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Catalytic photochemical enantioselective α -alkylation with pyridinium salts†

Santhivardhana Reddy Yetra, D Nathan Schmitt D and Uttam K. Tambar D*

We have developed a chiral amine catalyzed enantioselective α -alkylation of aldehydes with amino acid derived pyridinium salts as alkylating reagents. The reaction proceeds in the presence of visible light and in the absence of a photocatalyst via a light activated charge-transfer complex. We apply this photochemical stereoconvergent process to the total synthesis of the lignan natural products (–)-enterolactone and (–)-enterodiol. Mechanistic studies support the ground-state complexation of the reactive components followed by divergent charge-transfer processes involving catalyst-controlled radical chain and in-cage radical combination steps.

Introduction

Carbonyl compounds with α -stereocenters are key components of many biologically active molecules, including pharmaceutical drugs and secondary metabolites. To access this important class of compounds, chemists have developed numerous transformative concepts in asymmetric catalysis for the enantioselective α -alkylation of enolates. While these reactions are often categorized by the mode of catalysis and the enolate precursor, an underappreciated component of these processes is the identity of the alkylating reagents that are coupled with enolates. Commonly employed alkylating reagents include alkyl halides and sulfonates (2, Scheme 1A).

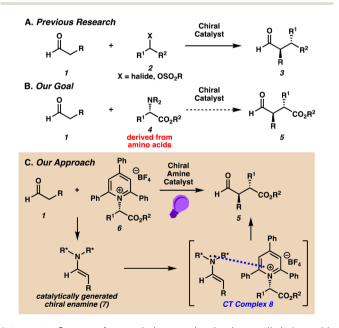
Recent elegant examples of photochemical enantioselective α-functionalizations of carbonyl compounds have been developed with alkyl halides.3-6 Notably, Melchiorre and co-workers pioneered the use of α-bromoketones and benzylic bromides as alkylating agents via light activated charge-transfer complexes.5 We were interested in developing a complementary approach for the catalytic enantioselective α -alkylation of aldehydes 1 based on renewable and sustainable sources of alkylating reagents. We identified amino acid derived substrates 4 as ideal reagents for enantioselective alkylations (Scheme 1B), as they possess several inherent advantages over traditionally used alkyl halides with respect to abundance, stability, versatility, and ease of preparation.7 In light of the poor electrophilicity of amino acid derivatives in enolate alkylations, we were motivated to devise a strategy for the activation of this class of substrates.

Department of Biochemistry, The University of Texas Southwestern Medical Center, 5323 Harry Hines Boulevard, Dallas, Texas 75390-9038, USA. E-mail: Uttam. Tambar@utsouthwestern.edu

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We report a catalytic photochemical enantioselective α -alkylation of aldehydes with amino acid derived pyridinium salts as alkylating reagents (Scheme 1C). These compounds are air and moisture stable crystalline solids that can be easily purified and stored for extended periods of time. Moreover, pyridinium salts can be generated on preparative scale from the facile condensation of amino acid derivatives and pyrylium salts.§

We hypothesized that pyridinium salts could form groundstate encounter complexes with catalytically generated electron rich chiral enolate equivalents. 9,10 Our approach was supported by an early report from Katritzky on the α -benzylation of



Scheme 1 Strategy for catalytic enantioselective $\alpha\text{-alkylation}$ with amino acid derivatives.

diethylmalonate with pyridinium salts of benzylamine, which he postulated proceeds through the formation of light activated charge-transfer (CT) complexes. ¹¹ More recently, Melchiorre has demonstrated the formation of CT complexes between enamines and alkyl halides. ⁵ In our case, the generation of chiral enamine 7 from the condensation of aldehyde substrate 1 and a chiral amine catalyst could form CT complex 8 with pyridinium salt 6, which would then undergo stereoselective C–C bond formation in the presence of visible light.

Our proposal to utilize pyridinium salts in enantioselective α -alkylations is motivated by their storied history as radical precursors. $^{12-16}$ More recently, pyridinium salts have been utilized in deaminative transformations through activation by photoredox catalysis 17 or the formation of CT complexes with electron rich molecules. 18 Despite the widespread application of pyridinium salts as radical precursors, the use of these substrates in catalytic enantioselective transformations is rare, 19 and enantioselective reactions with prochiral enolate equivalents is unprecedented.

