About the Cover:

Research endeavors are represented on the exploration of organofluorine chemistry at the multidisciplinary interface of biology and chemistry. The background is the tubulin-bound “REDOR–Taxol” molecule and a macrocyclic analog mimicking it at the β-tubulin binding site for Taxol. See Ojima, 6358. Professor Iwao Ojima is the recipient of the 2013 ACS Award for Creative Work in Fluorine Chemistry. View the article.
Exploration of Fluorine Chemistry at the Multidisciplinary Interface of Chemistry and Biology

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ABSTRACT: Over the last three decades, my engagement in “fluorine chemistry” has evolved substantially because of the multidisciplinary nature of the research programs. I began my research career as a synthetic chemist in organometallic chemistry and homogeneous catalysis directed toward organic synthesis. Then, I was brought into a very unique world of “fluorine chemistry” in the end of 1970s. I started exploring the interface of fluorine chemistry and transition metal homogeneous catalysis first, which was followed by amino acids, peptides, and peptidomimetics for medicinal chemistry. Since then, I have been exploring the interfaces of fluorine chemistry and multidisciplinary fields of research involving medicinal chemistry, chemical biology, cancer biology, and molecular imaging. This perspective intends to cover my fruitful endeavor in the exploration of fluorine chemistry at the multidisciplinary interface of chemistry and biology in a chronological order to show the evolution of my research interest and strategy.

INTRODUCTION

The extraordinary potential of fluorine-containing biologically relevant molecules in peptide/protein chemistry, medicinal chemistry, chemical biology, pharmacology, drug discovery as well as diagnostic and therapeutic applications was recognized by researchers who are not in the traditional fluorine chemistry field, and thus a new wave of fluorine chemistry has been rapidly expanding its biomedical frontiers. In fact, the importance of fluorine in bioorganic and medicinal chemistry has been demonstrated by a large number of fluorinated compounds approved by the FDA for medical use. According to our survey in 2008, 138 fluorine-containing drugs have received FDA approval for human diseases (of which 23, however, have been discontinued from the market), while 33 are currently in use for veterinary applications. These statistics make fluorine the “second-favorite heteroatom” after nitrogen in drug design.

Small atomic radius, high electronegativity, nuclear spin of 1/2, and low polarizability of the C−F bond are among the special properties that render fluorine so attractive. These atomic properties translate widely into equally appealing attributes of fluoroorganic compounds. Higher metabolic stability, often increased binding to target molecules, and increased lipophilicity and membrane permeability are some of the properties associated with the replacement of a C−H or C−O bond with a C−F bond in biologically active compounds. Because of the recognized value of fluorine, it is now a common practice in drug discovery to study fluoro-analogues of lead compounds under development. It should be noted that in 2006 the best and the second best selling drugs in the world were Lipitor (atorvastatin calcium) (by Pfizer/Astellas; $14.4 billion/year) and Advair (USA)/seretide(EU) (a mixture of fluticasone propionate and salmeterol) (by GlaxoSmithKline; $6.1 billion/year), which contain one and three fluorine atoms, respectively. These huge successes of fluorine-containing drugs keep stimulating research on fluorine in medicinal chemistry for drug discovery. As such, it is not an exaggeration to say that every new drug discovery and development today explores fluorine-containing drug candidates without exception.

Although medicinal chemists have been introducing fluorine into bioactive molecules on the basis of experience and intuition, it is only recently that experimental and computational studies have been conducted to better understand how the introduction of fluorine into small drug molecules results in higher binding affinities and selectivity. An understanding of how the replacement of H with F affects the electronic nature and conformation of small molecules is crucial for predicting the interaction of fluoroorganic molecules with proteins and enzymes. In addition, 19F NMR has found numerous applications to molecular imaging and promoted the development of molecular probes for imaging. The sensitivity of 19F NMR spectroscopy, along with large 19F−1H coupling constants and the virtual absence of 19F in living tissues, makes incorporation of fluorine into bioactive compounds a particularly powerful tool for the investigation of biological processes. Also, applications of 18F-PET (positron emission tomography), a powerful in vivo imaging technology in oncology, neurology, psychiatry, cardiology, and other medical specialties, have already become an essential part of medical care. In addition, 18F-PET has emerged as an important tool in...
drug development, especially for accurate measurements of pharmacokinetics and pharmacodynamics.

There is a strong demand for developing new and efficient synthetic methods as well as expanding the availability of versatile fluoro-containing synthetic building blocks and intermediates to promote medicinal chemistry, chemical biology, and molecular imaging research. The limited availability of fluorochemicals for bioorganic and medicinal chemistry as well as pharmaceutical and agrochemical applications is mainly due to the exceptional properties and hazardous nature of fluoro and fluorochemical sources. Also, in many cases, synthetic methods developed for ordinary organic molecules do not work well for fluorochemicals because of their unique reactivity. Therefore, the new and efficient synthetic methods applicable to organofluorine compounds, including $^{19}$F radiotracers, need to be continuously developed.

This Perspective was commissioned on the occasion of my receiving the 2013 ACS Award for Creative Work in Fluorine Chemistry. Accordingly, I would like to review the research strategy and programs in my laboratory in the last three decades in perspective, which would be useful to foresee the future directions in organofluorine chemistry at the multi-disciplinary interface of chemistry and biology.

