Marquette University e-Publications@Marquette

Chemistry Faculty Research and Publications

Chemistry, Department of

8-1-2009

Recent Applications of Acyclic (Diene)iron Complexes and (Dienyl)iron Cations in Organic Synthesis

William Donaldson Marquette University, william.donaldson@marquette.edu

Subhabrata Chaudhury Marquette University

Accepted version. *European Journal of Organic Chemistry*, Volume 2009, Issue 23 (August 2009). pp 3831-3843. DOI: 10.1002/ejoc.200900141 © 2009 Wiley-VCH Verlag. Used with permission. This is the pre-peer reviewed version of the article, which has been published in final form.

Recent Applications of Acyclic (Diene)iron Complexes and (Dienyl)iron Cations in Organic Synthesis

William A. Donaldson Department of Chemistry, Marquette University Milwaukee, WI

Subhabrata Chaudhury

Department of Chemistry, Marquette University Milwaukee, WI

Abstract:

Complexation of (tricarbonyl)iron to an acyclic diene serves to protect the ligand against oxidation, reduction and cycloaddition reactions while the steric bulk of this adjunct serves to direct the approach reagents to unsaturated groups attached to the diene onto the face opposite to iron. Furthermore, the Fe(CO)₃ moiety can serve to stabilize carbocation centers adjacent to the diene (i.e. pentadienyl-iron cations). Recent applications of these reactivities to the synthesis of polyene, cyclopropane, cycloheptadiene and cyclohexenone containing natural products or analogs will be presented.

Keywords: Diene ligands, Iron, Synthetic methods, Regioselective nucleophilic addition.

Introduction

While Reihlen and co-workers were the first to prepare an acyclic (butadiene)(tricarbonyl)iron (**1**, Figure 1) in 1930,[1a] the structure of this compound was not proposed until 1958 by Hallam and Pauson who were also the first to note that complexation of butadiene to iron protected the ligand towards catalytic reduction and cycloaddition reactions.[1b] Their structural assignment was eventually corroborated by X-ray crystallography in 1963.[1c] At about this same time, acyclic (pentadienyl)iron(1+) cations (**2**) were first reported by Pettit and co-workers.[2] Complexes of these types as well as the corresponding cyclic counterparts (**3**, **4**) have found great utility in the synthesis of natural products. Numerous reviews have appeared concerning the use of complexes of type **3** and **4**.[3] Similarly, reviews on the chemistry of complexes of type **1** and **2**, covering up to 1999, have appeared.[4] For this reason this review will focus on chemistry related to complexes **1** and **2** from 2000 forward.

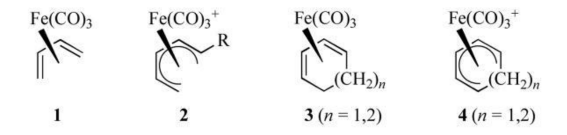


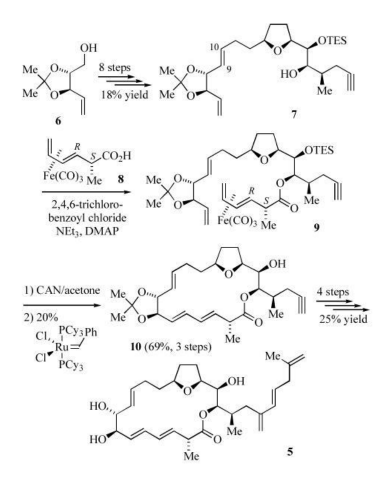
Figure 1 Structures of diene- and dienyl-iron complexes.

Use of Fe(CO)₃ as a Protecting and Stereodirecting Group

Synthesis of amphidinolide E

Amphidinolide E (**5**, Scheme 1) is a member of a family of macrolides isolated from the *Amphidinium* species of dinoflagellates.[5] Va and Roush have recently reported a synthesis of **5** which utilized $Fe(CO)_3$ to protect a 3,5-hexadienoic acid against conjugation.[6] The synthesis begins with conversion of protected 4-penten-1,2,3-triol **6** into the tetrahydrofuranyl alcohol **7** in eight steps. Key steps in this sequence included a Johnson orthoester Claisen

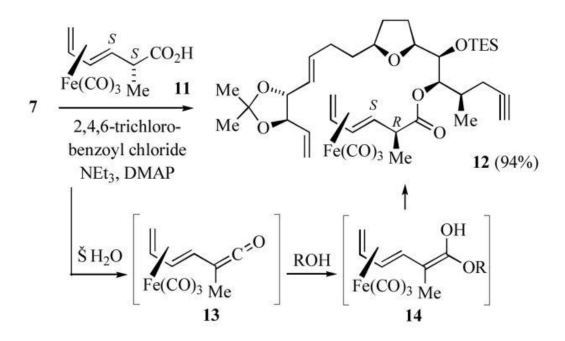
rearrangement to form the C9–C10 olefin and a [3+2] annulation[7] using an allylsilane to form the *cis*-tetrahydrofuranyl ring. Attempts at esterification of **7** with 2-methyl-3,5-hexadienoic acid were unsuccessful and generally led to recovery of **7** and the *conjugated* diene, 2-methyl-2,4-hexadienoic acid. Alternatively, esterification of (2*S*,3*R*) (2-methyl-3,5-hexadienoic acid)Fe(CO)₃[8] (**8**) with **7** cleanly gave **9**. In this case, iron serves as a protecting group such that the diene does not undergo isomerization. Oxidative decomplexation of **9**, followed by ring closing metathesis[9] with Grubbs' 1st generation catalyst afforded the macrolide ring **10** exclusively as the 3*E*,5*E*,9*E*-stereoisomer. Completion of the synthesis involved hydrostanallation of the alkyne, conversion to the 2-alkenyl iodide, cleavage of the protecting groups and Pd-catalyzed coupling.

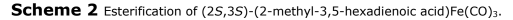


Scheme 1 Synthesis of amphidinolide E.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

During the course of this work, Va and Roush discovered that the esterification of the diastereomeric (2S,3S) (2-methyl-3,5hexadienoic acid)Fe(CO)₃ (**11**) with **7** proceeded with complete inversion of the C2 methyl bond to afford **12** (Scheme 2).[10] These authors propose that the esterification of **11** proceeds via dehydration to generate the ketene intermediate **13**; addition of the alcohol generates the ketene hemiacetal **14**. Protonation of **14** occurs via the *s*-trans conformer and on the face opposite to the sterically bulky Fe(CO)₃ group. Notably, the *relative* configurations at C2,C3 of **9** and **12** are the same (i.e. 2S,3R compared to 2R,3S), and thus it is likely that the transformation of **8** to **9** proceeds via the enantiomeric ketene (*ent*-**13**).

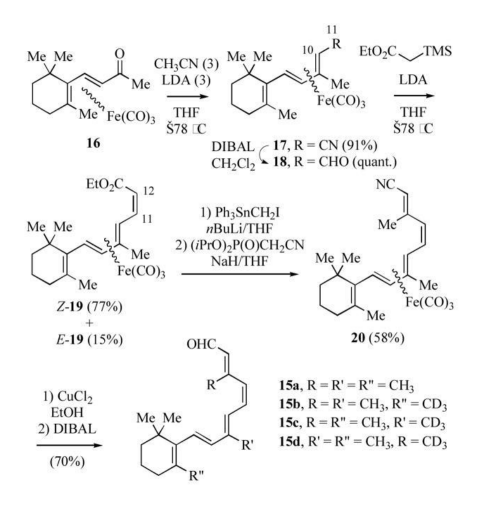




Stereoselective synthesis of 11Z-retinal

Ito and co-workers have reported a highly stereoselective synthesis of 11*Z*-retinal (**15a**, Scheme 3), the chromophore of the visual pigment rhodopsin, which utilizes Fe(CO)₃ complexation to facilitate generation of the 11*Z*-olefin.[11a–b] The synthesis begins with a nitrile aldol reaction of (β -ionone)Fe(CO)₃ (**16**) with acetonitrile. This reaction proceeds with migration of the iron fragment to give **17**.

1,3-Migration of the tricarbonyliron group have previously been observed.[4g,12] The presence of a terminal electron-withdrawing substituent (e.g. –CN) and the use of excess nucleophile generally lead to the more thermodynamically stable (diene)iron complex. Reduction of **17** gives the trienal **18**, which upon Peterson olefination with ethyl trimethylsilylacetate affords a separable mixture of *Z*- and *E*-**19** (77:15). Notably, Wittig or Horner-Emmons olefination of **18** gave only the *E*-stereoisomer. Conversion of *Z*-**19** into nitrile **20**, followed by decomplexation and nitrile reduction gave **15a**. Nakanishi's group has recently used this route to prepare the isotopically labelled 11*Z*retinals **15b d**; examination of the labelled retinals by solid state ²H NMR spectroscopy provided information on the orientation of these molecules in the rhodopsin binding pocket.[11c]



Scheme 3 Stereoselective synthesis of 11*Z*-retinal via organoiron chemistry.

