

Recent trends in the structural revision of natural products

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Covering 2011 to 2017

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This article reviews recent reports on the structural revision of natural products. Through a critical assessment of the original and revised published structures, the article addresses why each structure was targeted for revision, discusses the techniques and key discrepancies that led to the proposal of the revised structure, and offers measures that may have been taken during the original structure determination to prevent error. With the revised structures in hand, weaknesses of original proposals are assessed, providing a better understanding on the logic behind structure determination.

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1 Introduction

Human medicine has relied on bioactive natural products and their derivatives for the treatment of a wide variety of diseases. In fact, a substantial proportion of all FDA approved drugs are natural products or their derivatives.¹ The biological activities of these molecules are governed by their three-dimensional structures and the functional groups present.^{2, 3} Failure to establish the correct structure can ruin a drug development program, as exemplified by the TIC10 case. This drug candidate, identified from a free NCI database, was found to be active against a plethora of cancer cell lines.⁴ Scientists from Scripps proved that the structure from the NCI database used for the TIC10 patent was incorrect and they subsequently filed a patent with the correct structure. This led to a legal disagreement that is still ongoing.⁵ Hence, knowledge of the correct structure of a natural product, especially if it is a starting point for the development of a drug, is crucial.

In the past, structure determination was an arduous process occupying years of effort. Studies to identify molecular structures relied on degradation, derivatization, and characteristic reactivities of the functional groups, all based on synthetic chemistry.⁶ X-ray crystallography, available since the 1920's,^{7, 8} has been considered the gold standard for structure elucidation. However, it is limited to samples with sufficient material that can be crystallized, is not always suitable for establishing absolute configuration, and does not always lead to accurate prediction of bonding relationships as demonstrated by the revision of diazamide B.⁹ In the late 1960's, NMR spectroscopy started to establish itself as an indispensable tool for structure determination prompting a renewed era of drug discovery.¹⁰ In recent years, computational prediction of NMR and other spectroscopic properties has made spectral interpretation more accurate.^{11, 12} According to Koji Nakanishi (1991): "Until the mid-1960s, structure determination was an art that could be likened to solving a complicated detective case, but with the spectacular advancement in spectroscopy it has become less inspiring, and since the mid-1980s, in most cases, structure determination has become rather routine."¹³ However, it is important to note that although structure determination benefits from the vast array of spectroscopic techniques now available, there are still challenges inherent to identifying novel and complex natural products that enrich the elucidation process. Use of spectroscopic techniques to identify a molecule is an inverse problem, meaning that one uses

observations about acquired data to discard all solutions but the most logical one.¹⁴ Thus, individual bias can potentially result in a proposed structure that is a mere figment in the eyes of the beholder rather than a product of critically testing all possible hypotheses.

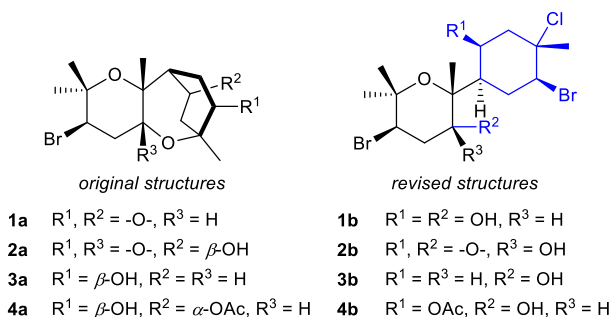
Previous reviews have evaluated important cases of erroneous structures and their revisions, bringing awareness to the surprisingly large number of structural misassignments reported in the literature. Nicolaou and Maier emphasized the value of total synthesis in confirming the structure of natural products,^{6,15} whereas Lawrence highlighted the contribution of biosynthetic rationalization of natural products to structure determination.¹⁶ A tabulated review presented by McPhail covered marine natural product revisions from 2005–2010.¹⁷ In the current publication, we review 23 unique structural revision cases reported from 2012 to 2017, each showcasing different approaches including computational chemistry, NMR spectroscopic methods, empirical rules, techniques in X-ray crystallography, and biosynthetic studies that revealed and resolved structural errors.

2 Structural revision based on computational prediction of NMR spectroscopic data

The field of natural products has advanced considerably with the development of technology that has resulted in the ability to compare experimentally derived NMR spectroscopic data with predictions generated by computational chemistry.¹⁸⁻²⁰ These computational methods have become increasingly accessible with the development of user-friendly software and increased computing speed. In addition to aiding the structure elucidation of new natural products, computational methods have found great utility in helping to uncover and ultimately revise errors in previously reported compounds. The case studies included in this section exemplify the efficacy of computational prediction towards correcting errors in molecular structures of natural products.

2.1 Aldingenins A–D

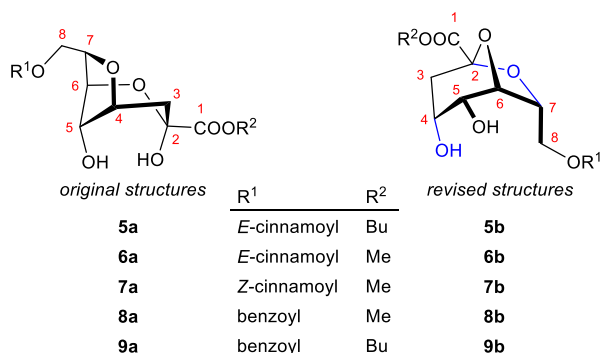
The red algal genus *Laurencia* has been the source of many isoprenoid natural products, including the chamigrene sesquiterpene (–)-elatol,²¹ the halogenated sesquiterpenes rhodolaureol and rhodolauradiol,²² and the brominated bisabolene derivatives aldingenin A–D (**1–4**).^{23, 24} The molecular masses of **1–4** were determined using HREIMS and the structures (**1a–4a**) were based on analysis of 1D and 2D NMR spectra.²⁵ Comparison of NMR spectroscopic data of synthesized aldingenins B and C with natural isolates did not agree with the proposed structures **2a** and **3a**.^{25, 26} Studies on aldingenin C (**3**) and D (**4**) using computer-assisted chemical structure elucidator (CASE),²⁷ revealed that they were in fact the known compounds caespitol (**3b**) and 5*S*-acetoxycaespitol (**4b**).²⁶ Based on this realization, Kutateladze and coworkers used a newly developed strategy to calculate spin-spin coupling constants named relativistic force field (*rff*), using an optimized basis set (DU8c) and scaled Fermi contacts to identify the correct structures for aldingenins A (**1b**) and B (**2b**).²⁸ Not surprisingly, the newly proposed structures bear chlorine atoms, as do caespitol (**3b**) and 5*S*-acetoxycaespitol (**4b**). It is unclear how the MS signatures of natural products with two bromine and one chlorine atoms could be confused with those possessing only one bromine. One possible explanation is that electron impact ionization led to weak molecular ion peaks for these non-aromatic halogenated compounds.²⁹



2.2 Decurrensides A-E

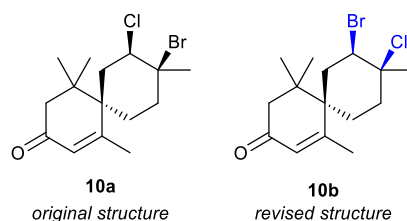
After decurrensides A–E (**5–9**) were isolated from the Chinese herb *Solidago decurrens*,³⁰ the proposed structure **8a** was synthesized; however, the NMR spectroscopic data did not match the natural product.³¹ This discrepancy led to the revelation that the depicted structures **5a–9a** did not match the compound names. More specifically, the paper proposed **5–9** to be 2,6-hemiacetal-3-deoxy-D-*altro*-2-octulosonic acids (**5a–9a**) but named them 3-deoxy-D-*manno*-2-octulosonic acids. Further studies based on computation of δ_C and the proton spin-spin coupling constants determined that both the original name and depiction of **5–9** were incorrect and instead should be revised to 2,6-anhydro-3-deoxy-D-*manno*-2-octulopyranosonates (**5b–9b**).³² DFT-based computation of δ_C for **8a** using the gauge-including atomic orbital (GIAO) method showed poor correlation with the experimental values, including a deviation of more than 10 ppm for the hemiacetal carbon. In the original proposal **5a–9a**, a spin system involving four consecutive 5 Hz spin-spin coupling constants was reported between H₂-3/H-4/H-5/H-6/H-7. Relativistic force field and *DU8c* based coupling constant calculations on the proposed structure predicted $J_{H-6,H-7}$ to be 0.2 Hz, which was drastically smaller than the reported value of 5 Hz. When computational predictions were made for α - and β -2,5-hemiacetals of the methyl ester of 8-*O*-benzoyl-3-deoxy-D-*altro*-2-octulosonic acid, both 2,5- and 2,6-hemiacetals of 3-deoxy-D-*manno*-2-octulofuranosonic acid, and the two ketal forms of 3-deoxy-D-*manno*-2-octulofuranosonic acid, none of the generated values for these structures matched the experimentally acquired spectral data for **8**. The core structure 2,6-anhydro-3-deoxy-2-octulopyranosonic acid was found to closely match experimental spectral data of **8** through comparison of computationally derived δ_C and *rff*-calculated *J*-coupling values. In support of the revision, **5b–9b** contain a common spin system with four consecutive protons all having dihedral angles of 40–50° between vicinal neighbors resulting in the expected sequential 5 Hz coupling constants. Ultimately, *rff* based calculations of nuclear spin-spin coupling constants and

prediction of δ_C led to the conclusion that structures **5–9** consist of a 2,6-anhydro-3-deoxy-D-*manno*-2-octulopyranosonate core with different substitution in the C-8-*O*-acyl and C-1 ester.



