Engineering strategies for rational polyketide synthase design

Maja Klaus and Martin Grininger

Covering: mid 1990s to 2018

Over the last two decades, diverse approaches have been explored to generate new polyketides by engineering polyketide synthases (PKSs). Although it has been proven possible to produce new compounds by designed PKSs, engineering strategies failed to make polyketides available via widely applicable rules and protocols. Still, organic synthetic routes have to be employed whenever new polyketides are needed for applications in medicine, agriculture, and industry. In light of the rising demand for commodity products from feedstock and for fast and cheap access to pharmaceutical compounds, the need for harnessing PKSs to produce such molecules is more urgent than ever before. In this review, we focus on a multitude of approaches to engineer modular PKSs by swapping and replacing PKS modules and domains, which we analyze in the light of recent structural and biochemical data. We conclude with an outlook on possible strategies on how to increase success rates of PKS engineering in future.

1. Introduction

Polyketide synthases (PKSs) are responsible for the synthesis of polyketide natural products; among them many compounds with high bioactivity, such as environmental toxins (e.g. aflatoxin), antibiotics (e.g. erythromycin and tetracycline), anti-neoplastics (e.g. daunorubicin) and immunosuppressants (e.g. rapamycin). PKSs use simple acyl-CoA building blocks and assemble them into complex compounds with molecular weights of up to several kilodaltons (kDa).

PKSs can be classified into three different types. Type I PKSs are of highest synthetic capability, and are comprised of large multidomain proteins. Further, type I systems can be subdivided into the iterative (iPKSs) and modular PKSs (modPKSs), whereby the modPKSs occur as cis-AT and trans-AT PKSs. In the following, we focus only on type I systems and use the term PKSs for referring to type I cis-AT PKSs. iPKSs perform synthesis in a recursive manner, during which the catalytic domains of a single polypeptide repeatedly condense acyl-CoA precursor units until the specific length of the compound is achieved. In contrast, modPKSs occur as large assembly line-like complexes and successively condense precursor building blocks to the final natural compound. The 6-deoxyerythronolide B synthase (DEBS) is the prototypical example for a large pool of naturally occurring modPKSs, in which the arrangement of modules essentially defines the sequence of functional groups in the final compound (Fig. 1). Each module comprises a set of essential domains for C–C bond formation, namely the catalytic

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domains ketoacyl synthase (KS) and acyl transferase (AT), as well as the acyl carrier protein (ACP) domain. Within a module, the collaborative function of the AT, ACP, and KS domains results in a non-reduced polyketide chain. Optionally, further processing domains (ketoreductase (KR), dehydratase (DH), and enoylreductase (ER)) are used to catalyze the step-wise reduction of the β-keto group. The flexible ACP domain shuttles the polyketide intermediate both within a module and across modules. Physical linkage between individual modules within a modPKS is either achieved through covalent linkers or non-covalent linker domains (docking domains) at their C- and N-termini.5,6

The high pharmaceutical importance of polyketides has initiated intensive research on their biosynthetic generation. Whereas conventional polyketide synthesis through organic chemical routes can be highly intricate, polyketide biosynthesis is remarkably simpler, since it is built on the controlled step-wise condensation of small acyl-CoA substrates and their subsequent chemical modification. The strict correlation of the assembly line architecture of modPKSs and the final polyketide

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**Fig. 1** Architecture of the 6-deoxyerythronolide B synthase (DEBS). DEBS consists of three polypeptides (DEBS1, DEBS2, and DEBS3) each harboring two modules, which are connected by covalent linkers (colored lines). Modules across polypeptides interact through docking domains at the C- and N-termini (black taps). Additionally, a loading didomain (LDD) for starter unit selection and a thioesterase domain (TE) for product release are present. DEBS uses propionyl-, methylmalonyl-CoA and NADPH to produce 6-deoxyerythronolide B (6-dEB) the precursor for the antibiotic erythromycin. The polyketide intermediates of each module are depicted. Domain annotation: KS – ketosynthase, AT – acyltransferase, KR – ketoreductase, DH – dehydratase, ER – enoylreductase, and ACP – acyl carrier protein. Figure adapted from (ref. 58).

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Maja Klaus graduated 2015 from the Leibniz University Hannover with a M. Sc. in Life Science. During her Master’s thesis with Prof. Chaitan Khosla at Stanford University she started to work on the engineering of protein–protein interactions within polyketide synthases. In summer 2015, she joined Prof. Martin Grininger’s lab at the Goethe University Frankfurt to pursue a PhD in chemistry, where she continues her research on harnessing the modularity of polyketide assembly lines in collaboration with Prof. Khosla.

