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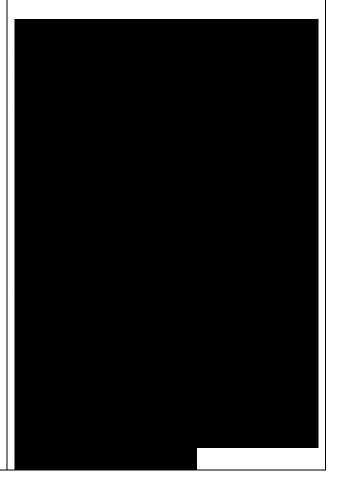
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Proline-catalyzed asymmetric reactions	Phản ứng bất đối xứng xúc tác bằng
1 h 09	Proline
1. Introduction	1.Giới thiệu

Several strategies available are enantioselective (chọn lọc đối quang, chọn lọc đối phân) catalysis, including heterogeneous catalysis, Br0nsted or Lewis acid and base catalysis, homogeneous transition-metal catalysis, and biocatalysis. One remarkable molecule, the amino acid proline, has become a crucial component in examples of all of the catalytic strategies listed above. Proline can be a ligand in asymmetric transition-metal catalysis, a chiral modifier in heterogeneously catalyzed hydrogenations, and, most importantly, proline itself can be an effective organocatalyst several of powerful asymmetric transformations, such as the aldol, Mannich, and Michael reactions.

In the focus of this review are prolinecatalyzed asymmetric reactions. Covered are those reactions that are either catalyzed by proline alone, or by proline in combination with cocatalysts such as metal salts. Selected non-enantioselective proline-catalyzed reactions have also been included. However, the present review does not cover the use of important proline-derived auxiliaries or catalysts including the Endershydrazones or the Corey-Bakshi-Shibatacatalyst, which have been reviewed elsewhere 1'2 Further-more. other remarkable and useful amino acid-based catalysts such as MacMillan's iminium catalysts and Miller's peptide catalysts are beyond the scope of this review.3

Đã có một số phương pháp để tiến hành xúc tác chọn lọc đối quang, bao gồm xúc tác không đồng nhất, xúc tác axit và bazo Br0nsted hoăc Lewis, xúc tác kim loại chuyển tiếp đồng nhất, và xúc tác sinh học. Một phân tử đáng chú ý, axit amin proline đã trở thành một thành phần quan trong và tiêu biểu trong tất cả các chiến lược xúc tác liệt kê ở trên. Proline có thể là một phối tử trong xúc tác kim loại chuyển tiếp bất đối xứng, chất cải biến chiral trong quá trình hydro hoá xúc tác không đồng nhất, và quan trọng nhất, chính proline có thể là chất xúc tác hữu cơ hiệu quả của một số quá trình chuyển đổi bất đối xứng mạnh, chẳng hạn như các phản ứng aldol, Mannich, và Michael.



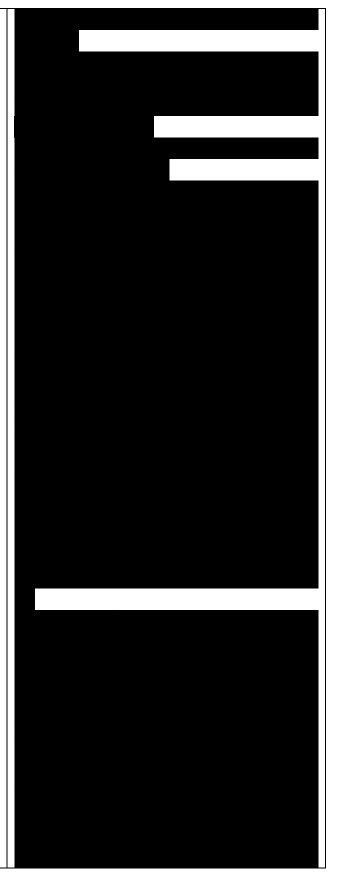
1.1. Proline—a universal asymmetric catalyst?

There are several reasons why proline has become an important molecule in asymmetric catalysis. Not least is the

Figure 1. Modes of action in proline-catalysis.

fact that proline is an abundant chiral molecule that is inexpensive available in both enantiomeric forms. Additionally, there are various chemical reasons that contribute to proline's role in catalysis. Proline is bifunctional, with a carboxylic acid and an amine portion. These two functional groups can both act as acid or base and can also facilitate chemical transformations in concert, similar to enzymatic catalysis. While enzymes typically use several different functional groups in their catalytic machinery, bifunctional asymmetric catalysis has become a very successful strategy in the laboratory.4 In addition, proline is a chiral bidentate ligand that can form catalytically active metal complexes (Fig. 1).