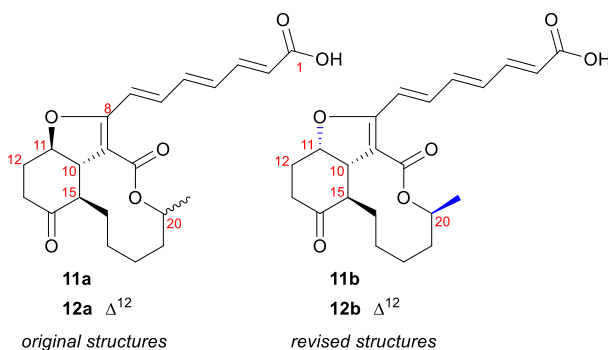
2.3 Tristichone

Tristichone C (**10**) is a chamigrane-type sesquiterpene obtained from the marine red alga *Laurencia tristicha*.³³ Its structure (**10a**) was elucidated using HRMS and NMR spectroscopic analysis, but no experimental evidence was provided to support the relative positions of the bromine and chlorine atoms. Recently, Kutateladze *et al.* developed a quadratic scaling correction to be used in conjunction with their *rff* method to take into account the effect of spin-orbit contribution on predicted δ_C .³⁴ The correction parameters were obtained from an experimental training set of more than 400 δ_C for carbon attached to S, Cl, Br or I. Among the 119 molecules chosen in this study, the structures of 16 were subsequently revised. Misassignments ranged from incorrect stereochemistry to incorrect carbon skeletons. The predicted chemical shift for the chlorine-bearing carbon atom of **10a** deviated by 6.6 ppm from the experimental value, but the constitutional isomer **10b** resulted in a much better fit. Interestingly, this computational prediction is in perfect agreement with a previous empirical study in which the δ_C for a series of polyhalogenated compounds similar to **10a** and **10b** were reported.³⁵



2.4 Glabramycins B and C

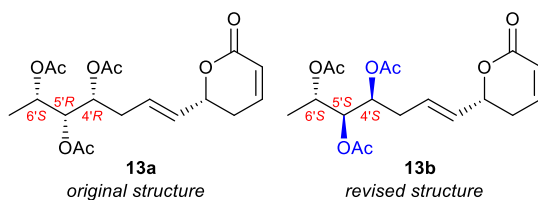
Antibiotic glabramycins B (**11**) and C (**12**) were isolated from *Neosartorya glabra*, a fungus obtained from soil samples.³⁶ In the original proposal for glabramycin B (**11a**), an *anti* relationship between C-10, C-11, and C-15 was proposed based on the triplet resonance of 9.6 Hz observed for H-10 whereas the configuration at C-20 was not determined. The decalactone polyketide structure of glabramycins B (**11**) and C (**12**) are closely related to the natural product dictyosphaeric acid, where C-11 exhibits a *syn* configuration relative to C-10.³⁷ Prompted by this difference in the configuration at C-11, the proposed structures of glabramycins and their possible diastereomers at position C-11 and C-20 were subjected to δ_C DFT calculation. Coupling constants between H-10 and H-11 and between H-11 and H-12 were also predicted computationally for all the diastereomers.³⁸ Calculated δ_C and J values favored **11b** and **12b** as the correct structures of glabramycins. This result was recently confirmed through total synthesis of **11b**.^{39,40} The original proposal lacked a key literature precedent: the structure determination of dictyosphaeric acid, which has the same decalactone moiety as glabramycins, utilized a J based configuration analysis to explain the large coupling constant observed for the eclipsed configuration between H-10 and H-11.^{37,41}



2.5 Synargentolide A

Syncolostemon argenteus, a South African plant of the mint family *Lamiaceae*, produces the 5,6-dihydro- α -pyrone synargentolide A (**13**).⁴² To determine its absolute configuration, the compound was fully hydrolyzed and transformed into 5- and 6-membered acetonides. First, the 5-membered ring acetonide tethering alcohols at positions 4' and 5' was used to deduce the relative configurations of these positions, as well as the configuration of the free alcohol at position 6' using derivatization of its α -methoxy- α -trifluoromethylphenylacetate ester. To relate the configuration of the alcohol at C-6' with the two other stereocenters, δ_C of the two methyl groups on the 6-membered acetonide between alcohols 4' and 6' were compared with each other (Fig. 1A). Since different

chemical shifts were obtained for the methyls of the acetonide, the authors concluded that the acetonide adopted a chair conformation which translates to a *syn* configuration of the corresponding 1,3-diol and the overall structure **13a**. Yet when **13a** was synthesized,⁴³ the NMR spectroscopic data did not match those originally reported. In a subsequent study, the 4'*R*,5'*R*,6'*R* isomer was synthesized and proposed as the absolute configuration of **13** based on visual comparison of ¹H NMR spectral features.⁴⁴ However, extraction of chemical shifts and *J* coupling constants by non-linear fitting of the spectra obtained from the natural product and the synthetic diastereoisomer highlighted slight differences disproving the revision.⁴⁵ Further predictive calculation of the coupling constants by DFT computation of four possible diastereoisomers suggested the configuration to be **13b**. The simulation took advantage of the spin-spin option in the program Gaussian 09⁴⁶ which calculates the scalar coupling constant as a summation of Fermi contact, spin-dipolar, paramagnetic spin-orbit, and diamagnetic spin-orbit.⁴⁷ The reason for the erroneous stereochemical assessment in the original proposal is uncertain. The acetonide strategy has been extensively studied from the conformational point of view⁴⁸ and suffers no exception when the acetonide is substituted at C-5', as in this case.⁴⁹ Also, it was shown that substituents at C-4' or C-6' exhibiting weak steric hindrance, like nitrile or alkyne derivatives, may produce an *anti* acetonide. This could mislead configurational assignment since *anti* acetonides, which normally adopt a twist-boat conformation, can flip to a chair conformation (Fig. 1B).⁵⁰ However, the methyl group of **13** was not shown to be small enough for this to happen. A possible alternative hypothesis is that epimerization was triggered by the harsh hydrolysis conditions (0.5 M NaOH in MeOH/H₂O for 2.5 days, followed by acidification and heating up to 100 °C for 2 min) or during acetonide formation in acetone with amberlyst 15.⁴²



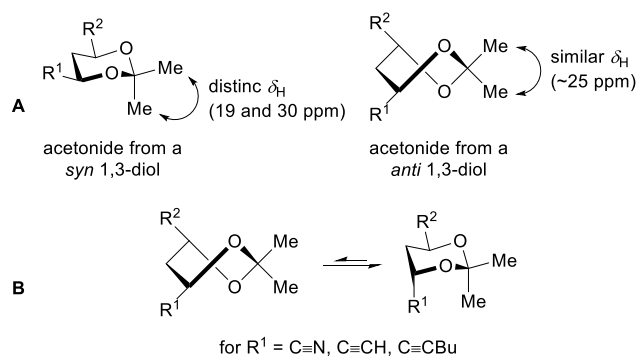
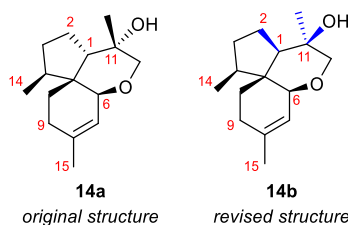


Fig. 1. (A) Acetonide methodology for configuration assignment of 1,3-diols. (B) Misleading configuration assignment for acetonide from *anti* 1,3-diols

2.6 Cordycepol A

Cordycepol A (**14**), along with analogues B and C, were isolated from *Cordyceps ophioglossoides*, a fungus that colonizes other fungal species.⁵¹ NOESY cross-peaks between H₃-14/H-1, H₃-15/H-1 and H₃-14/H-9 supported the relative configuration of the spiro[4.5]decane ring system of **14a** (Fig. 2). However, the structures of **14** and its analogues were recently revised.⁵² The calculated $J_{H-1, H-2b}$ and $J_{H-1, H-2a}$ as well as chemical shifts of C-1, C-2, C-6, C-7, C-10, and C-12 derived using the *rff/*DU8c parametric method for **14a** significantly diverged from the experimental NMR spectroscopic data. In contrast, the calculated parameters of the revised structure **14b** very closely match the experimental values. The original proposal contains a NOESY correlation between H₃-15 and H-1⁵¹, which should be at least 6.4 Å apart based on computational predictions. It is possible that H-1 was wrongly assigned since several ¹H NMR signals were reported in the same range (δ_H 1.00–1.50), but in the absence of public access to free induction decay (FID) files as proposed by Pauli,⁵³ this cannot be confirmed.



