

Decarbonylative Approach to the Synthesis of Enamides from Amino Acids: Stereoselective Synthesis of the (Z)-Aminovinyl-D-Cysteine Unit of Mersacidin

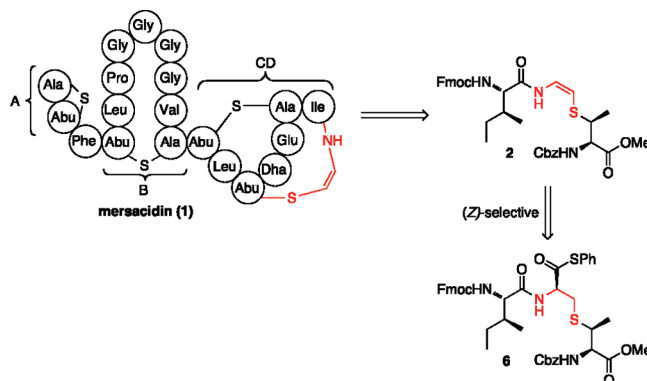
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Received December 20, 2011

ABSTRACT



The Pd- and Ni-promoted decarbonylation of amino acid thioesters proceeds smoothly to yield enamides. The synthesis of the (S)-(Z)-AviMeCys subunit of mersacidin, an MRSA-active lantibiotic, via this approach, is described.

The prevalence of antibiotic resistance poses a significant public health problem and continues to undermine the achievements that have led to the development of our current antibiotic arsenal. Among these agents, “last line” glycopeptide antibiotics, such as vancomycin, have arguably received the most attention in light of the onset of methicillin-resistant *S. aureus* (MRSA), a particularly problematic nosocomial infection. However, the emergence of new resistance mechanisms, coupled with the paucity of novel antibiotics belonging to new structural classes, further underscores the need to revitalize the search for alternative therapeutics.

To address this need, we have initiated a broad research program based on the chemical synthesis and mechanistic study of several peptide antibiotics that inhibit bacterial peptidoglycan (PG) biosynthesis via mechanisms that are orthogonal to that utilized by vancomycin.¹ One

such candidate is mersacidin (**1**), a type B lantibiotic produced ribosomally by *Bacillus* sp. strain HIL Y-85,54728 and whose *in vivo* activity against MRSA is comparable to that of vancomycin.^{2,3} The discovery that mersacidin’s bioactivity is uninhibited by vancomycin despite also forming a 1:1 stoichiometric complex with lipid II suggests that its mode of action is distinct.

Its unique binding site as well as inhibition of the transglycosylation reaction make it an attractive target for development of novel therapeutics because (i) transglycosylation occurs on the cell surface so cellular penetration of the antibiotic is not a requirement for activity and (ii) the transglycosylation reaction utilized in PG biosynthesis is unique to bacterial cells. With the ultimate goal of delineating

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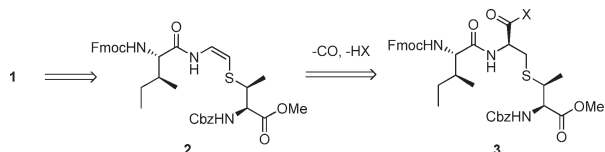
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the exact mode of action of mersacidin and its variants, as well as developing novel analogues with increased efficacy, we have embarked on the total synthesis of **1**.

The presence of the sensitive *S*-[(*Z*)-2-aminovinyl]-3-methyl-D-cysteine (AviMeCys) linkage in the CD ring system, as well as the unnatural amino acid (2*S*,3*S*)- β -methylcysteine (β -MeCys) and (2*S*,3*S*,6*R*)- β -methyl-lanthionine (β -MeLan) fragments in the A, B, and CD ring systems, renders mersacidin a challenging synthetic target. The AviMeCys subunit is present in other bioactive peptides including epidermin,⁴ thioviridamide,⁵ and the recently discovered microbisporicin, now considered the most potent lantibiotic to date.⁶ Still, at the time we initiated our studies, no synthesis of the AviMeCys functionality had been reported.⁷

Although little is known about the function of the enamide subunit, it appears to have a critical role for the biological activity of other agents.^{8,9} As a result, a wide range of methodologies has arisen to address the issues of chemo- and stereoselectivity associated with their chemical synthesis. Given that enamides possess intrinsically moderate reactivity, they may also serve as intermediates for accessing other functionality.^{10,11}