While all of these criteria apply for all amino acids, proline is a secondary, cyclic, pyrrolidine-based amino acid. A unique consequence of this property is the increased pKa value of its amine compared to primary amino acids. proline's Another consequence ofpyrrolidine portion is bicyclo[3.3.0]octane ring system ('open book structure') of its metal complexes. The most important difference to other is proline's amino acids effective

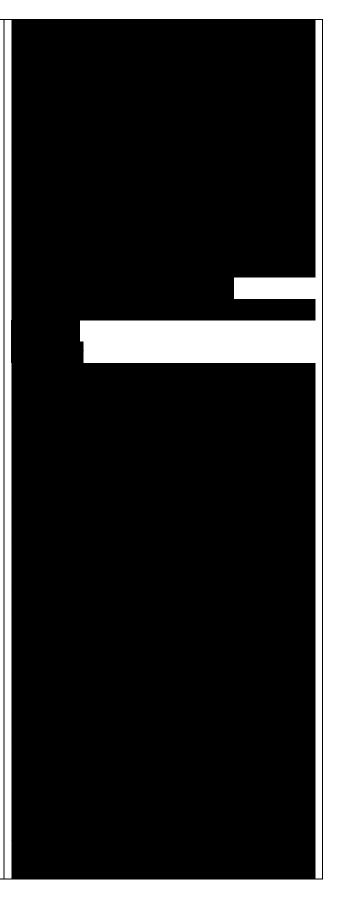


Lewis-base-type aminocatalysis—a catalysis that facilitates iminium- and enamine-based transformations.5 Proline's unique nucleophilic reactivity is primarily a consequence of the pyrrolidine portion, which forms iminium ions and enamines with carbonyl compounds more readily than most other amines, including cyclic ones such piperidine.6 The carboxylate further contributes to proline's aminocatalysis by acting as a general Br0nsted cocatalyst.

2. The Hajos-Parrish-Eder-Sauer-Wiechert reaction

2.1. Background

The 1960s witnessed a strong interest in efficient and economic steroid syntheses. This development was fueled by the commercial success of steroidal contraceptive agents (in 1969, million American women were taking the pill) and the promise of other pharmaceutically active steroids such as the 'wonder drug', cortisone. The best way to synthesize steroids at the time was the Marker process, a sequence of reactions that led from diosgenin, a plant steroid isolated from Mexican wild yams, to cortisone and other important steroids such as norethindrone.7 Early on, alter-native synthetic schemes that would not require the use of a potentially rare resource were envisioned. example, racemic tetrahydroindandione 1 and octahydronaphtalene- dione 2 (the Wieland-Miescher ketone) have been resolved and used in asymmetric steroid total syntheses (Scheme 1).8



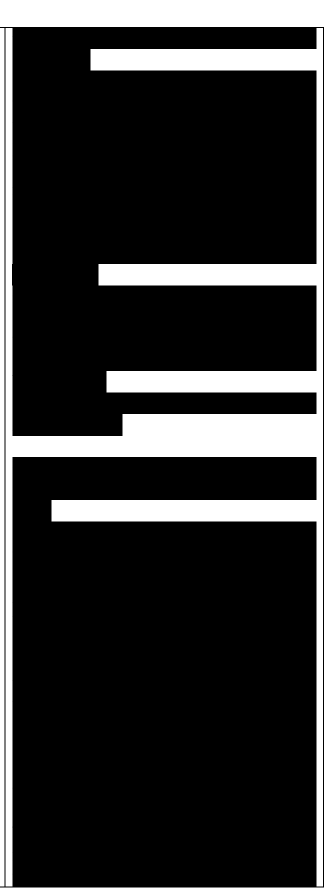
Ketones 1 and 2 can be made from symmetric monocyclic triketones 3 and 4 via intramolecular aldol condensation (Scheme 2). These reactions can be catalyzed by amines such as pyrrolidine,9 and Spencer et al. convincingly demonstrated that they involve enamine intermediates similar to certain enzymatic reactions.10

Contemporaneous to these experiments were Yamada's studies on asymmetric synthesis with amino acids. For example, asymmetric Robinson-annulations have been

Scheme 3. Yamada's asymmetric Robinson-annulation.

developed that are based on preformed proline-derived enamines (Scheme 3).11

Asymmetric enamine catalysis was first realized with the discovery of the proline-catalyzed asymmetric intramolecular aldol reaction by two industrial groups in the early 1970s. Hajos and Parrish at Hoffmann La Roche reported proline-catalyzed intramolecular aldol reactions triketones such as 3 and 4 to give aldols 5 and 6 in good yields and ees (Scheme 4).12 Acid-catalyzed dehydration furnished aldol condensation products 1 and 2 in a second step (Eqs. (1) and (2)). As shown by Eder, Sauer, and Wiechert at Schering, the aldol condensation products can also be obtained directly from triketones 3 and 4 if the cyclization



is performed in the presence of proline (10-200 mol%) and an acid-cocatalyst (Eqs. (3) and (4)).13

2.2. Scope and applications

The asymmetric proline-catalyzed intramolecular aldol

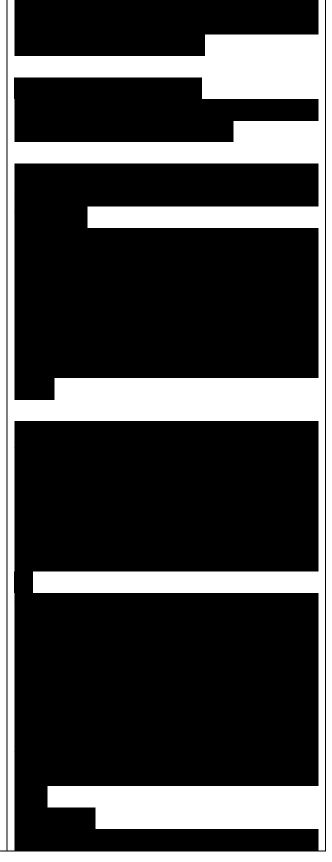
Figure 2. Selected products from Hajos-Parrish-Eder-Sauer Wiechert reactions.

cyclization, also termed the Hajos-Parrish-Eder-Sauer- Wiechert reaction,14 has been applied to several substrates since its invention over 30 years ago.15 A small selection of products obtained using proline-catalyzed intramolecular aldolizations is shown in Fig. 2

The Hajos-Parrish-Eder-Sauer-Wiechert reaction has not only been used in steroid syntheses but also in several other natural product total syntheses.17 The reaction has been studied using polymer-bound (S')-proline as the catalyst18

and Agami et al. described conceptually related proline- catalyzed enantiogroup differentiating aldol-cyclodehydrations of acyclic diketones (Table 1). 9 When compared to the Hajos-Parrish-Eder-Sauer-Wiechert reaction, the efficiency and enantioselectivity of Agami's desymmetrization reaction are generally lower.