Martin Grininger studied chemistry at the Universities Linz and Graz (Austria), and conducted his diploma thesis in natural compound synthesis in the lab of Prof. Alois Fürstner. He then worked with Prof. Dieter Oesterhelt at the Max-Planck-Institute of Biochemistry for his PhD, studying flavin sequestration and storage in Archaea and Bacteria. After becoming a project group leader at the same institution, he focused on multidomain fatty acid synthases and contributed to the current structural understanding of these proteins. He also held a visiting professorship at the University Vienna in Biological Chemistry. In 2012, he received a Lichtenberg Professorship of the Volkswagen Foundation hosted by the University Frankfurt. He is co-speaker of the research focus MegaSyn, which develops rules for engineering megasynthases for custom compound synthesis.
structure, known as the principle of co-linearity, stimulated efforts to engineer modPKSs by swapping entire modules or single domains to generate new polyketides.\textsuperscript{7-9} Recently, a broadly applicable mix-and-match strategy for engineering the related family of non-ribosomal peptide synthetases (NRPSs) has been presented,\textsuperscript{18} which uses defined exchange units as interchangeable synthetic modules. Concepts of comparable broad applicability for harnessing PKS modularity have not yet been established. The extensive scaffold of modPKSs comprises a wealth of permanent domain–domain and module–module interactions and samples a multitude of transient domain–domain interactions during ACP-mediated substrate shuttling. This complexity needs to be further untangled in structural and functional studies before programmable engineering becomes possible.

Besides swapping domains and modules, another PKS engineering strategy is the modification of active sites to induce different substrate specificity or higher substrate promiscuity. Such an approach preserves the structural and conformational properties of the scaffold, and only adapts active sites to the new requirements. PKS engineering by site-directed mutagenesis is inherently modPKS comprises a wealth of permanent domain–domain and module–module interactions and samples a multitude of transient domain–domain interactions during ACP-mediated substrate shuttling. This complexity needs to be further untangled in structural and functional studies before programmable engineering becomes possible.

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2. The PKS domain–domain and module–module interaction network

Engineering of PKSs requires detailed knowledge about structural arrangements and the interactions of domains and modules. While our understanding of the architecture of PKS modules and their arrangement into assembly lines is still limited, we have insight into structures of subregions or evolutionary related proteins. Recent progress in the characterization of the transient interactions during the catalytic cycle of PKS and an improved understanding of the vential synthetic progress in modPKS further helps to guide precise engineering approaches.

2.1. The PKS structural scaffold

The earliest X-ray structural information on the modPKS, the KS–AT condensing unit and the KR domain, was available from studies of the modPKS DEBS.\textsuperscript{18-20} When the X-ray crystal structure on porcine FAS was revealed in 2008, it not only gave new insights into mechanisms of fatty acid synthesis, but also spurred significant interest in the field of modPKS (Fig. 2A, mFAS).\textsuperscript{28} The mammalian fatty acid synthase (mFAS) is evolutionary related to PKSs, and many subregions from mFAS and modPKSs displayed structural consensus, including the non-catalytic linkers in the KS–AT didomain (the KS–AT linker and post-AT linker) and the non-catalytic KSKR domain.\textsuperscript{18-17} This comparison led to the appealing model that mFAS is simply a fully reducing PKS, which can accept deletions of modifying domains within the processing wing to build the partially reducing and non-reducing PKS folds. Notwithstanding the still valid tight structural relationship of folds, recent studies revised this view and show decisive structural differences between the processing parts of PKSs and mFAS. Particularly, the first almost complete X-ray structural model of the iterative Mycobacterium smegmatis mycocerosic acid synthase (MAS)-like PKS gave important insight,\textsuperscript{23} but also earlier models on separate proteins and the KR–ER didomain provided major contributions to the structural understanding of PKSs.\textsuperscript{20,22-25} The DH domain is most important in the different organizations of the processing part of mFAS and PKSs. In contrast to a V-shaped arrangement in mFAS, DHs of PKSs form an overall elongated dimeric arrangement inducing a different relative positioning of the modifying domains.\textsuperscript{21,22} The embedment of the ER domain in the PKSs fold is currently disputed. The ER is dimeric in the MAS-like (iterative) PKS, resting upon the extended β-sheet of the DH dimer by forming a small and variable interface (Fig. 2A, MAS-like model). The relevance of a dimeric ER arrangement in modPKSs is unclear,\textsuperscript{26} since the ER domains from modPKSs, structurally characterized by X-ray crystallography as individual proteins and as a part of KR–ER didomain construct, appeared to be monomeric.\textsuperscript{51,23} The ER domain may occur in different oligomeric states in PKSs, \textit{i.e.} as a monomeric ER in modPKSs and dimeric ER in iPKSs, consistent with ER–KR linkers of modPKSs being generally shorter (usually <8 amino acids) than those of iPKSs (>17). A short linker may restrain ER conformational variability in modPKSs and prevent ER dimerization at the twofold axis.\textsuperscript{26} SAXS data have been reported, which support both dimeric and monomeric ER in modPKSs.\textsuperscript{21,23} Accordingly, two models for the structural appearance of fully reducing modPKSs exist today: a MAS-like PKS model with the ER domains dimerizing at the C2-axis (Fig. 2A), and a SPNS M2 derived model, in which monomeric ERs are swung out and thereby free space at the protein’s central region (Fig. 2B).