Scheme 1. Retrosynthetic Analysis of the Enamide Unit



Methods that have been developed for the synthesis of enamides include the Curtius rearrangement,¹² Peterson olefination,¹³ vinyl transfer,¹⁴ hydrometalation of alkynes,¹⁵ radical decarboxylation,¹⁶ and radical addition to ynamides.¹⁷ Most recently, several reports of transition-metal-catalyzed cross-coupling reactions between amides and activated

olefins have appeared.^{18–21} Yet, despite its simplicity, this approach is hampered the need for strongly basic conditions, which may not be suitable for all substrates. In addition, these methods may also require access to activated and/or stereodefined olefins.

Given our recent report of the concise synthesis of orthogonally protected β -MeCys and β -MeLan,²² we envisioned the direct synthesis of this AviMeCys unit via decarbonylation of an activated lanthionine precursor (Scheme 1). In our efforts toward the chemical synthesis of mersacidin, we now wish to report our advances in this area, specifically the expeditious synthesis of the AviMeCys unit directly from the β -MeLan unit.²³

The concept of a decarbonylative pathway to enamides is not novel, although its documentation is limited,^{24–26} probably because it has traditionally been viewed as an undesirable side reaction. As a result, we took a systematic approach for optimization of this strategy.²⁷

Our initial approach consisted of evaluating amino acid chlorides in light of Bubner's observation that an enamide byproduct was isolated during the attempted Stille cross-coupling of Fmoc-Pro-Cl.^{24,28} Like Bubner, we also observed the lack of enamide formation in the absence of an organostannane. In search of additives that could enhance reactivity, we found that *n*Bu₄I was very efficient at promoting the catalytic decarbonylation of Fmoc-Pro-Cl, while only traces of the desired product were observed with NaI, presumably an artifact of its low solubility in organic solvents. The efficiency of this process is highlighted by the observation that Fmoc-Pro-OH could be activated *in situ* with diphenylphosphoryl chloride to give high yields of the pyrroline product. Unfortunately, when these conditions were applied to the synthesis of cysteine-derived enamides, only the enamine byproduct arising from β -elimination (i.e., desulfurization) was observed. CuI was added in an effort to sequester thiol byproducts and suppress the potential for poisoning of the Pd catalyst. However, this only increased the amount of the elimination byproduct that was generated. Furthermore, only decomposition was observed in the attempted decarbonylation of Fmoc-Ser(OBn)-Cl and Fmoc-Val-Cl.

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(28) We note that ref 27, highlighting a qualitatively analogous approach using Pd catalysis, appeared during the course of our study. Our method requires lower reaction temperatures (rt with most Ni catalyst systems) and utilized simple phosphine ligands. In addition, we also observed the desired (*Z*)-selectivity for cysteine- and serine-based substrates in contrast to the results reported in ref 27.

Table 1. Scope and Limitations of the Decarbonylative Synthesis of Enamides from *N*-Protected Amino Acids