2.3. Mechanism While realizing that their "results may be



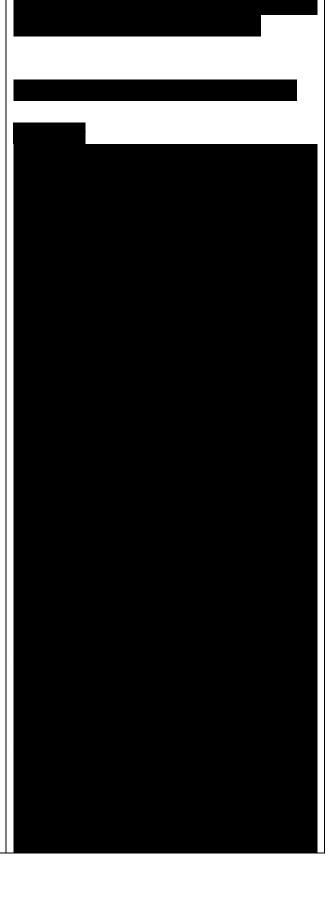
considered an example of a simplified model of a biological system in which (S')-proline plays the role of an enzyme", Hajos and Parrish initially rejected the aldolase-type enamine mechanism in their seminal work.12 According to the proposed alternative mechanism, one of the enantiotopic ring carbo- nyl-groups is activated as a carbinolamine, which C-C-bond-forming undergoes nucleophilic substitution reaction reaction with a side-chain enol (A in Fig. 3). This model is consistent with the low 180-incorporation into the product, an observation made if the reaction was conducted in the presence of 180-labeled water. However, the Hajos- mechanism has been rejected by Jung because it involves retention of configuration in an SN2-like process.20 Jung and later Eschenmoser et al.21 first discussed a 'one proline- mechanism' involving a side-chain enamine intermediate and Agami et al. proposed model B in which a second proline molecule is involved.22 Kinetic studies and an observed nonlinear effect in asymmetric catalysis supported the involvement of two proline molecules in the enantioselectivitydetermining step.23 Heterogeneous catalysis involving concerted a bifunctional acid/base- mechanism (C) has also been suggested as a possible mechanism on the basis ofthe observation that proline is often not soluble completely in organic solvents.24 Using quantum mechanical calculations, Houk et al. recently proposed a new model (D) that readily explains the observed

enantioselectivity.25 This elegant model is also consistent with the original Spencer-mechanism of the pyrrolidine-catalyzed intramolecular aldolization.

3. The direct intermolecular aldol reaction

3.1. Background

The direct intermolecular aldol reaction between two carbonyl compounds is central to sugar metabolism. Class I aldolases catalyze this process by using an enamine mechanism.26 Several early bioorganic studies appeared in which simple small molecule amines and amino acids served as aldolase models.27 Aldolase-like catalytically active amines, amino acids. and amine/antibody systems have been studied by Reymond et al.28 In addition, catalytic antibodies have been generated which also use an enamine mechanism.29 These important studies in particular have taught us the potential of enamine catalysis asymmetric synthesis. Lessons learnt from the aldolase antibodies, the Haios-Parrish-Eder-Sauer-Wiechert reaction, and the discovery of non-proteinogenic (xây dựng protein, sinh protein), metal complex-catalyzed direct asymmetric aldol reactions, 30 led to the development of the first proline-catalyzed direct asymmetric aldol reaction.31 Initially, it although proline was shown that, typically reacts unproductively with aldehydes, the intermolecular reaction between a ketone and an aldehyde is possible if a large excess of the ketone donor is used. For example, acetone (20 vol.%, ca. 27 equiv.) reacts with isobutyralde- hyde in DMS0 to give the



corresponding aldol in excellent yield and ee (Scheme 5).

Scheme 5. Highly enantioselective proline-catalyzed intermolecular aldol reaction.

3.2. Scope and applications

Several other aldehydes have been used in proline-catalyzed aldol reactions with acetone (Table 2). In general, aromatic aldehydes furnish aldols with ees of around 70% and in varying yields (54-94%). Higher enantioselectivities and yields were obtained when a-branched aldehydes were used and tertiary aldehydes gave exceptionally high ees of up to >99%. The only significant sideproduct in these reactions (and also in the Hajos-Parrish-Eder-Sauer- Wiechert reaction) the aldol-condensation is product.