In order to reach consensus on the appearance of individual PKS modules, structural characterization of a variety of intact modules is required. Until today, detailed structural information on a complete modPKS module is missing, with the exception of medium resolution (7.3–9.5 Å resolution) cryo-electron microscopic (cryo-EM) data on the partially reducing module 5 of picromycin synthase (PIKS M5). The PIKS M5 structure revealed an arched conformation,\textsuperscript{27} with a relative arrangement of the KS and AT domains that contradicts X-ray crystallographic studies on the mammalian FAS\textsuperscript{28-30} and PKS KS–AT structures.\textsuperscript{18,19,21} In all these structures, the AT domains were consistently found to be embedded as bulge of the KS fold framed by a N-terminal KS–AT linker that constitutes most of
the KS–AT interface, and a post-AT linker that wraps back to interact with the KS domain.\(^\text{18}\) In the PIKS M5 model, this structural integrity is dissolved inducing the overall arch-shaped appearance. The PIKS M5 model provides a structural basis that can explain modPKS function; however, before the PIKS M5 model can be accepted as a relevant model for the functional mode of modPKSs, several of its structural features need yet to be reconciled with a large amount of data collected over the last decades.\(^\text{31,32}\) A low-resolution model of partially reducing PKS modules was additionally derived from SAXS analysis of DEBS.\(^\text{33}\) Herein, a similar architecture of the KS–AT fold was observed as found in mFAS and PKSs. Thus, two different models for partially reducing modPKS modules can be derived today based on the analysis of PIKS M5 and DEBS modules (Fig. 2A).

While several models can be generated for the overall scaffold of a single PKS module, data on the architecture of a whole assembly line have not been presented to date. In general, modules can be connected by covalent ACP–KS linkers or through non-covalently interacting, \(\alpha\)-helical docking domains (Fig. 2B). While several excised docking domains were structurally solved,\(^\text{5,34–36}\) no high resolution structure of a bimodule was obtained so far.

### 2.2. Interactions and processes within modular PKSs

ACP domains are responsible for substrate shuttling in PKSs. They are generally loosely attached via unstructured linkers to the catalytic body of (type I) PKSs and interact transiently with the catalytic domains. Owing to the high conformational variability, ACP domains often remain unresolved in structural studies leading to paucity of details to the process of ACP-mediated substrate shuttling. The primary insight into the mode of substrate shuttling in PKSs was again provided by the homologous mFASs. For example, it was shown that swiveling and swinging motions of the condensing and processing part of mFAS occur and presumably assist the mobile ACP domain in substrate shuttling.\(^\text{27–39}\)

Despite being loosely attached, the ACP domains are differently constrained in their conformational space in iterative and modular proteins. While in mFAS and iPKSs the ACP is free to move, solely restricted by terminating domains such as TE, the linkage of ACP to the KS of the downstream module constrains its conformational variability in modPKSs. In covalently connected modules the average length of an ACP–KS linker is 18 residues in length (in ER containing modules).\(^\text{26}\) For anticipating a mode of action of ACPs in modPKS assembly lines, the ER arrangement is therefore of decisive impact (see above for the discussion about modPKS scaffolds). A monomeric ER would allow a central positioning of ACP and grant intuitive, easy access to each of the integral catalytic domains, whereas an ER dimer would crowd the C2-axis and force the ACPs toward the peripheral lining of the assembly line (Fig. 2B).

The pivotal steps for catalytic progress along the PKS assembly line are the ACP–KS mediated chain translocation and chain elongation reactions. For a successful chain translocation reaction, two adjacent modules need to interact to bring the upstream ACP and the downstream KS into close proximity. While part of this interaction is mediated through covalent linkers or docking domains, specificity is conferred through the

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**Fig. 2** Different models for assembly line PKSs in comparison to mFAS. (A) Schematic representation of mFAS (based on PDB 2VZB),\(^\text{29}\) a MAS-like PKS (based on PDB 5BP1 and 5BP4),\(^\text{30}\) a model for the partially reducing DEBS M3 based on SAXS analysis,\(^\text{23}\) and a model of the partially reducing pikromycin synthase module 5 (PIKS M5) based on cryo-EM analysis.\(^\text{27}\) (B) Graphical representation of a PKS module assembly using the example of the proposed model for the spinomycin synthase module 2 (SPNS M2).\(^\text{23}\) The SPNS M2 model was assembled from PDBs 5BP1 (KS–AT), 3SLK (ER–KR), and 3BP4 (DH) with KR of 3SLK superimposed on KR of 3BP4. PKS modules can either be connected through covalent ACP–KS linkers or through non-covalently interacting docking domains at the C- and N-termini. Additionally, chain translocation is mediated through specific interactions between the upstream ACP (ACP\(_{n−1}\)) and the downstream KS (KS\(_n\)), shown for one of the two ACPs in module\(_{n−1}\). Domain coloring: KS (blue), AT (light green), KR (dark green), DH (orange), ER (yellow), ACP (magenta), linker regions such as the KS–AT linker (dark gray), and only for mFAS non-catalytic pseudo-methyltransferase domain (light gray).
interaction of the ACP and the KS–AT fold itself. Using DEBS as a model system, the ACP–KS interactions were analyzed in a comprehensive study combining in vitro analysis of chimeric ACPs and in silico docking simulations. Generation of different chimeras revealed that during chain elongation, ACP_A docks with loop 1 to the KS_A–AT_A linker, while engaging with KS_B (of protomer B) for chain elongation.0 Two distinct recognition sites on the KS–AT linker were found to interact with a minimal epitope in ACP loop 1. In a similar manner, the upstream ACP docks with the first ten N-terminal residues of ACP helix 1 to KS_A–AT_A linker to mediate chain translocation with KS_B.40,41 Although the ACPs dock into the same deep cleft of the KS–AT fragment, the position and orientation is distinct in the event of chain translocation and chain elongation.