The rationale for Z-selective Peterson olefination rests on approach of the anion of trimethylsilylacetate to **18** in its *s*-transconformer (about the C10–C11 bond) on the face opposite to the sterically bulky (tricarbonyl)iron moiety. Of the two synclinal transition states of lowest presumed energy, **TS–1** and **TS–2** (Figure 2), only **TS–1** avoids the steric repulsions between the bulky TMS group and the (diene)Fe(CO)₃ group. *Syn*-elimination from the resultant β silylalcohol,[13] as is known for anionic conditions, results preferentially in the 11*Z*-stereoisomer.

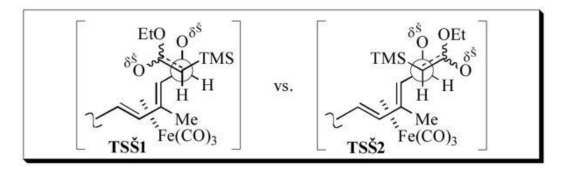
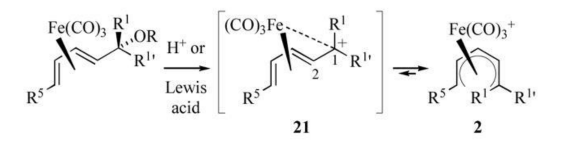


Figure 2 Rationale for Z-selective Peterson olefination of 18.

Reactivity of *In situ* Generated Transoid (Pentadienyl)iron(1+) Cations

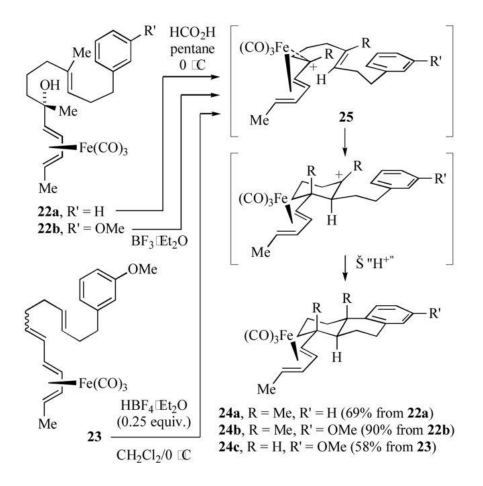
Acyclic (pentadienyl)iron(1+) cations **2** are most commonly prepared by ionization of (pentadienol)- or (pentadienyl ether)iron complexes under protic or Lewis acid conditions (Scheme 4).[2, 4] Ionization of the hydroxyl group occurs with anchimeric assistance from iron to generate the transoid pentadienyl cation **21**; subsequent isomerization of **21** occurs with retention of configuration about the C1–C2 bond.[14] In certain cases, the *in situ* generated transoid pentadienyl iron cation can undergo attack by weak nucleophiles present in the reaction mixture. These reactions generally proceed via attack at C1 on the face opposite to iron.

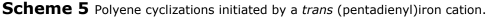


Scheme 4 Preparation of acyclic (pentadienyl)iron cations.

In situ generated transoid (pentadienyl)iron cations as initiators for polyene cyclization

Both the groups of Pearson[15] and of Franck-Neumann[16] reported polyene cyclizations initiated by *in situ* generated transoid (pentadienyl)iron cations (Scheme 5). These cyclizations may be terminated by attack of fluoride ion, formate ion or pendant electronrich aromatic groups. For example, reaction of dienol complexes **22a** or **b**,[16b] or the conjugated triene **23**[15c] under either protic or Lewis acidic conditions resulted in the diastereoselective formation of the octahydrophenanthrene skeletons **24a–c**. The relative configurations of **24a/b** were determined by X-ray crystallography, while the relative configuration of **24c** was assigned on the basis of extensive NMR spectral analysis of the free ligand (prepared by oxidation of **24c** with excess Me₃NO). The cyclizations were found to occur in a diastereoselective fashion; initial C-C bond formation occurred on the transoid (pentadienyl)iron cation (**25**) on the face opposite to the sterically bulky Fe(CO)₃ group.

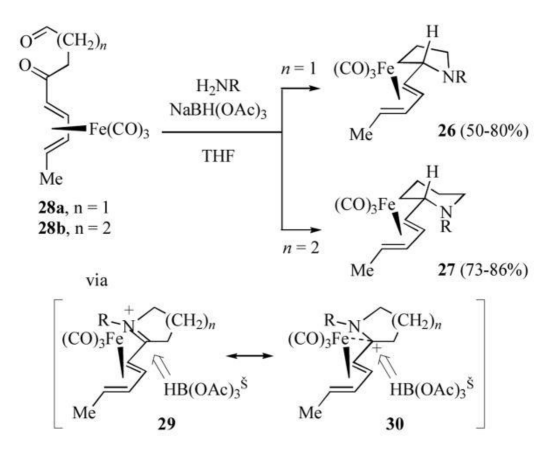




Diastereoselective Preparation of Dienyl Pyrrolidines and Piperidines

Cox and co-workers have reported on the diastereoselective preparation of dienyl pyrrolidine and dienyl piperidine complexes (**26** and **27** respectively, Scheme 6) by the reductive amination of ketoaldehydes **28a** and **28b**.[17] These reactions are proposed to proceed via reductive amination at the aldehyde, followed by generation of the iminium complex **29** (an alternative resonance contributor is the transoid pentadienyl iron cation **30**). The iminium ion/pentadienyl cation complex is preferentially oriented in the *s*-trans conformer about the diene-to-iminium carbon so as to minimize repulsion between the diene and the substituent R on nitrogen. Approach of hydride to the face opposite to iron (followed by rotation about the diene to pyrrolidine/piperidine bond) generated the products

with excellent diastereoselective control (ψ -exo diastereomer).[18] The relative configurations of **26** and **27** were confirmed by X-ray crystal structures of one example each.



Scheme 6 Reductive amination of diene ketoaldehyde complexes.

Preparation of Organoiron Nucleoside Analogs

Schmalz and co-workers reported on the preparation of organoiron containing nucleoside analogs by reaction of the dienyl ether complexes **31a** or **b** (prepared in 5 steps from a-methyl glucopyranoside) with silylated nucleobases in the presence of trimethylsilyl triflate (Scheme 7).[19] This reaction presumably proceeds via intermediacy of the transoid pentadienyl cation **32**. Nucleophilic attack on **32** occurs predominantly on the face opposite to the sterically bulky Fe(CO)₃ group to afford *exo*-**33a** or **b** as the major product, along with a lesser amount of the diastereomeric *endo* complex. Complexes *exo*-**33a** and *exo*-**33b** were found to be cytotoxic

against cultivated BJAB tumor cells (IC₉₀ = 30 and 20 μ M respectively). The cytotoxicity of *exo*-**33b** was attributed to its ability to induce apoptosis by DNA fragmentation. Notably, the free ligand of complex *exo*-**33b** exhibited considerably diminished cytotoxicity (IC₉₀ > 100 μ M), indicating a critical, but as yet undetermined, role for the metal.

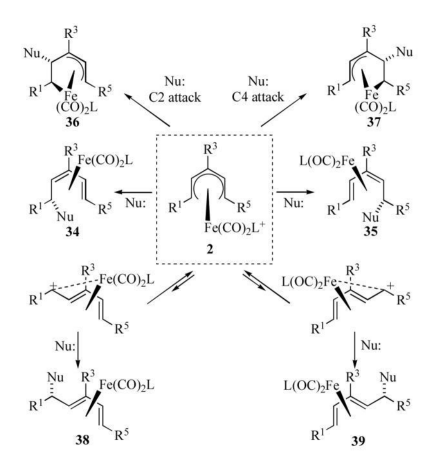
 α -methylglucopyranoside 5 steps NHTMS TDSO TDSO OMe Н Η TMSOTf OTMS e(CO)₃ CH₂Cl₂ $(CO)_3$ then H₂O 32 31a, R = H(26%)workup **31b**, $R = CO_2Et (32\%)$ NH_2 TDSO TDSO H' Н VH-Fe(CO)₃ Fe(CO)₃ exo-33a (2.3:1, 96%)endo-33a exo-33b (3.6:1, 86%)endo-33b

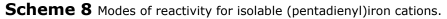
Scheme 7 Preparation of organoiron nucleoside analogs (TDS = thexyldimethylsilyl).

