Synthesis of ramariolide natural products and discovery of their targets in mycobacteria†

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Ramariolides A–D are natural products with antibacterial activity. To exploit their cellular mechanism, we here devise the first total synthesis and prepare a photoprobe for target identification. Antibacterial testing against several pathogenic strains including Mycobacterium tuberculosis revealed the highest potency for ramariolide A. Chemical proteomics unraveled binding to essential proteins for amino acid anabolism.

Treatment of bacterial infections by classical antibiotics is severely impaired due to the development of multiple resistant strains.1,2 In particular tuberculosis caused by Mycobacterium tuberculosis, responsible for millions of deaths worldwide,3 represents a major challenge for drug development due to its largely impermeable membrane barrier.4 Thus, even antibiotic-sensitive strains require the application of several different antibiotics including combinations of synthetic pharmaceuticals such as isoniazid as well as natural products such as rifampicin.5,6 Natural products have always been a rich source of antibiotic discovery addressing a wealth of different targets.7 Although the discovery of new natural product antibiotics is declining,8 their detailed study still represents a major inspiration for the exploration of unprecedented pathways and the identification of druggable targets. In 2012 Anderson and co-workers reported the isolation of ramariolides A–D (Fig. 1) from the coral mushroom Ramaria cystidiophora and highlighted their potent antibacterial activity.9 All ramariolide compounds belong to the group of butenolides with ramariolide A and B exhibiting a unique spirocyclic acetal functionality. Despite their promising bioactivities and interesting structural features no total synthesis and no investigation of their in-depth mode of action have been reported up to now.

Here, we present the first total synthesis of all four natural products as well as their structural analogs for target identification.

Structural examination of the individual natural products ramariolide A (1a), B (2), C (3a) and D (3b) revealed long chain fatty acids (4a, 4b) as commercially available precursors (Scheme 1). These were converted into x-bromo ketones in excellent yields (5a, 5b) in three consecutive steps.10,11 Transformation to the corresponding phosphonium-ylides was quantitative12 and enabled olefination using maleic anhydride yielding the butenolide moiety.11 However, the reaction was challenged by partial homo-olefination as well as polymerization caused by the acceptor-substituted ylide. Although dilution could not prevent this side reaction completely, the butenolides were obtained in good yields favoring the desired (Z)-isomer (7a–c) in a 3 : 1 ratio.14 Subsequent reduction15 revealed racemic ramariolides C (3a) and D (3b) both matching published spectral data.9 Ramariolide A was obtained by epoxidation of the allylic alcohol using Sharpless conditions favoring the anti-diastereomer in a 87 : 13 ratio.16,17 Use of (+)-tartrate ester gave rise to the enantienriched natural product in 55% ee. As the optical rotation of the synthetic product differed from reported values, we utilized chiral europium salt-assisted NMR titration to determine absolute values (Fig. S1, ESI†).18