2.3. Models for the origin of vectorial synthesis

Vectorial synthesis is the term given to the step-by-step synthetic progress along a directional scaffold; i.e. for modPKSs, the biosynthesis of polyketide compounds in a linear manner. While vectorial synthesis has emerged as a rather robust feature in modPKSs, its molecular foundation is unclear. Recently, a turnstile mechanism has been postulated, in which a not yet discovered molecular signal prevents the KS domain from iterative use.44 The reopening of the turnstile for reloading necessitates a vacant module as prerequisite. Such a state does not appear during synthesis in iPKSs (and FASSs), which implies that vectorial synthesis is a unique feature of modPKSs. An alternative explanation for the origin of vectorial synthesis is the kinetic coupling of catalytic steps along the assembly line, due to the serial arrangement of modules. As iPKSs and modPKSs are tightly evolutionary connected,44 kinetic coupling of catalytic processes would facilitate our understanding of vectorial biosynthesis as a result of the PKS architecture rather than an inherent feature of modPKSs. For example, the recently characterized iterative MAS-like PKS, producing an eicosenoic acid derivative, is assigned phylogenetically to modPKSs.21,45 Moreover, modPKSs have been demonstrated to also synthesize in iterative fashion as part of assembly lines (borrelidin synthase (BORS) module 5 (ref. 46) and aureothin producing PKS (AURS)45), and PKS modules have been turned into iteratively synthesizing units by mutation (PIKS M5 (ref. 48) and DEBS M3 (ref. 41)).

3. Engineering module–module and domain–domain interactions – early and recent efforts

Engineering of module–module and domain–domain interactions in modPKSs can be classified by different criteria. The often referred to mix-and-match approach considers the swapping of entire PKS modules from one synthase into another, while domain–domain swapping is focusing on single domains e.g. AT domains or KR domains to alter the product spectra of a given PKS. Within those categories, the studies differ in whether they were carried out in vivo, and thus relying solely on differences in product spectra, or in vitro, shedding light on enzyme kinetics and enzyme affinities. In this section, we highlight engineering approaches of modPKSs and evaluate them based on the available structural data.

3.1. Module–module exchanges

The discovery of modPKSs inspired the chemical biological community to harness their modularity for the rapid generation of new natural products.49,50 One of the biggest challenges in combining intact PKS modules to chimeric PKS assembly lines is to preserve their catalytic integrity. Early on, the necessity of preserving linker regions between adjacent modules to retain functional proteins was noticed.51–54 Although preservation of native interaction sites helped in generating productive assembly lines, the overall production rates were usually lowered in the engineered systems.

The first truly combinatorial approach of generating bimodular chimeric PKSs, consisting of intact modules from different PKS sources, was carried out in an in vitro study recombining 14 modules of 8 PKS clusters.4 Using docking domains derived from DEBS, a total of 154 bimodular chimeric PKSs were assembled. About 50% of the chimeric PKSs yielded detectable product albeit with lower product yields as their natural bimodular reference system.4 Later on, this approach was extended towards chimeric trimodular PKSs with similar results.55

Until now, chimeric PKSs have been constructed by either fusing proteins or by non-covalently connecting individual modules with docking domains derived from natural PKS sources. In vitro characterization of chimeric bimodular PKSs using fusion proteins, in which the non-naturally interacting modules were connected by a covalent linker, revealed similar $k_{cat}$ values for wild type and chimeric PKSs.56 A more thorough analysis showed that linker regions and specific ACP–KS interactions play equal roles for the turnover rate.57 Recently, the finding that linker regions alone are able to connect heterologous modules was challenged in a study creating a library of bimodular and trimodular chimeric PKSs in vitro by employing docking domains to mediate communication between chimeric modules. Surprisingly, and in contrary to the earlier in vivo results, the turnover rates of all chimeric PKSs were drastically diminished, which was assigned to impaired chain translocation due to the non-native ACP–KS interactions.58

As module–module communication is necessary to generate productive chimeric PKS assembly lines, questions arise towards the affinity of these interactions in native proteins. So far only a few studies have addressed this issue. The dissociation constant of docking domains or modules connected by docking domains was found to be $K_D$ 70–130 mM (ref. 35) and $K_D$ 1–2 mM (ref. 59), respectively. Titration of different DEBS polypeptides and following Michaelis–Menten fitting resulted in $K_{so}$ values of 2.5–4 μM, indicating that the interaction efficiency of the naturally interacting polypeptides is rather weak.69

Overall, efforts in engineering PKSs via module replacement revealed the importance of linker regions (covalent linkers or
matching docking domains) to mediate communication between heterologous modules. The nature of the non-native ACP–KS interaction during chain translocation appears as another factor influencing turnover in chimeric PKSs. A mere module–module exchange employing docking domains to facilitate communication across heterologous modules had limited success. Covalent linking of modules seems to allow the design of catalytically more active PKS chimera, but is done at the expense of modularity.