Results

We initiated our studies by coupling hydrocinnamaldehyde 9 with Katritzky salt 10a derived from the ethyl ester of glycine as the alkylating agent (Table 1). In the presence of MacMillan's amine catalyst A, 2,6-lutidine, and purple light (390 nm) in CH₂Cl₂, we observed trace amounts of product 11a in 58% ee (entry 1). Interestingly, the deaminated byproduct of Katritzky salt 10a and the corresponding 2,4,6-triphenylpyridine were formed, suggesting the formation of a light activated CT complex. We reasoned that the radical philicity of intermediates generated upon charge transfer between the enamine of aldehyde 9 and pyridinium 10a may not be matched for the desired C-C bond forming event.20 To test this hypothesis, we subjected electron-deficient Katritzky salt 10b derived from the 2,2,2-trifluoroethyl ester of glycine to the reaction conditions (entry 2). Gratifyingly, we obtained α -alkylation product 11b in 36% yield and 60% ee, presumably via a more electron-deficient α-carboxy radical. In a Lewis basic medium such as DMA, the desired product was formed in 40% yield and 92% ee (entry 3). Although other amine catalysts B-D also furnished product 11b (entries 4-6), catalyst A was still best for enantioselectivity.

With enantioselectivity optimized, we focused on improving the yield of the reaction by employing additives that could enhance the ground-state complexation of the reaction components. Ultimately, the addition of stoichiometric NaI resulted in an increase in yield to 65% (entry 7), presumably through the formation of a multicomponent CT complex.²¹ Inclusion of water to help solubilize NaI further improved the yield to 75% while maintaining the enantioselectivity at 92% ee, which represented the optimal conditions for the reaction (entry 8). Interestingly, while α-bromoketones have been demonstrated to be competent acceptors in CT complexes with catalytically generated enamines,⁵ the corresponding 2,2,2-trifluoroethyl ester of α-bromoacetic acid (10c) was not a suitable alkylating agent under our optimized conditions (entry 9). This result highlights a unique advantage of pyridinium derived

Table 1 Optimization studies^a

10b, $R = CH_2CF_3$				11b, R = CH ₂ CF ₃	
Alkylating				Yield ^b	ee ^c
agent	Catalyst	Solvent	Additive	(%)	(%)
10a	A	CH ₂ Cl ₂	_	5	58
10b	A		_	36	60
10b	A	DMA	_	40	92
10b	В	DMA	_	22	5
10b	\mathbf{C}	DMA	_	55	15
10b	D	DMA	_	52	23
10b	A	DMA	NaI	65	92
10b	Α	DMA	Nal, H ₂ O	75	92
10c	A	DMA	NaI,	<5	_
			H_2O		
10b	A	DMA	NaI,	80	46
			H_2O		
10b	A	DMA	NaI,	62	92
			H_2O		
10b	A	DMA	NaI,	18	92
			H_2O		
10b	A	DMA	•	<5	_
			H_2O		
Me	O. Me				
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N Me Bn	$\checkmark_{N} \searrow_{t\text{-B}}$	u `N' `	отмѕ	Br \\O	CH ₂ CF ₃
	Ĥ				2 3
				10c	
	Alkylating agent 10a 10b 10b 10b 10b 10b 10c 10b 10b 10b	Alkylating agent Catalyst 10a A 10b A 10b A 10b B 10b C 10b D 10b A 10c A 10c A 10b A 10b A 10c A 10b A	Alkylating agent Catalyst Solvent 10a	Alkylating agent Catalyst Solvent Additive 10a	Alkylating agent Catalyst Solvent Additive (%) 10a

^a Reaction conditions: 9 (0.30 mmol), 10 (0.1 mmol), catalyst (20 mol%), 2,6-lutidine (0.1 mmol) NaI (0.1 mmol), $\rm H_2O$ (1.0 mmol), 4 °C, 24 h. ^b Isolated yield. ^c Enantiomeric excess determined by chiral HPLC analysis of a lactone derivative (see ESI). ^d Reaction conducted at 23 °C. ^e Irradiation with 370 nm Kessil lamp. ^f Irradiation with 427 nm Kessil lamp. ^g No light.

alkylating agents over the more traditionally used alkyl bromides in enantioselective α -alkylations.

