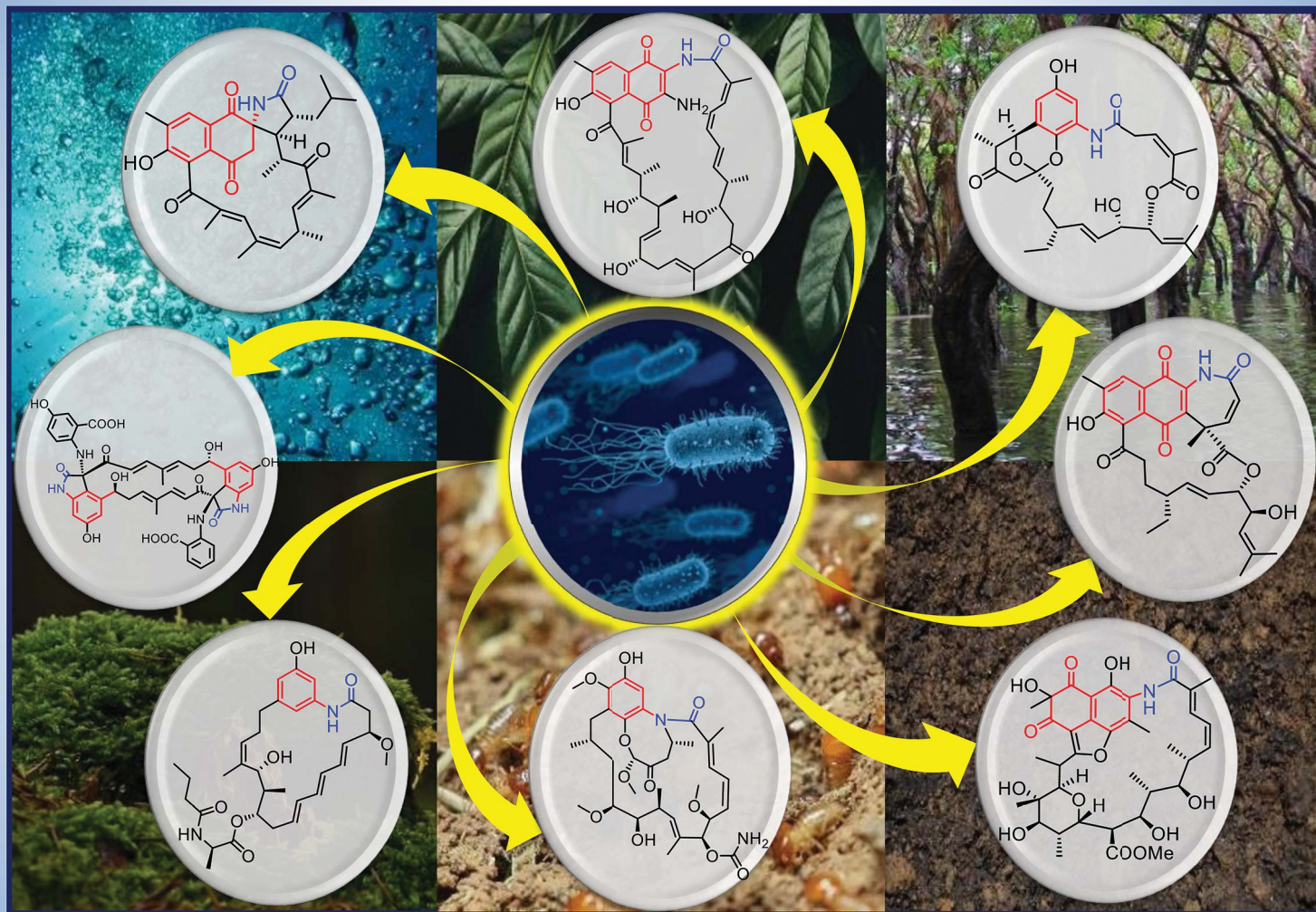


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REVIEW ARTICLE

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Modifications, biological origin and antibacterial
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Modifications, biological origin and antibacterial activity of naphthalenoid ansamycins

Natalia Skrzypczak and Piotr Przybylski *

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Structural division of natural naphthalenoid ansamycins, regarding the type of the core and length of the ansa chain, and their biosynthetic pathways in microorganisms are discussed. The great biosynthetic plasticity of natural naphthalenoid ansamycins is reflected in their structural variety due to the alterations within ansa bridge or naphthalenoid core portions. A comparison between the biological potency of natural and semisynthetic naphthalenoid ansamycins was performed and discussed in relation to the molecular targets in cells. The antibacterial potency of naphthalenoid ansamycins seems to be dependent on the ansa chain length and conformational flexibility – the higher flexibility of the ansa chain the better biological outcome is noted.

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1. Introduction

The term “ansamycin” was first suggested by Prelog and Oppolzer, to describe compounds being closely related to those discovered by Lüttringhaus (ansa compounds), and derives from the name for the characteristic ansa chain structural motif (ansa in Latin means “handle”).^{1,2} The discovery of the first naphthalenoid ansamycin antibiotics called rifamycins, was made by Professor Piero Sensi and his research group from Milan (Lepetit Research Laboratories).^{3–5} The basket-like

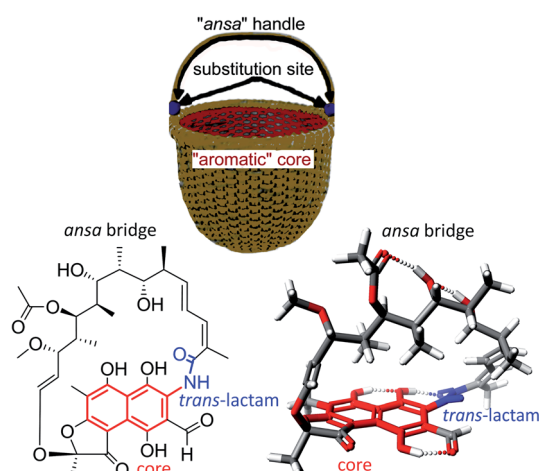


Fig. 1 Structural analogy between the basket and 3-formylrifamycin SV, a member of naphthalenoid ansamycins.

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structure of ansamycins (Fig. 1 and 2), consisting usually the aromatic core and flexible ansa bridge, has been shown to display some structural flexibility influencing biological properties, characteristic for chameleonic-type drugs.^{6–8} Taking into account recent achievements with mutasynthesis and isolation of novel natural scaffolds among macrolactams from bacteria strains, the term “ansamycins” can be used in a broader sense. Natural ansamycins can be described as a family of macrolactams, bearing relatively rigid central cores as benzenoid, naphthalenoid or atypical ones as *e.g.* alicyclic or heterocyclic ones, originated from AHBA or other-type small semisynthetic precursors (mutasynthons), and produced in as a result of cooperation between polyketide (PKS type I) and amide synthases (Fig. 2). The important role of naphthalenoid rifamycin-type antibiotics in medical therapy as antibacterial agents, especially against *M. tuberculosis*, encouraged us to report about their recently discovered natural sources, biogenesis, modifications, and structure–activity relationships.

2. Structural division of naphthalenoid ansamycins and their natural sources

Ansamycins can be roughly divided into three main groups called benzenoid, naphthalenoid, and atypical ones and further based on the length of the ansa chain. Funayama and Cordell divided ansamycins into two main subgroups and highlighted common biosynthetic paths of these natural products.⁹

The structural classification of ansamycins can be performed in a more detailed manner *e.g.* taking into account oxidized/reduced structure of the rigid core or the regioisomers of the quinone/iminoquinone moieties, as shown in Fig. 2. Thus, in an addition to the Funayama and Cordell classification, we propose several groups of naphthalenoid ansamycins, which can be divided into those having naphthalene, naphthalene-1,3(2*H*,4*H*)-dione, naphthalen-1(2*H*)-one, 2,3-

dihydronaphthalene-1,4-dione, 3,4-dihydronaphthalen-1(2*H*)-one, 1,4-naphthoquinone, 1,2/1,4-iminonaphthoquinone or naphthalen-1(4*H*)-one cores and lactam ansa bridges of different length (Fig. 2). Some of biosynthetic, biological and analytical aspects of naphthalenoid ansamycins have been discussed in earlier reviews published between the years 1972 and 2011.^{9–23} Since that time many biological relevant natural naphthalenoid ansamycin scaffolds have been isolated from bacteria or semisynthetically modified, revealing differences in their modes of action. Isolation of novel naphthalenoid ansamycins of unique structure, in terms of the new core type or the length and structure of the ansa bridge, indicated a great structural diversity among these natural products and shed new light on alternative biosynthetic pathways realized in microorganisms. Comparison of structural division between benzenoid and naphthalenoid ansamycins indicates a greater structural variety of the latter than the former ones. Irrespectively on the type of the core or the length of the ansa bridge, the main natural source evidenced for naphthalenoid ansamycins are bacteria strains (Fig. 2; bottom), similarly as for the benzenoid ansamycins, which has been demonstrated. Different bacteria strains producing naphthalenoid ansamycins can be, however, derived from soil (*Streptomyces* sp.; *Amycolatopsis* sp.; *Micro-monospora* sp.), water (*Streptomyces* sp.; *Salinispora* sp.; *Micro-monospora* sp.) as well as from higher plants as *e.g.* trees or shrubs *Manglietia hookeri* (*Streptomyces* sp. CS).

3. Comparison between biosynthetic pathways of naphthalenoid and benzenoid ansamycins

Ansamycins are synthesized in nature *via* the polyketide pathway from AHBA, being a product of the modified shikimate pathway, and acetate with propionate units, irrespective of the type of the rigid core, as evidenced by experiments with isotopically labeled biosynthetic precursors (Fig. 3).^{12,24–27} The major difference



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transformation of ansamycins cores, based on modern synthetic methods (click chemistry).



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research interests are focused on modification and determination of SAR for natural products and their congeners, and with tautomerization, atropisomerization and zwitterionization processes. His current studies are related to cascade approaches enabling modification of lactone and lactam macrolides.



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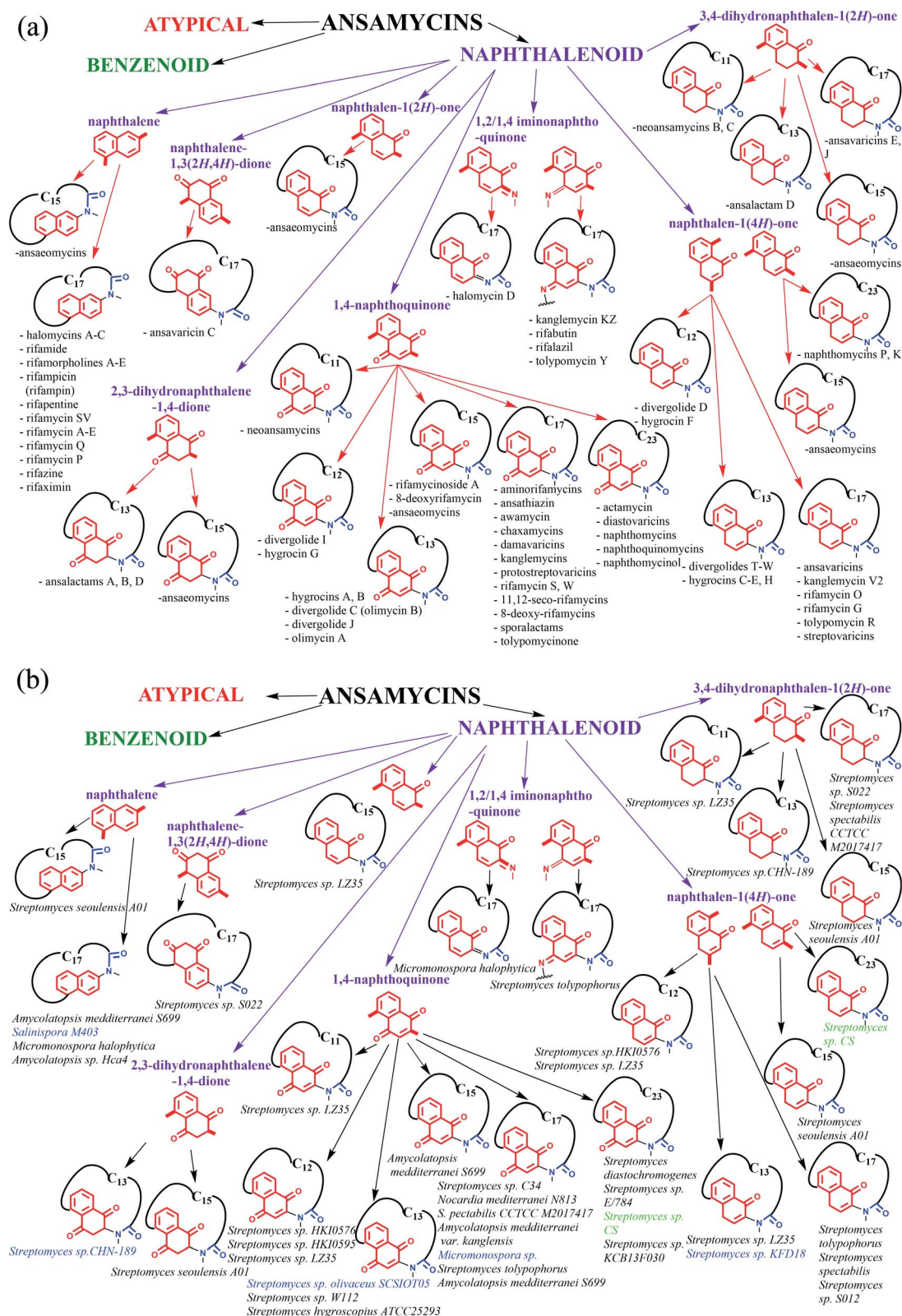


Fig. 2 Structural division of naphthalenoid ansamycins regarding the core type and the length of the ansa chain (top) and natural sources of these ansamycins (bottom): soil-derived bacteria (black); marine bacteria (blue), bacteria associated with the higher plants (green).

substitution) and the central core (type), as shown in Fig. 3. Studies with isotopically labeled biosynthetic units *e.g.* malonyl, methylmalonyl, glycolate, and methyl (methionine), required for

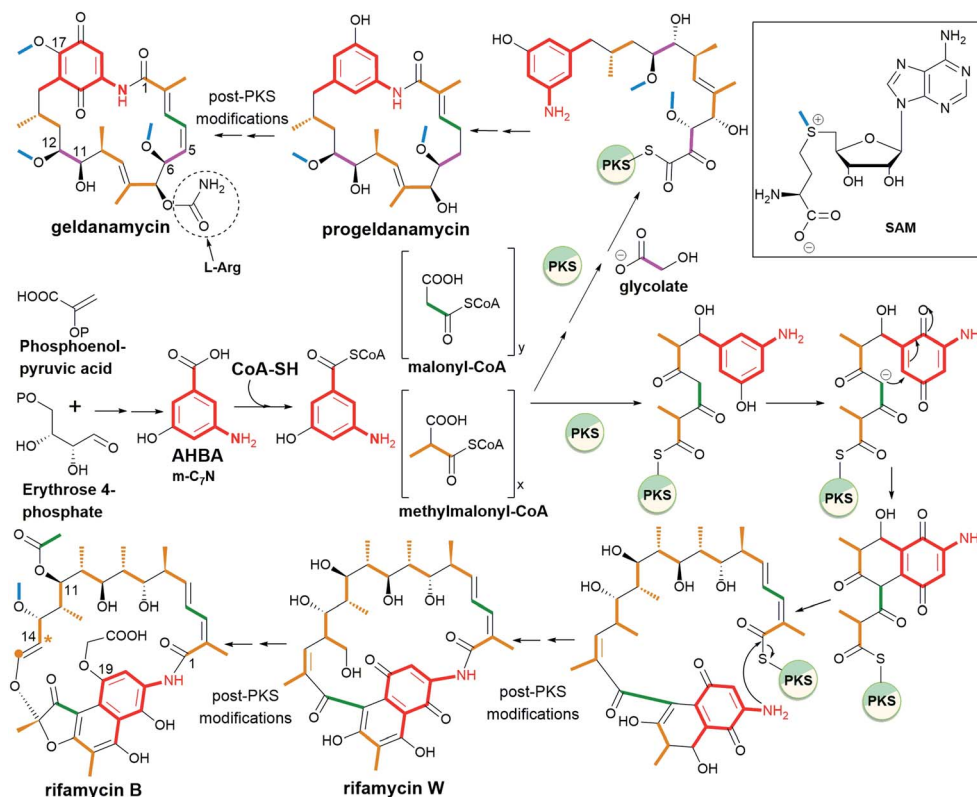


Fig. 3 Biosynthetic pathways of the two representatives of ansamycins: geldanamycin (benzenoid – benzoquinone- C_{15}) and rifamycin B (naphthalenoid – naphthalene- C_{17}).