3.2. Domain–domain exchanges

Another strategy of engineering PKSs for producing novel compounds is the exchange of individual domains. One of the most common ways of altering the final polyketide product is to insert AT domains with altered specificity to incorporate a different starter or extender unit and thus change the entity at the α-carbon. Altering the stereochemistry of the final product can also result in novel biological activities making KR domain exchanges or reductive loop swaps of particular interest. The other domains (KS, ACP, and DH) were only rarely employed in domain exchanges. In the following, we highlight successful examples of engineering projects and analyze their approaches in the light of available structural information. For a recent comprehensive overview of engineering approaches, we refer to Barajas et al.61

3.2.1. AT domain exchanges in loading modules.

One of the easiest ways of inserting a new chemical moiety into a polyketide product is to exchange or mutate the AT domain of the loading module to incorporate different starting unit. For example, the AT of the loading didomain of DEBS has been exchanged with the respective AT of the avermectin synthase (AVES) loading didomain, which increased the diversity of erythromycin analogs produced in vivo.62 In another study, the complete exchange of loading didomains of the same type (AT–ACP form) of the tylactone synthase (TYLS) into the platenolide
synthase was reported. It was even possible to exchange loading domains of different architecture (AT-ACP from DEBS vs. KS-AT-ACP from oleandomycin synthase (OLEAS)). While an exchange of whole loading modules interferes in ACP-KS interaction during translocation and thus may decrease product yields, a mere AT exchange affects the specific interaction of AT–ACP during transacylation. As both interactions are important for proper turnover, an adaption of the non-native module–module or domain–domain interfaces seems necessary to avoid kinetic penalties.

3.2.2. AT domain exchanges in elongating modules. The exchange of single extender AT domains within a given PKS system is the most commonly used engineering strategy to alter the polyketide product at a certain position. The first AT exchange was carried out in 1996 and similar approaches followed in these early years after PKS discovery. In most of these studies, the novel polyketide product was formed, albeit at lower yields. For receiving an initial understanding about the structural integration of extender AT domains in the overall PKS fold, limited proteolysis experiments were conducted. Those experiments indicated that linker regions especially downstream of the AT domain are important for a successful AT exchange. While it was shown that the post-AT linker is neither required for acylation of the AT domain nor for the transacylation from the AT to the ACP domain, its presence was recognized as mandatory for KS–ACP catalyzed chain elongation. The impact of linker regions on the overall protein stability became apparent upon obtaining the first high resolution structures. A conserved sequence in the C-terminal part of the post-AT linker (PAL2cons.) folds back onto the KS and also the KS–AT linker interacts with the post-AT linker (Fig. 3B and C). Recently, a first systematic analysis of junction sites was conducted by exchanging ATs of DEBS as well as of lipomycin synthase (LIPS). Based on sequence alignments of different AT domains, optimal fusion sites for AT domain exchanges were derived. Best tolerated AT exchanges were achieved by swapping the AT with the adjacent KS–AT linker (KAL) and the N-terminal part of the post-AT linker (PAL1) (construct KAL–AT–PAL1). In agreement with previous studies, this strategy is successful, as it preserves the unit of KS with the conserved N-terminal part of the post-AT linker (PAL2cons.). Thus, there is good evidence that the best fusion sites for AT domain exchanges lie in the KS–AT and post-AT linker regions.

AT domain exchanges are a good example for how structural information can guide PKS engineering in defining appropriate exchange sites.

3.2.3. Implication of ACP–AT interactions on PKS engineering. Only few studies have addressed the impact of non-native ACP–AT interactions on PKS turnover. As the interaction of the ACP with the AT is transient and presumably weak, mapping of the ACP–AT interface has been difficult in the past. Recently, the first X-ray structural model of an ACP–AT complex from the trans-AT vicenistatin synthase (VINS) revealed an interaction of ACP helix II with the AT. While this study provided the first high-resolution 3D-structure, previous studies already employed docking simulation and alanine scanning mutagenesis to map the respective ACP–AT interaction interface. Studies of the ACP–AT interaction in the cis-AT PKS DEBS, the trans-AT disorazole synthase (DSZS), and the iterative acting enediyne synthase (PKSE DynE8) revealed overlapping – yet not identical – interfaces, in which the ACP interacts with the AT via regions on helix I and helix II (and additional loop regions depending on the different models). A combination of docking simulation and alanine-mutagenesis identified residues responsible for the ACP–AT interaction of the trans-AT kirromycin synthase (KIRS). Similar to previous studies, residues of ACP helix I, loop I and helix II were predicted to be involved in AT recognition. Based on this knowledge a non-native AT–ACP interaction between the trans-AT KirCII and DEBS ACP6 was engineered towards improved transacylation properties.