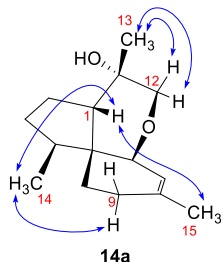
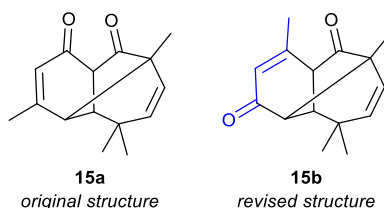


Fig. 2. NOESY correlations reported for the original cordycepol A structure (**14a**).

2.7 Meridane

A complex mixture of $7\beta,9\alpha$ -longipinene diesters extracted from the leaves and stem of *Stevia lucida* were subjected to hydrolysis, oxidation, acid-catalyzed dehydration, and rearrangement.⁵⁴ This yielded meridane, proposed as **15a** based on NMR spectroscopic evidence and a rational reaction mechanism of formation. Shortly thereafter, the structure **15a** was disproved based on comparison of experimental and computed δ_C using (GIAO mPW1PW91/6-311+G(d,p)).⁵⁵ Close examination of the reaction conditions and other possible rearrangement pathways led to the alternative structure **15b**, supported by computed δ_H , δ_C , J_{H-H} and J_{C-H} . It is difficult to envision what could have been done to avoid this misassignment, since all HMBC correlations equally support the original and the revised structure.



2.8 Brassicenes

The plant pathogen *Alternaria brassicicola* infects most *Brassica* species and produces brassicene diterpenes **16–23** that are phytotoxic.^{56–58} These molecules belong to the fusicoccane terpenoid family which has been extensively studied for their interesting biological activities.^{59–63} Initial computational studies predicted chemical shifts deviating 10–11 ppm from observed data for C-7 of **16a** and C-10, C-13 and C-18 of **22a**, disfavoring the proposed structures.^{56, 64} Also, the original proposal placed H-17 and H-18 6.8 Å apart, which did not meet expectations for an observed NOE correlation (**22a**, Fig. 3). Recently, **22** was re-isolated and revised to **22b** by pairing quantum-chemical predictions with extensive NMR spectroscopic analysis, resulting in stronger agreement

between calculated and experimental δ_C along with favorable NOE distances supporting correlations between H-18 and H-17 (Fig. 3).⁶⁴ The structure was further validated using X-ray crystallography and electronic circular dichroism spectroscopy (ECD) for absolute stereochemistry. Subsequently, structures of other brassicene diterpenes **17–21** and **23** were also revised based on δ_C calculations.⁶⁴ In the original studies,^{56, 57} important HMBC correlations to support the structures were missing. For example, an isolated spin system of brassicene G (**19**), composed of the methyl H₃-18 and the methine H-12, was proposed based on an HMBC correlation of H-12 with C-1, C-10, C-11 and C-13, but the more important HMBC correlations from H₃-18 were not reported.

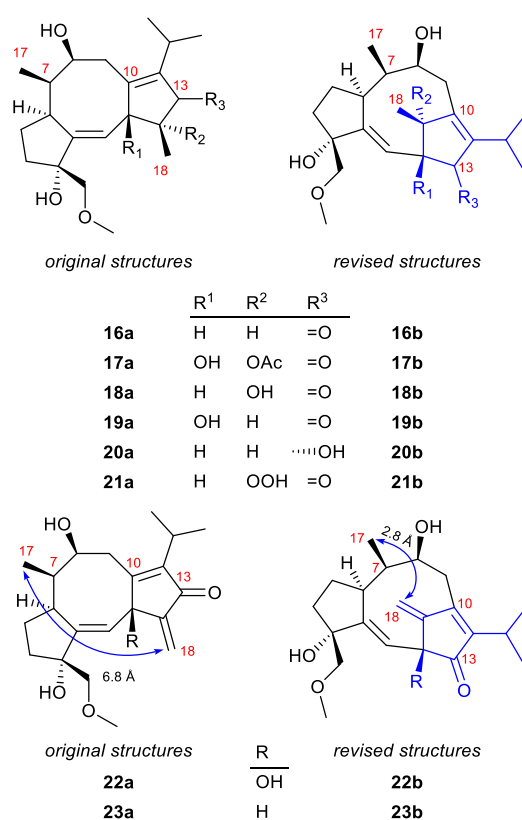


Fig. 3. Original and revised structure showing a key NOE enhancement.

2.9 Type C Polycyclic Polyprenylated Acylphloroglucinols

Polycyclic polyprenylated acylphloroglucinols (PPAPs) are a class of compounds featuring a highly oxygenated bicyclo[3.3.1]nonane substituted by prenyl side chains.⁶⁵ These compounds are found exclusively in plants from the Clusiaceae family, which includes St-John's wort (*Hypericum perforatum*), well known for diverse medicinal properties.⁶⁶ There are three categories of PPAPs, types A, B, and C, distinguished by the position of the acyl moiety relative to the quaternary center

of the prenyl side chain (Fig. 4).⁶⁵ A series of the rare type C PPAPs were isolated from the seeds of *Garcinia subelliptica* with structures **24a–26a** elucidated by conventional MS and multidimensional NMR spectroscopy.⁶⁷ However, no experimental evidence was provided to support whether the prenyl side chain was located at C-5 (type A) or C-1 (type C). In a recent examination of *Hypericum cohaerens*, the structures of **24–26** were revisited.⁶⁸ Three-dimensional models of both type C (**24a**) and type A (**24b**) were subjected to a molecular mechanic conformational search followed by DFT geometrical optimization, frequencies and thermochemical computation, and δ_C prediction, all at the mPW1PW91/6-31G(d,p) level of theory. The predicted spectroscopic data for **24b** were in closer agreement with experimental values, especially for positions 1 and 6, which were the most impacted by the acyl group. The same rationale was used to revise co-isolated PPAPs to **25b–26b**.⁶⁸ It is worth noting that while the 3D structures for **24a** and **24b** provided in the supplemental information of the revision were correct, the structures of **24a** and **24b** portrayed in the main manuscript were depicted incorrectly with an α -H at R³ rather than a β -H.⁶⁸ The structures included in the current review are based on the correctly portrayed 3D structures.

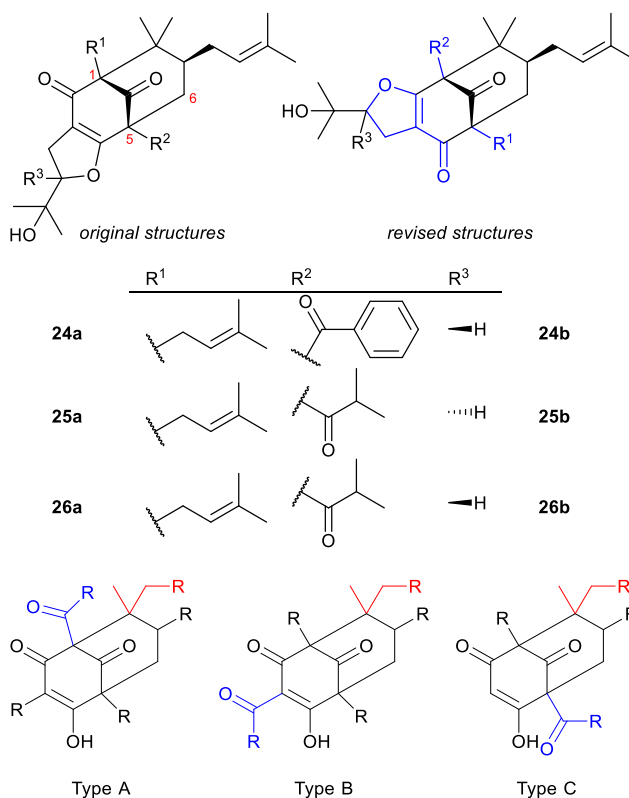


Fig. 4. Categories of polycyclic polyprenylated acylphloroglucinol are based on the relative position of the acyl moiety (blue) and the quaternary center (red).

3 Structural revision based on calculation of chiroptical properties

A major challenge often encountered by chemists attempting to elucidate structures of natural products is determination of relative and absolute configuration.⁶⁹ The chiroptical properties of these chemicals are often the only parameters available to distinguish some types of isomers. Advances in computational tools have elevated the empirical interpretation of chiroptical properties to a true predictive methodology aiding in assignment of configuration.⁷⁰ These new methods have contributed to corrections of erroneous structures, as demonstrated in the following cases.