Next, we pursued conversion of ramariolide A to B by recatellisation of the epoxide to an oxetane moiety applying either alkaline or Bronsted/Lewis acidic conditions.19 Since only the formation of the epimeric spirocycle of ramariolide B (15) was observed, we rationalized that the inverted stereochemistry
Staphylococcus aureus against strains with ramariolide A exhibiting the most pronounced potency. In contrast to control substances, followed by esterification of the secondary alcohol (Scheme 1). Satisfyingly, the modified structure retained the alcohol function were unsuccessful. Instead, a sequence of oxidation and subsequent stereoselective reduction utilizing zinc-(μ)-borohydride yielded the correctly configured substrate. After screening a variety of reaction conditions, the use of camphorsulfonic acid finally led to formation of 2, which was confirmed by spectral data as well as X-ray structural analysis. Interestingly, the previously isolated spirocyclic epimer was confirmed by spectral data as well as X-ray structural analysis. All standard procedures for inversion of the alcohol function were unsuccessful. Instead, a sequence of oxidation and subsequent stereoselective reduction utilizing zinc-(μ)-borohydride yielded the correctly configured substrate.22 After screening a variety of reaction conditions, the use of camphorsulfonic acid finally led to formation of 2, which was confirmed by spectral data as well as X-ray structural analysis. Interestingly, the previously isolated spirocyclic epimer was validated as a side product by NOE experiments (Fig. S2, ESI†), supporting an acyclic intermediate (7a) (Fig. S3, ESI†). All natural products were tested against a series of pathogenic bacterial strains to determine their minimal inhibitory concentration (MIC) of bacterial growth (Table S1, ESI†). Interestingly, all ramariolides were active against diverse Gram-positive strains with ramariolide A exhibiting the most pronounced potency against Staphylococcus aureus USA300 (30 μM) as well as mycobacteria including Mycobacterium tuberculosis H37Rv (25 μM). Since antiminocobacterial compounds are rare, we were interested in unraveling their molecular targets responsible for growth inhibition. Therefore, a ramariolide-based probe exhibiting a terminal alkyne was designed, which allowed for modification with functionalized azides via click chemistry after target binding.23 Inspired by the terminal olefin-containing ramariolide D, we utilized this blueprint for designing an alkyne analog. The synthesis followed the established procedure yielding probe 1c without the need for alkyne protection. In contrast to control substances, 1a and 1b, terminal alkynylated compound 1c did not maintain antimicrobial activity suggesting that the altered bond geometry negatively interferes with target binding (Table S1, ESI†). Validation of the compound structure for alternative modification sites revealed the hydroxy functionality of ramariolide A as a synthetically accessible position for the introduction of an alkyne tag. Although ramariolide A contains protein-reactive epoxide as well as Michael acceptor moieties, the mode of target binding could be driven by (non-covalent) reversible interactions. The natural product was thus also provided with a diazirine UV-crosslinker located next to the alkyne moiety.25 Diazirines covalently attach to amino acids in physical proximity to the binding pocket upon UV irradiation thus enabling mass spectrometric (MS) target identification under protein denaturing conditions.26 Corresponding probe 8 was accessible by esterification of the secondary alcohol (Scheme 1). Satisfyingly, the modified structure retained antymycobacterial activity, an important prerequisite for following ABPP studies. Mycobacterial cells were grown to mid-log phase, harvested and incubated with DMSO or probe 8 in PBS for 1.5 h. UV-irradiation for 10 min at 365 nm and 4 °C was performed to account for covalent as well as reversible target binding. Cells were lysed at 4 °C by sonication and soluble as well as insoluble fractions were separated by centrifugation. Protein fractions were clicked to biotin azide, enriched on avidin beads and released by tryptic digest (Fig. S4, ESI†). Cleaved peptides were modified by dimethyl labeling using light, medium and heavy isotope reagents.28 A label switch was performed throughout the biological replicates and differentially labeled samples were pooled prior to LC–MS/MS analysis. Identified proteins were ranked in corresponding volcano plots (log2 ratio of 8 to DMSO treatment (Fig. 2A) against statistical significance (−log10 p-value). Proteins enriched by a factor >2 with a p-value of 0.05 or below were considered hits (Table S2, ESI†). Evaluation of the soluble protein fraction revealed five hits as essential proteins for mycobacterial growth, while none were detected in the insoluble fraction (Fig. S5, ESI†). These hits comprise the 30S ribosomal proteins S4 (RpsD) and S5 (RpsE), the
covalent binding mode we repeated gel-free ABPP using probe E. coli with natural substrates for enzyme binding by blocking their thus intriguing to speculate that ramariolide A might compete † labeling of purified protein was successful in contrast to heat indicating a covalent mode of binding. Furthermore gel based molecular size (Fig. 2B) upon induction of protein expression, chemistry resulted in a strong fluorescent band of predicted 30 Mc2155 soluble fraction preincubated with 100 M. smegmatis vs. E. coli-D-thiogalactopyranosid. These results suggest a specific interaction of ramariolide A with a natively folded protein (Fig. S8, ESI†). Based on the covalent binding mode we repeated gel-free ABPP using probe 8 against DMSO (Fig. 2C) as well as competition with unmodified ramariolide A (Fig. 2D) without UV crosslinking. Importantly, Ask was consistently among the most significantly enriched proteins in both experiments. Although these findings highly interefere of ramariolide natural products with amino acid anabolism in mycobacteria, binding to other target enzymes cannot be excluded and is under further evaluation.

In summary we provided a first total synthesis of the family of ramariolides and verified their antimicrobial activity against Gram-positive strains. Molecular targets in mycobacteria were identified with customized probes. Essential members of the bacterial amino acid metabolism were found as interactors and validated by in situ and in vitro experiments providing strategies for future drug development.

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Notes and references