3.2.4. Reductive loop swaps/KR exchanges. KR domains are responsible for the NADPH-dependent reduction of the β-keto group and if present also for the epimerization of the α-substituent. The specificity of KR domains for their cognate ACPs has been suggested to be relatively low, therefore KR domains may be more tolerant in domain swaps than AT domains.

The KR domain has been successfully exchanged in several cases in partly reducing β-modules, particularly for studying the stereochemistry of the reduction reaction that configures the β-hydroxy functionality and the α-alkyl group in case other units than malonyl-CoA are accepted for elongation. These engineering successes can be attributed to the early availability of high resolution 3D-structures, which allowed for a better understanding of the KR from a structural point of view. Although KR domains are well described, which has led to a general classification scheme by their reduction and epimerization activity, they have, however, been shown to be inefficient in the simultaneous alteration of alkyl- and hydroxyl-group configurations in a heterologous context.

More recently, a dimerization element (DE) was identified, which is placed N-terminally to 50% of PKS modules that contain KR as the only processing domain. The DE turned out to be responsible for the stability of the respective KR domains. As a current rule for KR engineering in partially reducing PKSs (KS–AT–KR–ACP), DE domains of the host PKSs should be preserved in KR exchanges, while a DE–KR unit should replace KR in PKSs not natively carrying a DE structural motif (Fig. 4A).

Swaps of complete processing wings (or reductive loops) were among the earliest achievement of PKS engineering, due to the structural integrity of this unit. A similar strategy has led to the successful production of adipic acid. Here, the processing wing of the borrelidin synthase module 1 (BORS M1), which contains only a KR domain, has been replaced with the fully reducing processing wing of the second module of the spinosyn synthase (SPNS M2). The DH domain turned out to be problematic owing to restricted substrate specificity, but when swapped with the DH-domain of BORS M2 the expected product was produced. Not surprisingly, the definition of junction sites for reductive loop insertions underlay the same rules as AT
exchanges pointing again at the importance of preserving the KS–PAL2cons. interface (Fig. 4B).

3.3. Non-rational engineering strategies

Besides the aforementioned rational engineering attempts, non-rational engineering can also result in the generation of new polyketides. In nature, homologous recombination is an often used process for the generation of diversity. Based on homologous recombination in *S. cerevisiae*, chimeras of PPKS M5 and DEBS3 were successfully generated in *in vivo*, of which some had measurable *in vitro* activity indicative of a stable protein fold.98 Although it was possible to rapidly generate a library of chimeric PPKSs, stability of the resulting folds could not be ensured under the experimental conditions. Recently, a similar approach used homologous recombination within the rapamycin synthase (RAPS) to generate several rapamycin derivatives with altered biological activity.99 Interestingly, the best chimeras revealed junction sites in the KS or AT domain or in the linker upstream of the ACP domain. The approaches use homologous recombination as a tool to mimic evolutionary pressure for receiving chimeric PPKSs with sufficient catalytic fitness. Such strategies can identify junction sites that may eventually support rational engineering strategies.

4. Roadmap for another 20 years of PKS engineering

Below we highlight several aspects of PKS research that should be considered to come closer to the goal of custom polyketide synthesis by designer PKSs.

4.1. Continuing efforts in the structural analysis of PKSs

In recent years, a wealth of structural data at high resolution has given insight into the principles of polyketide biosynthesis.10 There is still the clear need to further analyze PKS in terms of their structural and conformational dynamic properties with a dedicated focus on understanding module–module interactions in PKS assembly lines. The technological progress in cryo-EM will foster efforts in the characterization of particularly the larger assemblies.99 As a single-particle method, cryo-EM can additionally give insight into conformational dynamics when particles are classified during manual and automated sorting.110,111 or initial electron density maps are analyzed for principle components.112 In addition to the static snapshots provided by cryo-EM or X-ray crystallography, the dynamic character of PKS necessitates the use of spectroscopic methods. These methods can deliver...
a continuous spectrum of protein conformational dynamics in real-time and add valuable information on the internal dynamics of these domains.\textsuperscript{99} As conformational motions in proteins are stochastic and synchronization of populations are hardly possible, spectroscopic methods will be particularly powerful when applied at the single-molecule level.

4.2. Characterizing selected PKSs in vitro

The access to several PKSs by recombinant production in high yield, high protein quality and by fast purification protocols has been achieved in \textit{Escherichia coli} and in \textit{Streptomyces}.\textsuperscript{94} On the basis of this methodological foundation, selected PKSs have been established as model systems \textit{in vitro} and characterized to considerable detail to date; \textit{i.e.} the modPKSs DEBS, PIKS, and SPNS, and the iPKS PKSA and MSAS.\textsuperscript{95–99} One needs to further analyze these proteins in quantitative terms \textit{in vitro} by classical enzymatic methods, as well as to enlarge the toolbox with a set of new enzymes. The carefully selected PKSs can function as testbeds to probe the precise impact of domain swaps and mix-and-match approaches. A tight network of quantitate data on the selected proteins will help understanding structure-function-relationships in PKSs to establish a toolkit of methods and engineering rules that are broadly applicable for the PKS protein family.