3.1 Brevianamides

Diketopiperazine alkaloids are a rich class of cyclic dipeptide molecules exhibiting a range of biological properties.⁷¹⁻⁷⁴ Brevianamide M (**27**) and its oxidation product **28** were isolated from *Aspergillus versicolor* along with a related diketopiperazine dimer and oxepin-containing alkaloids.⁷⁵ The relative configuration of the two stereocenters in **27** was determined by X-ray crystallography to be (2*R*,13*R*) or (2*S*,13*S*). Hydrolysis of the natural products in aqueous solution gave L-phenylalanine suggesting an *S*-configuration at position 13, which led to the proposal of a (2*S*,13*S*) configuration for **27a**. However, in contrast to the positive optical rotation calculated for the proposed configuration **27a** using four different DFT quantum chemical models, the molecule showed an experimental optical rotation of -147.7° , suggesting a (2*R*,13*R*) configuration.⁷⁶ After careful analysis of this discrepancy, the hydrolysis process was questioned since it could cause epimerization at C-13 of **27b** to form the low energy diastereomer **27c** that could hydrolyze to give L-phenylalanine (Fig. 5). As expected, DFT studies predicted the epimer **27c** to be energetically more stable than **27b** with a predicted conversion rate of more than 92%. Since **28** was isolated using the same procedure as **27**, it was proposed to share the same absolute configuration. The reassignments were confirmed by agreement between experimental data and DFT computed optical rotation, predicted VCD spectra, and the theoretical prediction of positive and negative cotton effects for ECD spectra.⁷⁶

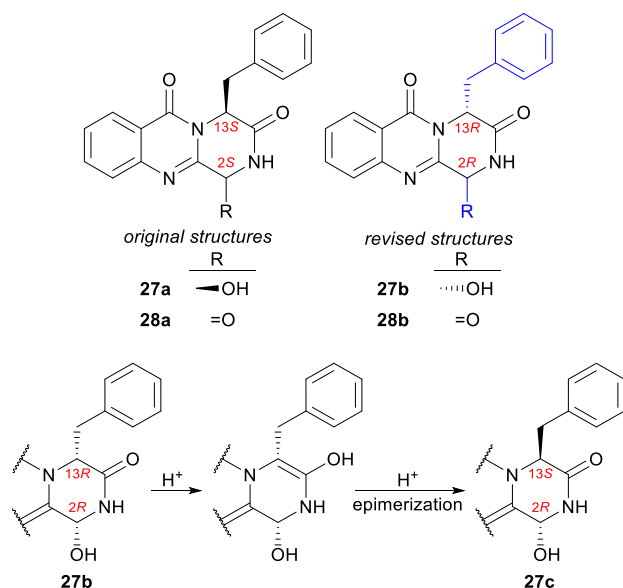
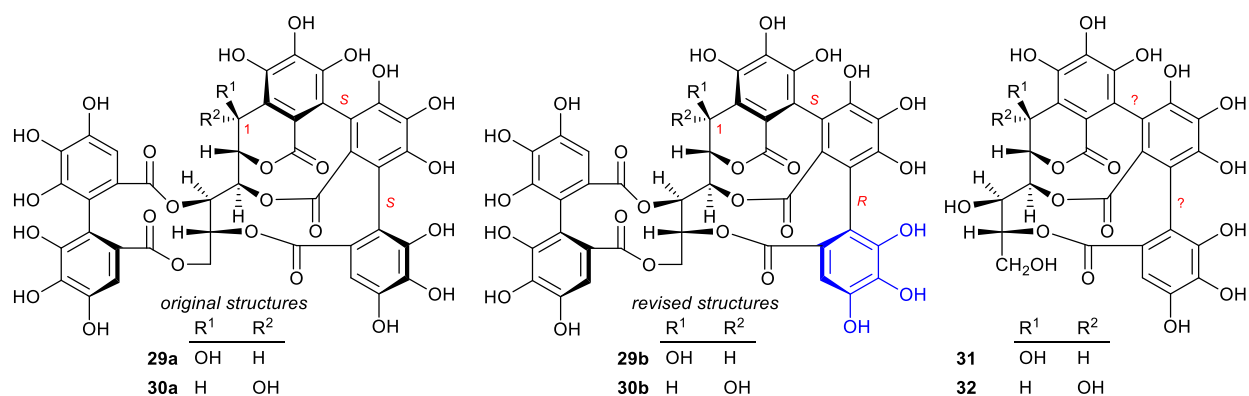


Fig. 5. Hypothetical epimerization reaction of **27b**.

3.2 Castalagin and vescalagin

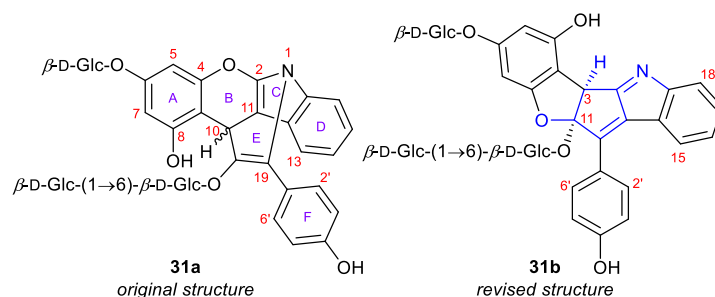
Castalagin (**29**) and vescalagin (**30**) are epimeric ellagitannins exhibiting antioxidant, antiviral, and inhibitory activity against DNA topoisomerase II.^{77, 78} They were initially isolated from the wood of chestnut (*Castanea sativa*) and oak (*Quercus sesseliflora*) but their exact atropisomerism configuration could not be determined at that time.⁷⁹⁻⁸¹ The configuration for the nonahydroxytriphenoyl group was proposed to be (*S,S*) based on a comparison of the circular dichroism spectrum with the one obtained for a model compound of known configuration.⁸² In a follow-up molecular mechanics study, it was proposed that an (*S,R*) configuration at the nonahydroxytriphenoyl group would be more stable.⁸³ The discrepancy in proposed configurations was resolved by predicting the ECD spectra of hydrolyzed analogues castalin (**31**) and vescalin (**32**) using TDDFT calculation.⁸⁴ The hydrolysis was necessary since strong Cotton effects from the (*S*)-hexahydroxydiphenoyl ester overlapped with the small Cotton effects of the triphenoyl group. The experimental ECD spectra obtained for **31** and **32** were in agreement with the predicted spectra of the (*S,R*) configuration. This led the authors to conclude that castalagin and vescalagin could be revised to **29b** and **30b**, respectively. To confirm the ECD spectroscopy results, DFT prediction of δ_{H} and δ_{C} were performed⁸⁴ and found to be in good agreement with experimental values for **29** and **30**.⁸⁵



3.3 Nudicaulin

The flower petals of *Papaver nudicaule* owe their yellow color to a group of glycosides called nudicaulins.⁸⁶ After extensive NMR spectroscopic analysis, nudicaulin I was suggested to possess an unprecedented 10*H*-1,10-ethenochromeno[2,3-*b*]indole moiety (**33a**). The downfield chemical shift of C-2 led the authors to propose that this position was attached to one of the oxygens of ring A, leading to the asymmetry needed to explain the two distinct ¹H NMR signals of H-5 and H-7 within ring A. Similarly, the relatively upfield chemical shift of C-19 was interpreted as an indication for a bond between C-19 and the indole nitrogen. However, if **33a** was stable enough to be isolated, it would be expected to experience severe strain at C-2 and C-11 which would limit the aromaticity to isolated phenyl rings rendering the molecule colorless. This prompted the revision of **33a** to **33b** using NMR spectroscopic data, chemical derivatization, and comparison of experimental and predicted chiroptical properties.⁸⁷ The evidence included HMBC correlations from H-3 and H-18 to N-1 and ROESY correlations between H-2'/6' and H-15. The ECD spectrum of **33b** was predicted with TDDFT using CAM-B3LYP functional and conductor-like polarizable continuum model (CPCM) for solvent effects of methanol. However, prediction at this level of theory was unsatisfactory and the size of the molecule precluded the use of a higher level of theory. Simplification of the molecule to a partially hydrolyzed derivative with one glucose unit at C-11 was evaluated but this prediction was also unsuccessful. Interestingly, modeling a hypothetical chemically unstable form of **33b** with a hydroxyl in place of the sugar accurately predicted the experimental spectrum. While this advanced demonstration is an example of how complex a structural elucidation with quantum-chemical computation can be, the authors missed a simpler solution: since the relative stereochemistry between the aglycon and the sugar moieties was determined by

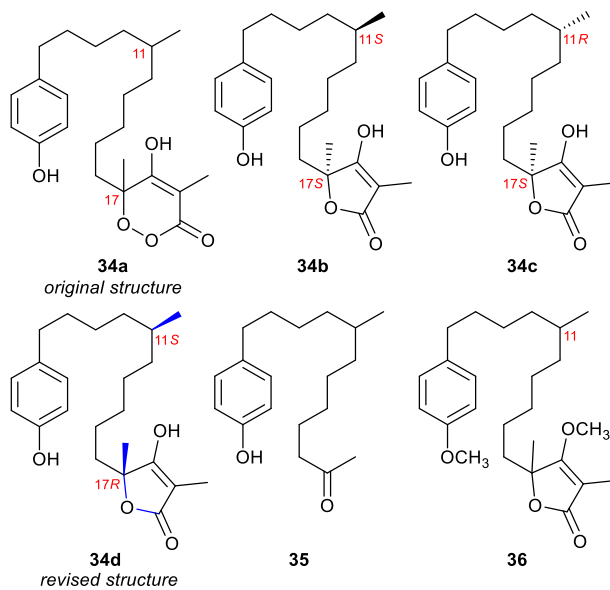
semi-quantitative ROESY, the absolute configuration of **33b** could have been deduced from the hydrolyzed glucose unit.