entry ^a	amino acid thioester	MLn ^b	CuX	<i>t</i> (°C)	Time (h)	product	yield ^c	<i>E</i> : <i>Z</i>
1	Fmoc-Phe-SPh (4a)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		90%	7.1:1
2	Cbz-Phe-SPh (4b)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		82%	4.9:1
3	Cbz-Phe-SPh (4c) ^d	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		77%	7:1
4	Boc-Phe-SPh (4d)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		92%	5.6:1
5	Phth-Phe-SPh (4e)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		33%	<i>E</i> only
6	Phth-Phe-SPh (4e)	120% Ni(PPh ₃) ₄	CuTC	rt	22		94%	<i>E</i> only
7	Fmoc-Trp(Boc)-SPh (4f)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		81%	1:1.4
8	Cbz-Asp(OtBu)-SPh (4g)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		82%	1:3.7
9	Fmoc-Asp(SPh)-OMe(4h)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		trace	---
10	Fmoc-Asp(SPh)-OMe(4h)	120% Ni(PPh ₃) ₄	CuTC	rt	22		93%	---
11	Cbz-Pro-SPh (4i)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		33%	---
12	Fmoc-Pro-SPh (4j)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		41%	---
13	Fmoc-Pro-SPh (4j)	120% Ni(PPh ₃) ₄	CuTC	50	5		89%	---
14	Tos-Pro-SPh (4k)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		46%(68%) ^e	---
15	Boc-Ser(Bn)-SPh (4l)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		42%	<i>Z</i> only
16	Fmoc-Cys(Me)-SPh (4m)	120% Ni(PPh ₃) ₄	CuMeSal	rt	1		91%	<i>Z</i> only
17	Fmoc-Cys(tBu)-SPh (4n)	120% Ni(PPh ₃) ₄	CuMeSal	50	3		53%	<i>Z</i> only
18	Fmoc-Cys(Trt)-SPh (4o)	120% Ni(PPh ₃) ₄	CuMeSal	50	5		0%	---
19	Fmoc-Val-SPh (4p)	15% Pd[P(OEt) ₃] ₄	CuMeSal	50	3		trace	---
20	Fmoc-Val-SPh (4p)	120% Ni(PPh ₃) ₄	CuTC	50	5		73%	---
21	Fmoc-Ile-SPh (4q)	120% Ni(PPh ₃) ₄	CuTC	50	5		67%	1.2:1
22	Boc-α-Me-Val-SPh (4r)	120% Ni(PPh ₃) ₄	CuTC	50	5		0%	---
23	Fmoc-β-Ala-SPh (4s)	120% Ni(PPh ₃) ₄	CuTC	rt	22		82%	---
24	FmocNHPh-CH ₂ CH ₂ CH ₂ COSPh(4t)	120% Ni(PPh ₃) ₄	CuMeSal	rt	20		94% ^f	---
25	BocNH(CH ₂) ₁₀ COSPh(4u)	120% Ni(PPh ₃) ₄	CuTC	rt	22		95%	---
26	Boc-Ala-Phe-SPh (4v)	120% Ni(PPh ₃) ₄	CuMeSal	rt	12		89%	17:1
27	Fmoc-Phe-Cys(Me)-SPh (4w)	120% Ni(PPh ₃) ₄	CuTC	35	18		74%	<i>Z</i> only

^aFor reaction conditions, see Supporting Information. ^bPrepared from either Ni(COD)₂ and PPh₃ or Pd(OAc)₂ and P(OEt)₃. ^cIsolated yields. ^dPrepared in situ with DCC/PhSH. ^eBased on recovered starting material. ^fMixture of isomers (External:Internal (*E*)/Internal (*Z*) = 10:5.8:0.1).

The isolation of dehydroalanine as a byproduct from decarbonylation reactions with Cys-derived acid chlorides, coupled with the lack of generality of this process, prompted us to examine milder activating groups for the amino acid derived esters. Although a range of common activating groups, including -OSu, -OCOR, -OOBt, -OAt, and -OPy, gave only traces of product at best, the 2-pyridyl thioester showed promise. While trace amounts of the desired enamide were observed under Mukaiyama's

conditions,²⁵ added DPEPhos as a ligand with Ni(COD)₂ as the precatalyst yielded 15% of the vinyl sulfide with 37% of the product resulting from sulfur abstraction.

Even more promising was thioester activation,²⁹ which allowed for the efficient decarbonylation of Fmoc-Pro-SPh (**4j**). When these conditions were applied to Fmoc-Cys(Me)-

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SPh, high yields of the decarbonylation product were observed, albeit as the trapped thioaminal, along with reduced amounts of the undesired desulfurization byproduct.

Inspired by Liebeskind's work on the Cu(I) carboxylate activation of thioesters,²³ we sought to limit the propensity for SPh deinsertion by taking advantage of the inherent thiophilicity of Cu(I). Although simple salts, such as Cu(I) and Cu(OAc), showed limited activity, high quantities of the desired enamide product **5** were isolated, along with a significant amount of the sulfur-abstraction product, when copper(I) 3-methylsalicylate (CuMeSal) was used. Similar yields were observed with an SⁱPr ester suggesting that steric hindrance may not play a significant role in the transition state leading to thioaminal formation. Under optimized conditions for decarbonylation of Fmoc-Cys-(Me)-SPh, thioaminal formation was kept to a minimum through use of PPh₃ as a ligand, providing the desired aminovinyl sulfide in 91% yield after 1 h at rt.

