Regio- and Stereoselective Isomerizations of Allenamides: Synthesis of 2-Amido-Dienes and Their Tandem Isomerization-Electrocyclic Ring-Closure

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ABSTRACT



A regio- and stereoselective isomerization of allenamides is described, leading to preparations of de novo 2-amido-dienes and a tandem isomerization- 6π -electron electrocyclic ring-closure.

Synthesis of conjugated dienes via an allene isomerization, while a thermodynamically favored process, is not trivial kinetically. The required 1,3-H-shift constitutes a fourelectron $[2\pi + 2\sigma]$ process that would call for an antarafacial approach if proceeding through a concerted and anti-Hückel [or Möbius] transition state.^{1,2} Although impossible in an allylic system, it is relatively more feasible for an allenic system because of the presence of orthogonally oriented p-orbitals of the sp-hybridized central allenic carbon [Scheme 1]. The orthogonal *p*-orbital at C3 [in blue] introduces a formal phase change required for an anti-Hückel transition state, or formally allows a six-electron $[2\pi + 2\sigma + 2\pi]$ process when the second set of allenic π -electrons becomes involved. Nevertheless, the calculated^{2a} ΔE_{act} value remains high at 77.7 kcal mol⁻¹ and consequently, concerted or not, most thermal isomerizations of allenes take place at high temperatures,^{3,4} thereby rendering it difficult to control E/Z ratios of the resulting dienes. There are more practical approaches would involve stepwise processes promoted by acid, base, or metal, but their examples are limited and the level of stereo- and regiochemical control needs to be improved.^{3,5}

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Given that most dienes can be prepared from an array of stereoselective transformations, synthesizing conjugated dienes from structurally more challenging allenes through a kinetically demanding and stereochemically undistinguished isomerization does not appear to be a logical first choice. However, our efforts with the chemistry of allenamides⁶ allowed us to envision a much greater potential in constructing amidodienes through isomerizing allenamides^{7–9} because there are no consistent approaches for synthesizing amido-dienes.^{10–12} Of the two major methods for preparing amido-dienes,¹⁰ the one involving acid-mediated condensations suffers from functional group tolerance with the metal-mediated amidative cross-coupling^{13,14} suffering from limited access to halo-

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dienes [Scheme 1]. In contrast, substituted allenamides are quite accessible through α -alkylations of parent allenamide^{15,16} or amidative cross-couplings of allenyl halides.¹⁷ Their isomerizations can prove to be an invaluable entry to amido-dienes. We communicate here a regio- and stereoselective isomerization of allenamides in the synthesis of 2-amido-dienes and a tandem isomerization- 6π -electron electrocyclic ring-closure.

Screening through various thermal conditions [entries 1-7 in Table 1] including several solvents distinctly revealed that

Table 1. Thermal vs Acidic Conditions

on-Pr ↓		Conditions: Thermal or Acidic		N <u>E</u> 2 n-Pr 2 2		
		acid	temp	time	yield	
entry	solvent	[10 mol %]	[°C]	[h]	$[\%]^{a,b}$	$E:Z^{c}$
1	CH ₃ CN	_	25	16	0	_d
2	CH ₃ CN	_	55	16	51	$\geq 20:1$
3	CH_3CN	-	85	16	88	$\geq 20:1$
4	CH ₃ CN	-	115	16	91 [78]	16:1
5	THF	-	115	16	51	9:1
6	ClCH ₂ CH ₂ Cl	-	115	16	79	7:1
7	Tol	-	150	16	55	4:1
8	CH_2Cl_2	$HNTf_2$	25	$5 \min$	0	_e
9	CH_2Cl_2	PTSA	25	1	66	2:1
10	CH_2Cl_2	$4-NO_2PhCO_2H$	25	16	81	15;1
11	CH_2Cl_2	$PhCO_2H$	25	16	85 [55]	18:1
12	CH_2Cl_2	PPTS	25	16	77	15:1
13	CH_2Cl_2	CSA	25	10 min	95 [74]	18:1
^a NMR yields ^b Isolated yields in the bracket ^c Determined by ¹ H NMP						

^{*a*} NMR yields. ^{*b*} Isolated yields in the bracket. ^{*c*} Determined by ¹H NMR. ^{*d*} Allenamide **1** was recovered. ^{*e*} Allenamide **1** decomposed.

isomerization of achiral allenamide **1** was the most effective at 115 °C in CH₃CN [sealed tube], leading to 2-amido-diene 2^{18} in 78% isolated yield and 16:1 ratio [entry 4] in favor of the *E*-geometry [assigned later]. While there appears to be a solvent effect on the *E/Z* ratio [entries 5–7], we found that with the exception of HNTf₂ and PTSA [entries 8–9], a range of Brønsted acids were equally effective and more facile at RT in providing 2-amido-diene **2** with excellent *E/Z* ratio [entries 10–13].

Generality of this α -isomerization could be established as shown in Table 2. Key features are: (1) An array of chiral allenamides **5**–**7** could be employed to construct *de novo* 2-amido-dienes **8**–**10** with comparable yields and *E/Z* ratios under thermal [higher temperature at 135 °C] or acidic conditions [entries 2–11]; (2) unsubstituted 2-amido-dienes **8d** and **9c** could also be accessed in good yields [see R = H in entries 7 and 9]; (3) allenamide **11** containing an acyclic

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Table 2. Isomerization of Allenamides at the α -Position



^a Unless otherwise noted, CH₂CN was the solvent for thermal conditions. and CH₂Cl₂ was the solvent when using 10 mol % of CSA at rt. For all reactions, concn = 0.10 M.^b Isolated yields.^c Determined by ¹H NMR. ^d Temp started at -78 °C. ^e ClCH₂CH₂Cl was used. ^f MS (4 Å) was used.

amide is also feasible for the isomerization; and (4) a singlecrystal X-ray structure of 10b was attained to unambiguously assign the *E*-configuration [Figure 1].