3.4 Plakinidone

Sponges belonging to the genus *Plakortis* are well known to produce molecules having cyclic five- or six- membered endoperoxide rings.⁸⁸⁻⁹¹ Plakinidone (**34**), isolated from *Plakortis angulospiculatus*, was proposed to have a *p*-hydroxyphenyl and a six membered per lactone ring flanking the two ends of a ten-carbon alkyl chain (**34a**).⁹² Attempts to synthesize an analogue having the same per lactone moiety were unsuccessful which brought into question the validity of **34a**.⁹³ Careful NMR spectroscopic analysis resulted in the proposal of more plausible **34b** composed of a tetrone acid moiety instead of the per lactone. This compound and the 11*R*,17*S* stereoisomer **34c** were synthesized and their optical rotations were compared to the value for the isolated compound, leading to the conclusion that the correct configuration was 11*S*,17*S* (**34b**).⁹³ A subsequent study challenged this finding due to the unstable nature of **34** and proceeded to conduct a thorough study of the chiroptical properties of re-isolated **34**.⁹⁴ The *S* configuration at C-11 was unambiguously determined by comparing the optical rotation values of the degradation product **35** and the corresponding synthetic compound **36**, obtained from a 6-step sequence starting from (*S*)-citronellol. The other chiral center was determined as 17*R* from comparison of experimental VCD and ECD spectra with those predicted from quantum chemical computation leading to the revised structure

34d. The difference in molecular masses of the originally proposed **34a** and revised **34d** was attributed to air oxidation of **34** which resulted in a molecular ion peak at m/z 374 for **34d** instead of m/z 390 for **34a**.⁹³

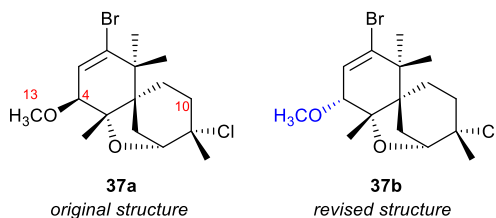


4 Structural revision based on the crystalline sponge method and X-ray crystallography

X-ray crystallography is a powerful tool for structure determination; however, it requires high quality diffracting crystals in sufficient quantity.^{48, 95} An alternative that has recently been proposed, the crystalline sponge method, utilizes a porous metal framework composed of tris(4-pyridyl)1,3,5-triazine and a metal salt to hold the target molecule of interest for crystallographic analysis, eliminating the need to obtain crystals.⁹⁶ Both methods have found use in structural revision studies, as demonstrated in the subsequent case studies.

4.1 Cycloelatanene A-B

Cycloelatanene A and B, two C₁₆ chamigrenes, were isolated from the Australian marine alga *Laurencia elata*.⁹⁷ A preliminary chemical profiling using HPLC-NMR spectroscopy resulted in the isolation of the two C-4 epimeric cycloelatanenes, described as A (**37a**) and B (**37b**). Their structures were proposed from NMR spectroscopic analysis with the relative configuration established by selective 1D NOE irradiations. In a recent study, the structures were revisited using the crystalline sponge method,⁹⁸ which is suitable for rigid, nonpolar molecules that are smaller than the cross-sectional area of pores of the host complex.⁹⁹ X-ray diffraction analysis of the molecules trapped inside the ordered cavities of the crystalline sponge [(ZnI₂)₃ (tpt)₂ (cyclohexane)_x] (tpt = 2,4,6-tris-(4-pyridyl)-1,3,5-triazine) confirmed a rigid tricyclic framework for the cycloelatanenes and ultimately structure **37a** was revised to **37b** and **37b** was revised to **37a**. A close evaluation of the reacquired 1D NOE spectroscopic data showed key correlations between H-10 and H₃-13 for **37b** and between H-10 and H-4 for **37a**, further supporting the revision (Fig. 6).



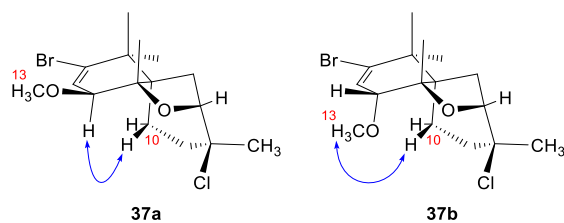
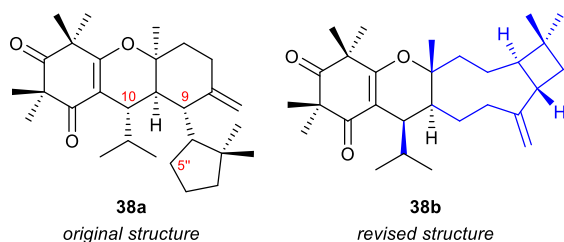
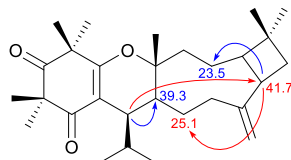


Fig. 6. 1D NOE NMR signal enhancement⁹⁷ favoring **37b** as the correct structure of cycloelatanene A and **37a** for cycloelatanene B.

4.2 Myrtucommulone K

Myrtus communis L, a Mediterranean evergreen shrub, has been traditionally used as an antiseptic, disinfectant, and hypoglycemic agent.¹⁰⁰ A variety of bioactive compounds have been isolated from *M. communis* including the cytotoxic phloroglucinol derivative myrtucommulone K (**38**).¹⁰¹ Structure **38a** was derived through MS and NMR spectroscopy, relying on two critical HMBC correlations from H-10 to C-9 and from H-9 to C-5". When **38** was later re-isolated, X-ray crystallographic analysis led to revised structure **38b**.¹⁰² The MS, ¹H, and ¹³C NMR spectroscopic data originally published for **38a** were identical to those obtained for **38b**, prompting the structural revision of **38**. Recently, a biomimetic synthesis of **38** was reported with agreement between NMR spectroscopic and X-ray crystallographic data of synthetic and natural **38b**.¹⁰³ Some originally reported HMBC correlations cannot be rationalized in light of the new structure since the hydrogens and the carbons are separated by 4–6 bonds.¹⁰¹ However, due to the wide spectral window necessary to record a full HMBC spectrum for this compound (0-220 ppm in the indirect dimension), it is likely that the resulting low resolution led to confusion in the assignment of carbons with similar chemical shifts (Fig. 7).





38b

Fig. 7. Improbable HMBC correlations observed in the original report (red) of **38a** transposed on the revised structure and possible correlations with which they were confused (blue). Numbers are ^{13}C chemical shifts.

5 Structural revision based on NMR spectroscopic analysis and empirical rules

Quantum chemical computation of spectroscopic parameters as a tool for structural revision is in vogue but it requires powerful computing resources. Sometimes, a careful analysis of NMR chemical shifts and coupling constants of a series of analogues is sufficient to assess the precise configuration of a compound. Identifying potential structure misassignments can be enabled by application of accepted empirical rules. The following proposed revisions illustrate how an understanding of basic principles of NMR spectroscopic analysis and fundamental organic chemistry facilitate correction of structural errors.