Reactivity of Isolable Cisoid (Pentadienyl)iron Cations

The acyclic (pentadienyl)iron(1+) cations **2** can act as excellent organometallic electrophiles toward a wide variety of nucleophiles. Nucleophilic attack can take place on the cisoid form of the pentadienyl

cation at either termini to afford the $E_{,Z}$ -diene complexes **34** or **35**, or on the internal atoms of the ligand (C2/C4) to afford complexes 36, or **37** (Scheme 8). Alternatively, since the transoid form exists in equilibrium with the cisoid form, nucleophilic attack on the transoid pentadienyl cation generates E, E-diene complexes **38** or **39** as a single diastereomer. The regioselectivity for nucleophilic attack depends on the nature of the substituents present on the pentadienyl ligand as well as the "spectator" ligand L, the nature of the nucleophile, and even the nucleophile counter ion. While not all of these factors are well understood, a few generalities can be made. [20] In general for tricarbonyl-ligated cations 2 (L = CO), weak neutral nucleophiles (e.g. H₂O, alcohols, aryl amines, electron-rich aromatics, allyl silanes) reaction proceeds via the higher energy (and thus more reactive) transoid pentadienyl forms to afford products 38/39. Reaction of more reactive organocadmium reagents, organocuprates, phosphines and alkyl amines proceeds via attack at the terminal carbons of the cisoid conformer to give products 34/35. These reactions are believed to be under frontier orbital control. If the pentadienyl ligand bears a terminal electron-withdrawing group (e.g. $R^1 = CO_2Me$), reaction with methyl lithium, alkenyl Grignards, potassium phthalimide and stabilized carbon nucleophiles proceeds by attack at C2/C4, and this regioselectivity is believed to be due to charge control (i.e. nucleophilic attack at the pentadienyl carbon bearing the greatest partial positive charge). For cations in which the substituents are neither strongly electron-withdrawing or electon-donating, nucleophilic attack frequently does not occur in a regioselective fashion. There are considerably fewer cases of acyclic (pentadienyl)iron cations bearing a phosphine ligand (i.e. $\mathbf{2}$, $L = PR_3$), however in these cases the regioselectivity is generally improved over that of their corresponding $Fe(CO)_3$ cations due to the greater stability/decreased reactivity of the $Fe(CO)_2PR_3$ cations.





Synthetic Studies on Diterpenes with a 3-Methyl-1,3Zpentadienyl Side-chain

Heteroscyphic acids A and B are novel clerodane-type diterpenes isolated from cultured cells of the liverwort *Heteroscyphus planus*, whose structural assignments (**40a/b**) were based on their MS and NMR spectral data (Figure 3).[21a–c] In particular the 12*Z*-stereochemistry for **40b** was assigned on the basis of NOEs between Me-16 and H-12 and between H-14 and H11. While no biological activity was reported for **40a** or **b**, these compounds are nonetheless structurally related to the clerodane caseargrewiin D[21d] (**41**) which exhibits both antimalarial and antitumor activity.

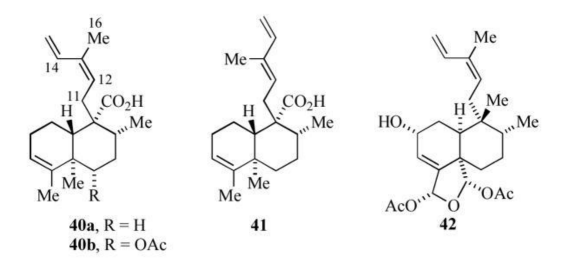
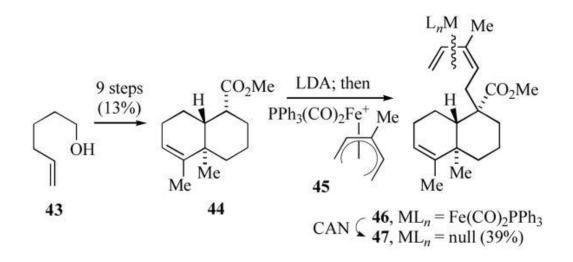


Figure 3 Proposed (**40a**) and revised (**41**) structures for heteroscyphic acid A, and structure for caeswaregiin D (**42**).

Donaldson's group envisioned introduction of the 3-methyl-1,3Z-dienyl side-chain by nucleophilic addition to a (3methylpentadienyl)iron cation.[22] To this end, 5-hexen-1-ol (43) was transformed into the decahydronaphthalene ester 44 (Scheme 9); the fused bicyclic skeleton was formed by a Mn-mediated oxidative radical cyclization.[23] Generation of the ester enolate anion from 44 and addition to the Fe(CO)₂PPh₃ ligated pentadienyl cation **45** gave complex **46**. This was produced as a mixture of diastereomers due to nucleophilic attack at one or the other pentadienyl terminal carbons of the symmetrical cation. Decomplexation of **46**, followed by purification by $AqNO_3$ impregnated silica gel qave **47** as a single diastereomer. It was surprising to note that the NMR spectral data for the dienyl sidechain of **47** (confirmed as *Z* by comparison of its NMR spectral data to that of other known diterpenes possessing a 3-methyl-1,3Zdienyl group) did not match well with that reported for the heteroscyphic acids A and B. In fact, the chemical shifts reported for heteroscyphic acids A and B are more consistent with those observed for a number of diterpenes possessing a 3-methyl-1,3E-dienyl group, and thus it was suggested that the heteroscyphic acids have this geometry for the sidechain (c.f. **41**, Figure 3). This methodology might prove useful for the introduction of the 3-methyl-1,3Z-dienyl sidechain in **42**.





Synthetic Studies on Macrolactin A

Macrolactin A (**48**, Figure 4) is a polyene macrolide aglycon originally isolated from a taxonomically unidentified marine bacterium.[24] More recently, other members of this family of 24membered macrolides have been isolated from *Bacillus* sp. Sc026, *Bacillus* sp. PP19-H3, and *Actinomadura* sp.[25] Initial screening revealed that **48** displayed antibacterial, antiviral and antitumor activity. The complex structure of macrolactin A presents several synthetic challenges, including four sp³ asymmetric centers and three conjugated dienes. Several groups have reported synthetic studies,[26] including total syntheses by the groups of Smith,[27b] Carreira,[27c] and Marino.[27d]

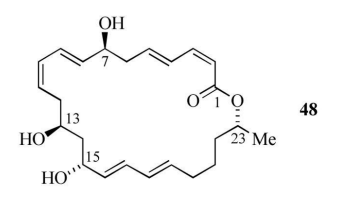
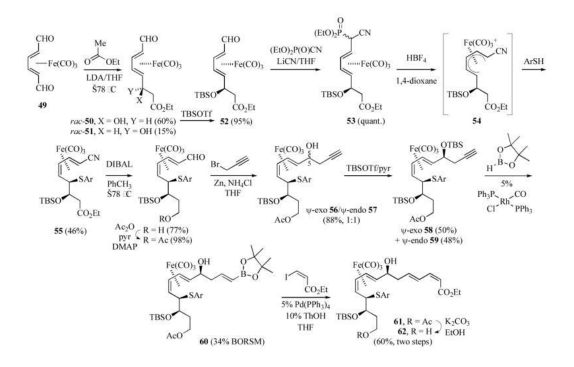


Figure 4 Structure of macrolactin A (48).

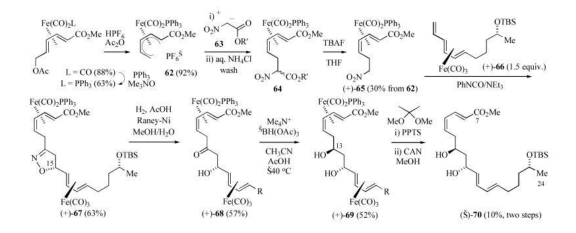
Takemoto's group has prepared the C1–C15 segment of macrolactin A, in racemic form, utilizing the $Fe(CO)_3$ group as a mobile chiral auxiliary.[28] The synthesis begins with the achiral (2,4hexadiendial)Fe(CO)₃ complex **49** (Scheme 10). Condensation of **49** with the enolate anion derived from ethyl acetate proceeded in a diastereoselective fashion to afford a separable mixture of predominantly the ψ -exo β -hydroxyester rac-**50** along with the ψ endo alcohol rac-51. Reaction of the derived TBS ether 52 with diethyl phosphorocyanidate gave the crude cyanophosphate 53 as a mixture of diastereomers, which were used in the next step without further purification. Protonation of 53 with HBF₄ in the presence of 4fluorobenzenethiol afforded the E_{z} -dienylnitrile complex 55, along with a minor amount of the corresponding *E*,*E*-diene complex. This 1,2-migration of iron presumably proceeds via the intermediacy of the cisoid (pentadienyl)iron cation 54. The success of this reaction was highly dependent on the solvent and acid used; use of BF_3 -etherate gave greatly diminished yields of 55 at the expense of a variety of other nucleophilic addition products. Similarly, attempts to use hydride nucleophiles (Et₃SiH or NaBH₃CN) in the in situ formation of 54 were unsuccessful. Treatment of **54** with 6 equivalents of DIBAL, followed by quenching with aqueous NH₄Cl, resulted in reduction of the nitrile and ester to an aldehyde and 1° alcohol respectively. After protection of the 1° alcohol as an acetate, addition of the organozinc reagent prepared from propargyl bromide and zinc in the presence of NH₄Cl gave an equimolar mixture of the diastereometric dienol complexes ψ exo **56** and ψ -endo **57**. Separation of the diastereomers was possible after protection as their TBS ethers 58/59. Rh-catalyzed hydroboration of ψ -exo **58** with pinacolborane gave the crude Evinylboronate 60 in modest yield. Pd-catalyzed coupling of 60 with ethyl (Z)-3-iodopropenoate afforded a mixture of acetate **61** and alcohol 62. Hydrolysis of 61 afforded the (2Z,4E,8E,10Zpentadecatetraenyl complex 62 in 60% overall yield from 60. Unfortunately, while the 1° alcohol of 62 could be oxidized with IBX, attempts at coupling the resultant aldehyde with an alkenylzirconium reagent, to generate the C15-C16 bond, were unsuccessful.