producing geldanamycin and rifamycin B, clearly revealed key differences in biosynthetic pathways between these ansamycins. The three malonyl and eight methylmalonyl units are incorporated into the rifamycin B scaffold, where seven of methylmalonyl units are inserted into the ansa bridge portion and the one is required for cyclization of the naphthalenic core (orange, Fig. 3). In contrast to that, only four methylmalonyl, two glycolate and one malonyl units are inserted into the molecular “handle” of the geldanamycin basket by PKS I (Fig. 3).²⁸ At the early stage of rifamycin B biosynthesis realized by bacteria, the closing of the naphthalene core *via* an intramolecular conjugate addition of Michael-type takes place (Fig. 3). Post-PKS I tailoring of rifamycin B and geldanamycin is based on cyclization catalyzed by amide synthase with the formation of rifamycin W or progeldanamycin, respectively. Rifamycin B post-PKS I transformations, apart from the lactam formation and methylation *via* SAM, involve acetylation of hydroxyl at C(11), attachment of 3-phosphoglyceric or 3-phosphohydroxypyruvate to the phenol group at the C(19), oxygen insertion between vinyl and naphthalene portions and the C_1 unit lost at the C(14) position (asterisk, Fig. 3). In turn, post-PKS I modifications of progeldanamycin are concerned with oxidation of C(4)–C(5) positions, three *O*-methylations with SAM, and carbamoylation at C(7). Another difference between the above biosynthetic pathways is monooxygenation at C(17) and oxidation of *p*-phenol moiety into the quinone for geldanamycin, and the reduction of the quinone to the phenol form for rifamycin B. It should be noted that quinone or quinol forms of the

above ansamycins are important for better molecular recognition with their different targets in cells. The absolute configurations of contiguous stereogenic centers in the polyketide chains of naphthalenoid and benzenoid ansamycins, can be predicted

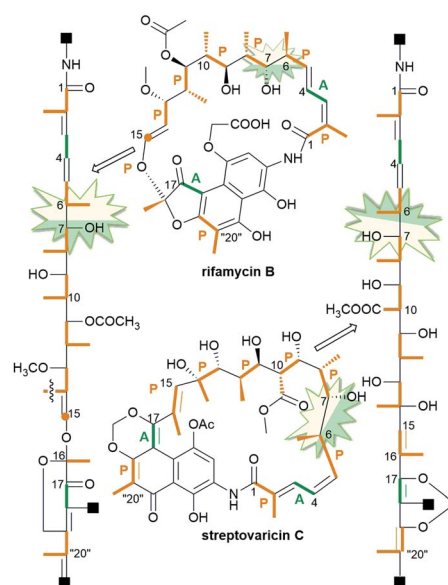


Fig. 4 Model of expected stereochemistry within the ansa chains of naphthalenoid ansamycins according to Funayama *et al.* or Celmer *et al.*; *p*-propionate (orange) and A-acetate (green) units.



using a model developed by Funayama *et al.*, analogous to the Celmer model referred to macrolactones (Fig. 4).^{29,30} As it can be deduced from Fig. 4, rifamycin B and streptovaricin C belonging to naphthalenoid ansamycins are biosynthesized using a common amide-head biosynthetic pathway; with the exception of the C(6) and C(7) positions, the absolute configurations of the stereogenic centers along the ansa chain of rifamycin B appear to be identical to those along the ansa chain of streptovaricin C. Most of the ansamycins, having seventeen carbon atoms in the polyketide chain, show the same structural sequence along the chain, *i.e.*, PAPP PPPAP, where P and A are propionate (methylmalonyl) and acetate (malonyl) units, respectively. Kanglemycins are the exception to this rule because they have an extra C₁ unit.^{31,32} In contrast to the naphthalenoid-C₁₇ ansamycins, those having a longer “handle” of the molecular basket, as *e.g.*, actamycin with C₂₃-ansa chain, show a different structural sequence along the bridge: AAAPAPAAPPAP.

4. Structural aspects of naphthalenoid ansamycins influencing their physico-chemical and biological properties

Naphthalenoid ansamycins possess a different number of double bonds within the ansa bridge (Fig. 5). The number and position of double bonds along the ansa chain together with intramolecular interactions, as *e.g.* H-bonds, have an impact on the possibility to adopt various conformations of the ansa “handle” of naphthalenoid ansamycins in response to environmental factors such as the nature of cell barriers or steric demands of the molecular target. Due to the existence of the two structural portions, a rigid naphthalenoid core and the flexible-

ansa chain bearing stereogenic centers, the relative arrangement of them contributes to atropisomerism (from Greek “a” meaning “not” and “tropos” meaning “turn”). This special case of stereoisomerism, dependent on the altered arrangement of the whole ansa bridge relative to the plane of the rigid core for naphthalenoid ansamycins, was also observed for the other medicinal relevant biomolecules as benzenoid ansamycins, alkaloids, purines or diaryl or aryl-heterocyclic drugs, where both axial and planar chiralities were well evidenced.^{33–36} Excellent, experimentally indicated examples of stereoisomerism (atropisomerism) among naphthalenoid ansamycins, induced by plane or axis, are streptovaricins (Fig. 6). Single crystal X-ray studies showed that one of the boronic esters of streptovaricin C (Fig. 5) has an opposite helicity of the ansa bridge to that present in most natural streptovaricins and other naphthalenoid ansamycins (Fig. 6).^{37,38} Furthermore, it was evidenced that for various rifamycins (including rifabutin, rifapentine, rifaximin, and rifampicin) the conformation of the ansa bridge and its arrangement relative to the central core are dependent on the phase (solid/liquid) or polarity and nature (protic/aprotic) of the solvent used.^{39–42} The conformation flexibility of the ansa chain for ansamycins, *e.g.* rifamycins, is not always concerned with atropisomerism, however, is crucial for chameleonic properties of this-type macrolactams, reflected in a facile adaptation to the changing cell environment during

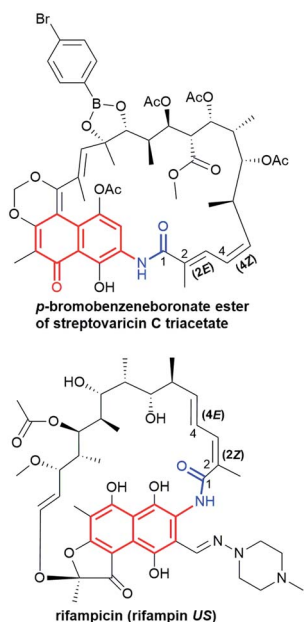


Fig. 5 Different configuration of double bonds within ansa bridge for exemplary naphthalenoid ansamycins.

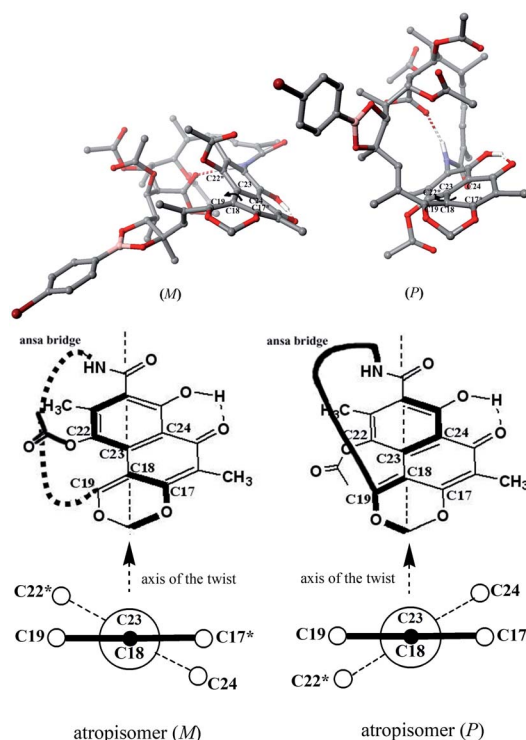


Fig. 6 Newman projection of the axial stereoisomerism of streptovaricin C derivatives, implied by the two different orientations of the ansa bridge (bottom) relative to the aromatic core, visualized on molecular models (top) via Scigress (EU 3.1.8), Fujitsu. Models of streptovaricins for DFT optimizations were assumed using X-ray structure of *p*-bromobenzenboronate ester of streptovaricin C triacetate.³⁷



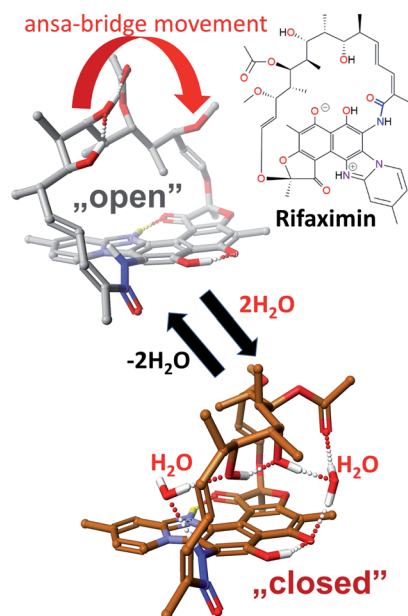


Fig. 7 Key conformers for a total functionality of rifamycins: an “open” ansa bridge conformer of lower lipophilicity and crucial for binding with bacterial RNA polymerases (top), and “closed” ansa-bridge conformer of higher lipophilicity needed to overcome natural cell barriers, visualized by Scigress (EU 3.1.8), Fujitsu.⁴⁴

transport of the drug to the target site.⁴³ Recently, it was demonstrated that interactions of naphthalenoid ansamycins (rifamycins) with water molecules lead to the formation of a “closed”-type structure (Fig. 7), with polar groups hidden toward the core, which increases the lipophilicity and enable diffusion of these antibiotics in bacterial cell membranes.⁴⁴ However, a more hydrophilic rifamycin structure in solution, with polar groups exposed to the solvent, *i.e.*, in the so-called “open” structure (Fig. 7), is necessary for the rifamycins to target bacterial RNA polymerases.^{45–47} Hence, a smooth conversion between these two-type structures with a displacement of the whole ansa bridge (Fig. 7), *i.e.* chameleonic nature, is highly important to the presence of attractive biological activity of naphthalenoid ansamycins.⁴⁴

Another structural factor of naphthalenoid ansamycins is their zwitterionization as a result of the presence of both acidic and basic groups in scaffolds, which affects lipophilicity and water solubility in the cell environment. This phenomenon was the best researched for naphthalenoid rifamycins, where the proton of the phenol group was transferred to the most basic centre in the molecule (Fig. 7 and 8).^{44,48–50} It was indicated that despite unfavorable low water-solubility of the non-ionic form of rifamycins, the optimization of this physico-chemical parameter is realized by the proton transfer yielding zwitterions, at the active participation of the protic solvent. In turn, aprotic solvents favoured non-ionic form of rifamycins, better adapted to lipophilic cell barrier environment. This another chameleonic feature of rifamycins such as rifampicin or its various amine congeners was evidenced not only *via* 1D and 2D NMR and FT-IR spectroscopic studies in solutions but also was proved in crystal state where different solvents were

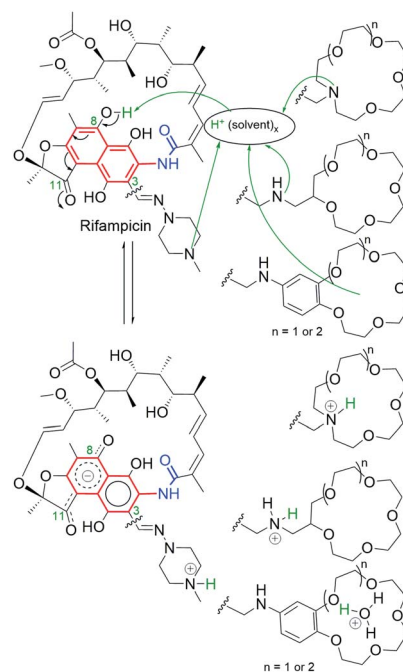


Fig. 8 Regioselective zwitterionization of rifampicin and its amine derivatives containing different-types of basic “centers”.

applied.^{44,48,51,52} X-ray and CP-MAS NMR solid state studies showed that crystallization of the rifampicin from protic solvents yielded zwitterionic form (water-soluble) of the antibiotic whereas crystallization from aprotic solvents afforded the non-ionic form (lipophilic) of this antibiotic.^{42,49,53} The other example of chameleonic ansamycin is rifaximin, where the zwitterion formation was accompanied by the equilibrium between more lipophilic (closed) and more hydrophilic (open) conformers, supported by water or other protic solvent molecules (Fig. 7).⁴⁴ Hence, the non-ionic structure of rifampentine, the antibiotic analogous to the above-mentioned rifamycins, in the solvate with methanol molecules⁵⁴ seems to be incorrect. The problem of ansamycin zwitterionization, influencing the molecular recognition of the antibiotic with the target in cells (bacterial RNA polymerases – RNAPs, Fig. 9) and lipophilic permeability efficiency, is often neglected in the literature.^{50,55,56} In the work of Campbell *et al.*, the structure of the non-ionic form of rifampicin (incorrect non-ionic pentahydrate crystal form – reinvestigated later as zwitterionic⁵¹) was used to build the model of interactions between this ansamycin and bacterial RNAPs (Fig. 9a).⁵⁷ Considering the presence of ansamycin zwitterionic form in water and close proximity of the protonated basic piperazine substituent at C(3) of rifampicin or its congeners and the negatively charged carboxylate of a highly conserved E445 of RNAPs (distance 3.4 Å), alternative binding models of rifamycins with RNAPs were proposed (Fig. 9b).^{51,53,58–60} In one of these alternative binding patterns concerning rifampicin, the protonated piperazine arm at a slightly altered conformation takes part in a very strong H-bond interaction (rifampicin) $N^+H \cdots H \cdots OOC(E445_{RNAP})$ of the salt-type character (H-bond donor ...acceptor ~ 2.6 Å; angle