Figure 3. Proposed mechanisms of the Hajos-Parrish-Eder-Sauer-Wiechert reaction.

a-Unbranched aldehydes turned out to be a difficult substrate class and did not provide the corresponding aldol products under standard conditions. Only homoaldol- addition- and condensation of the aldehyde or elimination of the crossaldol product to the a, P-unsaturated ketone were observed in DMSO. Using acetone or acetone/CHCl3 mixtures as solvents and 10-20 mol% of proline as the catalyst allowed isolation of the cross-aldol products in modest yields and good enantioselectivities (Table 3).32

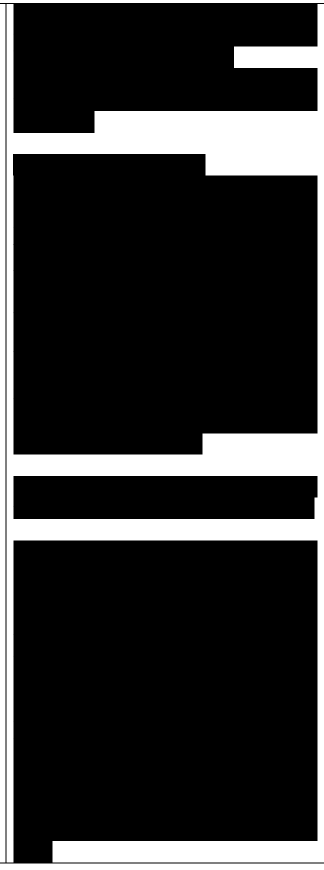


Table 3. Proline-catalyzed direct asymmetric aldol reactions using a-unbranched aldehydes as acceptor

Scheme 6. Catalytic asymmetric total synthesis of (S)-ipsenol.

Observed side-products are the crossaldol condensation products and the homo-aldol addition product of acetone.

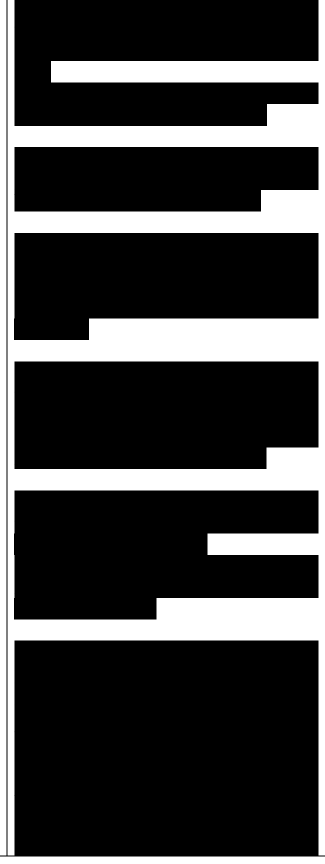
The proline-catalyzed aldol reaction of acetone with a-unbranched aldehydes has been used in a short synthesis of the natural pheromone (^)-ipsenol (Scheme 6).

Recently, the proline-catalyzed intermolecular aldol reaction with acetone has been applied to the highly dia- stereoselective synthesis of complex sugar derivatives (Scheme 7).33

Scheme 7. The proline-catalyzed intermolecular aldol reaction in the synth¬esis of complex sugars.

Table 4. The proline-catalyzed intermolecular aldol reaction using cyclic ketones as donors

A general limitation of the method is the scope of the ketone component. Since a large excess of the ketone is often required, small and inexpensive ketones such butanone. acetone. and cyclohexanone typically used. are Selected reaction products from prolinecatalyzed aldol reactions with ketones other than acetone are provided in Table 4. Other ketones such as 3-pentanone



and cetophenone have not been successfully used yet.

Excellent results have also been obtained with hydroxy¬acetone as the donor. In this case, anti-diols are formed in high regioselectivities, diastereoselectivities and enantio- selectivities (Table 5).

Several proline-catalyzed intermolecular aldol reactions have recently been successfully repeated, both with proline itself and with poly(ethylene glycol)-supported proline.34'35

In addition to serving as acceptors in proline-catalyzed aldol reactions, aldehydes can also act as donors under conditions. certain For example, acetaldehyde trimerizes in the presence of (^)-proline to give aldehyde 7 in low relatively yield but high enantioselectivity (Scheme 8).36

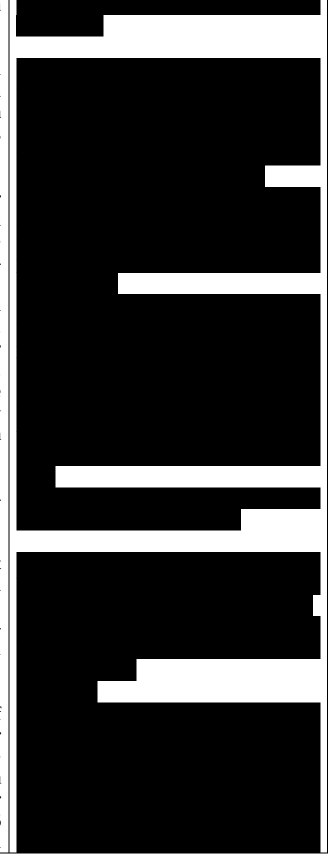
Furthermore, it was found that several aunbranched alde-

Table 6. The proline-catalyzed direct asymmetric intermolecular aldol reaction using aldehydes as donors

hydes react with activated nonenolizable ketones to give aldols in good yields and ees (Table 6).37