Scheme 10 Takemoto synthesis of the C1–C15 segment of macrolactin A (Ar = p-FC₆H₄).

Li and Donaldson have also applied diene-iron complexes to the synthesis of the C7–C24 segment of macrolactin A in enantiomerically enriched form (\geq 90% ee) (Scheme 11).[29a] Generation of the 8E,10Z-diene segment of macrolactin utilized nucleophilic addition to the enantiomerically enriched Fe(CO)₂PPh₃ ligated cation **62**. This cation was prepared by standard procedures from enantiomerically pure methyl 6-oxo-2,4-hexadienoate complex.[30] Addition of nitroacetate anion 63 proceeds at an internal pentadienyl carbon under kinetic control, however a brief work-up of the initially formed (pentenediyl)iron complex with aqueous NH₄Cl gave the E,Z-dienoate complex 64 as a mixture of diastereomers. This isomerization is proposed to proceed via protonation at the ester carbonyl, dissociation to the (pentadienyl)Fe(CO)₂PPh₃⁺ cation, followed by attack at the terminal position to generate the more thermodynamically E,Zdienoate complex. Cleavage of the trimethylsilylethoxy ester from 64 and subsequent decarboxylation generated the C7-C13 segment (+)-65. Generation of the nitrile oxide from (+)-65 using Mukiayama conditions[31] in the presence of 1.5 equivalents of the enantiomerically enriched triene complex (+)-**66**[32] (\geq 90% ee) gave

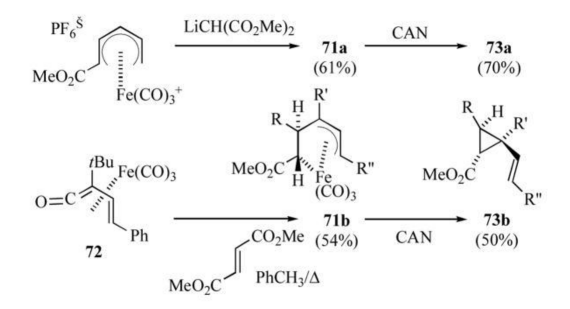
the bimetallic tetraene isoxazoline (+)-**67** as a single diastereomer. The selective formation of the ψ -exo diastereomer in this intermolecular cycloaddition is due to approach of the nitrile oxide on the less hindered face of the s-trans triene rotomer.[33] Reductive hydrolysis of isoxazoline 67, using commercially purchased Raneynickel, gave the bimetallic β -hydroxyketone (+)-**68**. Using this less reactive form of the catalyst, the two iron adjuncts serve to protect the diene segments against hydrogenation.[34] Diastereoselective reduction[35] of **68** gave the diol (+)-**69**. Generation of the acetonide followed by oxidative decomplexation with CAN gave the tetraene (-)-**70**. Oxidative removal of the two iron moieties was accompanied by cleavage of the acetonide group due to the acid generated under these reaction conditions. The diminished yield for this last step may be due to the lability of this tetraenyldiol as others have reported that removal of hydroxyl protecting groups from intact macrolactin A has proven to be difficult.[27b] In this synthesis of the C7–C24 segment, the ironcarbonyl adjuncts are responsible for *i*) stereoselective preparation for the C8–C11 E,Z-diene, ii) diastereoselective generation of the C23 alcohol by remote asymmetric induction, iii) introduction of the C15 stereocenter by a highly diastereoselective intermolecular nitrile oxideolefin cycloaddition, and iv) protection of the C8-C11 and C16-C19 dienes during reductive hydrolysis of the isoxazoline group.



Scheme 11 Synthesis of the enantiomerically enriched C7–C24 segment of macrolactin A ($R = CH_2CH_2CH_2CH(OTBS)Me$, $R' = CH_2CH_2TMS$).

Synthesis of Vinylcyclopropanes

(3-Pentene-1,5-diyl)iron complexes **71a**, bearing an electron withdrawing group at C1, have been prepared by addition of carbon nucleophiles to (pentadienyl)iron(1+) cations (Scheme 12).[36] Alternatively, the thermal reaction of (vinylketene)iron complex **72** with dimethylfumarate generated the (pentenediyl)iron complex **71b**.[37] Oxidation of either **71a** or **71b** with ceric ammonium nitrate gave the vinylcyclopropanecarboxylates **73a** or **b**.[36a, 37] Since this is formally an oxidatively induced-reductive elimination, the reaction generally proceeds with retention of configuration at the two centers undergoing C–C bond formation.



Scheme 12 Synthesis of vinylcyclopropanes via oxidative decomplexation of (3pentene-1,5-diyl)iron complexes (\mathbf{a} , $R = CH(CO_2Me)_2$, R' = R'' = H; \mathbf{b} , $R = CO_2Me$, R' = tBu, R'' = Ph).

Synthesis of 2-(2-Carboxycyclopropyl)glycines and Dysibetaine CPa

The selective activation of different glutamate receptors may depend on recognition of a particular conformer of this flexible molecule. For this reason, the synthesis and evaluation of a number of 2-(2'-carboxycyclopropyl)glycines (e.g. **74a–f**, Figure 5), as

conformationally restricted analogs of glutamate, has led to the discovery of ligands with mGluR specificity.[38] In particular the extended conformation, as exemplified by compounds **74a–d**, is believed to be a requirement for binding to the mGluR1 and mGluR2 receptors. Recently, Sakai and co-workers isolated a novel water-soluble cyclopropane containing betaine from *D. herbacea* which they termed dysibetaine CPa (**75**, Figure 5).[39] Compound **75** displaced kinate from the NMDA-type glutamate receptor with $IC_{50} = 13 \mu M$.

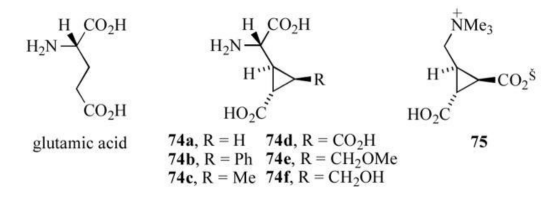
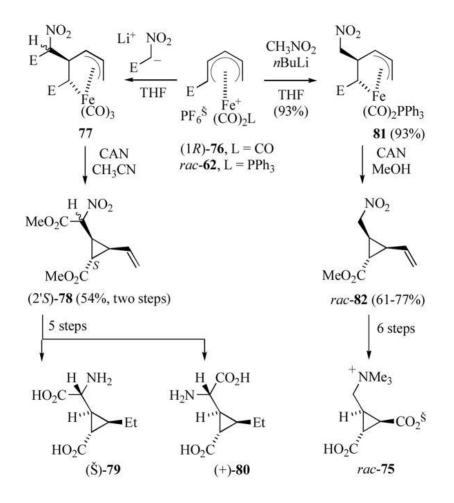


Figure 5 Structure of conformationally restricted glutamate analogs (**74**) and dysibetaine CPa (**75**).

Reaction of the enantiomerically enriched tricarbonyl-ligated cation (1R)-**76** (\geq 80% ee) with the anion generated from methyl nitroacetate gave (pentenediyl)iron complex **77** as a mixture of diastereomers at the nitroacetate carbon (Scheme 13).[36a] Decomplexation of the mixture of diastereomers afforded vinylcyclopropanecarboxylate (2'S)-**78** as an inseparable mixture of diastereomers at the nitroacetate carbon. Transformation of the diastereomeric mixture (2'S)-**78** into the individual 3-ethyl CCGs (–)-**79** and (+)-**80** required reduction of the vinyl and nitro groups, conversion of the amines into a separable mixture of diphenylmethylene imines,[40] hydrolysis of the separate diphenylmethylene imines and the methyl esters and finally generation of the free bases.



Scheme 13 Synthesis of 2-(2'-carboxycyclopropyl)glycines and dysibetaine CPa ($E = CO_2Me$).