5.1 2β -Hydroxynagilactone F and nagilactone I

The bioactive compounds 2β -hydroxynagilactone F (**39**) and nagilactone I (**40**) were originally isolated from the root bark of the Japanese evergreen *Podocarpus nagi*.^{104, 105} The proposed structures **39a** and **40a** included an α -hydroxy at C-2, which was established from the half-height width of the H-2 NMR signal and the assumption that ring A adopted a chair conformation (Fig. 8A).¹⁰⁶ Recently, **39** and **40** were re-isolated from Ethiopian *P. falcatus* and were found to be active against human colorectal adenocarcinoma.¹⁰⁷ Conformational analysis using MM2 force field suggested that a boat conformation at ring A for epimers **39a** and **39b**, with 2α - and 2β -hydroxy groups, respectively (Fig. 8B), would have lower energy than the originally proposed chair conformation.¹⁰⁷ This change in conformation required that the dddd H-2 signal ($J = 12.9, 10.0, 7.2, 5.1$ Hz) was axially positioned, thus supporting the β -oriented C-2 hydroxy group in revised **39b**. The ^1H and 2D-NOESY NMR spectra were analyzed to further support the revised orientation of the hydroxy group at C-2 and X-ray crystallographic analysis was used to confirm this new assignment. The absence of a NOESY correlation due to a 1,3-diaxial interaction between H-2 and the methyl group at C-10 should have raised red a flag on the validity of the original structure **39a**.¹⁰⁵

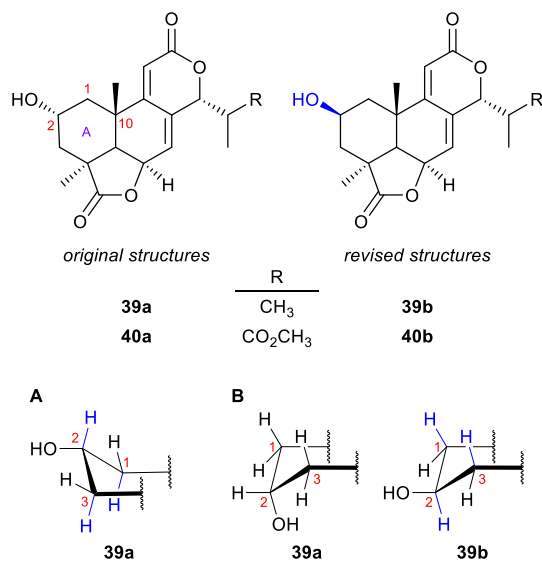
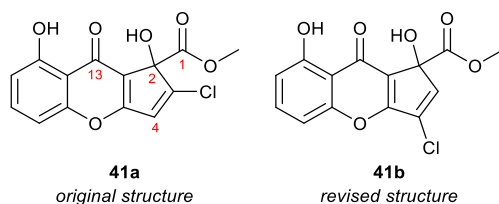


Fig. 8. (A) Chair conformation of ring A with 2 α -hydroxy group (**39a**). (B) Boat conformation of ring A with 2 α - (**39a**) and 2 β - (**39b**) hydroxy groups. Bonds in blue are near 180° torsional angles leading to strong J -coupling.

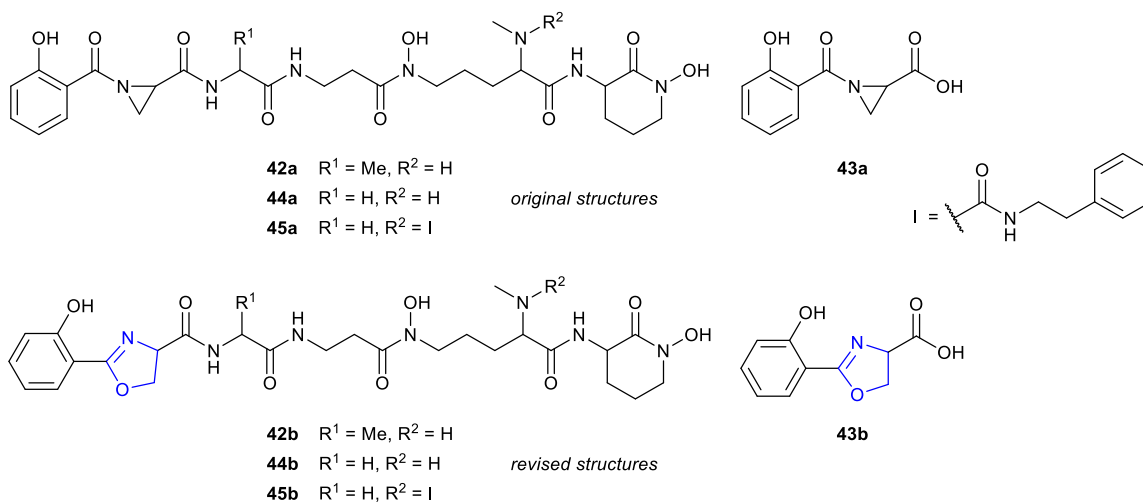
5.2 Coniothyrione

Coniothyrione (**41**) is a chlorinated cyclopentadienylbenzopyrone antibiotic produced by *Coniothyrium cerealis*, an ascomycete obtained from decaying livestock manure. A mechanistic assay involving interaction with the ribosomal protein small subunit D was used to guide the isolation of **41**.¹⁰⁸ Lack of HMBC correlation between olefinic H-4 and the methoxycarbonyl C-1 was used to support the proposed structure **41a**; however, the absence of a correlation does not affirm that they are necessarily more than three bonds apart.¹⁴ Furthermore, as the C-3/C-4 double bond is conjugated to a ketone at C-13, the assignment of C-3 at δ_C 127.2 and C-4 at δ_C 143.1 was unconvincing.¹⁰⁹ The structure was revised to **41b** by interchanging δ_C between C-3/C-4 along with the position of chlorine, whereas the absence of a three bond HMBC correlation from H-3 to C-1 was attributed to an unfavorable dihedral angle. Biosynthetic analysis was also used to support the revised structure **41b**. Although the biosynthetic claim was disputed, the revised structure **41b** was ultimately confirmed based on 1,1-ADEQUATE, $^1J_{CC}$ -edited HSQC-1,n-ADEQUATE, J -modulated ADEQUATE experiments and computational studies.¹¹⁰ In a recent study the structure was confirmed using ACD/Structure Elucidator in combination with DFT calculations.¹⁴



5.3 Madurastatins

Madurastatins are siderophores produced by bacteria and fungi that facilitate uptake of essential metals into bacterial cells.¹¹¹ It has been well established that siderophore-dependent iron sequestration is essential for the virulence and pathogenicity of some bacteria and fungi.¹¹² Antibacterial madurastatins A1 (**42**) and B1 (**43**) are produced by *Actinomadura madurae*,¹¹³ whereas madurastatin C1 (**44**)¹¹⁴ and MBJ-0035 (**45**)¹¹⁵ were isolated from *Actinomadura* sp. and *Streptosporangium* sp., respectively. Their structures (**42a–43a**), proposed based on NMR and mass spectral analysis, featured a unique aziridine ring thought to be essential for their antibacterial activity.¹¹³ The aziridine ring in the original structures **42a–45a** was derived from HMBC correlations between α -proton H-19 and β -methylene protons H₂-20 to carbonyls C-18 and C-21 (Fig. 9A).¹¹³ However, these HMBC correlations are equally justified by a dihydrooxazole moiety (Fig. 9B), which was recently observed in spoxazomicin C and D.¹¹⁶ An aziridine ring carbon has an expected δ_C of 33–44 ppm while dihydrooxazole δ_C would be expected further downfield. The experimental values for C-19 and C-20 in **42a** were 67.2 and 69.4 ppm, respectively, favoring the dihydrooxazole moiety.¹¹³ In a recent study, the revised structures were further confirmed by partial synthesis of an aziridine ring, a dihydrooxazole moiety, and comparison of δ_C at C-19 and C-20.¹¹⁷



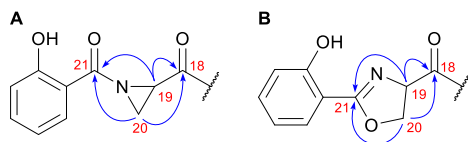
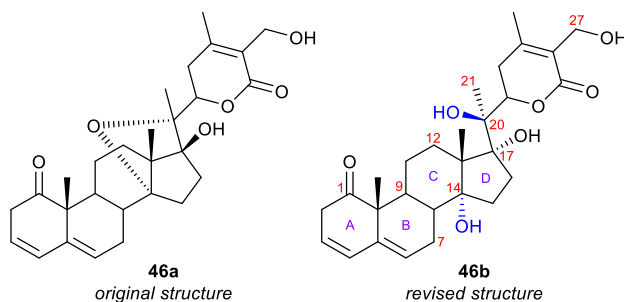


Fig. 9. HMBC correlations for aziridine moiety present in the original proposals (A) and dihydrooxazole present in the revised structures (B).