3.3. Mechanism

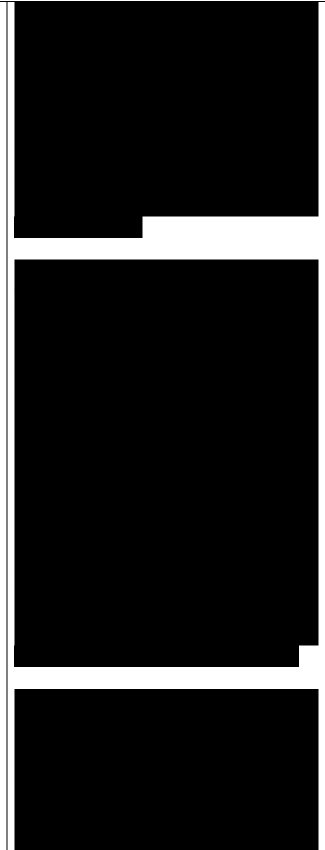
The originally proposed mechanism of the proline-catalyzed intermolecular aldol reaction31 was based on the established class I aldolase-mechanism that involves carbinolamine, imine or iminium, and enamine intermediates.26 The catalytically active functional



groups in class-I aldolases are an samino group of a lysine residue (một chuỗi có thứ tự các axit amin) and, depending on the enzyme subtype, a set of Br0nsted cocatalysts required for the various proton-transfers of the multi-step reaction mechanism. In the proline-catalyzed version, the catalytic amine is proline's pyrrolidine. The carboxylate could function as a multi-purpose Br0nsted cocatalyst for the proton-transfers (Scheme 9).

The enantioselectivity was explained with a transition state (E) that can be described as a metal-free version of the classical Zimmermann-Traxler model successfully explains (F),38which stereoselectivities of metal enolate aldol reactions. Furthermore, model E is similar to Houk's recently calculated transition state of the Hajos-Parrish-Eder-Sauer-Wiechert reaction (D).25However. according these to calculations, an N-H hydrogen bond does not lower the energy of the and model E transition state consequently been advanced to model G,5 which is superimposable to the calculated transition state of the prolinecatalyzed intermolecular aldol reaction (Fig. 4).

The proposed multi-step reaction mechanism (Scheme 9) has very recently been confirmed using density functional theory calculations.39a Moreover, the validity of transition state G has been demonstrated by using density functional theory predictions followed by experimental verification of



stereoselectivities of proline-catalyzed aldol reactions.39b

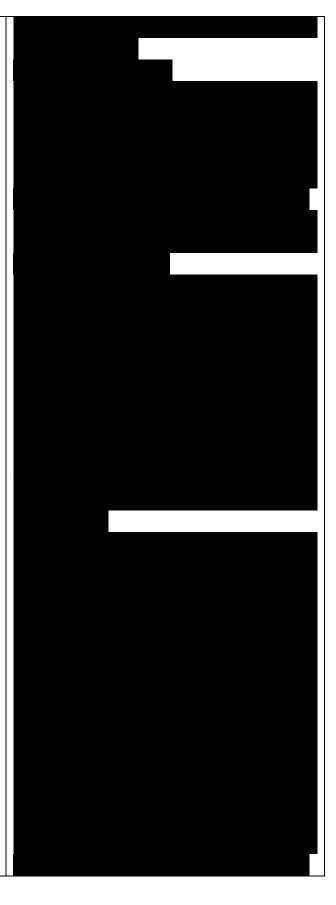
4. Mannich reactions

Proline-catalysis has recently been extended to the direct asymmetric three-component Mannich reaction of ketones, aldehydes, and amines to give (P-amino ketones in high yields and enantioselectivities (Scheme 10).40

Scheme 9. Originally proposed mechanism of the proline-catalyzed direct asymmetric aldol reaction.

typically were indirect and required the use of preformed imine- and enol equivalents.41 In contrast, the proline-catalyzed version constitutes a rare example of a catalytic asymmetric multi-component reaction. The substrate scope of this reaction has recently been explored.42 Various ketones can be employed in proline-catalyzed Mannich reactions with ^-anisidine (PMPNH2) and ^-nitrobenzalde- hyde with excellent results (Table 7).

Catalyst and amine-component have also been varied and so far proline seems to be the optimal catalyst while ^-anisidine turned out to be the most useful amine-component. A remarkable aspect of the reaction is its tolerance to a broad range of diverse aldehydes as substrates. Both aromatic and aliphatic aldehydes used. Aromatic can be aldehydes generally give the Mannich products in high ees yet modest yields.43 Most importantly, and in contrast to the proline- catalyzed aldol reactions and to all other catalytic asymmetric Mannich reactions, a-unbranched aldehydes were



Proline-catalyzed Table 7. direct asymmetric Mannich reactions varying the ketone component Proline-catalyzed Table direct 8. asymmetric Mannich reactions varying the aldehyde component efficient substrates in the prolinecatalyzed variant. Here, acetone or a chloroform/acetone mixture was used as the solvent instead of the commonly used DMSO (Table 8). Recently, ethyl glyoxylate has been added to the list of aldehydes that may be used.44 Functionalized a-amino acid esters obtained were in high stereoselectivities in such reactions if the preformed imine of ethyl glyoxylate was used. Both ketones and a-unbranched aldehydes could be utilized as donors to give the products in high enantioselectivities (Table 9). Because of the exceptionally high regio-, enantioselectivities diastereo-. and observed in the Mannich reaction with hydroxyacetone, reactions ketone as the donor component were studied with several different aromatic aldehydes and isobutyraldehyde (Table 10).42 Good yields and diastereoselectivities and excellent regio- selectivities were generally observed. Enantioselectivities were typically very high (up to >99%), yet dependent on the electronic nature of the aldehyde component. A good Scheme 11. Proposed mechanism of the proline-catalyzed direct asymmetric

three-component Mannich reaction.

correlation of enantioselectivities with Hammett (7p-values was observed, and a linear Hammett plot was obtained with a reaction constant p for the proline-catalyzed three-component Mannich reaction of 1.36 (R2=0.95) (Fig. 5).

