

Organocatalytic Multicomponent Synthesis of α/β -Dipeptide Derivatives

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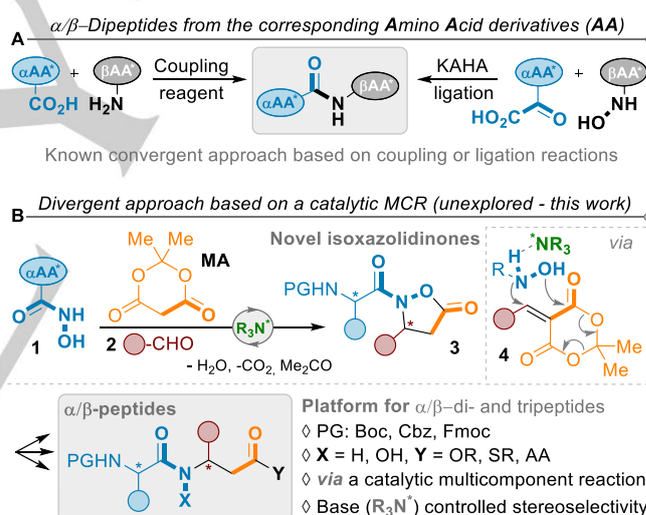
Dedication ((optional))

Abstract: A straightforward multicomponent Knoevenagel-aza-Michael-Cyclocondensation reaction involving readily available hydroxamic acid-derived from naturally occurring α -amino acids allows a diversity-oriented synthesis of novel isoxazolidin-5-ones possessing an *N*-protected α -amino acid pendant with good to high diastereoselectivities thanks to a match effect with a chiral organocatalyst. These diversely substituted heterocycles, easily isolated as a single diastereoisomer, proved to be versatile platforms for the formation of an array of α/β -dipeptide fragments.

Naturally occurring peptides currently hold a privileged place for the development of bio-inspired ingredients in pharmaceutical industry.^[1] In order to overcome the inherently sensitivity to proteolytic degradation of native peptides, while giving the opportunity to afford new properties and bio-tools to probe protein functions, synthetic chemists have tackled the elaboration of peptide analogues. In that peptidomimetic context, β -amino acids proved to be excellent building blocks for the elaboration of either β -peptides and hybrid α/β -polyamino acids having not only an improved stability towards peptidases but also displaying unique secondary structures and pharmacological activities.^[2] Furthermore, the α/β -dipeptide fragments are not only a key elements of naturally occurring products,^[3] but have met important successes in medicinal chemistry either in a linear form in case of cardiovascular diseases target for instance,^[2b] or to construct cyclic-peptide architectures in which the 1,4-diazepane-2,5-diones, and heterocycles derived thereof, proved to be privileged platforms in a large array of pharmaceutical applications.^[4] The attractiveness of these frameworks has fueled numerous research efforts dealing with the asymmetric synthesis of β -amino acids, and the elaboration of original derivatives is still of high added value.^{[3a],[5]}

The synthesis of the α/β -dipeptide motif essentially relies on the construction of the amide bond by means of coupling reagents, essentially stoichiometric, between suitably (orthogonal) protected and pre-elaborated α -amino acids (α AA) and β -amino

acids (β AA) derivatives (Scheme 1A).^[6] Bode and co-workers have made a major achievement in this field, by developing the α -ketoacid-hydroxylamine (KAHA) ligation between two pre-synthesized AA derivatives, one bearing a α -keto acid pendant and another possessing an hydroxylamine *N*-terminus functional group.^[7] Recently, Takemoto expended the portfolio of hydroxylamine-derived β AA to aspartic acid derivatives thanks to an organocatalytic aza-Michael addition to fumaric monoacid electrophiles.^[8] This elegant strategy, making use of hydroxylamine functions as key building blocks for the elaboration of β AA, was applied to the elaboration of α/β -dipeptides albeit with moderate diastereoselective ratio (dr).



Scheme 1. A new entry to β -amino acid derivatives.