With these promising results in hand, we proceeded to examine the substrate scope as summarized in Table 1. We were pleased to find that Pd[P(OEt)₃]₄ efficiently catalyzed the formation of conjugated enamides (entries 1–9).

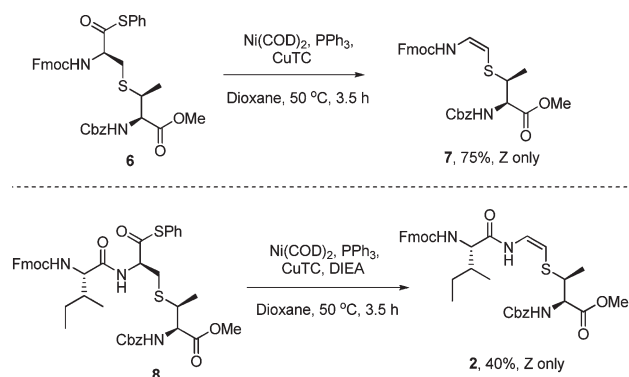
However, moderate yields were observed with amino acids such as Pro (entries 11–14) and Ser (entry 15). In spite of near-quantitative recovery of starting material, enamide yields plateaued to a similar level regardless of whether the reaction was run at or above rt. Moreover, added CuMeSal did not have any positive effect on the yield, while only slightly increased yields were observed when more Pd catalyst was added to the reaction mixture.

In most instances, the Ni-based approach proved to be superior and, in general, yielded enamides in good to excellent yields; some reactions proceeded very efficiently at rt (entries 23–27). This is further highlighted by the contrast between the good yields obtained for the Asp- and Val-derived enamides with Ni that were unattainable with the Pd catalyst (entries 9, 10, 19, 20). To the best of our knowledge, these conditions provide a route to enamides under nonoxidative, nonbasic conditions not reported previously. We did observe, however, that steric hindrance in the substrate may be a limiting factor for successful application of this methodology (cf., entries 18 and 22).

A series of thioesters (**4s–u**) were prepared in order to assess the relationship between the proximity of the protected nitrogen to the thioester and the overall efficiency of the decarbonylation reaction. The data reveal the possibility of a modest trend toward improved yields with increasing distance between the two functional groups. In general, excellent yields of alkene products were observed with simple alkyl thioesters (entries 23 and 24), an indication that this approach may be general for the synthesis of alkenes from thioester precursors.

The decarbonylation of dipeptide substrate **4w** returned an excellent yield of the desired (Z)-enamide. This result

Scheme 2. Synthesis of the AviMeCys Unit of Mersacidin



was critically dependent on the choice of Cu(I) salt and ligand (Table S2, Supporting Information). While reactions utilizing copper(I) thiophene-2-carboxylate (CuTC) reproducibly returned good yields of the desired product, reactions employing CuMeSal resulted in isolation of an alkene byproduct arising from a desulfurization process.

Product selectivity (i.e., enamide vs desulfurization) was also strongly influenced by the choice of ligand. Reactions employing PPh₃ provided the desired enamide in good yields (CuTC additive) while reactions using (*o*-Tol)₃P provide the desulfurization product exclusively. Bulky, as well as sigma-donating, ligands also returned higher yields of the desulfurization byproduct.

Under the optimized conditions, we were pleased to find that lanthionine **6** could be decarbonylated smoothly to yield aminovinyl cysteine **7** in 75% yield (Scheme 2). More importantly, the synthetically more relevant subunit **2** could be prepared in satisfactory yields directly from lanthionine **8** and can be used directly for the synthesis of mersacidin.

In conclusion, we have developed a mild alternative to the synthesis of enamides that makes use of readily available amino acid starting materials. This has allowed efficient access to the AviMeCys fragment of mersacidin. Future work directed toward the total synthesis of mersacidin as well as attempts to determine the function of the AviMeCys subunit will be presented in due course.

Acknowledgment. Financial support provided by the NIH (AI 059327) and Indiana University is gratefully acknowledged. P.G.-R. thanks Eli Lilly and Company for fellowship support. We also thank Professor Lanny Liebeskind (Emory University) and Professor Kenneth Caulton (Indiana University) for helpful discussions.

Supporting Information Available. Experimental details and spectral data for all new compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.

The authors declare no competing financial interest.