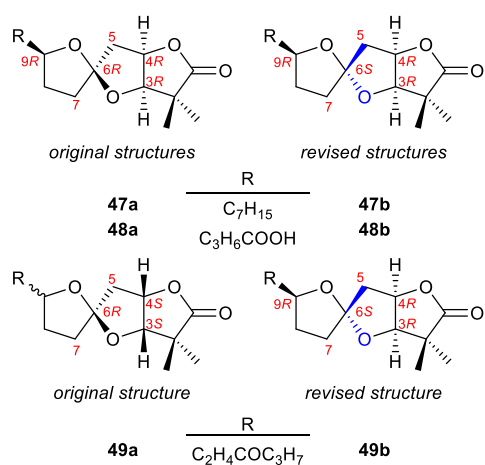
5.4 Coagulin

There are over 900 naturally occurring withanolides originating from numerous genera of the plant family Solanaceae that exhibit biological activities including anti-inflammatory, immunomodulatory, antifeedant, and antitumor.^{118,119} Withanolides are steroids that contain a characteristic highly oxygenated ergostane backbone. Comparison of ¹³C NMR spectroscopic data from known withanolides whose structures have been well established by X-ray crystallography and 2D NMR spectroscopy enabled revisions to 29 withanolide structures.¹²⁰ The observations suggested that the δ_C of C-7, C-9, C-12, and C-21 are shifted, some upfield and others downfield, by the presence of a hydroxy group at C-14 or C-17 via the γ -gauche effect and that the configuration of the hydroxy groups at C-14 and C-17 can be determined from the δ_C at C-9, C-12, and C-21. Coagulin (**46**), a product of the medicinal plant *Withania coagulans*,¹²¹ was suggested to contain an uncommon C-14/O/C-20 bridge (**46a**), based on the degree of unsaturation as determined by EIMS and the δ_C of three oxygenated quaternary carbons at C-14, C-17, and C-20. The proposed linkage for **46a** requires that the 17-hydroxy group be in a β -orientation, but it was revised to be in a α -orientation as indicated by the upfield chemical shift of C-12.¹²⁰ Further, the ¹³C NMR spectroscopic data for positions 1–22 of **46** were almost identical to those of a known analogue, withanolide K. Since **46** contained the same steroid nucleus as withanolide K, the structure was revised to 27-hydroxywithanolide K (**46b**). It should also be noted that the authors of the original paper mistakenly concluded the observed $[M - H_2O]^+$ to be the molecular ion of **46**, resulting in an inflated value for degrees of unsaturation.¹²⁰



5.5 Cephalosporolides and penisporolide

Cephalosporolides H (**47**) and I (**48**) and penisporolide B (**49**) are [5,5]-spiroacetal-*cis*-fused- γ -lactones originally described following their discovery from a marine-derived *Penicillium* fungus.¹²² The proposed stereochemistry for cephalosporolide H (**47a**) was established by NOESY correlations from H-5a to H-4 and from H-5b to H-7. Penisporolide B was shown to have structure **49a** which, if the lateral chain is neglected, is a diastereoisomer of **47a** and **48a**. In all cases, the relative configuration of the chiral centers at C-3, C-4 and C-6 was supported by NOESY data, but the configuration at C-9, which is relatively distant from positions 3, 4, and 6, was not discussed. Recently, NMR spectroscopic data of the four possible relative stereoisomers of analogues of **47**–**49** (R = Me) were examined.¹²³ By searching for significant patterns in δ_{H} and J values, a decision tree was formulated based on three NMR spectroscopic parameters to determine the relative stereochemistry for this compound class: (i) the scalar coupling constants of H-4 with H-3 and H₂-5; (ii) the difference in δ_{H} between the two H-5 protons, which allows determination of the configuration at C-6 relative to the orientation of H-3/H-4; and (iii) the difference in δ_{H} between both protons at position 8 which determines the configuration at C-9, based on arguments of steric compression.¹²⁴ To strengthen the model, the chemical shifts of C-3 and C-7 were also considered. The decision tree was used to confirm the assignment of 21 such molecules from different sources and to revise the structures of cephalosporolide H (**47a**) and I (**48a**) and penisporolide (**49a**) to **47b**, **48b** and **49b**, respectively.



5.6 Cryptospirolepine

The ‘Crews rule’ states that the structure of a molecule is expected to be especially difficult to solve when the ratio of the number of hydrogens to carbons is fewer than 1, since NMR spectroscopic methods are so dependent on detection through protons.¹²⁵ The polyaromatic alkaloid cryptospirolepine (**50**) obtained from the West African shrub *Cryptolepis sanguinolenta*, certainly falls into this category.¹²⁶ In addition to being proton-deficient, its yield was minimal, increasing the challenging nature of the original structure determination. The validity of structure **50a** was called into question when a sealed NMR tube containing degraded product **52** (Fig. 10) was evaluated by members of the same research group 12 years after its report.¹²⁷ In the originally proposed **50a**, vinyl proton H-13 was placed near a four-spin aromatic system based on a ROESY correlation with H-12. Additionally, a strong HMBC correlation was observed between H-12 and C-13a, which is unlikely as they are four bonds apart (Fig. 10).¹²⁶ Considering these discrepancies, the structure was revised to **50b**, based on long-range HRSQMBC correlations between both *N*-methyl groups and C-2 as well as a weak correlation from H-13 to C-2.¹²⁷ Further analysis of **50** by a 1,1-HD-ADEQUATE experiment established that vinyl C-13 is bonded to carbonyl C-2 and quaternary C-13a. In addition, spiro C-1 was shown to interact with C-13 through a $^2J_{CC}$ correlation. The revised structure **50b** justifies both degradation products as well as the questionable four bond HMBC correlation observed in the original proposal. The revision was recently confirmed by the residual dipolar coupling and residual chemical shift anisotropy analysis.¹²⁸

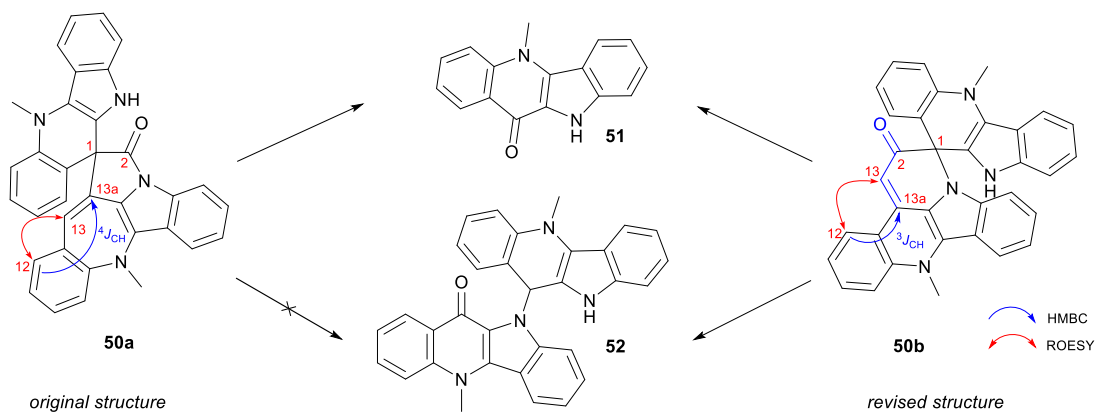


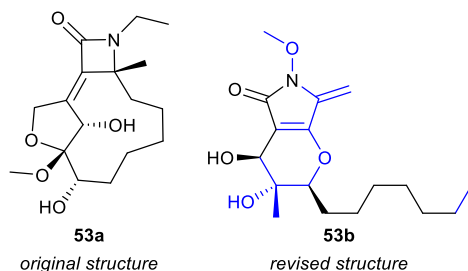
Fig. 10 Degradation products **51** and **52** contributed to structural revision of **50**.

6 Other cases

Sometimes, the solution to a structural elucidation problem can be found using traditional strategies such as incorporation of isotopes for biosynthetic studies or targeted degradation. These approaches offer inexpensive alternatives that can be used when sophisticated computational methods and analytical instrumentation are not available. The two last cases are examples falling into this category.

6.1 Phyllostictine A

Phyllosticta cirsii is a pathogenic fungus that infects the perennial flowering plant *Cirsium arvense* and produces phyllostictine A (**53**).¹²⁹ The molecule was reported to have an unprecedented *Z*- α -(dihydrofuran-3(2*H*)-ylidene)- β -lactam core (**53a**) which sparked interest about the possibility of a novel biosynthetic pathway. Recently, the structure was revised based on biosynthetic studies whereby feeding experiments incorporated ¹³C at either C-1 or C-2 of acetate. This revealed that the distribution of carbon was complex in proposed structure **53a** but logical in revised structure **53b**, which had an alternating pattern for the distribution of ¹³C atoms consistent with a hybrid polyketide synthase and nonribosomal peptide synthetase biosynthetic pathway (Fig. 11A).¹³⁰ Additionally, NMR spectra acquired for re-isolated **53** showed some major discrepancies in HMBC correlations with respect to **53a**. For example, an HMBC correlation from the methyl protons at C-5 to C-11 would have required 7-8 bond coupling if **53a** was correct (Fig. 11B). The new NMR spectroscopic data led to a revision of the structure to **53b**, which contains similarities to other known, fungal metabolites.^{131, 132}