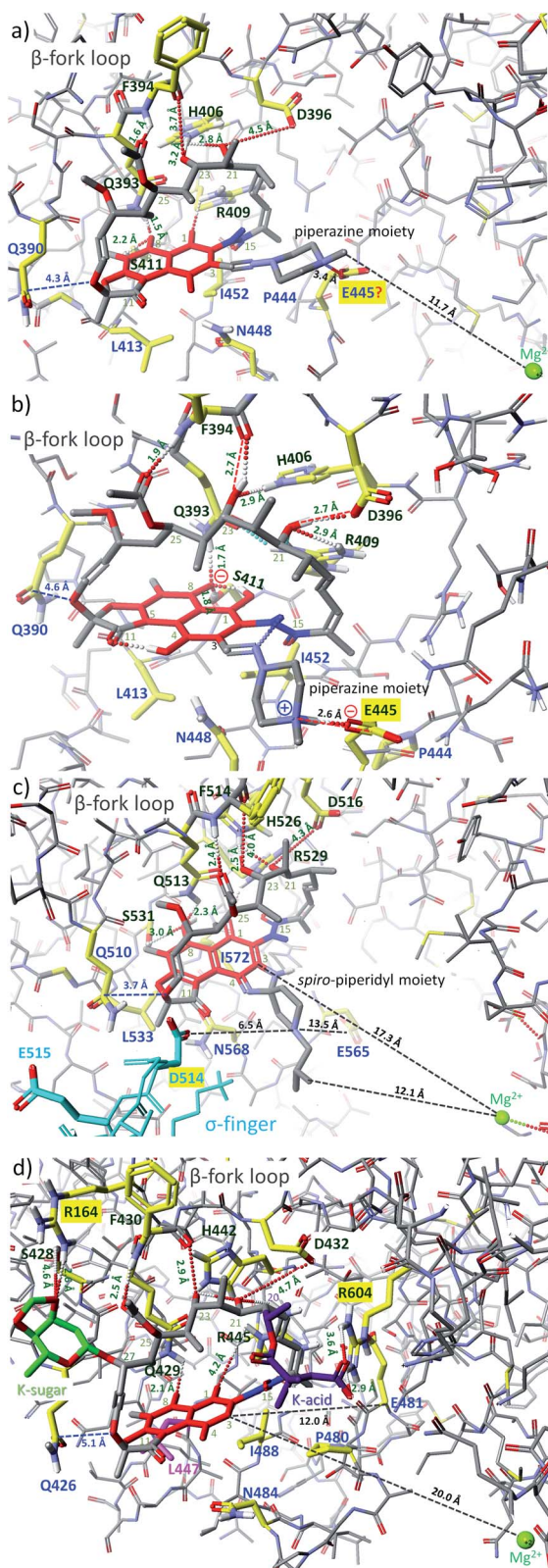


Fig. 9 Binding modes of: (a) rifampicin to RNAP of *Thermus aquaticus* (PDB 1I6V);⁵⁷ (b) rifampicin to RNAP of *Thermus aquaticus*⁵³ (alternative model built on the basis of MOG-PM6 calculations with RNAP structure – PDB 1I6V); (c) rifabutin to RNAP of *Thermus thermophilus* (PDB 2A6E);⁶¹ (d) kanglemycin A to RNAP of *Mycobacterium tuberculosis* (Mtb) mutant S447 → L447 (PDB 6DCF), resistant to rifampicin;³² visualized by Scigress (EU 3.1.8, Fujitsu). Interatomic distances are

~170°).⁵³ It should be mentioned that new models for rifampicin and its amine congeners explained the structure–activity relationships at changing C(3)-substituent of congeners having comparable lipophilicity and water solubility.

Bacterial RNAPs, as well-known targets in cells of naphthalenoid ansamycins, have identical key amino acids at analogous positions of the sequence, irrespectively on the bacteria strain (Fig. 9). Naphthalenoid ansamycin – rifabutin possesses spiro-piperidyl basic C(3)–C(4) arm, which is directed toward σ -finger of RNAPs (Fig. 9c),⁶¹ at nearly conserved interactions with β -fork loop, when compared to rifampicin. This model of rifabutin interactions, however, did not consider that C(3)–C(4) spiro-piperidyl arm possesses a basic nitrogen atom directed toward σ -finger of RNAPs, rich in acidic amino acids, as *e.g.* D514 or E515 (Fig. 9c). Following the above informations concerning binding of rifampicin with RNAPs, it was suggested that the C(3)–C(4) arm in the protonated form (*in vivo*), can be involved in H-bonding with D514 of the σ -finger.⁴⁴ In contrast to the above rifamycins varying at the core substitution, kanglemycins are examples of naphthalenoid ansamycins modified with K-acid and K-sugar at C(20) and C(27) positions of the ansa bridge, respectively (Fig. 9d). Due to this structure of kanglemycins, their location at RNAPs pocket is different than those of rifampicin or rifabutin antibiotics, taking into account the distance to Mg^{2+} and Q426/510/390 (Fig. 9d).³² The substitution of naphthalenoid-C₁₇ ansamycins at the ansa bridge with bulky substituents often results in hindered binding with bacterial RNAPs (one of the resistance mechanisms) and the markedly decreased or lack of antibacterial potency. Kanglemycins are, however, antibacterially potent, even against *M. tuberculosis* strains resistant to rifampicin, *inter alia* due to extra interactions of K-acid and K-sugar with R604 and R164 of RNAPs, respectively. It should be noted that in this altered binding model of kanglemycins, similarly to that of rifabutin one, there are lost interaction with amino acid at 447 positions of the sequence and weakened interactions between C(25) acetyl with the NH group of F430/514/394, when compared to rifampicin models (Fig. 9). Due to the altered importance of the interactions with amino acids of RNAPs, the binding pattern for kanglemycins can be good starting point at designing alternative antibiotics to rifampicin against multiresistant bacteria strains.

Recent NMR, X-ray, surface plasmon resonance (SPR), and site-directed mutagenesis studies on the affinity of rifamycins to viral targets have revealed the biological secret of rifampicin-like ansamycins bearing C(3)-arm, *i.e.* their binding ability with proteins of poxvirus, as *e.g.* viral trimeric protein D13, in a highly conserved region – F-ring at the central channel.⁶² It is a surprising result since known antibacterial antibiotics are

marked by dots or dashed lines; key amino acids taking part in intermolecular H-bonds are marked by yellow (dark green codes); amino acids building the pocket and involved in hydrophobic interactions are marked by yellow – dark blue codes; the core of ansamycin was marked by the red and the lactam by a dark blue; in the structure of RNAP from *Thermus thermophilus* (c) the σ -finger region is marked by light blue whereas in the structure of RNAP mutant from Mtb (d) the mutated amino acid is marked by pink.



usually not involved in inhibiting viral membranes assembly processes. The activity of rifampicin, which forms complexes with mutants of D13 (K_D s = 19–285 μ M), hinders molecular recognition between viral protein D13 and A17 and prevents virion formation. Hence, this alternative mechanism of action for rifamycins can be used for designing novel-type antiviral agents, based on the structure of naphthalene ansamycins.

The chameleonic nature of naphthalenoid ansamycins is important not only for transportation and binding with the target but can be essential from a bacteria resistance viewpoint, *via* increased/decreased affinity to efflux pump systems.^{44,63,64} The zwitterionization phenomenon for naphthalenoid ansamycins as rifamycins have also impact on their polymorphism in the crystal state, which is a very important problem discussed at potential pharmaceutical applications of this group of macrolactams in the treatment of bacterial infections (including tuberculosis).^{52,65–69}

Thus, the chameleonic features of naphthalenoid ansamycins, concerned with zwitterionic/non-ionic structures and different conformations (sometimes related to stereoisomerism), should be taken into regard at their discussions focused on: the transport, binding with the target (SAR discussions), and bacteria resistance mechanisms.

5. Biosynthesis of naphthalenoid ansamycins

5.1. 1,4-Naphthoquinone- C_{23} ansamycins

The naphthoquinone- C_{23} ansamycin subgroup, of the longest ansa bridge among ansamycins, is relatively structurally homogeneous, and its members differ from each other mainly with regards to the groups present in their ansa chains and at the C(3) position of the core. Novel naphthomycins L (1), M (2) and N (3) were found to be secondary metabolites of *Streptomyces* sp. CS, associated with the tree/shrub tissues of *Manglietia hookeri* from China (Fig. 10).⁷⁰ These natural products, in contrast to the earlier identified naphthomycins, have amine- or sulfur-containing substituents instead of chlorine (naphthomycin A),

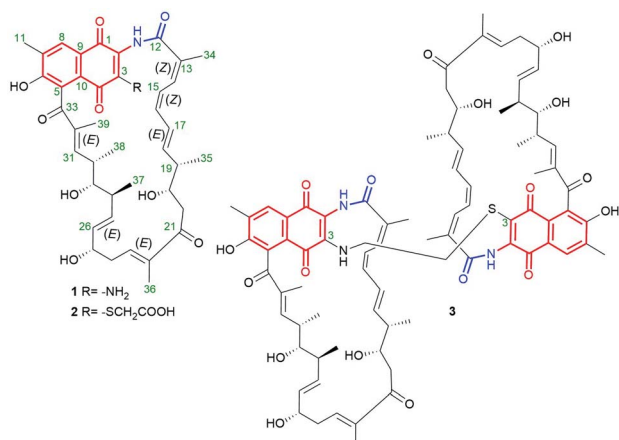


Fig. 10 Structures of new naphthomycins L (1), M (2) and N (3) from *Streptomyces* sp. CS.

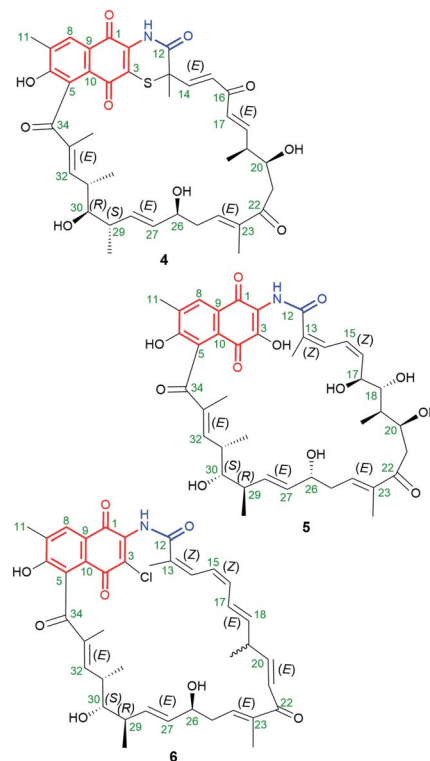


Fig. 11 Structures of new naphthomycins O (4), P (5) and Q (6) obtained from recombinatorial mutants of *Streptomyces* sp. CS/asm21-4.

hydrogen (naphthomycin E) or hydroxyl (naphthomycin D) at C(3) (Fig. 10). Ansamycin 3 is a dimeric-like scaffold bridged by one 2-aminoethanethiol unit. The basic biological precursors of these naphthomycins are: AHBA, malonyl or methylmalonyl units whereas the sulfur source in biosyntheses of 2 and 3 is unknown. Structures of 1–3 were elucidated from HR ESI MS and 1D and 2D NMR, and several unknown genes were found in their biosynthetic clusters.⁷⁰ Among these naphthomycins, only compound 1, bearing amine group at C(3), similarly as naphthomycin A, showed weak antifungal activities against several phytopathogenic fungi *Fusarium*, *Verticillium*, and *Phyricularia* at the lack of anticancer effects.⁷⁰ Natural naphthoquinone- C_{23} naphthomycins O (4), P(5), Q (6), were obtained as secondary metabolites of mutant *Streptomyces* sp. CS/asm21-4 strain, or of non-mutant strain *Streptomyces* sp., associated with Chinese medicinal plant *Maytenus hookeri*, producing also other naphthomycins (Fig. 11).⁷¹ These new naphthomycins 4–6 differ from each other by a number of hydroxyls and double bonds, including their stereochemistries within the ansa chain, and different substituents at C(3) of the naphthoquinone core. Different structures of 4–6, including altered stereochemistry at C(26), C(29) and C(30) as well as the fused/non-fused core with the ansa chain were indicated by 1D and 2D NMR. Compounds 4–6 were quite potent against *E. coli*, *S. typhimurium*, *S. aureus* (MICs = 25 μ g mL⁻¹) and weak active against fungi as *M. grisea* and *F. verticillioides* (MICs = 200 μ g mL⁻¹).⁷¹ Structural diversity of naphthomycins can be helpful in the future at their biological optimizations, especially against Gram-negative bacteria strains, although, the mechanism of their activity is still not revealed.



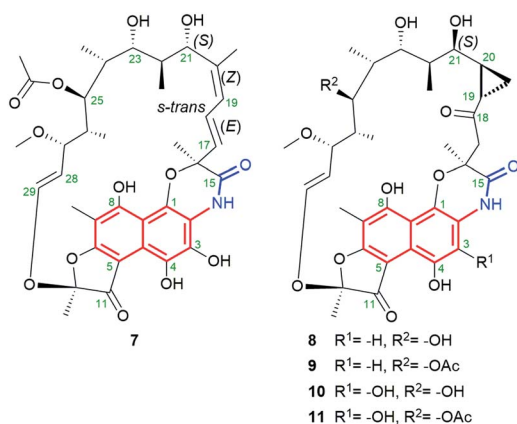


Fig. 12 Structures of new rifamorpholines 7–11.

5.2. Naphthalene-C₁₇ ansamycins

Natural naphthalene-C₁₇ ansamycins, called rifamorpholines A–E (7–11, Fig. 12), are produced by filamentous actinobacteria *Amycolatopsis* sp. Hca4 being associated with locust *Locusta migratoria* (Jiangsu Province, China).⁷² These natural rifamycins possess an extra fused ansa chain with the phenol group of the naphthalene core, as evidenced by X-ray studies of 7.⁷² Rifamorpholines 8–11, apart from the fused core with the ansa chain, possess ketone at C(18) and rare cyclopropane structural motif at C(19)–C(20). The natural production of rifamorpholines is based on the use of the gene cluster 19 having nearly the same organization (in particular a lack of *orf32c* and *orf37c*) as that producing rifamycins in *Amycolatopsis mediterranei* S699.⁷² The formation of demethyl desacetyl rifamycin S and transformation of the lactam *via* 1,6-cyclization yielding imino-quinone core

were postulated as intermediate steps at biosyntheses of 7–11 (Fig. 13). Next, the transformations of the iminoquinone intermediate are realized in two directions: one involving recovering the lactam and epimerization at C(21) (7), and the second recovering the lactam, formation of α,β -unsaturated ketone at C(18) instead of the C(17)–C(18) double bond, followed by epimerization at C(21) and insertion of a one carbon atom at the ansa bridge (8–11).⁷² The mechanism of cyclopropyl formation within the ansa bridge has not been presented up to now. As the result of the above transformations, all rifamorpholines gained C(21S) absolute configuration, as proved by 2D NMR and X-ray studies. Rifamorpholines 7–11 were tested against several bacteria strains, and 8 was found to be the most active of all ansamycins against *M. luteus* at MIC = 0.5 μ M. Importantly, the antibacterial activity of the most active 8 was identical to that of rifampicin against *M. luteus* and *S. aureus* strains, however, the mechanism of action of 8 was not discussed.⁷²

5.3. Naphthalene-1(4H)-one-C₁₇ ansamycins

Ansavaricin D (12, Fig. 14), a new streptovaricin analog lacking the characteristic naphthodioxine motif, has recently been isolated from *Streptomyces* sp. S012, collected from a soil sample (Zhongshan Botanical Garden, Nanjing, China).⁷³ Its structure was determined using NMR and HR-MS methods. Unfortunately, at a concentration below 100 μ M, 12 did not inhibit the secretion of SPI-1 effectors of *Salmonella enterica* serovar Typhimurium. In turn, novel kanglemycins A (13), V1 (14) and V2 (15) were obtained by metagenomic survey of *kng* gene cluster and the use of soil-derived microbiome as well *Amycolatopsis mediterranei* var. *kanglensis*.³² Kanglemycin V2 (15, Fig. 14), possessing methylenedioxy moiety bridging C(4) and C(11), β -O-3,4-O'-methylene digitoxose deoxysugar at C(27), and *gem*-dimethylsuccinic acid portion (K-acid) attached

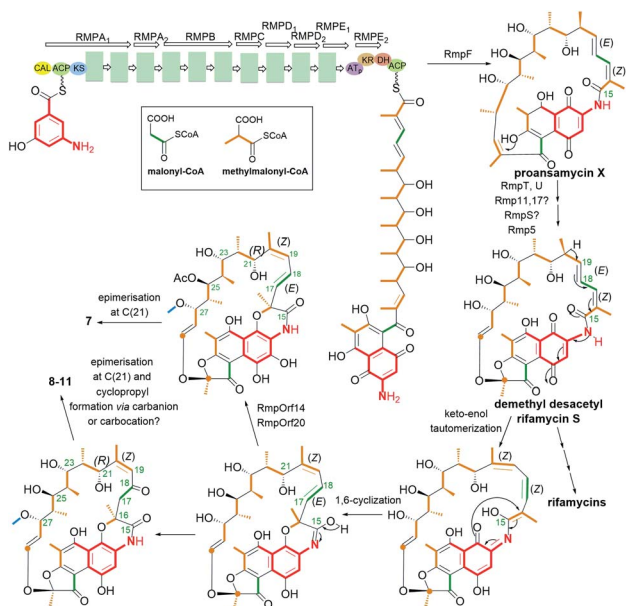


Fig. 13 Biosynthesis of the naphthalene-C₁₇ ansamycins – rifamorpholines containing the naphthalene core fused with the ansa bridge (7–11) and a unique cyclopropyl moiety (8–11).