Besides these convergent sequences, we tackled an alternative divergent approach in order to populate the chemical space within the biorelevant α/β -dipeptide series (Scheme 1B). We reasoned that the multicomponent Knoevenagel-aza-Michael-Cyclocondensation (KMC) reaction would take place between readily available hydroxamic acids **1**,^[9] derived from naturally occurring and enantiopure α AA, various aldehydes **2** and Meldrum's acid (MA) as a C2-synthon (first point of diversity).^{[10],[11]} Then, this strategy would afford a straightforward elaboration of novel isoxazolidin-5-ones **3**,^[12] as masked β AA with an α AA side chain, providing eventually versatile platforms to elaborate a library of α/β -dipeptides (second point of diversity).^{[13],[14]} In order to be synthetically useful, this sequence should demonstrate a significant functional group compatibility and furnish one stereoisomer. Thanks to the marked electrophilic

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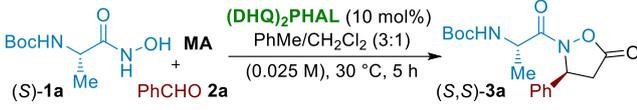
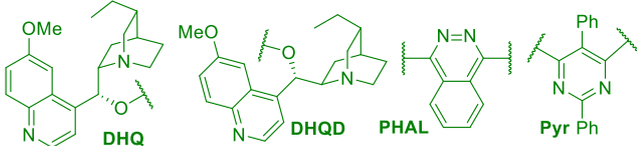
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reactivity of alkylidene **MA** intermediates **4**,^{[11],[15]} we postulated that (1) a facile domino addition process of hydroxamic acids **1** would occur and (2) the C-N bond formation between **1** and **4** would be under the influence of both a chiral nucleophile **1** and a Brønsted base organocatalysts to favor the formation one stereoisomer (Scheme 1B). We are delighted to report on the unprecedented diastereoselective multicomponent reaction (MCR) giving rise to versatile isoxazolidin-5-ones **3** thanks to the match influence of a suited organocatalyst, en route to the elaboration of various original α/β -dipeptide derivatives.

Table 1. Proof of principle and optimization.^[a]

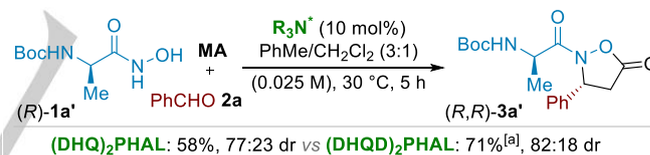



Entry	Deviation from the standard conditions	Yield [%]	dr ^[b]
1	Without catalyst	20 (55) ^[c]	57:43
2	Quinuclidine as catalyst	86	71:29
3	-	75	86:14
4	(DHQD) ₂ PHAL as catalyst	82	74:26
5	(DHQ) ₂ Pyr as catalyst	62	82:18
6	Quinine (QN) as catalyst	73	81:19
7	Quinidine (QD) as catalyst	66	48:52
8	MeCN (0.1 M) as solvent, 20 °C ^[d]	44	73:27
9	CH ₂ Cl ₂ (0.1 M) as solvent, 20 °C ^[d]	30	86:14
10	PhMe (0.1 M) as solvent, 20 °C ^[d]	traces	-
11	PhMe/CH ₂ Cl ₂ (3:1, 0.1 M) as solvent, 20 °C ^[d]	68	86:14

[a] Reaction conditions: **1a** (0.1 mmol), PhCHO **2a** (1 equiv), **MA** (1.5 equiv) in PhMe/CH₂Cl₂ (3:1, 0.025 M) at 30 °C for 5 h. Yields of both diastereoisomers determined by ¹H NMR with an internal standard. [b] dr determined by ¹H NMR on the crude product. [c] Bimolecular reaction from benzylidene Meldrum's acid **4a**. [d] **MA** (1 equiv), (DHQ)₂PHAL (20 mol%).