4.3. Understanding substrate shuttling

Conformational properties of the evolutionarily related mFAS proteins are stochastic and synchronization of populations are hardly possible, spectroscopic methods will be particularly powerful when applied at the single-molecule level.\textsuperscript{99} As conformational motions in proteins are stochastic and synchronization of populations are hardly possible, spectroscopic methods will be particularly powerful when applied at the single-molecule level.

4.4. Redefining module boundaries

Over the last two decades, mix-and-match approaches for PKS engineering were based on the traditional module definition in which the domains are ordered in a KS–AT–[DH–ER–KR]–ACP sequence. Although several studies implied that preservation of native interaction interfaces is crucial with regard to any engineering attempt (\textit{e.g.} ACP–KS linker,\textsuperscript{106} ACP–KS interaction\textsuperscript{52,58}), these traditional module boundaries were never challenged as suited junction sites in mix-and-match approaches. Recently, a study on the co-evolution of large PKS assembly lines revealed that the processing domains are more closely related to the downstream than to the upstream KS.\textsuperscript{107} A unit that may be better exchangeable in mix-and-match approaches than the genetically-encoded, “traditional” module could therefore be of AT–[DH–ER–KR]–ACP–KS domain sequence. From an engineering point of view, an exchangeable unit with such newly defined boundaries may indeed make sense, as it allows to preserve the important interface of the upstream ACP to the downstream KS,\textsuperscript{98} which was found to be crucial also in the closely related NRPSs,\textsuperscript{108} as well as in \textit{trans}-AT PKSs.\textsuperscript{109} The general applicability of the new module boundaries for PKS engineering remains to be shown.

4.5. Controlling vectorial synthesis

In the serial arrangement of modules in PKS assembly lines, reaction progress can occur via re-feeding of the intermediate to the integral KS, leading to repeated condensation and processing as encoded on the integral module (“stuttering”), or the translocation of the intermediate to the KS of the downstream module.\textsuperscript{11} Engineering of PKS assembly lines requires control of this decision to steer synthesis towards translocation (see Fig. 2B). The means to impose kinetic reaction control include the modulation in $k_{\text{cat}}/K_M$, the tuning of electrostatic properties of domain–domain interfaces, and the adjustment of the spatial arrangement of interacting domains.\textsuperscript{103,104} A recent model study has demonstrated that kinetic engineering can steer synthesis. On the example of the microbial FAS, selected mutations redirected the complex biosynthetic pathways of fatty acid synthesis towards the production of short fatty acids.\textsuperscript{105}

4.6. Modeling the PKS reaction networks

\textit{In silico} modeling of the reaction network underlying polyketide biosynthesis will be necessary to disclose and exploit enzymatic properties of PKSs. Owing to a set of comprehensive enzymatic assays in the labs, selected enzymatic functions have been already experimentally described and many more enzyme kinetic parameters of the catalytic processes will be available during the next years. \textit{In silico} models need to be implemented using these data to form a knowledgebase, to then, in a computational filtering process, give access to enzyme kinetic properties that are not directly experimentally accessible.\textsuperscript{109} Once established, reaction networks can be extended to represent effects of engineering, such as the modulated enzymatic constants resulting from the swapped domain and/or changed spatial constraints introduced by binding site design. Computational tools will add a new dimension to the quality of understanding of PKSs that goes beyond a conventional phenomenological analysis, and will eventually inform PKS engineering strategies.
5. Conclusions

In this review, we summarize and revise engineering approaches in the light of available structural and functional data. We present several models for PKS architectures and employ sequence and structural analysis to investigate the recent strategies conducted in PKS engineering.

There is still a strong demand for a highly detailed structural and biochemical characterization of PKSs especially with improvements in structure determination methods. Inclusion of complementary techniques that provide dynamic information will expand the view on programming PKS biosynthesis. One of the key elements in the rational design of chimeric PKSs is the definition of appropriate junction sites that lead to structurally intact protein folds to guarantee the productive interaction of modules and domains throughout the catalytic cycle. Since many of the important interfaces are still poorly described, we are convinced that a combination of experimental data, evolutionary analysis and in silico modeling will help overcome these shortcomings and pave the way for general applicable engineering rules to PKS systems.

6. Conflicts of interest

There are no conflicts to declare.