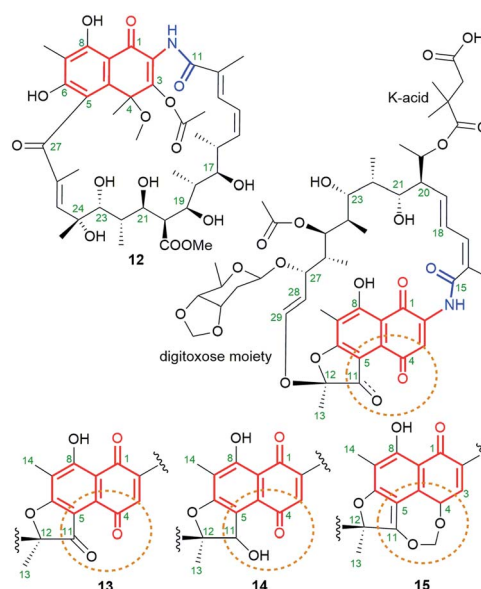


Fig. 14 Structure of new naphthalene-1(4H)-one-C₁₇ ansamycins: ansavaricin D (12) and kanglemycins: A (13), V1 (14), V2 (15).



at C(20), showed an attractive inhibitory activity, toward a rifampicin-resistant RNAP mutant (S447 → L447).³² Kanglemycins A and V1 (13, 14, Fig. 14), lacking methylene-dioxy bridge, are also natural congeners of rifamycins, possessing C(11) ketone or hydroxyl groups, and produced by *Amycolatopsis mediterranei* var. *kanglensis*.³² These natural compounds, shown to inhibit growth of the transcript at one step earlier than that shown for rifampicin. Binding affinity of 13 to RNAPs of rifampicin-resistant mutants (Fig. 9d), e.g. *E. coli*, was higher than rifampicin and the best for RNAP_{βD516V} (IC₅₀ (13) = 39 μg mL⁻¹; IC₅₀ (rifampicin) = 320 μg mL⁻¹). Hence, the greatest advantage of 13 was, the first of all, its potency against rifampicin-resistant *M. tuberculosis* strain and also against several other pathogens as: *S. pyogenes* (MIC = 0.031 μg mL⁻¹), *B. subtilis* (MIC = 1 μg mL⁻¹), *S. aureus* MRSA (MIC = 1 μg mL⁻¹). Different binding models of kanglemycins and rifampicin contributed to slightly lower activities of the former ones than rifampicin against most of tested strains, as e.g. standard *M. tuberculosis* strains (H37Rv; 1192/015).⁷⁴ The explanation of this result is probably lost interactions between the ansamycin and important amino acids of RNAP (as e.g. S447), stabilizing mainly the core at the RNAP pocket, in consequence of the presence of bulky digitoxose deoxysugar portion attached to C(27).

5.4. 1,4-Naphthoquinone-C₁₇ and 4-iminonaphthalen-1(4H)-one-C₁₇ ansamycins

Chaxamycins A–D (16–19, Fig. 15), isolated from the *Streptomyces* sp. C34 strain collected from the soil of Chaxa Lagoon (Atacama Desert, northern Chile), were structurally characterized by 1D and 2D NMR methods.^{75,76} X-ray structure of 16 showed collective intramolecular H-bond system between hydroxyls at C(21), C(23) and C(25), H-bonded phenol at C(8) with the quinone oxygen at C(1) as well as the presence of *trans*-lactam moiety.⁷⁵ The identical stereochemistry of the ansa chains of 16–19 was also independently confirmed by the total synthesis of the C(16)–C(28) portion.⁷⁷ Compound 19, being closely structural related to classical rifamycins, showed attractive potency against *S. aureus* MRSA (MICs = 0.06–0.25 μg mL⁻¹) and *E. coli* (MIC = 1.21 μg mL⁻¹), at almost no inhibitory

activity toward the ATP-ase function of human Hsp90.⁷⁵ Furthermore, studies involving docking and inhibition tests confirmed that Hsp90α is not molecular target for chaxamycin 19, as its mechanism of action is probably closer to that realized for classical rifamycins binding to RNAPs (Fig. 9). Despite potency of 19 was weaker against *S. aureus* than rifampicin, in contrast to the latter it showed attractive potency toward Gram-negative strain.⁷⁵

Novel protostreptovaricins I–V (20–24, Fig. 16) were obtained independently from mutant strain *Nocardia mediterranei* N813 and *S. peccabilis* CCTCC M2017417, collected from a soil sample (Wuhan University campus, China).⁷⁸ These ansamycins, being structural analogous to 16–18, differ from each other by the substituents at the ansa bridge at C(22) and C(26) as well as the substituent at C(6) of the core, as indicated by MS and NMR. The biosynthetic route of protostreptovaricins formation shares a common intermediate (prostrestreptovaricin) with the pathway yielding streptovaricins (Fig. 16). AntiSMASH analysis showed activity of cytochromes P450s, especially at hydroxylation processes being post-PKS tailoring stage of protostreptovaricins and streptovaricins. Notably visible role of P450 is at the oxidation of the methyl at C(25) into methylene hydroxyl group, which subsequently undergoes oxidation to the carboxylic group taking part in lactonization with the hydroxyl at C(22), as for streptovaricin F (25, Fig. 16).⁷⁸ The precursor of C(3) methyl unit is SAM, similarly as C(12) carbon atom of dioxamethylene bridge of streptovaricins, whereby the formation of this bridge is catalyzed by type-I O-methyl transferase (StvM1).⁷⁹ The

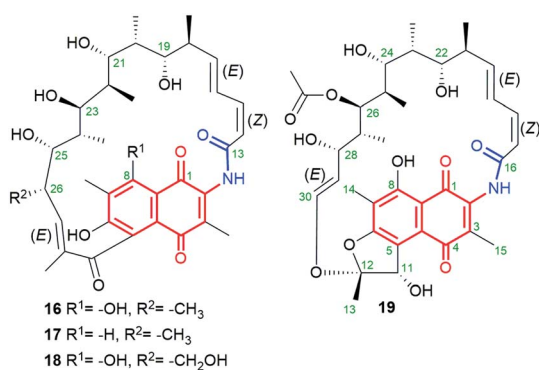


Fig. 15 Structure of new naphthoquinone-C₁₇ ansamycins called chaxamycins A (16), B (17), C (18) and D (19), isolated from soil-derived *Streptomyces* sp. C34 strain.

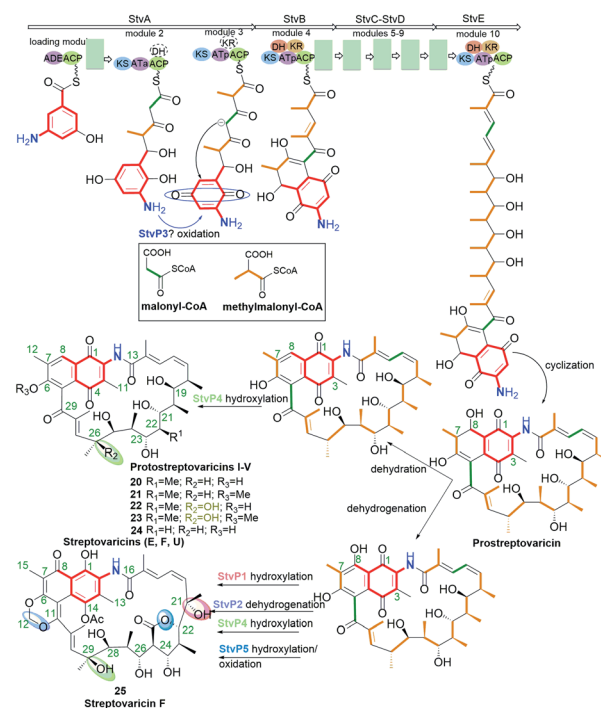


Fig. 16 Common main biosynthetic route, via prostrestreptovaricin intermediate, leading to natural protostreptovaricins I–V (20–24) and streptovaricins with the key role of cytochrome P450s at post-PKS I tailoring of the structure (StvP1–P5, highlighted by colors).

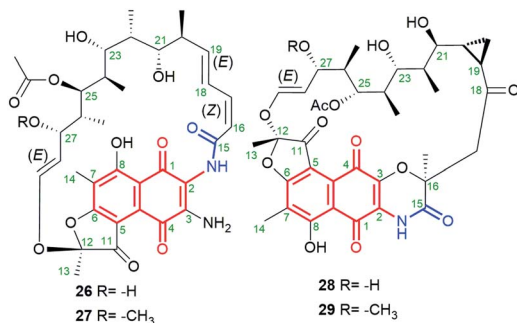


Fig. 17 Structure of new naphthoquinone- C_{17} ansamycins called aminorifamycins (26, 27) and sporolactams A (28) and B (29), isolated from Northeastern-Pacific marine-sediments bacteria of *Micromonospora* sp.

presence of hydroxyl group at C(26) is essential for enhanced antibacterial potencies of protostreptovaricins against *S. aureus* strains (standard and MRSA, MICs = 16 $\mu\text{g mL}^{-1}$), which was even better than those of some obtained streptovaricins.⁷⁸ However, the most potent of all studied new ansamycin derivatives was streptovaricin F (25, Fig. 16; MIC = 0.125 $\mu\text{g mL}^{-1}$), containing a tertiary hydroxyl group in the ansa chain at the analogous site [at C(29)] as that of the most active protostreptovaricin III (22), as well as one extra tertiary hydroxyl at C(21).⁷⁸

Novel natural aminorifamycins 26 and 27 (Fig. 17), possessing an amine group at C(3), were isolated from cultures of *Micromonospora* sp. RJA4480.⁸⁰ Stereochemistry of aminorifamycins was determined by X-ray structure of 26 as well as by 1D and 2D NMR studies both of them. Sporolactams A and B (28 and 29, Fig. 17), containing the core fused with the ansa bridge by C(3)-O-C(16) ether moiety and cyclopropane motif within the ansa bridge, were found also to be secondary metabolites of *Micromonospora* sp.⁸⁰ Sporolactams 28 and 29 showed very good antibacterial potency, especially against Gram-negative *E. coli* (MIC₉₀s = 0.4–1.8 μM) and Gram-positive *M. tuberculosis* (MIC₉₀s = 0.8–0.06 μM). However, ansamycins 26 and 27, bearing C(3)-amine substituent, showed the most attractive potency of all against *S. aureus* MRSA, *E. coli* and *M. tuberculosis* (standard and in macrophages) in the range far beyond expectations (MIC₉₀s = 0.0001–0.7 μM), making them a very attractive starting point to drug design against hard to control pathogens. Such a biological outcome clearly shows that for very attractive potency relatively high flexibility of the ansa bridge is desired.⁸⁰

New 11,12-*seco*-rifamycin W analogs 30–32 (Fig. 18), originating from common biosynthetic precursor – rifamycin W, were found in an agar fermentation extract of *Amycolatopsis mediterranei* S699.⁸¹ These naphthoquinone- C_{17} ansamycins possess included oxygen heterocycles as oxirane, tetrahydrofuran or tetrahydropyran into the ansa bridge. Such type structures of 30–32 were indicated by HR-MS and 1D and 2D NMR. It should be mentioned that 30 and 32 possess (13*Z*)-configuration, in contrast to 31 of (13*E*)-configuration (Fig. 18). Biosynthesis of 31, containing both the lactam and the lactone

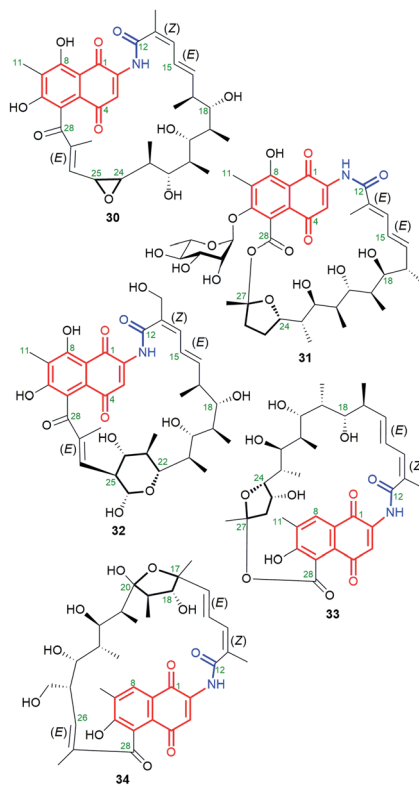


Fig. 18 Structure of new natural naphthoquinone- C_{17} ansamycins (30; 31 named rifamycin B, 33), derivatives of 11,12-*seco*-rifamycin W, as well as 8-deoxy derivatives obtained from genetic modified *A. mediterranei* ΔrifT strain (33, 34), all bearing oxygen heterocycles incorporated into the ansa bridge.

portions within the ansa bridge, first requires glycosylation with α -rhamnose saccharide at C(6) and next the Baeyer-Villiger oxidation (C27–C28 oxidative cleavage) and an intramolecular transesterification with one of hydroxyls of the ansa bridge. Biosyntheses of 30 and 32 occur directly from rifamycin W, however, detailed mechanisms were not displayed. Ansamycins of greater heterocyclic ring motifs decorating their ansa chains, as 31 and 32, exhibited strong inhibitory activities toward DNA topoisomerase (Topo I) and Topo II α at 50 μM , in contrast to weak effects noted for the other earlier-obtained 11,12-*seco*-rifamycins.⁸¹ In turn, the limited flexibility of the ansa bridges belonging to 30–32 contributed to the lack of typical for rifamycins antibacterial properties. A much higher plasticity in the production of structurally diverse rifamycin-like natural products was presented *via* deleting the *rifT* gene from *Amycolatopsis mediterranei* S699.⁸² This approach enabled to obtaining of 8-deoxy ansamycins as *e.g.* 33 or 34 (Fig. 18), bearing tetrahydrofuran moieties at different positions of the ansa bridge. Compound 33, lacking C(8) phenol and possessing tetrahydrofuran ring next to the lactone, in contrast to 30–32 and 34, showed moderate antibacterial potency against *S. aureus* (MIC = 10 $\mu\text{g mL}^{-1}$), and was inactive in cancer cells KG1.⁸²



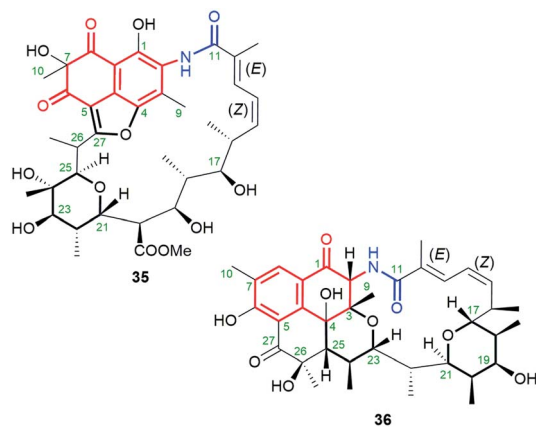


Fig. 19 Structure of new natural naphthalene-1,3(2H,4H)-dione- C_{17} (35, ansavaricin C) and dihydronaphthalen-1(2H)-one- C_{17} (36, ansavaricin E) ansamycins, isolated from botanical garden soil-derived (Nanjing, China) *Streptomyces* sp. S012 bacteria strain.