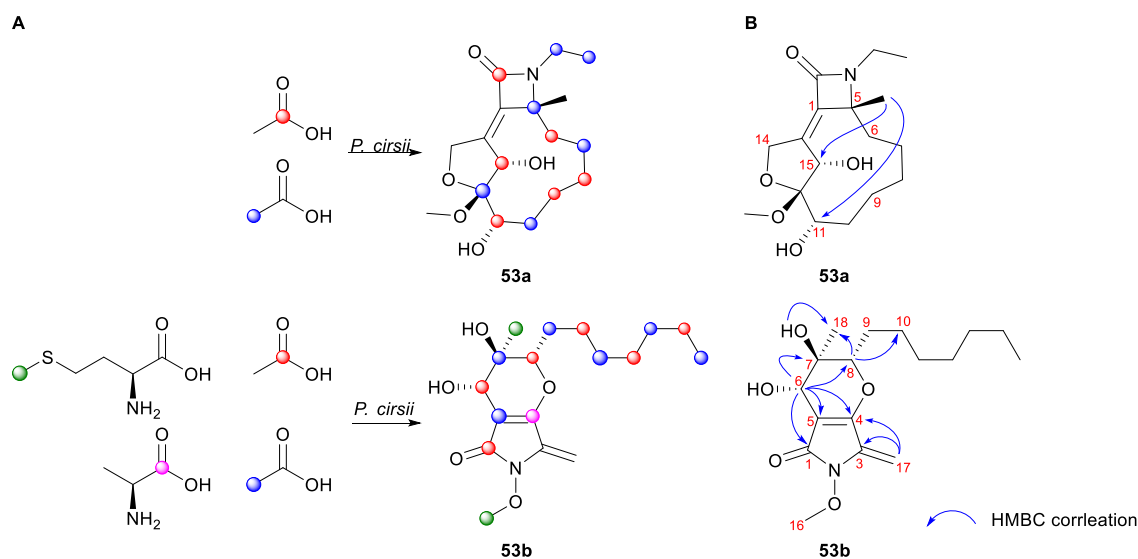


Fig. 11. Labeled carbon distribution (A) and relevant HMBC correlations¹³⁰ (B) projected in original (53a) and revised structure (53b) of phyllostictine A.

6.2 Poecillastrin C

Cytotoxic chondropsin-type macrolide lactams, like poecillastrin C (54), are potent natural products that have been obtained from different genera of deep-sea sponges in very small amounts, making it difficult to fully characterize the structures.¹³³ A challenging structural aspect of these molecules is establishing which of the two carbonyls of the β -hydroxyaspartic acid residue forms the ester linkage and which is a carboxylic acid functionality (i.e., position 1 or γ) (Fig. 12). Assignment of these carbonyl moieties was previously attempted with the related molecule chondropsin A.¹³⁴ The free carboxylic acid of chondropsin A was converted to a methyl ester and an NOE was observed between the newly formed *O*-methyl protons and the oxymethine proton β , which prompted the authors to suggest that C-1 formed the ester linkage. Subsequent structure determination of other chondropsin molecules, including 54, used the suggested ester linkage provided for chondropsin A, exemplified in 54a.¹³³ In a recent study, this strategy was questioned based on the fact that both the α -methine and β -methine protons are close enough to the *O*-methyl protons to produce observable NOEs.¹³⁵ Examination of additional NMR spectroscopic data suggested that the structure of poecillastrin C was as likely to be 54b as 54a. To determine whether position 1 or γ formed the ester bond, a hydride reduction of the lactam was used as a discriminatory tool due to its greater reactivity towards esters relative to carboxylic acids. If 54a was the correct structure, a reduction followed by an acid hydrolysis would lead to hydroxymethyl 55, keeping the γ -carboxylic acid intact while 54b exposed to the same treatment would convert to

hydroxymethyl **56** (Fig. 12). The absolute configuration was established by subjecting the acid hydrolysate of **54** to Marfey's derivatization and comparing it to synthetic standards of β -hydroxyaspartic acid treated with Marfey's reagent. The β -hydroxyaspartic acid residue liberated from poecillastrin C was found to be (2*R*,3*R*)-2-amino-3,4-dihydroxybutanoic acid, supporting the hypothesis that the γ -carboxyl group is esterified, consistent with revised structure **54b**.

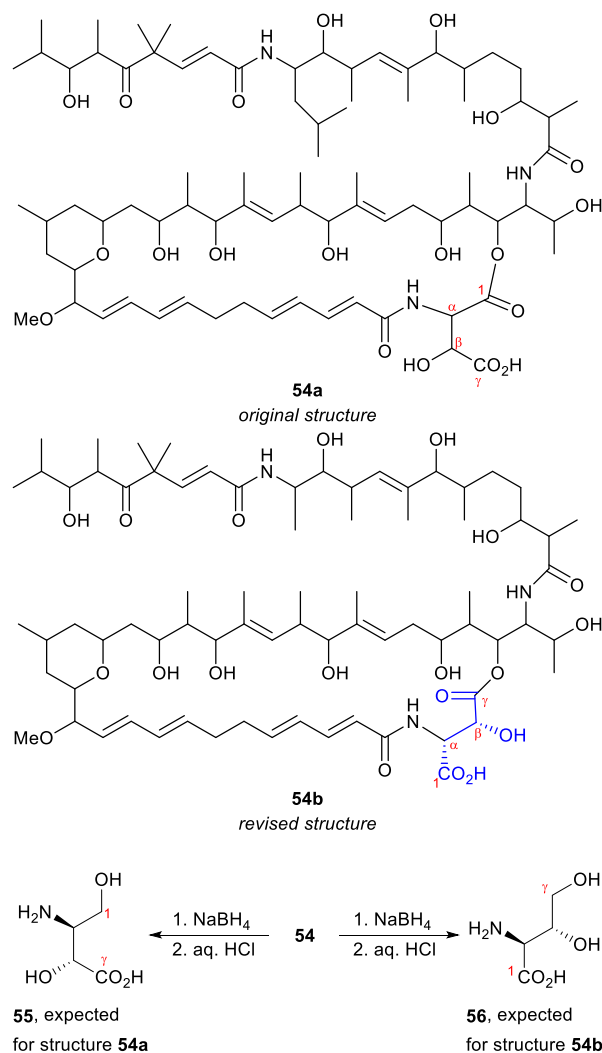


Fig. 12. Reduction and hydrolysis of poecillastrin C (**54**) to determine the orientation of the β -hydroxyaspartic acid moiety.

7 Conclusions

Despite the substantial advances made by modern spectroscopic methods and computational tools, natural product structure determination is still heavily influenced by the observations and conclusions of individual scientists solving structures. In fact, all incorrect structural assignments arise from the failure to recognize other possibilities, out of which the correct structure has the best agreement with the data. Avoiding this trap requires both the appropriate application of available tools and an adjustment in mindset. When one imagines a structural solution for a natural product and assembles the experimental dataset or modeling results to confirm the hypothetical structure, the goal of ruling out hypotheses, rather than proving them, become obscured. Thus, the correct approach is to imagine as many structural solutions as possible for a given case and assemble data and models to test the feasibility of each, leaving only the best hypothesis standing as the true structure. Many of the unique cases discussed herein exemplify the importance of realizing inconsistencies and considering alternative structures that falsify competing hypothesis and satisfy the data. There are several important lessons that can be learned from these cases of mistaken identity. For instance, the identification of unusual structural features should be carefully confirmed to avoid proposing an incorrect structure. In addition, a thorough reading of the literature can provide important information about existing precedents that can guide structural characterization. However, it is important to carefully assess and question literature precedents, as structural errors can be propagated. Extra care should be taken while using derivatization and hydrolysis techniques during structure elucidation as these chemical manipulations can lead to epimerization or unexpected side reactions, as seen with the brevianamides. When NMR spectroscopic data presents the challenge of signal overlap and unsatisfactory signal dispersion, meticulous interpretation in conjunction with computational modeling can aid in avoiding mistakes. Additionally, 2D band-selective NMR spectra that reduce the spectral window to focus on the region of signal overlap can enhance spectral resolution. *J*-based configurational analysis can be used to rule out errors, as seen with the glabramycins. However, for *J*-based analysis and with other NMR data as well, a carefully optimized NMR data acquisition setup with good sample shim is of utmost importance.

Comparison of experimental data to calculated spectroscopic or optical properties such as ^{13}C and ^1H NMR, ECD, VCD and optical rotation, can confirm the proposed structures and resolve errors. Computer-Assisted Structure Elucidation (CASE) algorithms have emerged as a promising tool

that generates a set of possible structures based on experimental NMR features provided. Finally, with computational tools being increasingly accessible, it is imperative that all natural product chemists utilize them as part of structure elucidation, especially in cases where ambiguities exist.

8 Conflicts of interest

There are no conflicts to declare.

9 Acknowledgement

We gratefully acknowledge Dr. William F. Reynolds for his valuable insights. We also thank the U.S. National Institutes of Health (ICBG U19-TW007401) for financial support.

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