID29 (European Synchrotron Radiation Facility, Grenoble, France) or Proxima 1 (SOLEIL, Saint-Aubin, France). Data were processed with either XDS [44] or iMosm [45] and scaled with XSCALE from the XDS package or SCALA from the CCP4 suite [46]. The crystal structures of SfapR alone and its complex with malonyl-CoA were solved by molecular replacement methods using the program AMoRe [47] and the effector-binding domain of B. subtilis FabR (PDB entry 2F41) as the search model. In all cases, the missing DNA-binding domains were manually traced from sigma A-weighted Fourier difference maps. The structure of the repressor-operator complex was solved by a combination of molecular replacement and single-wavelength anomalous diffraction (SAD) techniques using SeMet-labeled SfapR. The selenium substructure was determined with the program SHELXD [48] and further refined with the program SHARP [49]. Structures were refined with REFMAC5 [50] or BUSTER [51] (for the SeMet-labeled protein-DNA complex) using a TLS model and non-crystallographic symmetry restraints when present, alternated with manual rebuilding with COOT [52]. Models were validated through the MolProbity server [53]. Data collection and refinement statistics are reported in Table 1. Graphic figures were generated and rendered with programs Pymol [54] and 3DNA [55].

Isothermal titration calorimetry

ITC experiments were performed using the high precision VP-ITC system (MicroCal Inc., MA) and quantified with the Origin7 software provided by the manufacturer. All molecules were dissolved in 50 mM Tris-HCl, pH 8, 150 mM NaCl and the binding enthalpies were measured by injecting the SfapR solution into the calorimetric cell containing the 40 bp DNA solution. Heat signals were corrected for the heats of dilution and normalized to the amount of compound injected. Complementary DNA strands were heated to 90°C and annealed by a stepwise decrease to 25°C followed by 30 min on ice prior to use; DNA concentration was determined by absorption at 260 nm.

Accession codes

Crystallographic coordinates and structure factors were deposited in the Protein Data Bank, with accession codes 4a0x, 4a0y, 4a0z and 4a12.

Supporting Information

Figure S1 Fatty acid synthesis and phospholipid initiation steps in Bacillus subtilis. The FabH condensing enzymes initiates the cycles of fatty acid elongation by condensation of acyl-CoA primers with malonyl-ACP (3a). The resultant β-ketoester is reduced by the β-ketocarboxy-ACP reductase (4). Then, the β-hydroxyacyl-ACP is dehydrated to the trans-2 unsaturated acyl-ACP by β-hydroxyacyl-ACP dehydrase (5), which is finally reduced by enoyl reductase (6). Subsequent rounds of elongation
are initiated by the elongation-condensing enzyme FabF (3b) to generate an acyl-ACP two carbons longer than the original acyl-ACP at the end of each cycle. The long chain acyl-ACP end products of fatty acid synthesis are transacylated in three steps to glycerolphosphate, to generate phosphatidic acid (PA), a key intermediate in the synthesis of phospholipids. First, Phx catalyzes the synthesis of fatty acylphosphate from acyl-ACP (7); then, PhX transfers the fatty acid from the activated acyl intermediate to the 1-position of glycerol-3P (8) and finally, hyso-PA is acylated to PA by PlsC (9). Malonyl-CoA is generated from acetyl-CoA by acetyl-CoA carboxylase (1) and then is transferred to ACP by malonyl-CoA transacylase (2). Expression of the genes surrounded by shaded ellipses is repressed by FapR, whose activity is, in turn, antagonized by malonyl-CoA (enclosed in a rectangular box). R denotes the terminal group of branched-chain or straight-chain fatty acids.

(TIF)

Figure S2 Effect of malonyl-CoA and short-chain acyl-CoA thioesters on binding of SfFapR to PfapR. (A) Electrophoretic shift assay showing the binding of SfFapR to PfapR. Increasing amounts of SfFapR were incubated with 30 nM of double strand PfapR probe (left panel) or non-specific DNA (right panel). (B) Effect of malonyl-CoA on SfFapR binding to PfapR. The gel mobility shift assay was performed with two concentrations of SfFapR (100 nM or 500 nM), previously incubated at different concentrations of malonyl-CoA for 15 min at room temperature followed by addition of the probe to the SfFapR-malonyl-CoA complex. (C) Effect of malonyl-CoA analogs on SfFapR binding to PfapR. SfFapR (200 nM) was previously incubated, as described in b, with malonyl-CoA (lanes 3 and 4), acetyl-CoA (lanes 5 and 6), propionyl-CoA (lanes 7 and 8) or succinyl-CoA (lanes 9 and 10) before adding the PfapR probe to the reaction mixture. Gels were stained with SYBR GREEN and scanned at 530 nm.