5.5. Naphthalene-1,3(2H,4H)-dione- C_{17} and 3,4-dihydronaphthalen-1(2H)-one- C_{17} ansamycins

Naphthalenoid ansamycins containing the saturated and deoxygenated quinone core are rather rare in nature. This group ansamycins includes ansavaricins C (35, Fig. 19) and E (36, Fig. 19), produced by *Streptomyces* sp. S012.⁷³ Structures of these novel secondary metabolites, possessing identical 12E and 14Z configurations and decorating with tetrahydropyran and furan rings (35) or the two tetrahydropyran rings (36), were indicated by 1D and 2D NMR and HR-MS in negative and positive ion detection modes. The absolute configurations of C(7) and C(26) stereogenic centers in 35 were not proposed. The double fusion of the core at the C(23) and C(25) positions of the ansa chain was apparently responsible for the greater inhibitory activities observed for 36 than for 35 toward the secretion of SPI-1 effectors of *Salmonella enterica* serovar Typhimurium. Inhibitory activity of 36 at concentrations below 100 μ M in an anti-T3SS assay can be useful for its applications against resistant pathogens.⁷³ Ansavaricin J, showing nearly identical structure as 36, *i.e.* enriched by one phenol group at C(8) of the core, was isolated from the genetically modified *Streptomyces spectabilis* CCTCC M2017417 strain (Δ stvP5 cytochrome P450 gene deletion) producing also streptovaricins.⁸³ This compound obtained by genetic manipulation, which structure was proved by 1D and 2D NMR and HR-MS, showed only limited antibacterial effect toward *S. aureus* MRSA strains. In turn, much more biological attractive were ansavaricins F–I, named as “polyketides of ansamycin class”.⁸⁴ This description is, however, not precise because each of these ansavaricins F–I possesses cleaved the ansa chain and, hence these polyketides can be only precursors or metabolites of ansamycins (called ansavaricins). Open-ansa chain ansavaricins were potent inhibitors toward human DNA topoisomerases (Topo I and Topo II α) and one of them, *i.e.* ansavaricin H exhibited selectively cytotoxicity toward HeLa and MDA-MB-453 cancer cells at IC_{50} s < 50 μ M.⁸⁴ It is a surprising result that for ansavaricins belonging to classical ansamycins

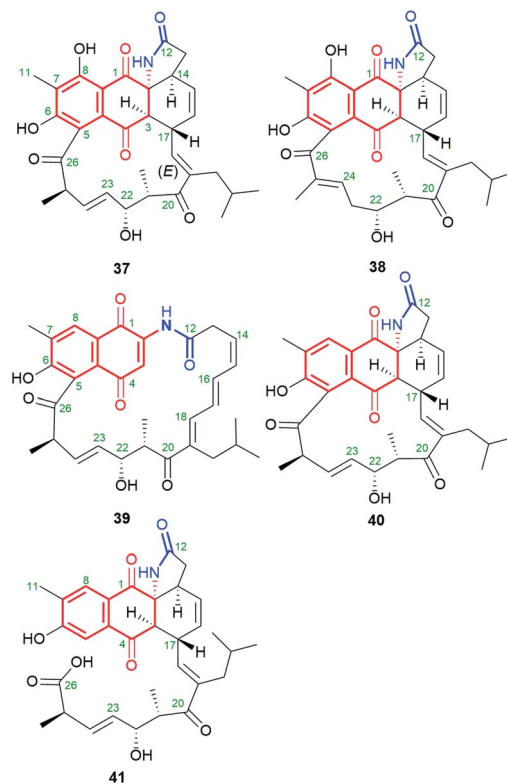


Fig. 20 Structure of new natural naphthalenoid- C_{15} ansaeomycins (37–40) and polyketide 41 with the cleaved the ansa bridge structure.

with the ansa bridge the biological potency was lower or none, in contrast to their open-ansa chain congeners.

5.6. Naphthalenoid ansamycins of C_{15} ansa chains

The naphthoquinone- C_{15} ansamycins, similarly to other ansamycins, are solely secondary metabolites produced by bacteria. The most number and structurally diverse group of ansamycins with C_{15} ansa chains (Fig. 2) are recently-discovered ansaeomycins (exemplary 37–40; Fig. 20).⁸⁵ These ansaeomycins were obtained through the expression of the *asm* gene cluster (cryptic red-ox gene) in a mutant of *Streptomyces seoulensis*. The activation of berberine bridge enzyme (BBE), *i.e.* oxidase *AsmF*, dictated the formation of natural ansamycins bearing C(8) phenol (37, 38, Fig. 21) whereas deletion of the *AsmF* gene contributed to a great structural variety of ansamycins structures, lacking C(8) phenol group (examples 39–41, Fig. 21).⁸⁵ The key steps in the natural production of structural diverse ansaeomycins are isomerization (double bond shifts) and conformational change as well as Diels–Alder reactions of the triple unsaturated moiety near the lactam (Fig. 21). Formation of polyketide 41 of cleaved the ansa bridge was postulated as a simple *seco*-dehydration process. All novel ansaeomycins were characterized by HR-MS, 1D and 2D NMR in detail. Among all ansaeomycins obtained, only those containing C(8) phenol (37 and 38) showed moderate anticancer effects in cell line K562. Surprisingly, antibacterial screening of novel ansaeomycin



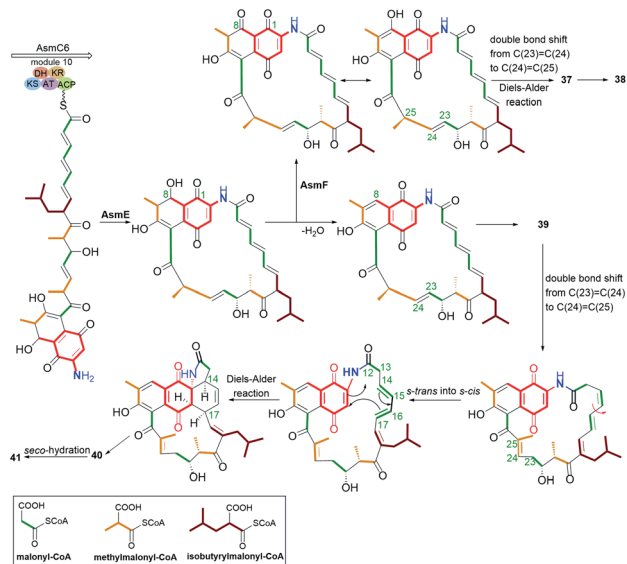


Fig. 21 Biosynthetic pathway of natural ansaeomycins (37–40) and polyketide 41 with the cleaved the ansa bridge structure.

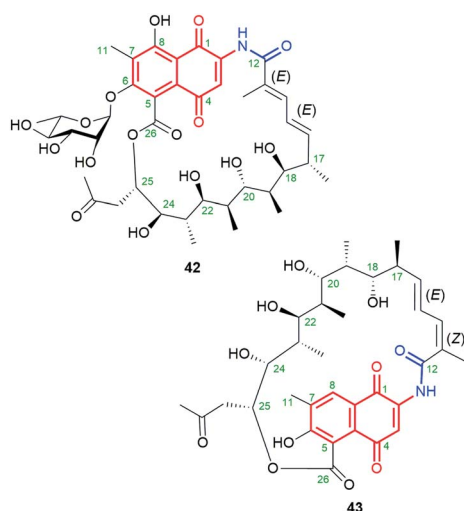


Fig. 22 Structure of new natural naphthoquinone- C_{15} ansamycins 42 (ansmitocinoside A) containing the lactone incorporated into the ansa bridge and L-rhamnose at the core and 43 with the unique propan-2-one motif attached at C(25) of the ansa chain.

scaffolds indicated potency of solely an open chain polyketide 41 toward *S. aureus* at MIC = 64 μM .⁸⁵ These results revealed that the polyketides with the cleaved ansa chain could be also an interesting starting point toward antibacterial potency optimization and that the limited flexibility of the ansa bridge abolishes the antibacterial properties of ansaeomycins.

Natural product 42 (rifamycininoside A, Fig. 22), belonging to naphthoquinone- C_{15} ansamycins, was isolated from *Amycolatopsis mediterranei* S699.⁸¹ It was postulated that 42 is formed on the common rifamycin biosynthetic pathway but with an extra oxidative decarboxylation and typical Baeyer–Villiger oxidation with C(25)–C(26) bond cleavage.⁸¹ The pattern of the

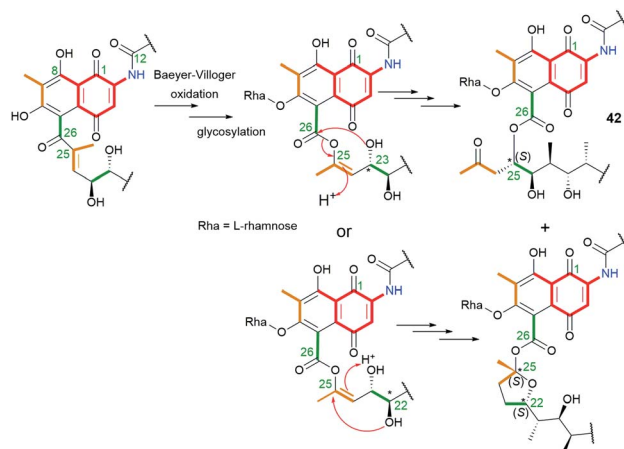


Fig. 23 Biosynthetic rearrangements after Baeyer–Villiger oxidations: on the top, leading to exclusion of the propan-2-one moiety from the ansa chain backbone, as for ansamycin 42 (ansmitocinoside A); on the bottom, leading to formation of tetrahydrofuran moiety, present also in ansamycins of longer C_{17} ansa chains (e.g. product 31).

cleavage within the ansa bridge, however, differs from those known contributing to formation divergolides R and S or hygrocins I and J,^{86,87} since Baeyer–Villiger oxidation is followed by rearrangement as shown in Fig. 23. As a result of this rearrangement an untypical propan-2-one motif, instead of usually occurring acetate, is being inserted at the ansa bridge structure.⁸¹ Subsequent glycosylation at C(6), followed by the rearrangement, yielded 42. It is worth noting, that the other type an intramolecular rearrangement with the nucleophilic attack of C(22) hydroxyl on the C(25) carbon (Fig. 23) yielded other ansamycins with tetrahydrofuran moiety, structurally analogous to those of longer C_{17} ansa chains (see e.g. 31, Fig. 18). Compound 42, showed good inhibitory activities toward the topoisomerases Topo I and Topo II α .⁸¹ The novel, untypical 8-deoxy rifamycin of the conserved 11,12-seco-rifamycin W scaffold (43, Fig. 22), classified to naphthoquinone- C_{15} ansamycins, has been obtained using the *Amycolatopsis mediterranei* S699 ΔrifT mutant strain.⁸² The structure of 43, is lacking one phenol group and the saccharide at the core as well as 13Z configuration is present, in contrast to 42. The presence of the propan-2-one motif in structures of 42 and 43 were proved using 1D and 2D NMR methods. Biological studies of 43 revealed moderate antibacterial activity against *Staphylococcus aureus* ATCC 25923 at MIC = 20 $\mu\text{g mL}^{-1}$.⁸² The limited potency of 43 can be a result of the lack of O(8)H phenol group in the core, important at binding with bacterial RNAPs (interaction with S411, Fig. 9), however, the binding pose of this compound at RNAPs has not been revealed yet.

5.7. Naphthalenoid- C_{13} ansamycins

Natural occurring ansamycins, consisting of an oxidized/reduced naphthalene moiety and bearing relatively short C_{13} ansa chain linked at several sites to the core are called ansalactams A (44), B (45), and D (46) (Fig. 24).^{88–90} Ansalactam 44 was extracted from marine sediment bacterium *Streptomyces* sp.



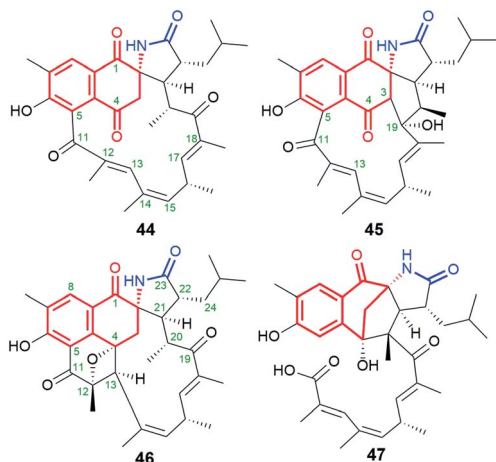


Fig. 24 Natural ansalactams A (44), B (45) and D (46), having C₁₃ ansa chain fused with the core as well as ansalactam C (47) containing cleaved the ansa bridge.

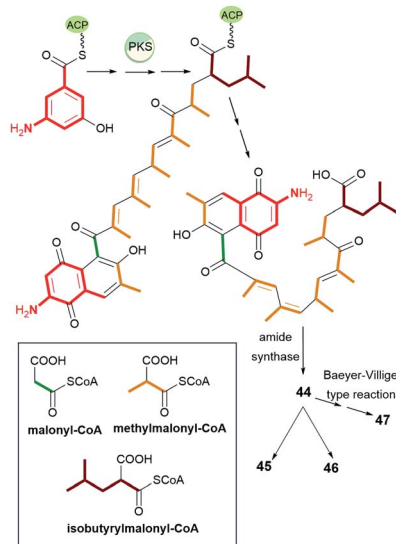


Fig. 25 The biosynthetic route leading to ansalactams A (44), B (45) and D (46), having C₁₃ ansa chain fused with the core and ansalactam C (47) with the cleaved ansa bridge.