The positive reaction constant is consistent with partial negative charge formation in the transition state (trang thái chuyển tiếp, trạng thái dịch chuyển) and with the proposed mechanism that involves nucleophilic addition of a proline-enamine to an imine (Scheme 11).

According to the mechanistic proposal, a proline-enamine reacts with an imine in the C-C-bond forming and enantio-selectivity-determining step. Both the imine and enamine intermediates are formed in situ from an aldehyde and a ketone in two separate pre-equilibria.

One of the intriguing aspects of the proline-catalyzed Mannich reaction is its stereoselectivity. Diastereoand enantioselectivities are opposite to those observed in proline-catalyzed intermolecular aldol reactions. This result was initially explained with transition states that involved (Z)-imines. However. (E)-aldimines strongly equilibria predominate with corresponding (Z)-imines. For example, only the (E)-aldimine can be detected 1H NMR-spectro- scopically in the reaction of ^-nitrobenzaldehyde with ^-anisidine in DMS0-d6.42b Therefore, although



(Z)-imines cannot be excluded, it seems more likely that (£)-imines are involved in the reaction mechanism. The currently preferred transition state models for the proline-catalyzed Mannich reaction (H) and intermolecular aldol reaction (G) are shown in Fig. 6.

The assumed transition states reflect the fact that enantio- faciality of the electrophile (imine si vs aldehyde re), but not that of the enamine, is reversed in aldol vs Mannich

Figure 6. Proposed transition states of proline-catalyzed aldol and Mannich reactions.

Scheme 12. Iminium and enamine catalysis of the Michael reaction. reactions and in order to allow for protonation of its lone pair, an (£)-imine

has to approach the enamine with its siface to avoid unfavorable steric interactions between the pyrrolidine and aromatic ring.

5. Michael reactions

The Michael addition is a particularly interesting reaction because proline-catalysis may proceed by both amino-catalytic pathways, iminium (a) and enamine catalysis (b) (Scheme 12); both reaction types have been realized.

5.1. Iminium catalysis of the Michael reaction

Yamaguchi et al. found the Michael



addition of malonates to a,P-unsaturated aldehydes to be catalyzed by secondary amines, including (S)-proline.45 For example, dimethyl malonate reacts with hex-2-enal in the presence of pyrrolidine or proline to furnish Michael adduct 8. It was noted that triethylamine and N-methyl proline are inactive and that lithium prolinate is superior to proline itself (Table 11).

Table 11. Initial study by Yamaguchi et al. on iminium catalysis of the Michael reactions

Table 12. Studied metal prolinates

Scheme 14. Merck's synthesis of substituted prolines.

Scheme 18. The enamine catalysis cycle. processes, it has been pointed out that the two-step procedures require significantly less catalyst and generally provide better yields and ees.58

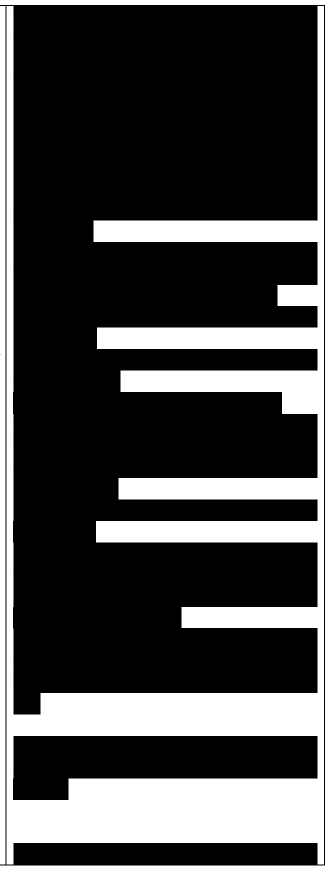
6. The direct electrophilic a-

Proline-catalysis in aldol, Mannich, and Michael reactions can be rationalized with a general enamine catalysis cycle (Scheme 18).5

Accordingly, carbonyl compounds react with proline to generate an enamine intermediate and water. This enamine

Table 18. The first direct catalytic asymmetric electrophilic a-amination of aldehydes

reacts with an electrophile X=Y, which

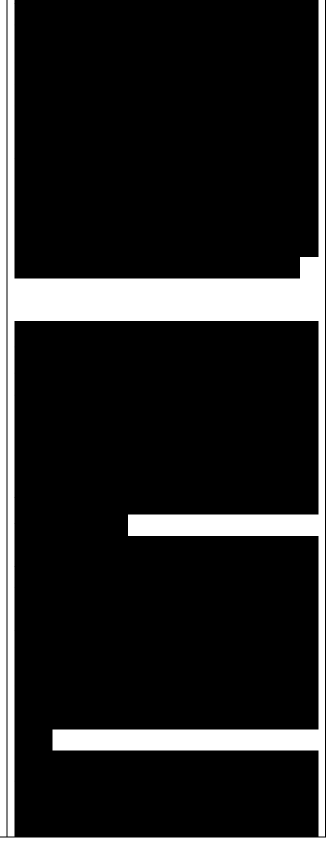


may be an aldehyde, an imine, or an activated olefin in an aldol, Mannich, or Michael reaction. Hydrolysis of the iminium intermediate then gives the product under regeneration of the proline catalyst. An alternative electrophilic species could be dialkyl a azodicarboxylate (X=Y:RCO2N=NCO2R). The overall transformation would result in an electrophilic a-amination of the carbonyl compound.61 The products of this reaction, if produced enantioselectively, could be useful precursors for various amino acid derivatives.