For the preparation of dysibetaine CPa, reaction of the dicarbonyl(triphenylphosphine)-ligated cation *rac*-**62** with the anion generated from nitromethane gave (pentenediyl)iron complex **81** in excellent yield (Scheme 13).[41] Oxidative decomplexation of **81** gave the vinylcyclopropanecarboxylate **82**. Transformation of **82** into *rac*-**75** required conversion of the vinyl functionality to an ester, subsequent reduction of the 1° nitro group, hydrolysis and exhaustive methylation.

Synthesis of the C9–C16 Segment of Ambruticin

Ambruticin (**83**, Figure 6) is a structurally unique carboxylic acid isolated from *Polyangium cellulosum var fulvum*, which exhibits potent oral antifungal activity against *Coccidioides immitis*, *Histoplasma capsulatum*, and *Blastomyces dermititidis*.[42] The complex structure of ambruticin presents several synthetic challenges, including a tetrahydropyranyl ring, a dihydropyranyl ring and a divinylcyclopropane ring. Several groups have reported synthetic studies,[43] including total syntheses by the groups of Kende,[44a] Jacobsen,[44b] Martin.[44c] and Lee.[44d]

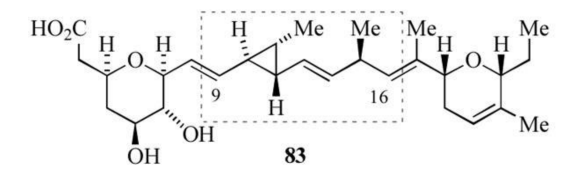
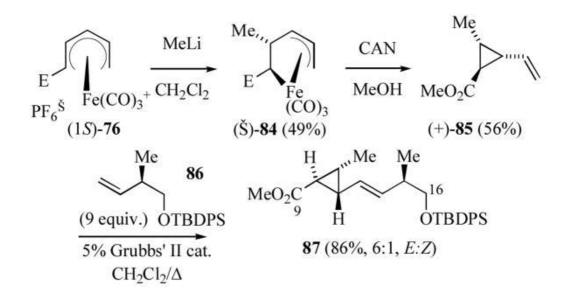
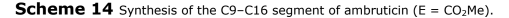


Figure 6 Structure of the antifungal agent ambruticin (83).

Reaction of (1S)-**76** in CH₂Cl₂ with a ethereal solution of methyl lithium gave the (pentenediyl)iron complex (–)-**84** along with a minor amount of tricarbonyl(methyl-3,5-hexadienoate)iron (Scheme 14).[45] It was found that use of CH₂Cl₂ as solvent was crucial to the success of this reaction. Use of either ether or THF gave reduced yields of the (pentenediyl)iron complex. Oxidative decomplexation of (–)-**84** cleanly gave the stereodefined vinylcyclopropanecarboxylate (+)-**85**. Cross metathesis of **85** with a nine fold excess of (*R*)-**86** in presence of 5 mol % of Grubbs' 2nd generation catalyst gave **87** as a mixture of *E*- and *Z*-isomers (6:1 ratio).[46]



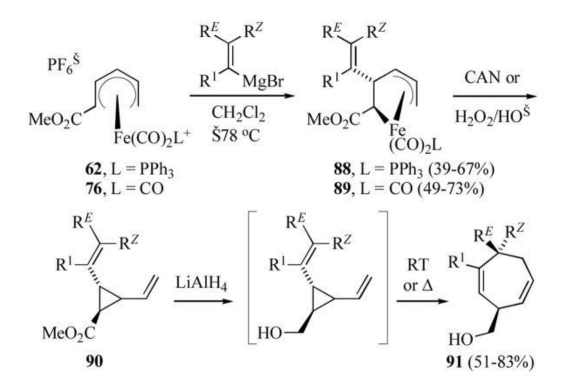


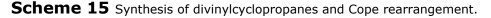
Synthesis of Divinylcyclopropanes and Cope rearrangement

Donaldson and co-workers have demonstrated that the reaction of (2-methoxycarbonylpentadienyl)iron(1+) cations 62 or 76 with alkenyl Grignard reagents gave primarily the corresponding (2-alkenyl-3-pentene-1,5-diyl)iron complexes 88 or 89 respectively (Scheme 15).[47] The yield of these products was dependent on the reaction media; use of methylene chloride gave the best results while THF or toluene led to diminished yields of 88/89. Nucleophilic attack on the face opposite to the metal was corroborated by X-ray crystal structure of the parent complex **89** ($R^1 = R^E = R^Z = H$).[47b] Oxidative decomplexation of 88/89 gave the divinylcyclopropane 90. In most cases CAN gave good yields of the 2,3-divinylcyclopropanecarboxylate, however for complexes with an electron rich 2-alkenyl group secondary oxidation of the resultant divinylcyclopropane product led to diminished yields. In these cases, oxidation with alkaline hydrogen peroxide provided superior yields, but led to mixtures of both *cis*- and trans-divinylcyclopropanes. Reduction of 90, followed by [3,3]sigmatropic rearrangement[48] afforded the 2,6cycloheptadienylmethanol **91**. While the temperature required for the Cope rearrangement varied depending on the alkenyl substituents and

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

olefin geometry, good overall yields were obtained from complexes **88**/**89**.





Synthesis of a Guianolide Skeleton

The guianolides are a family of sesquiterpenes characterized by a 5,7,5-fused tricyclic skeleton. The majority of these compounds possess a *trans*- γ -butyrolactone ring, but differ with respect to the oxygenation and oxidation state(s) of carbons 2–5, 8, 10, and 11.[49] Representative members of this family include chinesiolide B (**92**, Figure 7),[49d] cynaropikrin (**93**),[49e] and cladantholide (**94**).[49f]

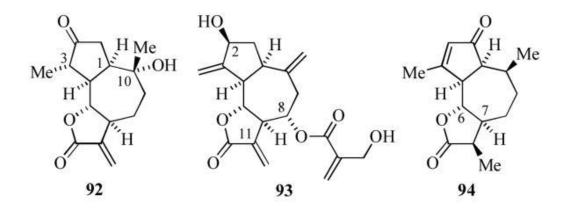
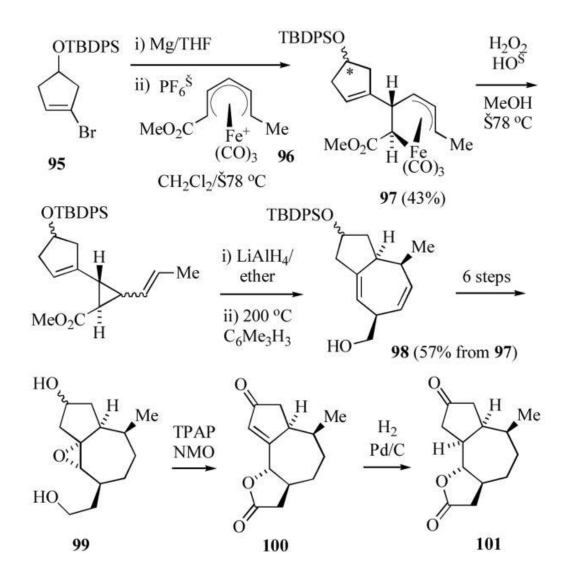


Figure 7 Representative guianolide natural products.

Donaldson and co-workers have applied organoiron methodology to the synthesis of the 5,7,5 ring system of the guianolides (Scheme 16).[50] Reaction of the Grignard reagent derived from known[49] cyclopentenyl bromide 95 with the $(dienyl)Fe(CO)_{3}^{+}$ cation **96**[50] gave the (pentenediyl)iron complex **97** as a mixture of diastereomers at the silvl ether carbon (*). Oxidative decomplexation, ester reduction and Cope rearrangement at elevated temperatures gave 98. The hexahydroazulene 98 was transformed into the epoxydiol **99** via *i*) selective hydrogenation of the less substituted olefin, ii) extension of the C3 sidechain by tosylation and cyanide displacement, *iii*) cleavage of the silvl ether, iv) epoxidation and finally, v) twofold reduction of the nitrile sidechain. Oxidation of **99** with catalytic TPAP and NMO (3.2 equiv) gave a single lactone **100**. This transformation presumably proceeds via oxidation of both the 1° and 2° alcohols, followed by β -elimination of the epoxide, generation of a lactol and further oxidation to the lactone. Reduction of 100 afforded **101**, which possesses the relative stereochemistry about the seven-membered ring of cladantholide.

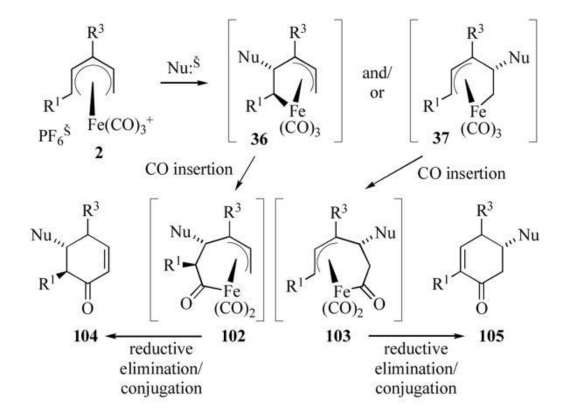


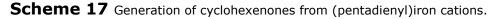
Scheme 16 Preparation of the 5,7,5 ring system of the guianolides.