CHN-189 and its stereochemistry was determined by 2D NMR and X-ray analysis of its reduced congener containing hydroxyls at C(1) and C(4).⁸⁸ Biosynthetic isotopic studies of ansalactams indicated their main precursors as: AHBA, one malonyl, six methylmalonyl and one isobutyrylmalonyl units (Fig. 25).^{88,89} The inclusion of the isobutyrylmalonyl unit into the ansalactam scaffolds is enabled by activity of β -ketoacyl-ACP-synthase, dehydrogenase and crotonyl-CoA carboxylase producing isobutyrylmalonyl-CoA in *Streptomyces* sp. CHN-189. The biosynthetic plasticity of 44–46 is well reflected in the ability of 44 to be converted into 45 or 46 via an extra cyclization involving the ansa chain and the core.⁸⁹ In addition to 45 and 46, another natural secondary metabolite, an ansalactam C (47, Fig. 24) with the cleaved ansa chain, is also produced by marine sediment

bacterium *Streptomyces* sp. CHN-189.⁹⁰ This metabolite obtained probably due to the Baeyer-Villiger type reaction from 44 can be further used in macrolactonization due to the possible approaching phenol by the carboxylic group. The above ansalactams 44–47 showed no anticancer effects but most of them were weak active against methicillin-resistant *S. aureus* at MICs = 31 $\mu\text{g mL}^{-1}$ (45, 46) and MIC = 62 $\mu\text{g mL}^{-1}$ (47). Thus, decreased flexibility of the ansa bridge or its cleavage contributed to limited antibacterial effects when compared to those of the most active rifamycins.

Another small subgroup of naphthoquinone C₁₃-ansa chain ansamycins includes those lacking the C(8) phenol and containing the lactone within the ansa chain and dihydro- or tetrahydro-2H-azepin-2-one ring next to the core (48–51, Fig. 26). Divergolide C (48), found in endophytic *Streptomyces* sp. associated with the mangrove tree *Bruguiera gymnorrhiza*,^{91,92} with its recently revised (15*R*) configuration, possesses structure identical to that of differently named “olimycin B”, isolated from the deep-sea bacterium *Streptomyces olivaceus*.⁹³ Bioinformatic analysis indicated that Baeyer-Villiger monooxygenase (*DivO*) contributes to the presence of the lactone moiety in the ansa chain whereas an alternative exclusion of C(17) tail from the main backbone of the macrocyclic system of divergolides is also a result of the activity of P450-dependent oxidoreductase *DivQ*. Thus, these enzymes are essential at shaping the naphthalenoid ansamycin scaffolds of great structural variety concerning different ansa chain lengths, isolated from *Streptomyces* sp. W112 and bacterial endophytes (Fig. 27).^{87,94,95} Divergolide J (49, Fig. 26), bearing a partial saturated azepinone ring, has been

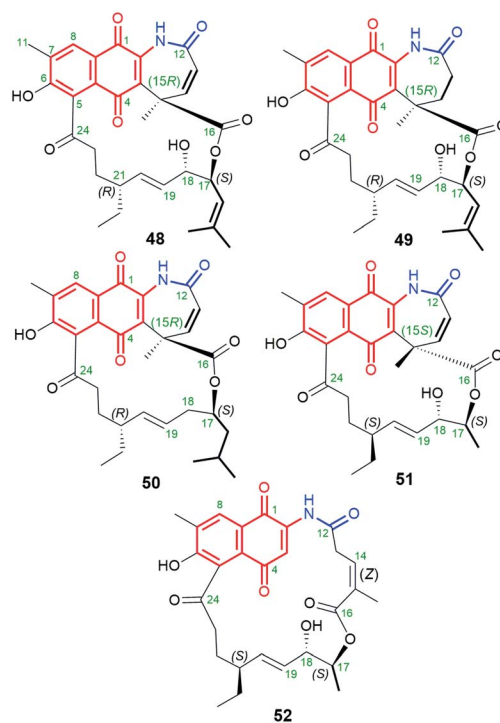


Fig. 26 Naphthoquinone-C₁₃ ansamycins, the so-called: divergolide C (48; also named-olimycin B), divergolide J (49), as well as olimycin A (50) and structurally related to them hygrocin B (51) and A (52).



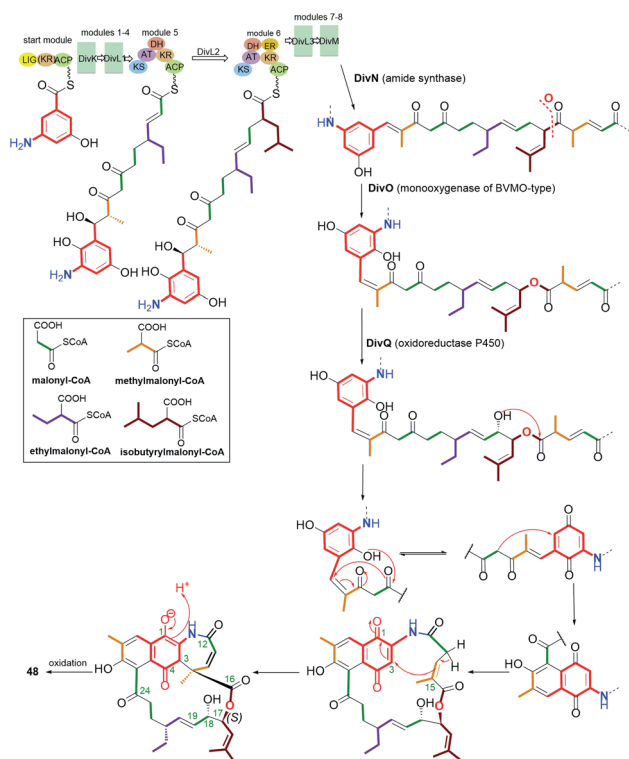


Fig. 27 Biosynthesis of divergolide C (**48**, olimycin B) with the pivotal role of monooxygenase *DivO* and oxidoreductase *DivQ*.

obtained from the same source and on the similar biosynthetic pathway as those of **48**. In turn, olimycin A (**50**, Fig. 26), isolated from *Streptomyces olivaceus* SCSIO T05, has the absolute configuration (15*R*), similarly as **48** and **49**, and contains, in contrast to them, the saturated C(17) tail.^{93,96} Interestingly, a biosynthetic insertion of an O atom into the scaffold of olimycin A is realized by luciferase-type monooxygenase *OvmM*, an analogous to those producing divergolides (e.g. **49**), and acting before amide synthase *OvmN*.⁹⁶ Inactivation of *OvmM* lead to production of an open-chain olimycins C and D.⁹⁶ Structural close to the above ansamycins **48–50**, are hygrocins A and B (**52**, **51**; Fig. 26), isolated from the *Streptomyces hygrosopicus* ATCC25293 and *Streptomyces* sp. LZ35.⁹⁷ Structure of hygrocine B (**51**), of the absolute configuration (15*S*), which is opposite relative to those of **48–50**, was evidenced by 1D and 2D NMR. The biosynthetic pathway of **51** and **52** was based on the formation of the common intermediate prohygrocin and requires the presence of the redox genes *hgc2* and *hgc4* of the *hgc* cluster.⁹⁷ The mutant strains $\Delta hgc1A$, $\Delta hgc1L2$ and $\Delta hgc5$ and $\Delta hgc11$ were shown to be able to produce structurally diverse compounds **51–53** (Fig. 28 and 29). Classes I, II and O Baeyer–Villiger monooxygenases (BVMOs) play a key role in the incorporation of the lactone in the natural scaffolds, whereas oxygenation of prohygrocin is realized by luciferase-like homologue, namely monooxygenase *Hgc3* (Fig. 28), of functionality similar to that producing olimycins (**48** and **50**).⁹⁷ Compound **49** (Fig. 26) was shown to be inactive toward fungi and bacteria.⁹⁸ Hygrocins **51** and **52** (Fig. 26) exhibit moderate potency against *Aspergillus fumigatus*, whereas the latter was

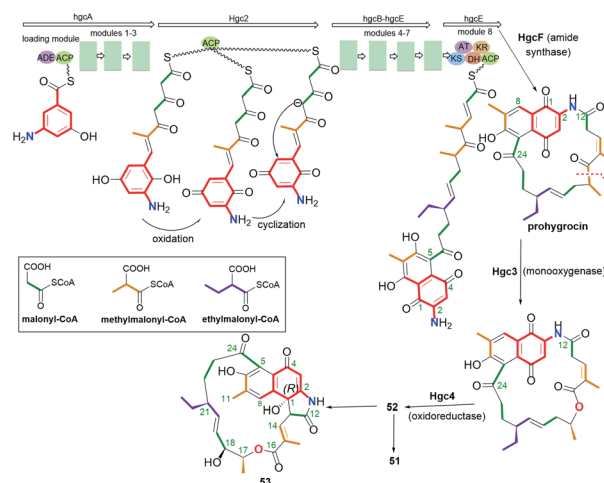


Fig. 28 Biosynthesis of naphthoquinone- C_{13} hygrocins A (**52**), B (**51**) and their structural analog **53** (hygrocin E) of untypically fused the ansa chain with the core at C(1).

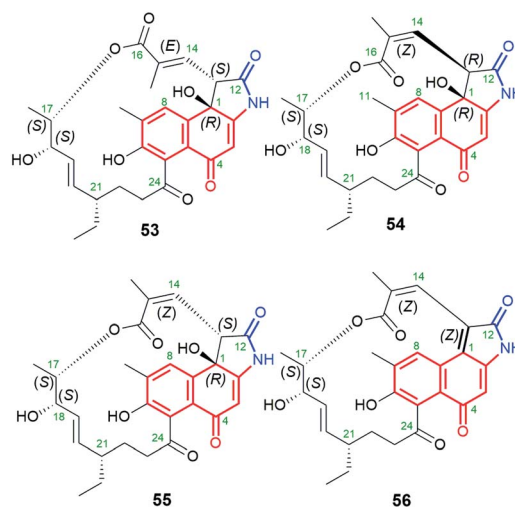


Fig. 29 Diastereomeric naphthalene-1(4*H*)-one- C_{13} hygrocins E (**53**), C (**54**), D (**55**), H (**56**), containing identical C(17)-methyl substituent within the ansa chain.

also active toward *S. pneumoniae*.⁹⁹ This result showed clearly that antibacterial potency is related to the not disturbed flexibility of the ansa bridge. In contrast to hygrocins **51** and **52**, hygrocine E (**53**, Fig. 29) bearing fused core at C(1) with the ansa chain, were tested toward anticancer potency in MDA-MB-231 and PC3 cell lines. These tests did not show, however, the activity up to $IC_{50} = 10 \mu M$.⁹⁷

Diastereomeric ansamycins of naphthalene-1(4*H*)-one- C_{13} group and decorated with 5-membered lactone moiety, similar as **53**, are hygrocins C (**54**), D (**55**) and H (**56**), produced by *gdmA*-disrupted *Streptomyces* sp. LZ35.^{86,100} These hygrocins are structural different regarding the absolute configuration of the stereogenic center at C(13) and the double bond C(14)=C(15), influencing the ansa chain conformation rigidity (Fig. 29).^{86,100} Structures of **54–56** were elucidated by HR-MS, 1D and 2D NMR

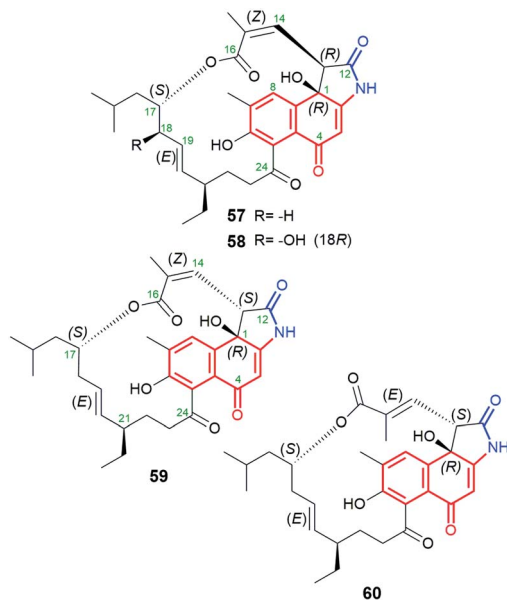


Fig. 30 Naphthalene-1(4H)-one-C₁₃ stereoisomeric divergolides T (57), U (58), V (59) and W (60), containing identical C(17)-isobutyl substituent within the ansa chain, being produced by marine-derived microorganisms, i.e. *Streptomyces* sp. KFD18.

as well as X-ray studies (structure of 54).^{86,100} It should be mentioned that hygrocin H (56), possesses an extra double bond at C(1), as result of dehydration process, in contrast to 53–55. Hygrocin H (56) showed a quite high cytotoxic potency in MDA-MB-231, PC3 and HeLa cancer cells (IC₅₀s = 0.5–5.0 μM), unlike to 53.^{86,100} This result suggested the impact of *E/Z* configuration at C(14) on anticancer activity, whereby the mechanism of this activity has not been revealed yet. This SAR among hygrocin is concerned probably with the fact that *E*-configuration limits flexibility of the ansa bridge.

A very similar in structure to the above hygrocin, are divergolides T (57), U (58), V (59) and W (60, Fig. 30), containing also C₁₃-ansa chains and isolated from marine-derived microorganisms *Streptomyces* sp. KFD18.¹⁰¹ Each of these compounds has an isobutyl group attached at C(17), whereas 59 and 60 have an inverted absolute configuration at C(13) relative to that of 57 and 58. Structures of 57–60 were confirmed by HR-MS, 1D and 2D NMR of them as well as by X-ray of 57. Interestingly, only 57 and 59 having (14Z) configuration, showed good potency in K562, A549, SGC-7901, and HeLa cancer cell lines, at IC₅₀s = 2.8–20.9 μM, and apoptosis-inducing activity.¹⁰¹ This result is in agreement with that obtained for the above-discussed hygrocin 53–56, where only those of the absolute configuration (14Z), i.e. of more conformational labile the ansa bridge, showed notable anticancer potency.¹⁰¹

5.8. Naphthalenoid-C₁₂ ansamycins

Polyketides, containing naphthalenoid core and C₁₂ ansa chain, are shown in Fig. 31. Divergolid D (61, Fig. 31)⁹¹ and hygrocin F (63, Fig. 31)⁹⁹ have common naphthalen-1(4H)-one core, whereas divergolid I (62, Fig. 31)⁹⁸ and hygrocin G (64, Fig. 31)⁹⁹ belong to the naphthoquinone-C₁₂ ansamycins (Fig. 2). All of them have

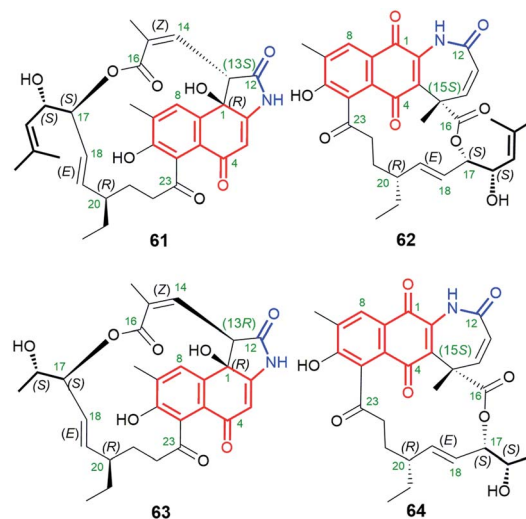


Fig. 31 Members of naphthalenoid-C₁₂ ansamycins group: divergolid D (61) and I (62) as well as hygrocin F (63) and G (64) of naphthalen-1(4H)-one or 1,4-naphthoquinone cores.

the lactone incorporated into the ansa bridge at C(16), enzymatically formed by a monooxygenases of BVMO-type. The stereochemistry of divergolid 62 has been revised from *R* into *S* at C(15), as evidenced by its total synthesis, 1D and 2D NMR, and comparison between X-ray structures and CD spectra of these ansamycins.^{93,102} It is worth to underlining that, apart from the above structural differences, compounds 61 and 63 also differ in the presence of opposite configurations at C(13). These hygrocin and divergolid possessing 2-propanol or 3-methylbut-2-en-1-ol C(17)-tails of identical stereochemistries were obtained by: the large-scale fermentation of *Streptomyces* sp. HKI0576 and HKI0595 associated with the mangrove tree *Bruguiera gymnorhiza* (61, 62),⁹¹ overexpression of *div8* in *Streptomyces* sp. W112 (61)⁸⁷ and isolation from *gdmA*I-disrupted *Streptomyces* sp. LZ35 (62, 63 and 64).¹⁰⁰ The biosynthetic pathway of these naphthalenoid-C₁₂ ansamycins, despite suspected to be analogous to those of other hygrocin or divergolid of naphthalenoid-C₁₃ group, was not however revealed or discussed. Compounds 63 and 61 (IC₅₀s ranging from 1 to 2 μM) showed high anticancer activities in MDA-MB-231, PC3, as well as LXFA629L, PANC-1, RXF486L and Saos-2 cancer cells.^{91,100} Neither ansamycin 62 nor 64 was found to exhibit essential antifungal or anticancer potencies. Among tested divergolid with C₁₂ and C₁₃ chains (divergolid I–N), divergolid I (62) of shorter C₁₂ ansa chain showed the strongest antibacterial activity (the best GIZs > 13 mm) against *B. subtilis*, *S. aureus* (MRSA), *E. faecalis* (vancomycin-resistant) and *Mycobacterium vaccae*.⁹⁸ Thus, the profile of biological potency of C₁₂ hygrocin and divergolid seems to be dependent on the type of the core, i.e. 1,4-naphthoquinone – antibacterial and naphthalen-1(4H)-one – anticancer.