We performed a series of control experiments to gain insight into the reaction (entries 10–13). Conducting the reaction at room temperature instead of 4 °C resulted in a loss of enantioselectivity (entry 10). Irradiation with various wavelengths of light resulted in diminished yields (entries 11–12), which confirmed the importance of activating the ground-state encounter complex at the appropriate wavelength. Furthermore, no product was observed in the absence of light (entry 13).

With optimal reaction conditions identified, we examined the substrate scope of the transformation (Table 2). The Katritzky salt of the trifluoroethyl ester of glycine was coupled with various aldehydes to furnish the desired products in

Table 2 Enantioselective α -alkylation with amino acid derived pyridinium salts^{α}

 a Reaction conditions: aldehyde (0.30 mmol), pyridinium salt (0.1 mmol), catalyst **A** (20 mol%), 2,6-lutidine (0.1 mmol), NaI (0.1 mmol), H₂O (1.0 mmol), DMA (0.1 M), Kessil lamp_{390 nm}, 4 $^{\circ}$ C, 24 h.

synthetically useful yields and greater than 90% ee (13a-d). We obtained products from linear aldehydes (13a-b) and branched aldehydes (13c-d). Although we did not observe any product with phenylacetaldehyde (13e), a carbamate functionalized aldehyde yielded the desired alkylation product (13f). Enantioenriched alkylation products were also formed from the coupling of hydrocinnamaldehyde and Katritzky salts derived from various natural amino acids, such as alanine (13g), valine (13h), phenylalanine (13i), methionine (13j), and tyrosine (13k). In addition, we generated the alkylated product derived from an unnatural amino acid (13l). The products were generated with high enantioselectivity but poor diastereoselectivity, presumably because of the formation of open-shell intermediates (vide infra).

To demonstrate the utility of this new mode of activation with other classes of substrates, we reacted hydrocinnamaldehyde **9** with Katritzky salts derived from various

Table 3 Enantioselective α -alkylation with other amine derived pyridinium salts^a

 a Reaction conditions: aldehyde (0.30 mmol), pyridinium salt (0.1 mmol), catalyst A (20 mol%), 2,6-lutidine (0.1 mmol), CH $_2$ Cl $_2$ (0.1 M), Kessil lamp $_{\rm 427~nm}$, 4 °C, 24 h.

amines (Table 3). Notably, these reactions were performed in CH2Cl2 in the absence of NaI and water. An electronwithdrawing group adjacent to the amine functionality was necessary for reactivity. For example, several 2-aminoacetophenone pyridinium salts were compatible substrates for the enantioselective transformation (15a-f). Through optimization, 427 nm irradiation performed as well as 390 nm for the aminoketone derived pyridinium salts, presumably because of different photophysical properties of the charge transfer complexes. With 427 nm being lower in energy, we chose to move forward with this wavelength. We also observed the desired product derived from aminoacetonitrile (15g). However, the Katritzky salt derived from 3-phenyl-1-propylamine did not yield alkylation product 15h under the reaction conditions. The requirement for an electron-withdrawing group in the pyridinium salt further demonstrates the importance of matching the radical philicities of intermediates generated upon light activated charge transfer.

The synthetic utility of our new catalytic method is highlighted by the enantioselective total synthesis of the lignan natural products (–)-enterolactone 17 and (–)-enterodiol 18 (Scheme 2).²² Under optimal conditions, 3-(3-hydroxyphenyl) propanal 16 and the pyridinium salt of racemic m-tyrosine (12l) reacted in a stereoconvergent process to form the α -alkylated product, which was subjected to reductive conditions without purification to furnish (–)-enterolactone 17 and its epimer (*epi*-17) in 46% yield over 2 steps. Although the diastereomeric lactones were obtained in high ee but poor dr, we recognized the opportunity to epimerize the mixture for the synthesis of more complex structures with high diastereoselectivity.

Scheme 2 Synthesis of lignan natural products via enantioselective α alkylation with pyridinium salts

Therefore, the diastereomers were subjected to LHMDS and TMSCl to yield (-)-enterolactone 17 in 81% yield, 12:1 dr, and 97% ee upon diastereoselective protonation of the enolate intermediate. Reduction with LiAlH4 resulted in the formation of (-)-enterodiol 18 in 70% yield and 97% ee as a single diastereomer. Optical rotations of the synthetic samples of the two natural products also confirmed the absolute stereochemistry of the alkylation products obtained in our enantioselective reaction.

