SYNSTORIES

- Making the Baeyer–Villiger (BV) Reaction Catalytic and Enantioselective with Chiral Brønsted Acid Using Aqueous H₂O₂ as the Terminal Oxidant

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CONTACT

Your opinion about SYNFORM is welcome, please correspond if you like: 
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Dear readers,

Organocatalysis continues to represent a major topic and a very competitive area of research in the arena of organic chemistry. It is therefore not surprising that three out of the four SYNSTORIES that constitute this new issue of SYNFORM are dedicated to new organocatalytic processes. Professor Kuiling Ding (P. R. of China) shows how to perform an old Baeyer–Villiger reaction in a brand new enantioselective manner using a chiral Brønsted acid as organocatalyst. Professor Benjamin List (Germany) describes how to use a simple yet challenging substrate like acetaldehyde as nucleophile in organocatalytic Mannich reactions. Finally, Dr. Matthew J. Gaunt (UK) reports how phenols can be oxidatively dearomatized in an enantioselective manner by means of organocatalysis. There is little doubt that organocatalysis is progressively broadening its scope, although much research remains to be done in order to see more real-life synthetic problems solved by means of this powerful technology. The fourth SYNSTORY covers an extremely exciting piece of bioorganic synthesis reported jointly by Dr. Arata Yajima (Japan) and Professor Yong Qin (P. R. of China) who were able to synthesize and characterize stereochemically a structurally challenging hormone of a fungus-like plant pathogen.

Enjoy your reading!

Matteo Zanda

Editor of SYNFORM
Making the Baeyer–Villiger (BV) Reaction Catalytic and Enantioselective with Chiral Brønsted Acid Using Aqueous H₂O₂ as the Terminal Oxidant


The Baeyer–Villiger (BV) reaction, discovered in 1899, represents one of the most well-known and widely applied reactions in organic synthesis. Although more than a century has gone by since its discovery, the BV reaction is far from being at the end of its development. The use of stoichiometric amounts of peracid as the oxidant suffers from disadvantages – such as expensive and hazardous (because of shock-sensitivity) reagents, with the simultaneous formation of one equivalent of corresponding carboxylic acid waste – which limit their practical application. Therefore, the use of aqueous hydrogen peroxide as the stoichiometric oxidant in the presence of a promoter has been the focus of attention from the viewpoint of green chemistry. On the other hand, the area of catalytic asymmetric BV reactions is also far from fully developed even though its first enantioselective version was realized in 1994 by Strukul and Bolm independently. Since then, a variety of chiral metal complexes (or organic molecules) have been developed for application as promoters in the enantioselective BV reaction of various ketones in stoichiometric or catalytic quantity, but only very few catalyst systems are compatible with the use of the environmentally benign and economically viable aqueous hydrogen peroxide as the terminal oxidant. Another challenging issue of the catalytic asymmetric BV reaction is the difficulty associated with enantioselectivity control. The state-of-the-art of this reaction is that no chemical catalysts are able to afford products in more than 86% ee in the catalytic BV oxidation of 3-substituted cyclobutanones, despite the fact that some enzymes demonstrate excellent enantiocontrol in the catalysis.