(TIF)

Figure S3 Conservation of FapR function in S. aureus. (A) RT-PCR results showing the expression of two genes from the fap regulon (pfx, fabH) in two different strains (RN4220, HG001). Relative levels of transcripts were measured by qRT-PCR. Expression levels were normalized using 16S RNA as an internal standard and are indicated as an n-fold change, expressed as the means and standard deviations of quadruplicate experiments. (B) SfFapR complements a B. subtilis fabR null strain. Cultures of strains GS415 (squares; ΔfapR, ΔfabHB-lac-Δfap/R, amyE::spc), GS373 (circles; ΔfapR, ΔfabHB-lac-Δfap/R, amyE::Pysl-BsfapR::spc) and GS416 (triangles; ΔfapR, ΔfabHB-lac-Δfap/R, amyE::Pysl-SfapR::spc) were incubated in LB medium in the presence of xylene 0.1% (w/v) at 37°C and growth was monitored. Samples were removed to assay β-galactosidase-specific activity at the indicated times. A single experiment representative of three repeats is shown. Solid lines, β-galactosidase-specific activity; dotted lines, OD525.

(TIF)

Figure S4 Alignment of the promoter regions of the genes and operons regulated by FapR. The -10 motif of RNA polymerase binding boxes are indicated with rectangles. The conserved inverted repeat sequence is shaded and a consensus sequence is indicated at the bottom line. Bases are numbered related to the first base of the translation initiation codon. (TIF)

Figure S5 Different roles of the amphipathic linker helix ζL in stabilizing the SfFapR-DNA and SfFapR-malonyl-CoA complexes. (A) Close view of the SfFapR-DNA complex. (B) Close view of the interactions between the EBD, the linker helix (green) and the DBD in the structure of the SfFapR-malonyl-CoA complex. In both cases, hydrophobic and hydrophilic side chains involved in inter-domain interactions are labeled. Malonyl-CoA binding triggers significant conformational changes in three main regions of the repressor: the connecting loop ζL - β1, the amphipathic helix ζH and the DBD. The side-chain of Arg110 from the EBD makes a key hydrogen bond with the main-chain carbonyl group of Glu77’ in the loop. In addition, the Glu77’ side-chain is hydrogen bonded to the main chain of His104 from the EBD, facilitating the formation of a strong salt bridge between the side-chains of Lys105 and Glu74’. The Ser102 side-chain makes a strong hydrogen bond with Val123, whereas its main chain interacts through hydrogen bonding networks with Ser72, Pro129 and a water molecule. Furthermore, the main-chain of Glu69 makes an electrostatic interaction with the side-chain of Lys127. These newly formed electrostatic/hydrogen-bond interactions allow the connecting loop ζL - β1 to firmly interact with the core of the EBD. As a consequence, the amphipathic helix ζH forms now an extended hydrophobic interface (Leu55, Ile59 and Val62), with both β1 (Ile83) and β2 (Ile94) strands of the EBD, and the α1-α2 connecting loop of the DDB (Pro25, Phe26 and Thr28). This structural arrangement is further stabilized by key electrostatic interactions. Arg56 makes a salt bridge with Asp96 located at the top of β2 and the side-chain of Asn66 makes a hydrogen bond with the carbonyl group of the main-chain of Leu82 on the β1 strand. Moreover, the side-chain of Tyr67 is oriented towards a hydrophobic pocket including residues Leu82, Val83 from β1, and Ile70, Ile73 and Phe78 from the connecting loop ζL - β1.

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Figure S6 Sequence alignment of FapR. Amino acid sequence and secondary structure elements (green for DBD, pink for EBD) of SfFapR. Residues that interact with the pantetheine group of malonyl-CoA and with DNA are underlined by open and closed circles, respectively. Wavy lines indicate disordered regions in the crystal structure. Colored amino acid positions indicate highly conserved residues in non-redundant FapR sequences from 67 different bacterial species (yellow, >80%; orange, >95%).

(TIF)

Figure S7 Structural superposition of the BfFapRA43-malonyl-CoA and SfFapR-malonyl-CoA complexes. Although the orientation of helix ζL is different (it is involved in crystal packing contacts in BfFapRA43), binding of malonyl-CoA promotes the same conformation of the loop connecting this helix with the first β-strand of the EBD.

(TIF)

Table S1 The fap regulon in B. subtilis and S. aureus.

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Table S2 Local base-pair step parameters of the DNA in complex with SfFapR.

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Table S3 Oligonucleotides used in this work.

(DOC)

Text S1 Supplemental information, including additional experimental procedures.

(DOC)

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48. Author Contributions

Conceived and designed the experiments: DA FS AB DdM PMA. Performed the experiments: DA GR MEG FS DdM PMA. Analyzed the data: DA GR MEG FS MD AB DdM PMA. Contributed reagents/materials/analysis tools: MD GE DdM PMA. Wrote the paper: DA ME DdM PMA.

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