At the onset, it was shown that the model MCR involving the *N*-Boc alanine hydroxamic acid **1a** (Boc- α -AlaNH₂OH), **MA** and benzaldehyde **2a** furnished after 5 hours the corresponding isoxazolidinone **3a** in 20% yield (determined by ¹H NMR) as 57:43 mixture of diastereoisomers namely with virtually no stereoselection (entry 1, Table 1). By performing the di-component reaction with benzylidene Meldrum's acid **4a**, the putative intermediate involved into the formal domino aza-Michael-cyclocondensation process (see Scheme 1), similar poor dr was obtained with a better 55% yield (entry 1). The facility with which this MCR takes place is worthy of note, likely thanks to the high electrophilic reactivity of benzylidene MA,^[15] but the ability of a catalyst to accelerate this process and promote a stereoselective sequence remains an issue.^[10] To our delight the use of 10 mol% of a Brønsted base like the achiral quinuclidine allowed to improve significantly the yield to 86% and dr (71:29,

entry 2). A large screening of organocatalysts (see SI) showed that the commercially available and chiral amine (DHQ)₂PHAL turned out to be the more efficient catalyst leading to good 86:14 dr in 75% yield in 5 hours (entry 3 versus entries 4-7). Early optimization endeavors (see SI) showed that less polar solvents like CH₂Cl₂ (entries 8-9) benefited to the level of stereoselection, while a mixture of PhMe/CH₂Cl₂ was required for solubility issue (entries 10-11). Noteworthy, isoxazolidone **3a** proved to be somewhat unstable during the purification by silica-gel column chromatography but also in the presence of the Brønsted base in solution (*vide infra*). Then, some decomposition events occur during longer or forcing reaction conditions leading to erratic outcomes. However, the use of more diluted conditions and a slight excess of **MA** prevent the decomposition of product **3a** (thanks to its high acidity, pK_a = 4.8 in H₂O)^{[11],[16]} even at 30 °C, a temperature allowing both a faster process and a decrease of the amount of catalyst from 20 to 10 mol% while giving improved yield of 75% and the same dr in 5 hours (entries 3 vs 11, Table 1). This shows the key role of **MA** to secure soft reaction conditions. Interestingly, it was observed a match/mismatch effect between chiral Cinchona-derived organocatalysts and (S)-Boc- α -AlaNH₂OH **1a** (Table 1, entries 3-4 and 6-7) which demonstrated that quinine versus quinidine derivatives were more competent to give (S,S)-isoxazolidinone **3a** with a good dr. Accordingly, it was proven that quinidine derived (DHQD)₂PHAL catalyst was able to transform the (R)-Boc- α -AlaNH₂OH enantiomer **1a'** into isoxazolidinone (R,R)-**3a'** in 71% yield and 82:18 dr contrary to (DHQ)₂PHAL catalyst giving **3a'** only in 58% yield and 77:23 dr (Scheme 2). Eventually, it was validated that this MCR took place with no racemization and provided the major isoxazolidinones **3a** and **3a'** as a single enantiomer (>99:1 er, see SI).



Scheme 2. Molecular diversity towards a versatile synthesis of **3a'** - the other enantiomer of isoxazolidinone **3a**. Yields of both diastereoisomers determined by ¹H NMR with an internal standard. [a] 47% isolated yield after column chromatography.

The scope and limitation of this novel MC-strategy was next addressed (Table 2). Pleasingly, carrying out the reaction for 24 hours on a larger scale secured the complete transformation of Boc- α -AlaNH₂OH **1a** into isoxazolidinone **3a** in 88:12 dr (99% of NMR yield for both diastereoisomers). Importantly, the two diastereoisomers of isoxazolidinone **3a** could be separated by flash column chromatography, as a general behavior in this series, leading to the major (S,S)-**3a** in 71% isolated yield (slight drop of the theoretical yield of 88%).^[17] Starting from isovaleraldehyde **2b**, a facile synthesis of the corresponding isoxazolidinone **3b**, having β Leu backbone (homologated β^3 -Leucine), occurred in excellent dr > 95:5 and 96% yield for the pure major stereoisomer. As a rule of thumb, in comparison to the use of aromatic aldehydes such as **2a**, the isoxazolidinones **3** derived from aliphatic aldehydes **2** turned out to be more stable which allows (1) an easy purification by chromatography and (2) to carry out the reaction in higher