Figure 1. X-ray structure of 2-amido-diene 10b.

Although our main interest resides in identifying a useful protocol for synthesizing 2-amido-dienes given its greater scarcity,^{10–12,19,20} we examined isomerizations of allenamides from the γ -position en route to more well-known 1-amidodienes.²¹ As shown in Table 3, isomerizations of two types of γ -substituted allenamides, those with a cyclohexylidene group [see 13-16 in entries 1-13], and those with an isopropylidene group [see 17-19 in entries 14-19] led to 1-amido-dienes 20–26 exclusively as *E*-enamides [assigned based on the *trans*-olefinic proton coupling constant].





^a Unless otherwise noted, CH₃CN was the solvent for thermal conditions, and CH₂Cl₂ was the solvent when using 10 mol % of PTSA or CSA at rt. In all reactions, concn was 0.10 M. ^b Isolated yields. ^c Only E isomers were observed. ^d Ninety percent starting allenamide recovered. ^e Seventy percent starting allenamide recovered. ^f Forty-four percent starting allenamide recovered. ^{*s*} Toluene was the solvent. ^{*h*} MS (4 Å) was used. ^{*i*} Decomposition. ^{*j*} NMR yields.

A keen observation here for the γ -isomerization is that acidic conditions appear to be more effective in general with the exception of 17 [entry 15]. In addition, thermal isomerizations at the γ -position required higher temperatures and/ or longer reaction times than those of α -isomerizations. This difference prompted us to explore a possible regioselective isomerization. As shown in Scheme 2, when heating alle-



namides 27a and 27b, containing both α - and γ -substituents, at 135 °C in CH₃CN, isomerizations occurred exclusively at the α -position, leading to 2-amido-dienes **28a** and **28b**²² in 71 and 94% yields, respectively, all in favor of the *E*-enamide [assigned by NOE^{18}]. Isomerization of allenamide 27c took place at RT when in contact with silica gel but again α -isomerization was favored. This regioselective isomerization are both mechanistically intriguing²³ and should be great synthetic value in constructing highly substituted 2-amido-dienes.

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The *E*-selectivity²³ attained from α -isomerization provides an excellent platform for the following important pericyclic transformation. As shown in Scheme 3, isomerization of





 α -allylated allenamide **29** under acidic conditions afforded 3-amido-triene **30** in 86% yield. With the *E*-selectivity, triene **30** is perfectly suited for a thermal 6π -electron electrocyclic ring-closure²⁴ to give cyclic diene **31**. Although only in 35% yield,²⁵ examples of cyclic 2-amido-dienes such as **31** are more rare.²⁶ Allenamide **32a** provided a good example of synthesizing

(22) There appears to be \sim 5% of 1-amido-diene from γ -isomerization.

(23) Without detailed studies, a rationale for lowering of the thermal activation barrier of 1,3-H-shift is the stabilization of the bi-radical intermediate provided by the nitrogen atom, assuming a radical intermediate is considered electron deficient. Based on the this model, this stabilization is direct when isomerizations take place at the α -position [see i], and "vinylogous" for isomerizations at the γ -position [see ii]. Thus, thermal isomerizations at the α -position.



As one reviewer suggested, it is also possible that the nitrogen atom mediates a polarized transition sate in which an increasing charge density at the β -carbon could develop, leading to an N-acyl iminium ion-like character with the migrating hydrogen behaving more like a proton. This charged transition state instead of a neutral one should possess a lower thermal activation barrier for the 1,3-H-shift. Finally, a rationale for the E-selectivity from the thermal α -isomerization is that the *pro-Z*-TS experiences a greater allylic strain than the *pro-E*-TS during the 1,3-H-shift, although we cannot rule out equilibration from Z- to E-enamide after the initial isomerization.

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cyclic 2-amido-diene **34a** via acid-promoted α -isomerization followed by ring-closure. Allenamide **32b** demonstrated that the thermal isomerization could be arrested with the *gem*-dimethyl group in triene **33b** impeding the ring-closure. Unfortunatedly, attempted ring-closure of **32b** at 200 °C led to an unidentified product instead of **34b**.

At last, this process could be rendered in tandem under thermal conditions to access cyclic 2-amido-dienes **34a**, **37**, and **38** in good overall yields directly from respective allenamides **32a**, **35**, and **36** [Scheme 4]. It is noteworthy that these 6π -



electron pericyclic ring-closures mostly took place at 135 °C, which implies an accelerated process. This feature is consistent with related ring-closures of 1,3,5-hexatrienes bearing a C3-donating group.^{27,28}

We have described here a regio- and stereoselective isomerization of allenamides, leading to preparations of *de novo* 2-amido-dienes and a tandem isomerization- 6π -electron electrocyclic ring-closure. Studies involving applications of these dienes and this new tandem process as well as mechanistic understanding of this allene-isomerization are underway.

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Supporting Information Available: Experimental procedures as well as NMR spectra, characterizations, and X-ray structural files are available for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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