Synthesis of Cyclohexenones

The (pentenediyl)iron complexes discussed in Schemes 12–16 are stable, isolable species. This is believed to be due to the fact that the presence of an electron withdrawing group attached to a carbon-metal σ -bond slows the rate of carbonyl insertion.[53] In contrast, (pentenediyl)iron complexes lacking an electron withdrawing group at C1 (e.g. **36** or **37**, Scheme 17) may be generated by nucleophilic attack on acyclic (pentadienyl)iron cations at the internal C2 position.[54] These complexes are generally unstable and undergo CO

insertion to generate the (acyl)iron complexes **102**/**103**. Reductive elimination of **102**/**103**, followed by conjugation of the olefin, gives cyclohexenones **104**/**105** respectively.

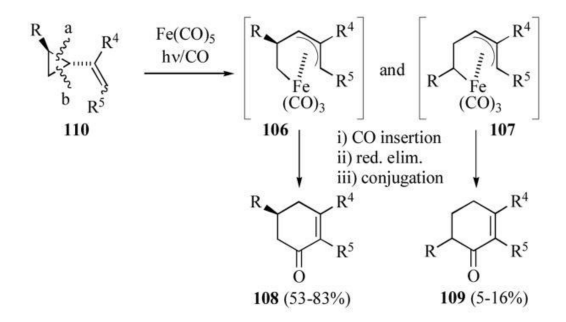




An alternative route to cyclohexenones is the photochemically initiated ring rearrangement carbonylation of alkenylcyclopropanes (Scheme 18).[55] While this reaction does not formally involve a (diene)iron complex or (dienyl)iron cation, it is nonetheless related by the presumed intermediates. This reaction is believed to proceeds via oxidative insertion of iron into one of the proximal vinylcyclopropane bonds (b or a) to generate (pentenediyl)iron intermediates **106** or **107** (respectively). Carbonyl insertion, followed reductive elimination and conjugation gives **108** or **109**. Isolation of **108** as the major cyclohexenone product indicates that insertion into the cyclopropane bond "b" is favored. Since the major product arises by cleavage of the less substituted vinylcyclopropanes bond "b", beginning with enantiomerically enriched (> 99% ee) vinylcyclopropane **110** (R = CH_2OBn , $R^4 = R^5 = H$) led to **108** in enantiomerically enriched form (>

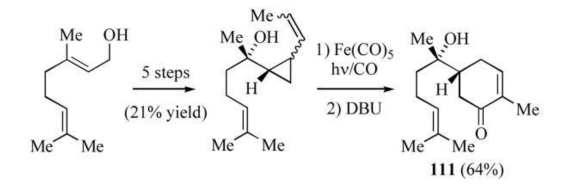
European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. DOI. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

95% ee). The enantiomeric excess and absolute configuration of the minor product **109** was not identified.



Scheme 18 Generation of cyclohexenones via iron-mediated carbonylation of alkenylcyclopropanes.

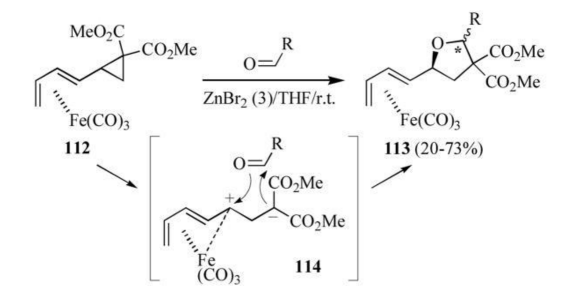
Taber and co-workers have applied this methodology to the enantioselective synthesis of (-)-delobanone **(111)** beginning with geraniol (Scheme 19).[55d]

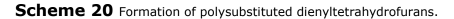


Scheme 19 Taber and co-workers synthesis of (–)-delobanone.

Miscellaneous

Christie and co-workers have reported the 1,3-dipolar cycloaddition of aldehydes with the racemic dienylcyclopropane complex **112** in the presence of $ZnBr_2$ (Scheme 20).[56] This reaction affords the dienylfurans **113** as a mixture of two (of the four possible) diastereomers. In all cases, the relative configuration at the iron-diene and the tetrahydrofuranyl carbon adjacent to the diene were found to be as indicated (i.e. ψ -exo), and thus the formed are due to the *cis*-and *trans*-2,4-disubstituted furan ring. at the dienyl at the indicated carbon (*) in moderate yield. The authors propose that this reaction proceeds via formation of the zwitterionic intermediate **114** which reacts with the aldehyde on the face opposite to the sterically bulky (tricarbonyl)iron adjunct.





Conclusions

Complexation of diene and dienyl ligands to iron facilitates the stereoselective preparation of conjugated *E,E-* and *E,Z-*1,3-dienes, trisubstituted cyclopropanes, 1,4-cycloheptadienes and cyclohexenones. These features of the (tricarbonyl)iron adjunct have been exploited by a number of research groups in the synthesis of polyene macrolides, optical pigment chromophores, heterocycles,

terpenes, conformationally restricted ligands for glutamate receptors, and antifungal agents.

Acknowledgments

WAD is grateful to his co-workers whose research accomplishments are described in this review. Their experimental skill, dedication, and keen observation turned ideas into reality. Financial support over the years from the National Institutes of Health (GM-42641) and the National Science Foundation (CHE-0415771) is gratefully acknowledged.

Biographies

• William Donaldson was born near Philadelphia, Pennsylvania. He received his B.A. degree in Chemistry from Wesleyan University (1977), and his Ph.D. in Organometallic Chemistry from Dartmouth College (1981) working with Prof. Russell Hughes, before conducting postdoctoral research with Prof. Myron Rosenblum at Brandeis University (1981–1982). Following a one-year position at Wesleyan University, he joined the faculty at Marquette University in 1983. His research has focused on the application of organoiron complexes to organic synthesis, as well as the synthesis of hydropyran natural products.

• Subhabrata Chaudhury is a Bengali from the southern part of Bengal. After receiving his B.Sc. from the University of Calcutta (1997) and M.Sc. degree from the Indian Institute of Technology, Kharagpur (1999), he joined the group of Professor Donaldson at Marquette University where he worked on the development of organoiron methodologies for the total synthesis of natural products. He obtained his Ph.D. degree in 2006 and was a postdoctoral fellow in the Department of Medicinal Chemistry at University of Kansas. Before he started his second period of postdoctoral research at the Department of Biophysics, Medical College of Wisconsin, he returned to Marquette and spent an interim period at Prof. Donaldson's laboratory. In September 2008 he moved to Scotland to join the group of Professor J. S. Clark at the University of Glasgow as a research associate. His research interests involve developing methodologies for the

preparation of organic building blocks using transition metal complexes and their application in organic synthesis.

References

- (a) Gruhl A, Hessling G, Pfrengle O, Reihlen H. Justus Liebigs Ann Chem. 1930;482:161–182.
 (b) Hallam B, Pauson P. J Chem Soc. 1958;168:642–645.
 (c) Mills OS, Robinson G. Acta Cryst. 1963;16:758–762.
- 2. (a) Mahler JE, Pettit R. J Am Chem Soc. 1963;85:3955–3959. (b) Mahler JE, Gibson DH, Pettit R. J Am Chem Soc. 1963;85:3959–3963.
- Choi TA, Czerwonka R, Forke R, Jaeger A, Knoell J, Krahl MP, Krause T, Reddy KR, Franzblau SG, Knoelker HJ. Med Chem Res. 2008;17:374– 385.Knoelker HJ. Curr Org Syn. 2004;1:309–331.Knoelker HJ, Braier A, Brocher DJ, Cammerer S, Frohner W, Gonser P, Hermann H, Herzberg D, Reddy KR, Rohde G. Pure Appl Chem. 2001;73:1075– 1086.Knoelker HJ. Chem Soc Rev. 1999;28:151–157.Stephenson GR. In: Handbook of Functionalized Organometallics. Knochel P, editor. Vol. 2. Wiley-VCH; Weinheim: 2005. pp. 569–627.Ong CW, Lia MC. Trends in Organomet Chem. 2002;4:47–58.Pearson AJ. In: Advances in Metal-Organic Chemistry. Liebeskind LS, editor. Vol. 1. JAI Press; Greenwich: 1989. pp. 1–49.
- 4. (a) Gree R. Synthesis. 1989:341-355.

(b) Gree R, Lellouche JP. In: Advances in Metal-Organic Chemistry. Liebeskind LS, editor. Vol. 4. JAI Press Inc; Greenwich, CT: 1995. pp. 129–273.