5.9. Naphthalenoid ansamycins of the shortest C₁₁ ansa chains

The naphthoquinone core is a predominant central motif in natural and the rarest ansamycin scaffolds of the shortest C₁₁



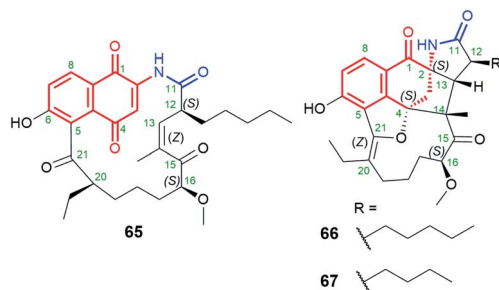


Fig. 32 Structures of neoansamycins A–C (65–67), produced by *nam7*-disrupted *Streptomyces* sp. LZ35 SR201*nam1OE* strain.

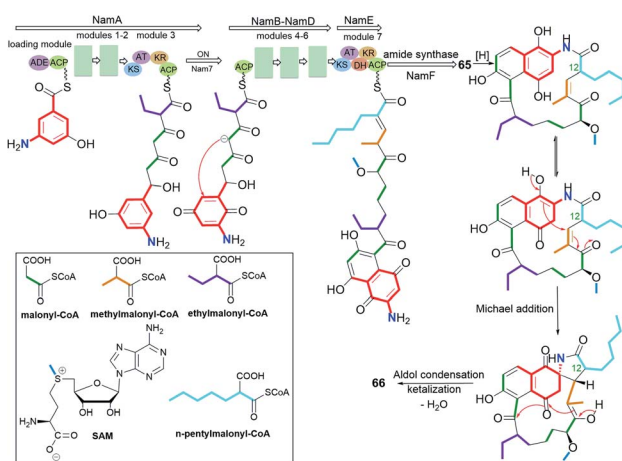


Fig. 33 Biosynthesis of neoansamycins A and B (65 and 66) by *nam7*-disrupted *Streptomyces* sp. LZ35 SR201*nam1OE* strain.

ansa chains of all known naphthalenoid ansamycins. Neoansamycin A (65, Fig. 32) is a type of natural 1,4-naphthoquinone- C_{11} ansamycin, whereas neoansamycins B and C (66 and 67, respectively; Fig. 32), formed from 65, belong to 3,4-dihydronaphthalen-1(2*H*)-one- C_{11} ansamycins (Fig. 33).^{103,104} The X-ray structure of 66 provided evidence for stereochemistry of this-type of novel neoansamycins.¹⁰⁴ These natural products were obtained through the activation of a cryptic ansamycin gene cluster in *Streptomyces* sp. LZ35.¹⁰⁴ Experiments with a mutant strain *Streptomyces* sp. LZ35, revealed that *nam7* hydroxylase is the key enzyme in switching between biosynthetic routes leading to the alternative formation of benzenoid ansamycins or neoansamycins in *Streptomyces* SR201*nam1OE* Δ *nam7*.^{103,104} The long alkyl chains at C(12) in neoansamycins A–C structures suggested that PKS module 7 is able to incorporate carbon skeleton not only of *n*-butylmalonyl-CoA but also *n*-pentylmalonyl-CoA unit (Fig. 33). Antiproliferative tests indicated clearly that potency of 66, bearing the ansa chain of almost no flexibility, is lower than that of 65 (IC_{50} (SW480) = 6.7 μ M). Surprisingly, the antibacterial screening of ansamycins revealed that both 65 and 66 showed potent activities towards *S. aureus* ATCC 25923 at low MICs = 3.125 μ g mL⁻¹.¹⁰⁴ This result suggested that the antibacterial and anticancer mechanisms of these compounds are totally different.

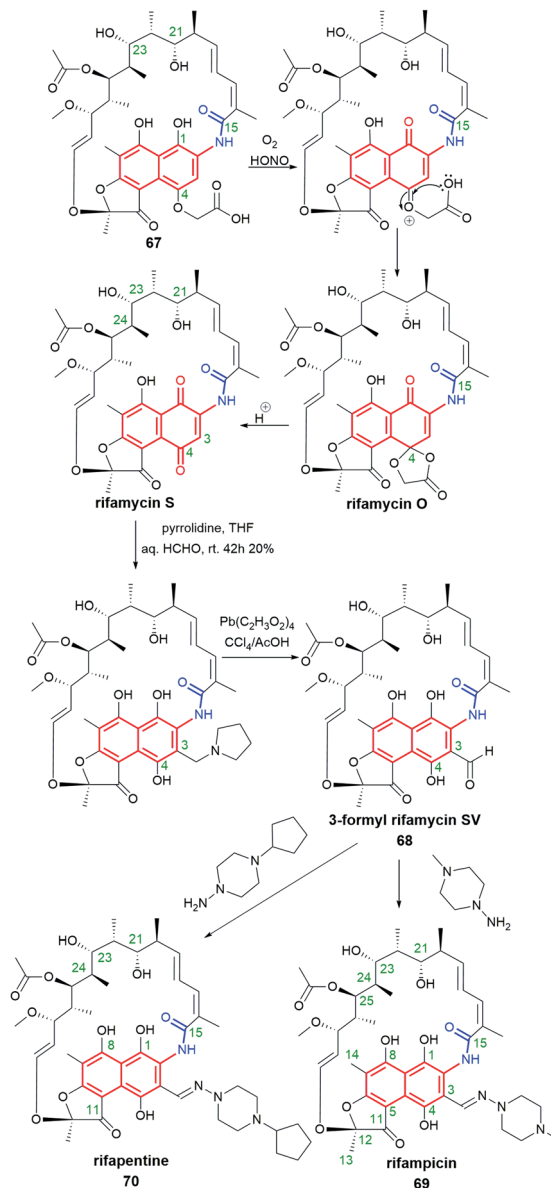


Fig. 34 Semisynthesis of potent antibiotics, rifampicin (69) and rifapentine (70), from natural rifamycin B (67) via intermediate 3-formylrifamycin SV (68), developed by Sensi *et al.*

6. Semisynthetic transformations of naphthalenoid ansamycins and their biological potency

Despite many natural-occurring naphthalenoid ansamycins revealed important antibacterial or other type activities, among them are also those possessing relatively weak biological activities, whose semisynthetic modifications help to improve their biological relevance. Here an excellent example is natural rifamycin B (67, Fig. 34), of weak antibacterial potency, which subjected to semisynthetic structural alterations *via* structure of 3-formyl rifamycin SV (68), yielded one of the best known naphthalene- C_{17} semisynthetic antitubercular antibiotic – rifampicin



(in US called also rifampin, **69**, Fig. 34).^{3,105} The other reason why semisynthetic transformations are desired among the naphthalenoid rifamycins is concerned with the resistance of bacterial strains, including multidrug-resistance (MDR) to semisynthetic antibiotics as **69** or its congeners.^{10,106} Resistance of different bacteria to naphthalenoid-C₁₇ rifamycins is realized *via e.g.* mutations of single amino acid residues of the target, ribosylation or glucosylation of hydroxyl groups of ansamycin, modification/hydrolysis of the C(3) arm of ansamycin, or *via* activation of the efflux pump systems in bacterial cells.^{107–109} In turn, modifications of rifamycin structures and/or the target performed by bacteria prevent stable binding of these ansamycins to the allosteric site of RNAPs, limits the permeability of the antibiotic through the natural cell barriers, or enhance affinity to efflux pump systems thereby decreasing the degree of accumulation of the antibiotic in bacterial cells.^{55,110} Recently it has been shown, that the loss of **69** antibacterial activity is a result of *N*-hydroxylation performed by flavoproteins such as rifampicin monooxygenase (RIFMO), co-working with NADPH, as evidenced by spectroscopic monitoring.¹¹¹ Furthermore, the discovered novel metabolites of **69**, being a result of the ansa bridge cleavage performed by monooxygenases (ROXs) from *Streptomyces venezuelae*, confirmed that structural alterations within naphthalenoid ansamycin scaffold are ‘double-edged swords’, not

only useful for human at designing novel antibiotics but also for bacteria as a defense response against newly developed antibiotics.^{112,113} Thus, in view of this still-emerging problem of bacteria resistance, in order to achieve chemical stability, improved binding mode with the target, and better balanced physico-chemical properties, semisynthetic modifications of naphthalenoid ansamycins have been widely performed, especially among natural naphthalene-C₁₇, 1,4-naphthoquinone-C₁₇ and 4-iminonaphthalene-1(4*H*)-one-C₁₇ scaffolds.

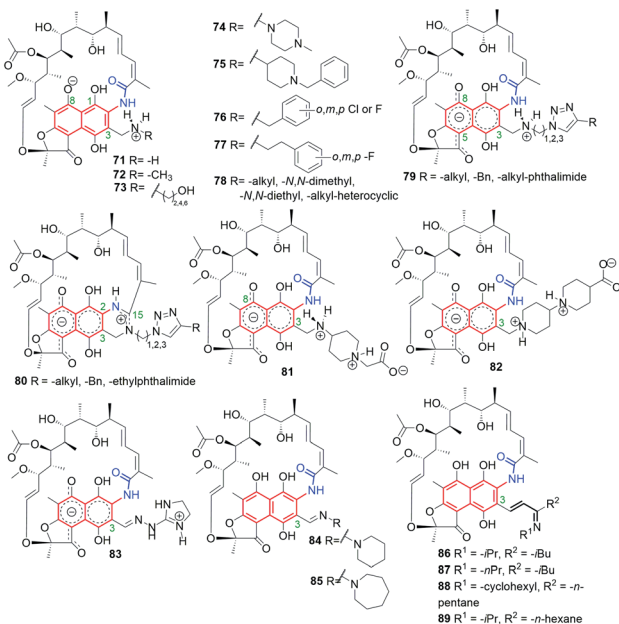
6.1. Naphthalene-C₁₇ ansamycins

It should be underscored here that only semisynthetic modifications of natural rifamycins have played a major role in pharmaceutical applications. A combined synthetic and genetic manipulation strategy with *A. mediterranei* S699 (AT6 mutant), targeting the ansa bridge, yielded rifamycin S and **69**, both lacking methyl at C(24).¹¹⁴ These new C(24)-desmethyl derivatives showed improved activity against rifampicin-resistant *M. tuberculosis*, containing RNAPs with single amino acid mutations H526 → T526 and S531 → L531 (Table 1), compared to **69** (MICs > 50 µg mL⁻¹). A such biological outcome suggested that the loss of C(24)-methyl, increasing the ansa bridge flexibility, and the presence of C(3)-arm of rifamycin-like derivatives,

Table 1 Antibacterial potency of semisynthetic naphthalene-C₁₇ ansamycins

Cmpd	MIC [µg mL ⁻¹]	
69 (ref. 48, 58, 60, 114 and 120)	MIC _{S. epidermidis} = 0.008 MIC _{OSDD321} > 50 MIC _{OSDD206} > 50 MIC _{OSDD55} > 50 MIC _{M. tuberculosis} H37RV = 0.005 MIC _{M. bovis} = 0.25	MIC _{S. aureus} MRSA = 0.0156 MIC _{S. aureus} MLSB = 0.03125 MIC _{E. faecalis} = 8 MIC _{S. pneumoniae} ≤ 0.25
Desmethyl- 69 (ref. 114)	MIC _{OSDD55} < 0.01 MIC _{OSDD321} = 0.05	MIC _{OSDD206} = 0.05
Desmethyl-rifamycin S ¹¹⁴	MIC _{OSDD55} = 0.1 MIC _{OSDD321} = 0.1	MIC _{OSDD206} = 0.05
74 (ref. 48)	MIC _{S. aureus} = 0.008–0.016 MIC _{S. epidermidis} = 0.008	
75 (ref. 58)	MIC _{M. tuberculosis} H37RV = 0.005 MIC _{M. bovis} = 0.125	MIC _{S. epidermidis} = 0.25 MIC _{S. aureus} = 0.25
75xHCl ⁵⁸	MIC _{M. tuberculosis} H37RV = 0.005 MIC _{M. bovis} = 0.125	MIC _{S. epidermidis} = 0.03–0.06 MIC _{S. aureus} = 0.125–0.25
79 (ref. 58)	MIC _{M. tuberculosis} H37RV = 0.5	MIC _{M. bovis} = 0.5
80 (ref. 58)	MIC _{M. tuberculosis} H37RV = 0.25	MIC _{M. bovis} = 0.25
81 (ref. 60)	MIC _{S. aureus} MRSA = 0.25 MIC _{S. aureus} MLSB = 0.5	MIC _{E. faecalis} > 8 MIC _{S. epidermidis} = 0.0625
82 (ref. 60)	MIC _{S. aureus} MRSA = 0.0078 MIC _{S. aureus} MLSB = 0.0078	MIC _{E. faecalis} = 1 MIC _{S. epidermidis} = 0.0078
83 (ref. 60)	MIC _{S. aureus} MRSA = 0.0156 MIC _{S. aureus} MLSB = 0.0078	MIC _{E. faecalis} = 1 MIC _{S. epidermidis} = 0.0078
84 (ref. 60)	MIC _{S. aureus} MRSA = 0.0156 MIC _{S. aureus} MLSB = 0.5	MIC _{E. faecalis} = 0.5
85 (ref. 60)	MIC _{S. aureus} MRSA = 0.0625 MIC _{S. aureus} MLSB = 0.5	MIC _{E. faecalis} = 0.5
100 (ref. 59)	MIC _{S. aureus} MRSA = 0.031–0.063 MIC _{S. aureus} = 0.031–0.063	MIC _{S. epidermidis} = 0.016
101 (ref. 120)	MIC _{S. pneumoniae} = 1	
102 (ref. 120)	MIC _{S. pneumoniae} = 2	
103 (ref. 120)	MIC _{S. pneumoniae} = 8	





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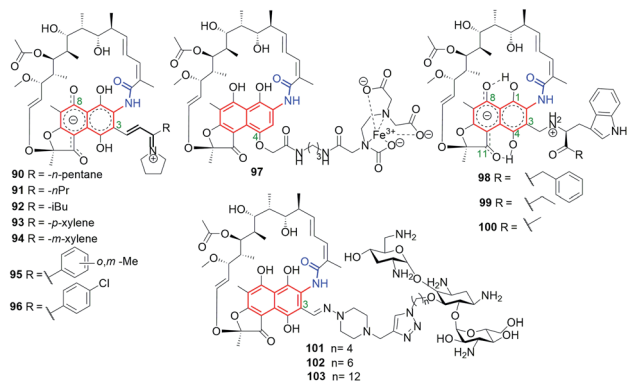


Fig. 36 Semisynthetic iminium analogs of **69** (90–96), and hybrid connections of rifamycins with EDTA (**97**), L-amino acid esters (**98–100**) and tobramycin (**101–103**).