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8. References

1. C. Hertweck, Angew. Chem., Int. Ed., 2009, 48, 4688–4716.
2. J. Staunton and K. J. Weissman, Nat. Prod. Rep., 2001, 18, 380–416.
3. K. J. Weissman, Nat. Chem. Biol., 2015, 11, 660–670.
4. J. Piel, Nat. Prod. Rep., 2010, 27, 996–1047.
5. R. W. Broadhurst, D. Nietlispach, M. P. Wheatcroft, P. F. Leadlay and K. J. Weissman, Chem. Biol., 2003, 10, 723–731.
6. J. Dorival, T. Annaval, F. Risser, S. Collin, P. Roblin, C. Jacob, A. Gruetz, B. Chagot and K. J. Weissman, J. Am. Chem. Soc., 2016, 138, 4155–4167.
7. P. F. Leadlay, Curr. Opin. Chem. Biol., 1997, 1, 162–168.
8. C. Khosla and R. J. X. Zawada, Trends Biotechnol., 1996, 14, 335–341.
9. R. McDaniel, M. Welch and C. R. Hutchinson, Chem. Rev., 2005, 105, 543–558.
10. K. A. J. BozhýyÜ, F. Fleischhacker, A. Linck, F. Wescbe, A. Tietze, C.-P. Niesert and H. B. Bode, Nat. Chem., 2017, 10, 275–281.
11. A. C. Murphy, H. Hong, S. Vance, R. W. Broadhurst and P. F. Leadlay, Chem. Commun., 2016, 52, 8373–8376.
12. F. Wang, Y. Wang, J. Ji, Z. Zhou, J. Yu, H. Zhu, Z. Su, L. Zhang and J. Zheng, ACS Chem. Biol., 2015, 10, 1017–1025.
13. L. Zhang, J. Ji, M. Yuan, Y. Feng, L. Wang, Z. Deng, L. Bai and J. Zheng, ACS Chem. Biol., 2018, 13, 871–875.
14. H. Jenke-Kodama, A. Sandmann, R. Müller and E. Dittmann, Mol. Biol. Evol., 2005, 22, 2027–2039.
15. S. Smith and S.-C. Tsai, Nat. Prod. Rep., 2007, 24, 1041–1072.
16. T. Maier, M. Leibundgut, D. Boehringer and N. Ban, Q. Rev. Biophys., 2010, 43, 373–422.
17. M. Grininger, Curr. Opin. Struct. Biol., 2014, 25, 49–56.
18. Y. Tang, C. Kim, I. I. Mathews, D. E. Cane and C. Khosla, Proc. Natl. Acad. Sci. U. S. A., 2006, 103, 11124–11129.
19. Y. Tang, A. Y. Chen, C.-Y. Kim, D. E. Cane and C. Khosla, Chem. Biol., 2007, 14, 931–943.
20. A. T. Keatinge-Clay and R. M. Stroud, Structure, 2006, 14, 737–748.
21. D. A. Herbst, R. P. Jakob, F. Zähringer and T. Maier, Nature, 2016, 531, 533–537.
22. A. Keatinge-Clay, J. Mol. Biol., 2008, 384, 941–953.
23. J. Zheng, D. C. Gay, B. Demeler, M. A. White and A. T. Keatinge-Clay, Nat. Chem. Biol., 2012, 8, 615–621.
24. J. Zheng, C. D. Fage, B. Demeler, D. W. Hoffman and A. T. Keatinge-Clay, ACS Chem. Biol., 2013, 8, 1263–1270.
25. D. Khare, W. A. Hale, A. Tripathi, L. Gu, D. H. Sherman, W. H. Gerwick, K. Hákansson and J. L. Smith, Structure, 2015, 23, 2213–2223.
26. A. T. Keatinge-Clay, Cell Chem. Biol., 2016, 23, 540–542.
27. S. Dutta, J. R. Whicher, D. a. Hansen, W. a. Hale, J. a. Chemler, G. R. Congdon, A. R. H. Narayan, K. Hákansson, D. H. Sherman, J. L. Smith and G. Skiniotis, Nature, 2014, 510, 512–517.
28. T. Maier, M. Leibundgut and N. Ban, Science, 2008, 321, 1315–1322.
29. G. Pappenberger, J. Benz, B. Gsell, M. Hennig, A. Ruf, M. Stihle, R. Thoma and M. G. Rudolph, J. Mol. Biol., 2010, 397, 508–519.
30. A. Rittner, K. S. Paitthankar, K. V. Huu and M. Grininger, ACS Chem. Biol., 2018, 13, 723–732.
31. A. Rittner and M. Grininger, ChemBioChem, 2014, 15, 2489–2493.
32. K. J. Weissman, Nat. Prod. Rep., 2015, 32, 436–453.
33. A. L. Edwards, T. Matsui, T. M. Weiss and C. Khosla, J. Mol. Biol., 2014, 426, 2229–2245.
34. C. D. Richter, D. Nietlispach, R. W. Broadhurst and K. J. Weissman, Nat. Chem. Biol., 2008, 4, 75–81.
35. T. J. Buchholz, T. W. Geders, F. E. Bartley III, K. a. Reynolds, J. L. Smith and D. H. Sherman, ACS Chem. Biol., 2009, 4, 41–52.
36. J. R. Whicher, S. S. Smaga, D. a. Hansen, W. C. Brown, W. H. Gerwick, D. H. Sherman and J. L. Smith, Chem. Biol., 2013, 20, 1340–1351.
37. E. J. Brignole, S. Smith and F. J. Asturias, Nat. Struct. Mol. Biol., 2009, 16, 190–197.