- (c) Iwata C, Takemoto Y. Chem Commun. 1996:2497–2504.
- (d) Donaldson WA. Aldrichimica Acta. 1997;30:17–24.
- (e) Donaldson WA. Curr Org Chem. 2000;4:837–868.

(f) Cox LR, Ley SV. Chem Soc Rev. 1998;27:301-314.

(g) Salzer A. In: Organometallics in Organic Synthesis. Werner H, Erker G, editors. Vol. 2. Springer-Verlag; Berlin, Germany: 1989. pp. 291–309.

- 5. (a) Kobayashi J, Ishibashi M, Murayama T, Takamatsu M, Iwamura M, Ohizumi Y, Sasaki T. J Org Chem. 1990;55:3421–3423.
 (b) Kubota T, Tsuda M, Kobayashi J. J Org Chem. 2002;67:1651–1656.
- 6. (a) Va P, Roush WR. Tetrahedron. 2007;63:5768-5796.
 (b) Va P, Roush WR. J Am Chem Soc. 2006;128:15960-15961.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

- 7. Micalizio GC, Roush WR. Org Lett. 2000;2:461-464.
- 8. For a previous preparation of 8 see: Wasicak JT, Craig RA, Henry R, Dasgupta B, Li H, Donaldson WA. Tetrahedron. 1997;53:4185–4198.
- 9. (a) Grubbs RH. Tetrahedron. 1998;54:4413–4450.
 (b) Schuster M, Blechert S. Angew Chem Int Ed Engl. 1997;36:2036–2056.
- 10. Va P, Roush WR. Org Lett. 2007;9:307-310.
- 11. (a) Wada A, Fujioka N, Tanaka Y, Ito M. J Org Chem. 2000;65:2438–2443.
 (b) Wada A, Hiraishi S, Takamura N, Date T, Aoe K, Ito M. J Org Chem. 1997;62:4343–4348.
 (c) Tanaka K, Struts AV, Krane S, Fujioka N, Salgado GFJ, Martinez-Mayrga K, Brown MF, Nakanishi K. Bull Chem Soc Jpn. 2007;80:2177–2184.
- 12. (a) Hafner A, von Philipsborn W, Salzer A. Angew Chem, Int Ed Engl. 1985;24:126–127.
 (b) Adams CM, Cerioni G, Hafner A, Kalchhauser K, von Philipsborn W, Prewo R, Schwenk A. Helv Chim Acta. 1988;71:1116–1142.
 (c) Takemoto Y, Ishii K, Ibuka T, Miwa Y, Taga T, Nakao S, Tanaka T, Oshishi H, Kai Y, Kanehisa N. J Org Chem. 2001;66:6116–6123.
- 13. (a) Hudric PF, Peterson D. J Am Chem Soc. 1975;97:1464–1468.
 (b) Ager DJ. Synthesis. 1984:384–398.
- 14. (a) Gresham DG, Kowalski DJ, Lillya CP. J Organomet Chem. 1978;144:71–79.
 (b) Dobosh PA, Gresham DG, Kowalski DJ, Lillya CP, Magyar ES. Inorg Chem. 1978;17:1775–1781.
- 15. (a) Pearson AJ, Alimardanov A, Pinkerton AA, Rouchard DM, Kirschbaum K. Tetrahedron Lett. 1998;39:5919–5922.
 (b) Pearson AJ, Alimardanov AR, Kerber WD. J Organomet Chem. 2001;630:23–32.
 (c) Pearson AJ, Ghidu VP. J Org Chem. 2004;69:8975–8978.
- 16. (a) Franck-Neumann M, Geoffroy P, Hanss D. Tetrahedron Lett. 1999;40:8487–8490.

(b) Franck-Neumann M, Geoffroy P, Hanss D. Tetrahedron Lett. 2002;43:2277–2280.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

17. (a) Williams I, Kariuki BM, Reeves K, Cox LR. Org Lett. 2006;8:4389– 4392.

(b) Williams I, Reeves K, Kariuki BM, Cox LR. Org Biomol Chem. 2007;5:3325–3329.

- 18. It should be noted that ψ-exo diastereomeric dienylpiperidenes have previously been prepared by intramolecular attack of tethered carbamate nucleophiles on in situ generated pentadienyl cations: Hachem A, Teniou A, Gree R. Bull Soc Chem Belg. 1991;100:625–626.Additionally, ψ-endo diastereomeric dienylpiperidone and 5-(dienyl)-2,3-dihydropyridinone complexes have been prepared by intramolecular attack of carbon nucleophiles or by cycloaddition of Danishefsky's diene with (dienylimine)iron complexes: Ripoche I, Bennis K, Canet JL, Gelas J, Troin Y. Tetrahedron Lett. 1996;37:3391–3392.Ripoche I, Canet JL, Gelas J, Troin Y. Eur J Org Chem. 1999:1517–1521.Takemoto Y, Ueda S, Takeuchi J, Nakamoto T, Iwata C. Tetrahedron Lett. 1994;47:8821–8824. As some of these reactions have previously been reviewed [4b,d] they are only mentioned here in the references.
- 19. Schlawe D, Majdalani A, Velcicky J, Hessler E, Wieder T, Prokop A, Schmalz HG. Angew Chem Int Ed. 2004;43:1731.
- 20. Donaldson WA, Shang L, Tao C, Yun YK, Ramaswamy M, Young VG., Jr J Organomet Chem. 1997;539:87–98.
- 21. (a) Nabeta K, Oohata T, Izumi N, Katoh K. Phytochem. 1994;37:1263–1268.
 (b) Nabeta K, Ishikawa T, Kawae T, Okuyama H. J Chem Soc, Chem Commun. 1995:681–682.
 (c) Nabeta K, Ishikawa T, Okuyama H. J Chem Soc Perkin Trans 1. 1995:3111–3115.
 (d) Kanokmedhakul S, Kanokmedhakul K, Kanarsa T, Buayairaksa M. J Nat Prod. 2005;68:183–188.
- 22. Chaudhury S, Li S, Donaldson WA. Chem Commun. 2006:2069–2070.
- 23. (a) Zoretic PA, Ramchandani M, Caspar ML. Syn Comm. 1991;21:915–922.
 (b) Snider B. Chem Rev. 1996;96:339–364.
- 24. (a) Gustofson K, Roman M, Fenical W. J Am Chem Soc. 1989;111:7519– 7524.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

(b) Rychnovsky SD, Skalitzky DJ, Pathirana C, Jensen PR, Fenical W. J Am Chem Soc. 1992;114:671–677.

- 25. (a) Jaruchoktaweechai C, Suwanborirus K, Tanasupawatt S, Kittakoop P, Menasveta P. J Nat Prod. 2000;63:984-986. (b) Nagao T, Adachi K, Sakai M, Nishijima M, Sano H. J Antibiot. 2001;54:333-339. (c) Kim HH, Kim WG, Ryoo IJ, Kim CJ, Suk JE, Han KH, Hwang SY, Yoo ID. J Microbio Biotech. 1997;7:429-434. 26. For previous synthetic studies see: (a) Benvegnu T, Schio L, Le Floc'h Y, Gree R. Synlett. 1994;7:505-506. (b) Tanimori S, Morita Y, Tsubota M, Nakayama M. Synth Commun. 1996;26:559-567. (c) Benvegnu TJ, Toupet LJ, Gree RL. Tetrahedron. 1996;52:11811-11820. (d) Benvegnu TJ, Gree RL. Tetrahedron. 1996;52:11821-11826. (e) Gonzalez A, Aiguada J, Urpi F, Vilarrasa J. Tetrahedron Lett. 1996;49:8949-8952. (f) Li S, Xu R, Bai D. Tetrahedron Lett. 2000;41:3463-3466. (g) Li SK, Rui X, Xiao XS, Bai DL. Chin J Chem. 2000;18:910–923. (h) Bonini C, Chiummiento L, Pullez M, Solladie G, Colobert F. J Org Chem. 2004;69:5015-5022. (i) Xiao X, Li S, Yan X, Liu X, Xu R, Bai D. Chem Lett. 2005;34:906-907. (j) Bonini C, Chiummiento L, Videtta V, Colobert F, Solladie G. Synlett. 2006:2427-2430. (k) Georgy M, Lesot P, Campagne JM. J Org Chem. 2007;72:3543. (I) Yadav JS, Gupta MK, Prathap I. Synthesis. 2007:1343–1348. (m) Yadav JS, Kumar MR, Sabitha G. Tetrahedron Lett. 2008;49:463-466. 27. (a) Boyce RJ, Pattenden G. Tetrahedron Lett. 1996;37:3501–3504. (b) Smith AB, III, Ott GR. J Am Chem Soc. 1998;120:3935-3948.
 - (b) Smith AB, III, Ott GR. J Am Chem Soc. 1998;120:3935–3948.
 (c) Kim Y, Singer RA, Carreira EM. Angew Chem Int Ed. 1998;37:1261–1263.
 (d) Marino JP, McClure MS, Holub DP, Comasseto JV, Tucci FC. J Am Chem Soc. 2002;124:1664–1668.
- 28. Fukuda A, Kobayashi Y, Kimachi T, Takemoto Y. Tetrahedron. 2003;59:9305–9313.
- 29. Li S, Donaldson WA. Synthesis. 2003:2064–2068.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