Hybrid connections of rifamycin scaffold have been also obtained *via* initial functionalization not only the C(3) but also the C(4) position (Fig. 36).^{59,120,123} The conjugate **97**, bearing a strongly complexing EDTA component, was proposed as an example of an “aggressive” agent, breaking polypeptide and polynucleotide chains near the areas of their binding to RNAP through the Fenton reaction.¹²³ Semisynthetic hybrids with L-amino acid esters, which selected examples were shown in Fig. 36 (**98–100**), revealed structure–activity relationship (SAR) in dependence of the bulkiness of the substituent at the ester portion.⁵⁹ The most potent of these compounds, derivative **100**, with a small substituent at the ester moiety of L-Trp, showed very good antibacterial potency, on the level comparable to that of **69** against *S. aureus*, *S. epidermidis* and MRSA strains (Table 1).⁵⁹ This result seems to be in line with that obtained for similar pharmacophore, *i.e.* tryptamine, attached at C(3) of rifamycin scaffold, which revealed also potent antibacterial activity.¹¹⁹ The molecular docking suggested that the loss of high antibacterial potency of rifamycin hybrids with L-amino acid esters in case of their bulky ester substituents is a result of steric hindrance with A447 of the RNAP binding pocket.

Click chemistry strategy was applied to obtain heterodimeric **101–103** rifamycin–tobramycin conjugates linked *via* triazole bridges (Fig. 36).¹²⁰ This type conjugates, in dependence on the length of the linker, were effective against Gram-negative *Pseudomonas aeruginosa* efflux-deficient mutants, resistant to doxycycline and chloramphenicol. Furthermore, one of this-type semisynthetic conjugates showed improved potency relative to tobramycin against *S. pneumoniae* strain (Table 1), in case when the linker was the shortest (**101**). Thus, rifamycin–tobramycin hybrids can be useful for overcome the resistance of a Gram-negative *P. aeruginosa* to chloramphenicol and doxycycline *in vitro* and *in vivo*.

6.2. 1,4-Naphthoquinone-C₁₇ and 4-iminonaphthalen-1(4H)-one-C₁₇ ansamycins

Semisynthetic C(25)-carbamate derivatives **104–106** (Fig. 37), decorated at C(3) a morpholine moiety, belong to 1,4-naphthoquinone-C₁₇ ansamycins.¹²⁴ These ansamycins were shown

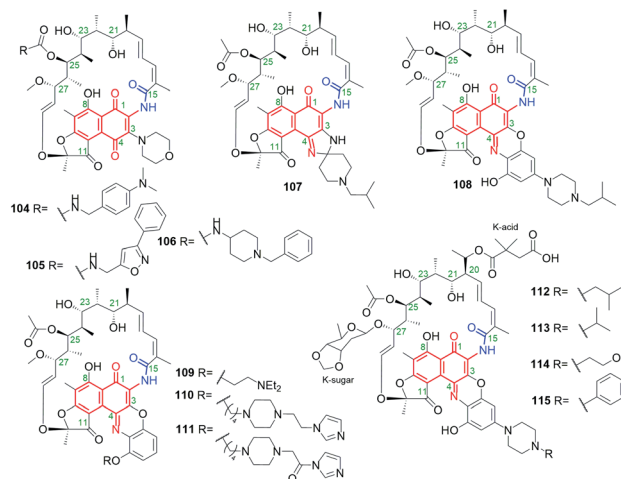


Fig. 37 Semisynthetic analogs of **69**, inspired by rifabutin (**107**) and rifalazil (**108**) structures.

to be even much more efficient than **69** against rapidly-growing mycobacteria such as *M. abscessus*, *M. chelonae*, *M. fortuitum* and *M. smegmatis* (Table 2). The most potent antimycobacterial potency showed piperidine derivative **106**, however, the explanation of this result, in view of the known binding modes of rifamycins with RNAPs (Fig. 9) and taking into regard bacteria resistance mechanisms *via* attachment of bulky groups to the ansa bridge, is difficult.

Rifabutin (**107**, Fig. 9 and 37) is capable of inhibiting bacterial RNAPs and is derived from natural rifamycin S, for which further natural transformations, specifically those yielding rifamycin O, are mediated by transketolase *Rif15* and the cytochrome P450 enzyme *Rif16*.^{127–130} High antibiotic activities of clinically-used **107**, against both Gram-positive and Gram-negative bacteria, created opportunity to produce alternatives relative to **69** and encouraged further exploration of semisynthetic transformations of iminoquinone ansamycins, assigned to the 4-iminonaphthalen-1(4H)-one-C₁₇ group (Fig. 2).^{109,130–132} Rifalazil (**108**) is an excellent example of a such semisynthetic ansamycin showing bactericidal potency against *M. tuberculosis*, *Chlamydia trachomatis*, *Clostridioides difficile* or *Helicobacter pylori* as stand alone antibiotic or with combination with the other antibiotics as *e.g.* vancomycin.^{10,133} Despite **108** helps to overcome resistance to other antibiotics, its toxic effects limited the clinical applications. Hence, novel semisynthetic benzoxazinorifamycins **109–111** (Fig. 37), being analogs of rifalazil and displaying high levels of affinity to RNAPs of wild-type and **69**-resistant mutants of *M. tuberculosis*, were obtained from rifamycin S.^{46,125} Compound **109** showed good metabolic stability. Structures of complexes between **109** or **110** and RNAPs indicated important interaction of C(3)–C(4) tails with a σ-finger of RNAPs (Fig. 9c), disturbing in this way the template DNA position in RNAP channels.^{44,46} Furthermore, derivatives **109–111** showed comparable abilities at inhibiting transcription relative to those of **69** or **108** in mutant *E. coli* and *M. tuberculosis* strains.⁴⁶ Antibacterial assays of **109–111** in aerobic (MABA) and anaerobic (LORA) conditions against *M.*



Table 2 Antibacterial potency of 4-iminonaphthalen-1(4H)-one-C₁₇ ansamycins

Cmpd	MIC or MIC ₉₀	
69 (ref. 124–126)	MIC _{M. abscessus} > 64 µg mL ⁻¹ MIC _{M. chelonae} > 64 µg mL ⁻¹ MIC _{90M. tuberculosis} ^{MABA} H37RV = 0.13 µM MIC _{90M. tuberculosis} ^{LORA} H37RV = 0.46 µM	MIC _{M. fortuitum} > 64 µg mL ⁻¹ MIC _{M. smegmatis} = 32 µg mL ⁻¹ MIC _{S. aureusS486L} > 64 µg mL ⁻¹ MIC _{S. aureusWT} = 0.00098 µg mL ⁻¹ MIC _{S. aureusH481Y} > 64 µg mL ⁻¹ MIC _{S. aureusH481Y} > 64 µg mL ⁻¹
13 (ref. 126)	MIC _{S. aureusS486L} = 0.25 µg mL ⁻¹ MIC _{S. aureusWT} = 0.016 µg mL ⁻¹	
108 (ref. 125)	MIC _{90M. tuberculosis} ^{MABA} H37RV < 0.004 µM MIC _{H. pylori} = 0.00025 µg mL ⁻¹	MIC _{90M. tuberculosis} ^{LORA} H37RV < 0.017 µM
104 (ref. 124)	MIC _{M. abscessus} = 8 µg mL ⁻¹	MIC _{M. fortuitum} = 1 µg mL ⁻¹
105 (ref. 124)	MIC _{M. chelonae} = 16 µg mL ⁻¹ MIC _{M. abscessus} = 8 µg mL ⁻¹	MIC _{M. smegmatis} = 0.06 µg mL ⁻¹ MIC _{M. fortuitum} = 1 µg mL ⁻¹
106 (ref. 124)	MIC _{M. chelonae} = 64 µg mL ⁻¹ MIC _{M. abscessus} < 0.03 µg mL ⁻¹ MIC _{M. chelonae} < 0.03 µg mL ⁻¹	MIC _{M. smegmatis} = 0.5 µg mL ⁻¹ MIC _{M. fortuitum} < 0.03 µg mL ⁻¹ MIC _{M. smegmatis} = 0.12 µg mL ⁻¹
109 (ref. 125)	MIC _{90M. tuberculosis} ^{MABA} H37RV = 0.02 µM	MIC _{90M. tuberculosis} ^{LORA} H37RV = 0.37 µM
110 (ref. 125)	MIC _{90M. tuberculosis} ^{MABA} H37RV = 0.08 µM	MIC _{90M. tuberculosis} ^{LORA} H37RV = 0.35 µM
111 (ref. 125)	MIC _{90M. tuberculosis} ^{MABA} H37RV = 0.07 µM	MIC _{90M. tuberculosis} ^{LORA} H37RV = 0.40 µM
112 (ref. 126)	MIC _{S. aureusS486L} = 1 µg mL ⁻¹ MIC _{S. aureusWT} = 0.0039 µg mL ⁻¹	MIC _{S. aureusH481Y} = 16 µg mL ⁻¹
113 (ref. 126)	MIC _{S. aureusS486L} = 1 µg mL ⁻¹ MIC _{S. aureusWT} = 0.0039 µg mL ⁻¹	
114 (ref. 126)	MIC _{S. aureusS486L} = 4 µg mL ⁻¹ MIC _{S. aureusWT} = 0.0039 µg mL ⁻¹	MIC _{S. aureusH481Y} > 64 µg mL ⁻¹
115 (ref. 126)	MIC _{S. aureusS486L} = 16 µg mL ⁻¹ MIC _{S. aureusWT} = 0.063	MIC _{S. aureusH481Y} > 64 µg mL ⁻¹

tuberculosis H37Rv showed that the most potent is compound **109**, even more active than **69** but lower active than **108** (Table 2). The most potent **109** inhibited the transcription process *in vitro*, catalysed by a wild-type *M. tuberculosis* RNAPs, at comparable IC₅₀s ~ 0.01 µM to those of **108**.¹²⁵ Furthermore, RNAPs of the mutants *M. tuberculosis* (D435 → V and S450 → L) were also better inhibited by **109** than by **108**. Semisynthetic derivatives of kanglemycin A (**13**, Fig. 14) described as **112–115** in Fig. 37, with modified end-group of C(3)/C(4) benzoxazino tails or at the K-acid portion at C(20) (the so-called Kangamides), showed attractive potencies against wild-type and **69**-resistant *S. aureus* strains (Table 2).¹²⁶ These studies indicated the presence of K-acid and the type of substituent at piperazine ring to be important for activity against **69**-resistant bacteria, whereas the introduced benzoxazino moieties were shown to improve *in vivo* efficacy.¹²⁶ The best antibacterial potencies against mutant *S. aureus* strains (H481 → Y) were noted for **112** and **113** ansamycins, which showed markedly improved properties relative to those of **13** and **108** (Table 2).

7. Conclusions

Our proposed division of natural and semisynthetic ansamycins (Fig. 2, top) based on the type of core and length of the ansa chain has neatly demonstrated the high levels of structural diversity of naphthalenoid ansamycins (23 structural subgroups). We also propose to use the term “ansamycins” as a family of macrolactams, bearing relatively rigid central cores as benzenoid, naphthalenoid, or atypical ones (e.g. heterocyclic,

alicyclic), originated from AHBA or other-type small semi-synthetic precursors (mutasynthons), and produced in a result of cooperation between polyketide (type I) and amide synthases. Due to the improved methods of extraction and genetic manipulations novel natural naphthalenoid ansamycin-like scaffolds, containing fused cores with heterocycles are available. Biosynthetic plasticity of naphthalenoid ansamycins is well reflected in the structural diversity of many naphthalenoid ansamycin congeners as divergolides, hygroscins, olimycins or 11,12-*seco*-rifamycin W analogs, bearing not only the lactam but also the lactone within the ansa bridge structure. Furthermore, there are also known examples of atypical fusion of the ansa chain with the core at C(1), as e.g. for hygrocin E or oligocyclic cores fused with the ansa chain, as noted for ansawaricin E and ansaeomycins. Among natural naphthalenoid ansamycins the shortest ansa chain is C₁₁ (neoansamycins A–C) whereas the longest is C₂₃ (naphthomycins L–Q).

The antibacterial potency of natural naphthalenoid ansamycins is moderate in most cases so the semisynthetic modifications of natural scaffolds are often required. One of exceptions are aminoansamycins, belonging to naphthoquinone-C₁₇ group, which antibacterial potencies are extremely high (even MICs = 0.0001 µM). It is interesting to note that mutasynthesis, often-used tool at optimization of anticancer activity of benzenoid ansamycins, has not been employed widely up to now in aim to modify the natural scaffolds of naphthalenoid ansamycins. Many natural ansamycin precursors containing cleaved ansa chain have been found in bacterial extracts as e.g. ansavaricins F–I, and despite most of



these precursors being relatively biologically inactive or of moderate potency, their macrolactamization and peripheral group modifications could provide alternatives of enhanced biological properties. The semisynthetic approach to modify naphthalenoid ansamycin natural scaffolds with the use of known 'starting points' as rifamycin SV, 3-formyl rifamycin SV, rifamycin S or kanglemycin A is predominant, in contrast to benzenoid ansamycins, where mutasynthesis is the main source of their structural diversity. It is noted also that some naphthalenoid ansamycins are called by different names for the identical structure as *e.g.*, divergolide C and olimycin B.

Overview on the biological properties of naphthalenoid ansamycins shows that these compounds (especially semi-synthetic derivatives) are predominantly powerful antibacterial agents, especially those of C₁₇ ansa chains, against both Gram-positive and Gram-negative bacteria, although notable other-type potency as anticancer was also found for *e.g.* hygrocin 53–56. The review revealed the following dependence that with increasing length of the ansa chain within the naphthalenoid ansamycins, the change in biological potency profile from the anticancer into the antibacterial occur. Furthermore, the profile of biological activity of structurally close C₁₂ hygrocin and divergolides, *i.e.* antibacterial or anticancer, seems to be dependent also on the type of the core. However, crucial for antibacterial potency of naphthalenoid ansamycins is conformational flexible ansa-bridge enabling adaptation of the molecule conformation (regulation of polarity/lipophilicity) to the outer environment during transportation to the target and enabling the molecular recognition of the ansamycin with the target. The best recognized molecular targets of naphthalenoid ansamycins are bacterial RNA-polymerases (RNAPs), which inhibition is related to antibacterial effects. Thus, target-directed designing of novel naphthalenoid ansamycin scaffolds, is reasonable way to obtain powerful antibacterials, even better than the clinically-used antibiotics as rifampicin (69), as demonstrated on semisynthetic C(3)-amine analogs, bearing rigid and basic amine arms, enabling zwitterionization. Interestingly, rifampicin-like derivatives possess the ability to interact with viral proteins, which are important at assembling viral membranes, therefore these compounds can be considered at designing novel antiviral agents.

8. Conflicts of interest

There are no conflicts to declare.

9. Acknowledgements

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