Very recently, this reaction has been realized for the first time.62 It was found proline catalyzes direct that the electrophilic a-amination of unbranched highly effectively aldehydes and enantioselectively. Because the produced a-hydrazino aldehvdes configurationally labile, they were in situ reduced to the corresponding alcohols (Table 18).

The potential of the produced amino alcohol derivatives as precursors for the asymmetric synthesis of a-amino acid derivatives was demonstrated with a straightforward synthesis of Evans-type oxazolidinone 18 via hydrogenation and work-up with phosgene (Scheme 19). The hydrogenation removes both Cbzprotecting groups and simultaneously cleaves the N-N bond.

The observed stereoselectivity can be explained with transition state(trang thái chuyển tiếp, trang thái dịch chuyển) J (Fig. 7), which again is superimposable



with Houk's transition states of the Hajos-Parrish-Eder-Sauer-Wiechert reaction (D) and Agami's diketonecyclization (D0. A comparison of the assumed preferred transition states of the proline-catalyzed enamine-involving intramolecular aldol reaction (D, D'),25 intermolecular aldol reaction (G).5Mannich reaction (H),42Michael reaction

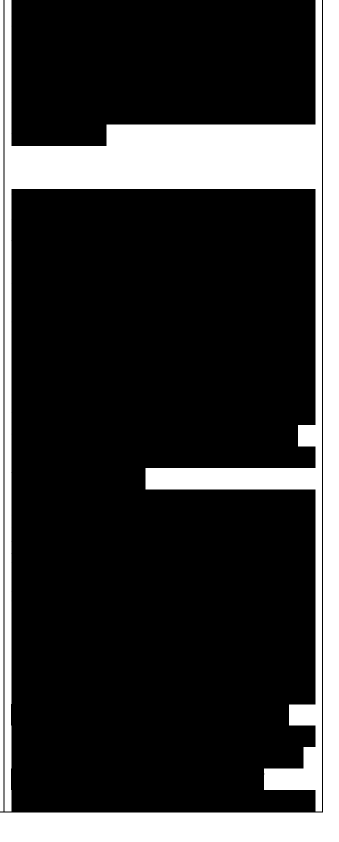
(I), ' and a-amination (J) reveals three important and general structural elements: (1) The assumed prolineenamine is always in a conformation in which the car- boxylate is anti to the enamine-olefin. (2) The enamine- olefin geometry is always (E). (3) The carboxylic acid protonates the electrophile to compensate negative formation. generalized charge A transition state (K) that combines these elements may be constructed (Fig. 7).

7. Miscellaneous proline-catalyzed asymmetric reactions

In addition to aldol, Mannich-, Michael and electrophilic a-amination reactions, proline has been used as a catalyst in several other asymmetric transformations such as allylic oxidations, transfer-hydrogenations, and Diels-Alder-type reactions. In these reactions, proline is not only used as an aminocatalyst but also as a chiral ligand in metal-mediated asymmetric processes.

Table 19. Dimerizations of unsaturated aldehydes (n.r.=not reported)
Figure 8. Possible intermediates.

7.1. Diels-Alder-type dimerizations of



a, P-unsaturated aldehydes

Asato and Liu et al.63 found that upon treating a,P-unsatu- rated aldehydes with (S)-proline in ethanol, cyclic optically active dimers were obtained (Table 19). Such dimerizations were known to occur under basic conditions, and to give products.64 racemic The prolinecatalyzed reactions furnish the products in encouraging enantioselec- tivity and it may be worthwhile to study other chiral amines as potential catalysts for this interesting reaction. Mechanistically, the reaction could proceed via dienamine 19 iminium and/or ion 20. These intermediates may undergo

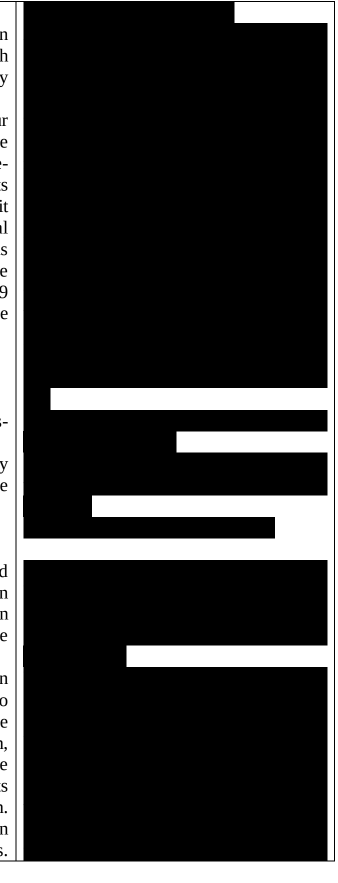
Table 20. Proline-catalyzed Baylis-Hillman reactions

a Diels-Alder-reaction followed by elimination and hydrolysis to give the observed products (Fig. 8).

7.2. Baylis-Hillman reactions

Very recently, a mixture of proline and imidazole (each 30mol%) has been found to catalyze Baylis-Hillman reactions of aldehydes with MVK (Table 20).65

Essentially, no asymmetric induction was observed (ees 5-10%). According to the proposed mechanism, proline activates MVK as the iminium ion, facilitating conjugate addition of the imidazole. The resulting enamine reacts with the aldehyde in an aldol reaction. The Baylis-Hillman product is then formed via elimination and hydrolysis.