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concentration.^[18] The MCR was applied successfully to *N*-Cbz- α AlaNH₂ **1b** (**3c-3d**, 64-85%, 83:17-89:11 dr)^[19] and *N*-Fmoc- α AlaNH₂ **1c** (**3e**, 86%, 93:7 dr) to provide the corresponding **3c-3e** with good to excellent stereoselectivities and yields. Then, various β hLeu-derived isoxazolidinones were constructed from isovaleraldehyde and Boc-protected proteinogenic α AA such as α Phe (**3f**, 71%, 89:11 dr), α Val (**3f**, 93%, >95:5 dr) and α Pro (**3i**, 63%, 75:25 dr) with yields ranging from 63% to 93%. It was also shown that *N*-Boc glycine hydroxamic acid afforded the corresponding isoxazolidinone **3h** in excellent 93% yield but low 58:42 er. In line with previous observations, this highlights a moderate capability of (DHQ)₂PHAL catalyst to promote an enantioselective process from an achiral α AANHOH.^{[10a],[20]}

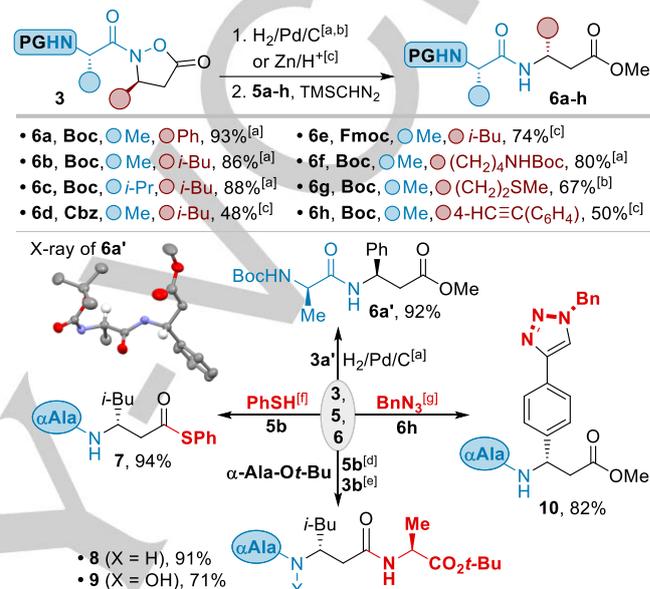
Table 2. Scope and limitations.^[a]

Reaction 1		Reaction 2	
From PG- α AlaNH ₂ 1a-1c		From <i>i</i> -BuCHO 2b	
• PG = Boc • PG = Cbz • PG = Fmoc		• PG = Boc • PG = Cbz • PG = Fmoc	
• Ph • <i>i</i> -Bu ^[b]		• Ph • <i>i</i> -Bu ^[b]	
• Bn (Boc- α Phe) • <i>i</i> -Pr (Boc- α Val) • H (Boc- α Gly) ^[d]		• Bn (Boc- α Phe) • <i>i</i> -Pr (Boc- α Val) • H (Boc- α Gly) ^[d]	
• R = F 3j , (99%), 64%, 92:8 dr • R = OMe ^[c] 3k , (91%), 51%, 88:12 dr • R = C \equiv CH 3l , (99%), 68%, 90:10 dr		• (Boc- α Pro), 3i • (91%), 63%, 75:25 dr	
• Ph • Bn • <i>i</i> -Pr • BnO		• Ph • Bn • <i>i</i> -Pr • BnO	
• MeS • BocHN		• MeS • BocHN	

[a] Reaction conditions: **1** (0.3 mmol), RCHO **2** (1 equiv), MA (1.5 equiv) in PhMe/CH₂Cl₂ (3:1, 0.025 M) at 30 °C for 24 h. Isolated yields of the pure major diastereoisomer after column chromatography. In parentheses, yields of both diastereoisomers determined by ¹H NMR with an internal standard. dr determined by ¹H NMR on the crude product. [b] 5 mol% of catalyst on 1 mmol scale gives **3b** (89%, 94:6 dr), **3d** (86%, 89:11), **3e** (80%, 94:6 dr), **3r** (47%, 94:6 dr). [c] 20 mol% of catalyst. [d] Absolute configuration was not determined for **3h**.