A new breakthrough in this area was realized very recently by Professor Kuiling Ding and his PhD student Senmiao Xu, as well as his Associate Professor Zheng Wang and Research Assistant Dr. Xue Zhang, at the Shanghai Institute of Organic Chemistry (SIOC), Chinese Academy of Sciences (P. R. of China). Senmiao Xu is also a joint PhD student with Dr. Xumu Zhang, a guest professor at SIOC, who is currently a professor at the State University of New Jersey (USA). The research was inspired by the facts that the BV reaction can be accelerated in the presence of strong Brønsted acids and that the reactivity of the peracid is dependent on the acidity of the corresponding Brønsted acid. “We envisioned,” explained Professor Ding, “that chiral peroxophosphoric acid, formed in situ from binol-derived phosphoric acid with hydrogen peroxide, might be usable for catalyzing the BV reaction in an enantioselective manner through intermediate 4. This concept also can be considered as an extension of our previous research on hydrogen-bond-promoted enantioselective reactions (Chem. Eur. J. 2004, 10, 5964; J. Org. Chem. 2006, 71, 2862).” The feasibility of the principle was demonstrated by the observation that a binol-derived phosphoric acid (10 mol%) can significantly accelerate the BV oxidation of 3-phenylcyclobutanone with aqueous H₂O₂ (30%), giving 3-phenyl-γ-butyrolactone in excellent yield (99%), albeit with very poor enantioselectivity (ca. 2% ee). Furthermore, no reaction occurred in the absence of phosphoric acid under otherwise identical experimental conditions. After screening a variety of binol-based phosphoric acids with diverse steric and electronic properties of the 3,3′-substituents and the backbone of the scaffold, catalyst 3, which features bulky pyren-1-yl groups at the 3,3′-positions with a 5,5′,6,6′,7,7′,8,8′-octahydro-1,1′-binaphthyl backbone turned out to be optimal. “The BV oxidation of a variety of 3-aryl-substituted cyclobutanones 1 with aqueous hydrogen peroxide in the presence of a catalytic amount of binol-derived chiral phosphoric acids (1–10 mol%) afforded the corresponding chiral γ-lactones 2 in high yields (91–99%) and good to excellent enantioselectivities (82–93%
Although the reactions of 3-alkyl-substituted or 3,3-disubstituted cyclobutanones gave the corresponding lactones with only moderate enantioselectivities (55–61% ee). When the catalyst loading was further reduced to 1 mol%, the oxidation of 3-(4-tolyl)cyclobutanone proceeded smoothly without any loss of enantioselectivity (93% ee).

“The present catalytic system possesses several advantageous features in terms of the following aspects,” said Professor Ding: “1) Although chiral phosphoric acids have been widely used in the catalysis of a variety of asymmetric transformations, including nucleophilic addition of aldimines, transfer hydrogenation, Diels–Alder reaction, Nazarov cyclization, multicomponent condensation and so on, this catalytic system represents the first example of a catalytic asymmetric (BV) oxidation with chiral phosphoric acid. 2) Both the activity and the enantioselectivity of the catalyst are the highest among the chemical catalysts (organometallic or organocatalysts) discovered so far for the asymmetric BV oxidation of 3-substituted cyclobutanones. 3) The use of aqueous hydrogen peroxide (30%) as the terminal oxidant merits green chemistry since water is the only byproduct formed.”

However, despite these very promising results, some challenges still remain for the further development of the procedure. “Although excellent enantioselectivities have been achieved for a broad range of 3-aryl cyclobutanones using the present procedure,” recognized Professor Ding, “further development of the catalytic system is still required to extend its synthetic utility to more challenging substrates, for example, more flexible 3-alkyl cyclobutanones or more sterically demanding 3,3-disubstituted cyclobutanones.” Some of their corresponding lactones are particularly useful in the synthesis of natural products.

“Detailed mechanistic studies for clarifying the substrate activation pathway and the origin of selectivity control in the catalytic process are critically important for the rational design of new-generation catalysts for the target reaction,” concluded Professor Ding.

**About the corresponding author**

**Kuiling Ding** was born in 1966 in Henan Province, P. R. of China. He received his BSc degree from Zhengzhou University (1985) and his PhD from Nanjing University (1990) under the supervision of Professor Yangjie Wu. He was a faculty member of Zhengzhou University from 1990–1998, and was promoted to Full Professor in 1995. In 1993–1994 he was engaged in postdoctoral research with Professor Teruo Matsuura at Ryukoku University (Japan). In the period from 1997–1998 he was a UNESCO research fellow with Professor Koichi Mikami at the Tokyo Institute of Technology (Japan). He joined the Shanghai Institute of Organic Chemistry in 1999, where he currently is a professor of chemistry. His research interests include the development of new chiral catalysts and methodologies for asymmetric catalysis.
According to Professor Benjamin List from the Max-Planck-Institute für Kohlenforschung of Mülheim (Germany), “even though acetaldehyde is structurally the simplest enolizable carbonyl compound, it can be difficult to use in chemical reactions. The undesired products stem mainly from self-aldolization pathways.” The few attempts at employing the molecule include a report on the self-aldolization of acetaldehyde by Barbas and co-workers in which very poor yields of an acetaldehyde trimer were isolated with respectable enantioselectivity, and Jørgensen’s finding of a high-yielding but racemic cross-aldol reaction. These results seemed to confirm the notion of acetaldehyde being uncontrollable. “However,” said Professor List, “we decided to see these results as encouragement instead: They showed that it is possible to (a) achieve decent enantioselectivities and (b) high yields in organocatalytic reactions of acetaldehyde. Now all we had to do was to combine these features in one reaction.”