That imidazole alone (in contrast to other nucleophiles such as DABCO) is not sufficiently reactive to induce the Baylis-Hillman process can be interpreted as evidence for the proposed iminium catalysis (Scheme 20). It will be interesting to note whether the use of other chiral amines or different reaction conditions may lead to a new catalytic asymmetric Baylis-Hillman variant.

Scheme 20. A mechanism of the proline/imidazole-catalyzed Baylis-Hillman reaction.

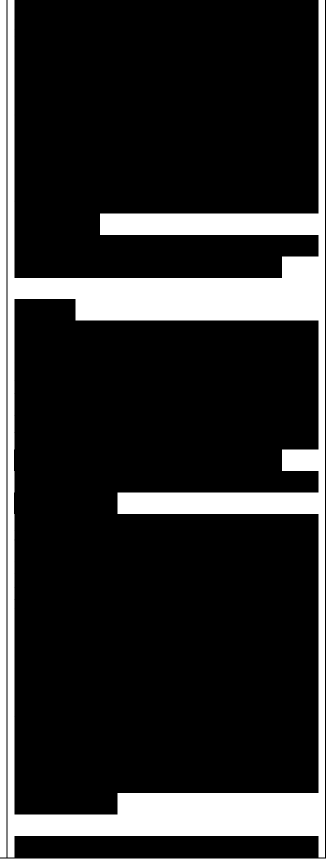
7.3. Reductions

Several reductions involving proline as the source for asymmetric induction have been developed. These include hydrogenations, epoxide reductions, borane and boranate reductions, and ruthenium-catalyzed transfer hydrogenations.

Ethyl acetoacetate has been enantioselectively reduced with

a Raney Cu-catalyst modified with (S)proline.66 Other heterogeneous catalysts in combination with proline have also been used, but typically reduce ketones with low enantioselectivities.67 Tungler et al. described the Pd/C-catalyzed reduction of isophorone in the presence of (S)-proline to give saturated ketone 22 in low yield and ca. 60% ee. The main side product results from a reductive amination of ketone 22 with proline. The reaction involve iminium may intermediate 21 (Scheme 21).68

An equimolar mixture of NaBH4 and



(S)-proline in THF reduces ketones to secondary alcohols in ees of up to 62%.69 Martens et al. used a mixture of borane with proline for the similar reactions and obtained enantioselectivities of up to >95%.70 Presumably, proline is initially converted to prolinol, which then forms an oxazaborolidine, derivative of the well-known CBS-reduction catalyst.

Racemic epoxides have been reductively cleaved with a mixture of zirconium tetrachloride, sodium borohydride, and (S)-proline to give enantioenriched alcohols (Scheme 22).71 Apparently, these reactions are not kinetic resolutions and the authors speculate that they may involve zirconium enolates.

Scheme 22. Proline-catalyzed reductive epoxide-opening.

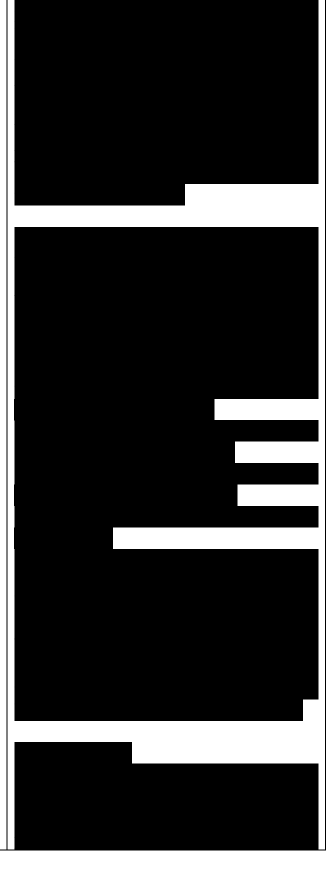
Table 21. Proline-catalyzed asymmetric transfer hydrogenation

Scheme 23. Proline-catalyzed asymmetric allylic oxidation.

An interesting Noyori-type asymmetric transfer hydrogena¬tion has been developed by Furukawa et al.72 The catalyst is prepared by mixing potassium prolinate with RuCl2 (p-cymene)2, and reduces aromatic ketones with isopropanol to give secondary alcohols in good yields and ees (Table 21).

7.4. Oxidations

One of the first catalytic asymmetric allylic oxidations was based on a copper catalyst combined with (S)-proline. Disclosed in a patent by Sumitomi



Chemical Co., Ldt.,73 this system was investigated by Muzart and Ferringa.74 An example is the oxidation cyclohexene with PhC03t-Bu in the presence of propionic acid and catalytic amounts of Cu(0Ac)2 and (S)-proline to give ester 24 in acceptable (Scheme enantioselectivity 23). Improved catalyst systems have recently been described.

8. Conclusions

It is remarkable that despite the diversity of reactions discussed in this review, from carbon-carbon ranging bondforming aldol-, Mannich-, and Michael reactions, to electrophilic aminations, transfer-hydrogenations, and allylic oxidations, the catalytically active species and source of asymmetry is a small and simple amino acid. While proline may not be the 'universal asymmetric catalyst' for all reactions, it clearly is a privileged molecule for enantio- selective synthesis. Not only is proline inexpensive, avail-able in both enantiomeric forms, stable, non-toxic, and a powerful catalyst for a number of asymmetric reactions; it also has a multifaceted mechanistic complexity hidden underneath its 'simple' structure. It would seem daring to expect anything less than the discovery of several new proline-catalyzed reactions in the future.