Based on our initial proposal of a light activated CT complex between the pyridinium substrate and a catalytically generated enamine (Scheme 1C), we performed a series of experiments to gain insight into the mechanism of the photochemical process. The subjection of either enantiomer of Katritzky salt 12g to the optimized reaction conditions with aldehyde 9 resulted in the stereoconvergent formation of the same major enantiomer of both diastereomers of product 13g (Fig. 1A). These experiments with both enantiomers of Katritzky salt 12g, in combination with the absence of product formation in the presence of 1 equivalent of TEMPO,23 are consistent with the formation of αcarboxy radical 19 as a common intermediate, which also accounts for the low diastereoselectivity in the transformation.

We were also interested in gaining insight into the role of NaI in the catalytic process. The use of 50 mol% NaI led to 58% isolated yield of the coupled product 11b. In addition, the pyridinium substrate 10b was not converted to the α-iodoester in the presence of NaI and absence of aldehyde.23 However, NaI caused a bathochromatic shift into the purple region of the absorption spectrum (Fig. 1B). Moreover, the inclusion of NaI had a profound impact on the equilibrium of enamine formation, presumably by affecting the equilibrium of CT complex 21 (Fig. 1B). Therefore, we believe NaI may affect the identity of the CT complex by forming a ternary complex with the catalytically generated enamine and pyridinium substrate.24

Next, we used the trans and cis isomers of the radical probe 22 as the aldehyde component to determine whether the enantioselective reaction proceeds through a radical chain or

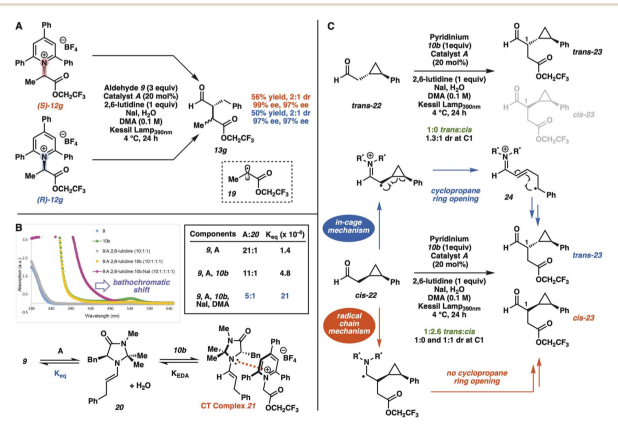


Fig. 1 Mechanistic experiments.

Scheme 3 Proposed mechanism.

in-cage radical combination process (Fig. 1C and S7†). ^{4g,5a} Radical probe *trans*-22 exclusively formed the alkylation product *trans*-23 as a mixture of diastereomers at C1, which is consistent with either a radical chain or in-cage radical combination. With radical probe *cis*-22, the expectation was that an in-cage radical process would exclusively furnish the thermodynamically stable alkylation product *trans*-23 *via* acyclic intermediate 24. ^{5a} Alternatively, the alkylation product *cis*-23 would form exclusively if the reaction proceeded through a radical chain. ^{4g} Surprisingly, starting from radical probe *cis*-22, we isolated both *trans* and *cis* isomers of alkylation product 23 in a 1:2.6 ratio.

These observations suggest that the catalytic enantiose-lective reaction may proceed simultaneously through two highly enantioselective processes (Scheme 3 and Fig. S7†): an in-cage radical combination mechanism (path a) and a radical chain mechanism (path b). Although we cannot rule out the possibility of a radical chain mechanism with cyclopropane ring opening as an off-cycle process, we believe the measured quantum yield of 4 may be more consistent with the coexistence of two distinct mechanisms.²³

Conclusions

In summary, we have developed a catalytic enantioselective alkylation of aldehydes with pyridinium salts derived from amino acids and other α -stabilized amines. The reaction is enabled by a visible light activated CT complex between electron-deficient pyridinium salts and electron-rich components of the reaction. The mild conditions are compatible with several functional groups, enabling the enantioselective synthesis of lignan natural products. Future studies will

examine the mechanism of this process in more detail. We anticipate this approach may be extended to the photochemical catalytic enantioselective alkylation of several classes of carbonyl compounds with pyridinium salts based on other modes of catalysis.

Data availability

The datasets supporting this article have been uploaded as part of the ESI. \dagger

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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