Recently, Professor List disclosed a breakthrough methodology that successfully exploits acetaldehyde as a substrate for highly enantioselective organocatalytic Mannich reactions leading to a wide range of β-amino aldehydes in enantiopure form.

Professor List and his group chose to use the Mannich reaction of N-Boc-imines they previously developed for their research efforts. “Not only had we already gained valuable experience on this highly efficient and enantioselective reaction from previous studies,” said Professor List, “but the β-amino aldehydes obtained as products would be valuable intermediates in β-amino acid syntheses as well as drug building blocks. We could quickly show that proline, an inexpensive, commercially available, and easy-to-handle catalyst, is able to produce the desired products in essentially enantiopure form. However,” he continued, “initial yields were extremely low and most of our efforts were then directed at devising reaction conditions that would lead to acceptable yields.”
Inspiration and a team effort in the lab ultimately led to success, and List and co-workers obtained the products in reasonable to good yields and excellent enantioselectivities. “Moreover,” Professor List said, “we were also able to develop several applications of our products, some of them offering shortcuts in the syntheses of newly approved drugs.”

“Our report,” said Professor List, “together with that of Hayashi and coworkers, who independently developed the corresponding cross-aldol reaction,’ describes the first useful applications of acetaldehyde as a nucleophile in organic synthesis. We are convinced of its enormous potential and are currently investigating several other reactions of acetaldehyde. For example, we have developed analogous organocatalytic asymmetric Michael reactions that provide the corresponding products in high enantioselectivities. The importance of our present work,” he concluded, “is underlined by the fact that pharmaceutical companies have already expressed their interest in its use.”

REFERENCES

Dearomatization of substituted phenols followed by a de-
symmetrization reaction to form chiral intermediates repre-
sents a popular strategy for synthesizing natural product mole-
cules. A further contribution in this area was recently reported
by Dr. Matthew J. Gaunt and his group at the University of
Cambridge (UK), who disclosed a catalytic enantioselective
single-reaction method for direct conversion of phenols into
highly functionalized chiral molecules. Gaunt and coworkers
carried out fast oxidation of para-substituted phenols to form
cyclohexadienones coupled with an amine-catalyzed intramo-
lecular Michael addition. Oxidation of the phenol ring occurs
rapidly without affecting the aldehyde side chain by using
methanol as the solvent and nucleophile and a hypervalent
iodine oxidizing reagent, PhI(OCOMe)₂. Minimal contact bet-
ween the iodine reagent and the chiral amine catalyst prevents
side reactions and the protic solvent methanol helps to control
the stereochemistry. The results are complex, non-racemic,
polycyclic molecules containing three new stereogenic centers
and an array of exploitable orthogonal functionality, directly from a
flat molecule that is devoid of architectural complexity.

Concerning what triggered the interest of Dr. Gaunt’s
group for this topic, he explained that “the elegance of these
stepwise tactics led us to speculate that a catalytic asymmetric
process that can directly transform an aromatic motif into the
non-racemic structure would provide a powerful strategy for
the rapid chemical synthesis of complex molecules. We de-
veloped a process that directly converts a para-substituted phen-
ol into a highly functionalized chiral product via oxidative
dearomatization and amine-catalyzed enantioselective de-
symmetrizing Michael reaction,” Dr. Gaunt continued. “This
one-step transformation reveals a complex structure, formed
with exquisite control of three new stereogenic centers and an
array of exploitable orthogonal functionality, directly from a
flat molecule that is devoid of architectural complexity.”

“We are currently investigating the application of this pro-
cess in the synthesis of natural products, in particular alkaloid,
terpene and polyketide structures,” concluded Dr. Gaunt. “We
have recently uncovered some exciting leads that will enable
us to generate complex non-racemic natural product struc-
tures directly from flat aromatic systems in a single step using
our catalytic enantioselective dearomatization strategy.”