- 30. Godula K, Barmann H, Donaldson WA. J Org Chem. 2001;66:3590-3592.
- 31. Mukaiyama T, Hoshino T. J Am Chem Soc. 1960;82:5339–5342.
- For an earlier synthesis of the C14–C24 segment of macrolactin A via organoiron chemistry see: Barmann H, Prahlad V, Tao C, Yun YK, Wang Z, Donaldson WA. Tetrahedron. 2000;56:2289–2295.
- 33. Le Gall T, Lellouche JP, Toupet L, Beaucourt JP. Tetrahedron Lett. 1989;47:6517–6520.
- 34. Franck-Neumann, et al. have reported that reduction of (diene)Fe(CO)₃ complexes using freshly prepared Raney-nickel results in decomplexation. Franck-Neumann M, Geoffroy P, Bissinger P, Adelaide S. Tetrahedron Lett. 2001;42:6401–6404.
- 35. Evans DA, Chapman KJ, Carreira EM. J Am Chem Soc. 1988;110:3560– 3578.
- 36. (a) Yun YK, Godula K, Cao Y, Donaldson WA. J Org Chem. 2003;68:901–910.
 (b) Motiei L, Marek I, Gottlieb HE, Marks V, Lellouche JP. Tetrahedron Lett. 2003;44:5909–5012.
- 37. (a) Saberi SP, Slawin AMZ, Thomas SE, Williams DJ, Ward MF, Worthington PA. J Chem Soc, Chem Commun. 1994:2169–2170.
 (b) Gibson SE, Saberi SP, Slawin AMZ, Stanley PD, Ward MF, Williams DJ, Worthington P. J Chem Soc, Perkin Trans 1. 1995:2147–2154.
- 38. (a) Shimamoto K, Ohfune Y. Synlett. 1993:919–920.
 (b) Ohfune Y, Shimamoto K, Ishida M, Shinozaki H. Biorg Med Chem Lett. 1993;3:15–18.
 (c) Pellicciari R, Marinozzi M, Natalini B, Constantino G, Lunei R, Giorgi Ge, Moroni F, Thomsen C. J Med Chem. 1996;39:2259–2269.
 (d) Jullian N, Barbet I, Pin JP, Acher FC. J Med Chem. 1999;42:1546– 1555.
 (e) Mazon A, Pedregal C, Prowse W. Tetrahedron. 1999;55:7057– 7064.
 (f) Collado I, Pedregal C, Mazon A, Espinosa JF, Blanco-Urgoiti J, Schoeep DD, Wright RA, Johnson BG, Kingston AE. J Med Chem. 2002;45:3619–3629.

(g) Collado I, Pedregal C, Bueno AB, Marcos A, Gonzalez R, Blanco-Urgoiti J, Perez-Castells J, Schoeep DD, Wright RA, Johnson BG, Kingston AE, Moher ED, Hoard DW, Griffey KI, Tizzano JP. J Med Chem. 2004;47:456–466.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. DOI. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

- Sakai R, Suzuki K, Shimamoto K, Kamiya H. J Org Chem. 2004;69:1180– 1185.
- O'Donnell MJ. In: Encyclopedia of Reagents for Organic Synthesis. Paquette LA, editor. Vol. 1. John Wiley & Sons; New York: 1995. pp. 293–294.
- 41. Siddiquee TA, Lukesh JM, Lindeman S, Donaldson WA. J Org Chem. 2007;72:9802–9803.
- 42. Connor DT, Greenough RC, von Strandtmann M. J Org Chem. 1977;42:3664–3669. Just G, Potvin P. Can J Chem. 1980;58:2173– 2175.
- 43. For an excellent review through 2004 see: Michelet V. Curr Org Chem. 2005;9:405–409.
- 44. (a) Kende AS, Mendoza JS, Fujii Y. J Am Chem Soc. 1990;112:9645–9646.
 (b) Liu P, Jacobsen EN. J Am Chem Soc. 2001;123:10772–10773.
 (c) Kirkland TA, Colucci J, Geraci LS, Marx MA, Schneider M, Kaelin DE, Jr, Martin SF. J Am Chem Soc. 2001;123:12432–12433.
 (d) Lee E, Choi SJ, Kim H, Han HO, Kim YK, Min SJ, Son SH, Lim SM, Jang WS. Angew Chem Int Ed. 2002;41:176–178.
- 45. Lukesh JM, Donaldson WA. Chem Commun. 2005:110–112.
- 47. (a) Wallock NJ, Donaldson WA. Org Lett. 2005;7:2047–2049.
 (b) Wallock NJ, Bennett DW, Siddiquee T, Haworth DT, Donaldson WA. Synthesis. 2006:3639–3646.
 (c) Pandey RK, Wang L, Wallock NJ, Lindeman S, Donaldson WA. J Org Chem. 2008;73:7236–7245.
- 48. (a) Hudlicky T, Fan R, Reed JW, Gadamasetti KG. Org React. 1992;41:1–133.
 (b) Hudlicky R, Kutchan TM, Nagvi SM. Org React. 1985;33:247–335.
- 49. (a) Fischer NH, Olivier EJ, Fischer HD. Progress in the Chemistry of Organic Natural Products. 1979:38.
 (b) Breitmaier E. Terpenes Wiley-VCH. Germany: 2006. pp. 37–39.For

a recent review on the synthesis of guianolides with a trans-lactone moiety see:

(c) Schall A, Reiser O. Eur J Org Chem. 2008:2353–2364.

⁽d) Zhang S, Wang J, Xue H, Deng Q, Xing F, Ando M. J Nat Prod. 2002;65:1927–1929.

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.

(e) Samek Z, Holub M, Drozdz B, Iommi G, Carbella A, Gariboldi P. Tetrahedron Lett. 1971:4475–4478.
(f) Daniewski WM, Danikiewics W, Gumulka M, Pankowska E, Krajewski J, Grabarczyk H, Wichlacz M. Phytochem. 1993;34:1639– 1641.

- 50. Gone JR, Wallock NJ, Lindeman S, Donaldson WA. Tetrahedron Lett. 2009;50:1023–1025.
- 51. Yong W, Vandewalle M. Synlett. 1996:911–912.
- 52. Gree R, Laabassi M, Moset P, Carrie R. Tetrahedron Lett. 1985;26:2317– 2318.
- 53. (a) Cotton JD, Crisp GT, Daly VA. Inorg Chim Acta. 1981;47:165–169.
 (b) Cotton JD, Crisp GT, Latif L. Inorg Chim Acta. 1981;47:171–176.
- 54. (a) McDaniel KF, Kracker LR, II, Thamburaj PK. Tetrahedron Lett. 1990;31:2373–2376.
 (b) Donaldson WA, Shang L, Tao C, Yun YK, Ramaswamy M, Young VG., Jr J Organomet Chem. 1997;539:87–98.
 (c) Chaudhury S, Donaldson WA. J Am Chem Soc. 2006;128:5984– 5985.
 (d) Chaudhury S, Li S, Bennett DW, Siddiquee T, Haworth DT, Donaldson WA. Organometallics. 2007;26:5295–5303.
- 55. (a) Aumann R. J Am Chem Soc. 1974;96:2361–2362.
 - (b) Schulze MM, Gockel U. J Organomet Chem. 1996;525:155–158.
 (c) Taber DF, Kanai K, Jiang Q, Bui G. J Am Chem Soc. 2000;122:6807–6808.
 - (d) Taber DF, Bui G, Chen B. J Org Chem. 2001;66:3423–3426.
 - (e) Taber DF, Joshi PV, Kanai K. J Org Chem. 2004;69:2268–2271.
 - (f) Taber DF, Sheth RB. J Org Chem. 2008;73:8030-8032.
- 56. Christie SDR, Cummins J, Elsegood MRJ, Dawson G. Synlett. 2009:257–259.

About the Authors

William A. Donaldson : Department of Chemistry, Marquette University, P. O. Box 1881, Milwaukee, WI 53201-1881, USA.

Fax: +1-414-288-7066

Email: william.donaldson@marquette.edu

European Journal of Organic Chemistry, Vol. 2009, No. 23 (2009): pg. 3831-3843. <u>DOI</u>. This article is © Wiley-VCH Verlag and permission has been granted for this version to appear in <u>e-Publications@Marquette</u>. Wiley-VCH Verlag does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Wiley-VCH Verlag.