An array of both aromatic aldehydes (**3j-3l**, 51-68%, 88:12 to 92:8 dr), linear and cyclic aliphatic aldehydes (**3m-3p**, 69-91%, 85:15 to > 95:5 dr) proved to be compatible with this multicomponent KaMC process, encompassing the construction of β hPhe **3n** and β hLeu **3o** precursors. Isoxazolidinones with ether **3q** (β hSer, 61%, 83:17 dr), thioether **3r** (β hMet, 41%, >95:5 dr) and NHBoc **3s** (β hLys, 44%, 89:11 dr) pendants could be isolated in diastereomeric pure form albeit in slightly lower yields

even with 20 mol% of catalyst, due to the instability of these aldehydes that results in an incomplete reaction. Although the homologous tryptophane derivative **3t** (β hTrp) could be constructed with excellent > 95:5 dr, a low conversion was observed leading to a moderate 19% isolated yield. Worthy of note, several MCR could be carried out on 1 mmol scale to give the isoxazolidinones **3b**, **3d**, **3e** and **3r** with rather similar yields and dr by means of only 5 mol% of (DHQ)₂PHAL as organocatalyst.



Scheme 3. Chemical transformations into α/β -dipeptides. Reaction conditions: [a] H₂, Pd/C (1 atm), *i*-PrOH, 30 °C for **6a-b**; 15 atm, RT for **6c**, 1 atm, 50 °C for **6f** and TMSCHN₂, MeOH, RT. [b] H₂, Pd(OH)₂/C (20 atm), *i*-PrOH/EtOAc (2:1), 60 °C for **6g**. [c] Zn (40-80 equiv), THF/H₂O/AcOH, 40 °C, and TMSCHN₂, MeOH, RT. [d] From **3b**, HCl- α AlaO*t*-Bu, *i*Pr₂NEt, DMAP (0.5 equiv), DMF, RT. [e] From **5b**, HCl- α AlaO*t*-Bu, *i*Pr₂NEt, EDC-HCl, HOBT, CH₂Cl₂, RT. [f] PhSH, DCC, HOBT, CH₂Cl₂, RT. [g] BnN₃, Na-Ascorbate, CuSO₄·5H₂O (20 mol%), *t*-BuOH/H₂O, RT.

Having an array of novel isoxazolidinones **3** in hand, decorated with various functionalities, the transformations of **3** into α/β -dipeptide derivatives was undertaken in order to probe the versatility of these masked β AA platforms (Scheme 3). By means of adapted palladium catalyzed N-O bond hydrogenolysis, followed by esterification of the obtained acids **5**, in order to facilitate the purification step, several *syn*- α/β -dipeptides-OMe **6a-6c** (86-93%),^[19] together with NHBoc **6f** (Boc- α Ala-Boc- β hLys, 80%) and SMe **6g** (Boc- α Ala-Boc- β hMet, 67%) derivatives were easily synthesized. This straightforward strategy was also applied to the formation of (*R,R*)-dipeptide **6a'** in 92% yield.^[19] On the other hand, by means of orthogonal Zn/AcOH deprotection conditions, the elaboration of *N*-Cbz and *N*-Fmoc α Ala- β hLeu dipeptides **6d** (48%) and **6e** (74%) respectively was allowed.^[10b] Product **6h**, displaying a sensitive but useful terminal alkyne moiety, was accessible in 50% yield under these reductive conditions (Zn/AcOH) and successfully transformed into triazole **10** in 82% yield, thanks to the copper-catalyzed 1,3-dipolar cycloaddition protocol. The readily available Boc- α Ala- β hLeu carboxylic acid **5b** (used as a crude product) was subjected to