Additionally, the research groups of Feringa, Hayashi and
Rovis have developed catalytic desymmetrization methods to
convert the cyclohexadienone motif into useful enantioenriched
molecules.”
Phenol oxidations to bicyclic frameworks

10 mol% catalyst
Ph-O-(OAc)$_2$
Nu-H
Ar = 2-naphthyl

99% ee, 12 examples

Proposed catalytic cycle

Proposed model for stereoselectivity
About the corresponding author

Following his BSc in Chemistry at the University of Birmingham (UK) Matthew Gaunt received his PhD in organic chemistry from the University of Cambridge (UK) under the guidance of Dr. Jonathan Spencer. Following postdoctoral studies with Professor Amos B. Smith III at the University of Pennsylvania (USA) he returned to Cambridge as a Research Fellow with Professor Steven Ley. He was appointed to the Faculty at Cambridge in October 2003 where he began his independent career and was recently appointed as a Royal Society University Research Fellow in October 2004. In October 2006 he was awarded tenure at Cambridge and promoted to Lecturer in Synthetic Organic Chemistry as a Philip & Patricia Brown Next Generation Fellow. Dr. Gaunt’s research program is centered around the development of chemical synthesis using enantio-selective catalysis, with specific interests in organocatalysis, metal-catalyzed C–H activation, cascade processes for the rapid synthesis of complex molecules and applications in chemical biology. The group’s research has been acknowledged by the award of DowPharma Prize for Creativity in Chiral Chemistry in 2005.
Synthesis and Absolute Configuration of Hormone α1


- Phytophthora, whose name translates as “plant-destroyer”, is one of the most destructive pathogens in the world. In the mid-1840s, late blight, the plant disease caused by a member of this fungus-like genus, destroyed potato crops in Europe and the United States and caused the Irish potato famine. The life cycle of Phytophthora species features characteristic biological events, including sexual reproduction. Each individual is bisexual, capable of producing both female (oogonia) and male (antheridia). There are two mating types, A1 and A2, with sexual reproduction requiring the interaction of both. After sexual reproduction, the oogonia develop into sexual spores called oospores, which can survive harsh conditions such as drying or freezing for months or years in the absence of a living host plant. “In 1929, Ashby proposed that sexual reproduction in Phytophthora was regulated by a hormone-like compound,” explained Dr. Arata Yajima from the Faculty of Applied Bio-Science, Tokyo University of Agriculture (Japan). “A factor secreted by the A1 mating type induces the formation of oospores in the A2 mating type, while a factor secreted by A2 induces the formation of oospores in A1. These factors are known as hormones α1 and α2, respectively,” he continued. “Recently, Professor Ojika from Nagoya University (Japan) and co-workers succeeded in isolating hormone α1 from 1830 L of culture broth of the A1 mating type of P. nicotianae. Surprisingly, hormone α1 was found to induce oospore formation not only in P. nicotianae but also in P. capsici, P. cambivora and P. infestans.” These results indicate that hormone α1 is a universal mating hormone in the heterothallic species of Phytophthora. “We became interested in synthesizing hormone α1 in order to confirm its structure and to investigate its biological activity,” said Dr. Yajima.

“This work might be one of the historical works on Phytophthora. More than 70 years after the first proposal of the existence of α1 by Ashby, we have succeeded in establishing the complete structure of α1. It goes without saying that the greatest contribution to solve the moldy old mystery, the structure of hormone α1, is Professor Ojika’s isolation and elucidation of the plane structure of α1. The remaining last piece of the mystery was its absolute configuration, and we put it on the mysterious picture.”