peptide-coupling reagents to give rise to the formation of either thioester **7** (94%), suited for further native ligation processes, or $\alpha/\beta/\alpha$ -tripeptide **8** in 91% yield. Interestingly, taking advantage of the reactivity of the N-EWG isoxazolidinone framework,^{[12], [17b], [7c]} the direct ring opening event of **3b** with *tert*-butyl-alanine furnished the $\alpha/\beta/\alpha$ -tripeptide **9**, similar to **8** but with a N-OH functionality known to be a valuable moiety to complex metal in bio-environment.^[21]

In summary, we have developed a multicomponent synthesis of novel isoxazolidin-5-ones possessing an N- α -amino acid. Thanks to the high reactivity of transient alkylidene Meldrum's acids, a smooth and diastereoselective domino addition reaction takes place upon the match influence of a commercially available quinine derived organocatalyst allowing to achieve good to excellent dr. The corresponding isoxazolidin-5-ones, easily obtained as single diastereoisomer after purification, proved to be versatile platforms for the diversity-oriented synthesis of an array of α/β -dipeptides, as useful fragment in medicinal chemistry.

Acknowledgements

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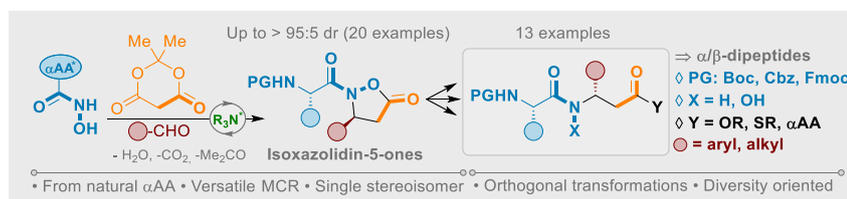
Keywords: amino acid • asymmetric synthesis • isoxazolidinone • Meldrum's acid • organocatalysis

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- [17] The N-O bond reductive cleavage could be easily achieved on the mixture of diastereoisomers **3a**, but it turned out to be much more difficult to separate the two diastereoisomers of the obtained dipeptide **6a** on column chromatography. This shows the clear advantage of working with isoxazolidinone **3a** whose pure major (S,S)-diastereoisomer could be isolated. This easy separation of the mixture of diastereoisomers of isoxazolidinones **3** is a general rule in this series.
- [18] In case of aliphatic aldehydes such valeraldehyde **2b**, the catalyzed reaction (optimized conditions) could be performed with more concentrated conditions with only a moderate impact on dr (**3b**, 0.1 M, 99% NMR yield, 94:6 dr; 0.2 M, 99% NMR yield, 93:7 dr).
- [19] CCDC 1983122, 1983125, 1983123, 1983124 contains the supplementary crystallographic data for respectively compounds **3c** (minor diastereoisomer), **6a** (the two diastereoisomers both originated from major and minor stereoisomers **3a**), **6a'**.
- [20] For a rare example of enantioselective catalytic C-N bond formation to alkylidene Meldrum's acids, see: E. Pair, C. Berini, R. Noël, M. Sanselme, V. Levacher, J.-F. Brière, *Chem. Commun.* **2014**, *50*, 10218.
- [21] a) H. C. J. Ottenheijm, J. D. M. Herscheid, *Chem. Rev.* **1986**, *86*, 697; b) R. Rani, C. Granchi, *Eur. J. Med. Chem.* **2015**, *97*, 505.

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Organocatalytic Multicomponent
 Synthesis of α/β -Dipeptide
 Derivatives

The diversity: A straightforward organocatalyzed multicomponent reaction (MCR) involving readily available hydroxamic acids-derived from naturally occurring α -amino acids (α AA) allows a stereoselective and diversity-oriented synthesis of novel isoxazolidin-5-ones as versatile platforms for the elaboration of α/β -dipeptide fragments.