According to Dr. Yajima, in the first synthesis the researchers suffered from the partial racemization of some asymmetric centers. “Because the NMR spectra of the stereoisomers of the synthetic intermediates and the final products were indistinguishable from each other, we could not realize the deterioration of stereoisomeric purity before we analyzed the corresponding MTPA esters of the final products,” he said. “In the second synthesis, we carefully chose the synthetic protocol to avoid any racemization, especially for C3 adjacent to the easily enolizable carbonyl group. Thus, we designed the second synthetic protocol toward hormone α1 as simple as possible.” The bioassay, carried out by Professor Qi’s group from the School of Pharmacy, Fudan University (P. R. of China), using synthetic samples from the two countries, Japan and China, showed the same result. “Only the synthetic (3R,7R,11R,15R)-isomer shows the oospore-inducing activity, which indicates that Phytophthora has some stereospecific hormone receptor,” said Dr. Yajima. “Since we did not synthesize 3S- or 15S-isomers, we still don’t know whether the stereospecificity of their hormone receptor is strict or not. But, 3S- or 15S-isomers of α1 will be synthesized and assayed, which will bring full information of the stereospecificity of the receptor in the near future. Now our attention is focused on the hormone receptor,” he said. “We are now trying to identify the receptor by using the synthetic α1 and its derivatives as a chemical probe. The information about the receptor will open the way to manage Phytophthora injury to potato or tomato crops.”
There are also interesting behind-the-scenes in this highly successful example of a collaborative project. “When I found the plane structure of hormone α1 in Professor Ojika’s paper, I was interested in it and the old story of α1,” said Dr. Yajima. “However, α1 has four asymmetric carbons on its linear carbon skeleton, and it seemed to be difficult to determine its absolute configuration by analytical methods, which meant that we had to synthesize the sixteen stereoisomers of α1 to compare their biological activity. In our preliminary work, we found that the stereoisomeric mixture showed a five times weaker oospore inducing activity,” he continued. “This indicated that some stereoisomers of α1 show the activity, and we could winnow the candidates down from sixteen to some extent. Thus, we started the synthesis of optically active α1. If the synthetic stereoisomeric mixture showed the same activity as the natural product, we might abandon the project of the asymmetric synthesis of α1.” Fortunately, during the synthetic studies, Professor Ojika brought a piece of good news. “They succeeded in determining the absolute configuration of two asymmetric centers by MTPA method, which limited the number of possible stereoisomers to four,” said Dr. Yajima. “This information encouraged us to complete the synthetic studies. We have prepared the four stereoisomers of α1, and only one isomer exhibited hormonal activity. How lucky we are! If the other isomers exhibited the activity, it would have been impossible to determine the absolute configuration of the natural product by this study. We owe our success to the stereospecificity of the Phytophthora’s hormone receptor!”

About the authors

This work is a collaborative work of three groups from Japan and P. R. of China. Dr. Yajima’s group and Professor Qin’s group synthesized the four stereoisomers of α1, and Professor Qi’s group assayed the synthetic samples. Originally, Professor Qin’s group and Dr. Yajima’s group were competitors. The two groups succeeded in synthesizing optically active α1 almost at the same time through totally different ways. The two groups became aware of each other’s activity when they both asked Professor Qi to bio-assay their samples. Finally, the groups followed the suggestion coming from Professor Sakagami from Nagoya University (Japan), and from an editor of Nature Chemical Biology and agreed to submit a collaborative paper.

Arata Yajima was born in Tokyo (Japan) in 1972. He received his BSc (1995) and PhD (2000) from Tokyo University of Science under the direction of Professor Kenji Mori, where he completed the total synthesis of phyto-cassane D. In 2000, he was appointed as a Research Associate at Tokyo University of Agriculture, and promoted to Assistant Professor in 2003. His current research interests are focused on the total synthesis of micro-organism regulators and rice phytoalexins and their chemical biology.

Yong Qin was born in Yunnan (P. R. of China) in 1967. He received his BSc (1989) from Yunnan University and MSc (1992) from the Chengdu Institute of Organic Chemistry and was appointed as Assistant and Associate Professor at Chengdu Institute of Organic Chemistry (1995–1996). He received his PhD (1995) from the Institute of Chemistry under the direction of Professors Zhitang Huang and Yaozhong Jiang. He joined the group of Professor Martin E. Kuehne at the University of Vermont (USA) as a postdoctoral fellow (1996–2000). He was appointed as research scientist at Triad Therapeutics, Inc. (USA) (2000–2003), before he moved to West China School of Pharmacy, Sichuan University, as a Professor and Vice Dean. His research interests are focused on the total synthesis of natural products, asymmetric synthesis and natural medicinal chemistry.

Jianhua Qi was born in Sichuan Province (P. R. of China) in 1967. He received his BSc (1987) from Sichuan Normal University (P. R. of China), his MSc (2000) and PhD (2002) from Nagoya University (Japan) under the direction of Professor Youji Sakagami. He stayed in Professor Sakagami’s laboratory as a post-doctoral fellow with a Japan Society for the Promotion of Science Fellowship (2002–2007), before he moved to the School of Pharmacy, Fudan University (P. R. of China), as a Professor. His research interests are focused on the isolation and structure determination of endogenous and exogenous bioactive substances, biological mechanisms, synthesis and biosynthesis of bioactive compounds.
FURTHER HIGHLIGHTS

SYNTHESIS

SYNLETT
Account on: Strategies for Constructing Diverse Chiral Environments in Multimetallic Bifunctional Asymmetric Catalysis
(by M. Shibasaki)

SYNFACTS
Synfact of the Month in category “Metal-Mediated Synthesis”: Suzuki–Miyaura Cross-Coupling of Trifluoroboratohomoenolates

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