1. EXPLORATION OF ORGANOFLUORINE CHEMISTRY BY MEANS OF TRANSITION-METAL CATALYSIS

My first encounter with “fluorine chemistry” was in the late 1970s, when I was a Group Leader for organometallic chemistry and homogeneous catalysis as well as organic synthesis at the Sagami Institute in Japan. At that time, the research council of the institute decided to add “fluorine chemistry” to one of its strategic research areas. To me, one of the most inspirational reports back then was the bold experiments done by Leland Clark. In this paper, Clark demonstrated the capability of perfluorocarbons (PFCs) to deliver oxygen to a living mouse placed deep in a beaker filled with PFCs, which clearly implied the extraordinary nature of the world of “fluorine”. Of course, fluoropolymers, especially “Teflons”, fluoroethenones, coolants/refrigerants, extinguishers, aerosols, etc., were the representative fluorochemicals commonly used in daily life. As for pharmaceutical drugs, only 5-FU was widely recognized followed by 9α-fluorohydrocortisone at that time.

From a strategic point of view based on synthetic organic chemistry, I reviewed and critically analyzed potential new methods that could introduce these fluoro groups efficiently and selectively to the desired molecules from readily available starting materials. Thus, we studied the hydroformylation of a variety of fluoro-olefins (eq 1) as one of our approaches to the functionalization of these building blocks by means of transition-metal catalysts. Then, we found unusually high regioselectivities and a remarkable dependency of the regioselectivities of the reaction on the catalyst metal species, which was unique in comparison with the hydroformylation of ordinary alkenes.

1.1. Remarkable Dependence of Regioselectivity on the Catalyst Metal Species. The hydroformylation of TFP was carried out with Co$_2$(CO)$_3$, Ru$_2$(CO)$_12$, Rh$_2$(CO)$_16$, and PtCl$_2$ (DIOP)/SnCl$_2$, which are typical hydroformylation catalysts, at 100 °C and 100 atm (CO/H$_2$ = 1) for the Co, Pt, and Ru catalysts and at 80 °C and 110 atm (CO/H$_2$ = 1) for the Rh catalyst (Scheme 1). The reaction of TFP catalyzed by Co$_2$(CO)$_3$ gave (trifluoromethyl)propanals (TFMPAs) in 95% yield, in which a “normal” (or linear) aldehyde, highly regioselective hydroformylation to the synthesis of fluoro-amino acids, (ii) a novel ureidocarbonylation process that gives S-((trifluoromethyl)dihydouracils in one step, and (iii) the hydroformylation-amidocarbonylation of fluoroolefins catalyzed by Co–Rh mixed-metal systems.

Although this program was initiated at the Sagami Institute in Japan, I moved to the State University of New York at Stony Brook in 1983 and continued the exploration and expansion of the scope of the research program.

![Scheme 1. Hydroformylation of TFP Catalyzed by Co, Pt, Ru, and Rh complexes](image-url)
CF$_3$CH$_2$CH$_2$CHO (3-TFMPA), was formed with high regioselectivity (93%). In sharp contrast with Co$_2$(CO)$_8$ Rh-carbonyl cluster Rh$_4$(CO)$_{16}$ exhibited extremely high catalytic activity and regioselectivity (96%) to give “iso” (or branched) aldehyde, CF$_3$CH$_2$CH$_2$CHO (2-TFMPA). The Pt catalyst, PtCl$_2$(DIOP)/SnCl$_2$, favored the formation of normal aldehyde ($n$/iso = 71/29), while Ru$_4$(CO)$_{12}$ gave isoaldehyde as the main product ($n$/iso = 15/85). In both cases, a substantial amount of hydrogenated product, CF$_3$CH$_2$CH$_3$, was formed (25–38%). Addition of PPh$_3$ to the Co, Ru, and Rh catalysts considerably decreased the catalytic activities but somewhat increased the isoaldehyde selectivity. The result made a sharp contrast to the cases of ordinary olefins, where the addition of PPh$_3$ increased normal aldehyde selectivity.

Since Rh$_4$(CO)$_{16}$ gave excellent regioselectivity for the formation of 2-TFMPA, several other Rh catalysts were employed and their catalytic activities as well as regioselectivities examined. The results clearly indicated that the Rh(I) complexes having chlorine as a ligand, such as RhCl($	ext{PPh}_3$)$_3$, were less active than HRh(CO)($	ext{PPh}_3$)$_3$. Rh–C, Rh$_4$(CO)$_{12}$, and Rh$_6$(CO)$_{16}$, but the regioselectivity was virtually the same in all cases examined. Consequently, it was concluded that the nature of the central metal of the catalyst played a key role in determining the regioselectivity of the reaction. It was noteworthy that the metal species dependency of the regioselectivity in this reaction was remarkable compared to that reported for propene.22

The reaction of PFS was carried out in a similar manner at 90 °C and 82 atm, using Co, Pt, Ru, and Rh catalysts.29–31 Rhodium catalysts exhibited high catalytic activity to give isoaldehyde, C$_5$F$_7$C($	ext{Me}$)$_2$CHO (3-PFPPA), with excellent regioselectivity (97–98%) in quantitative yields, while Co$_2$(CO)$_8$ gave normal aldehyde (3-PFPPA) as the major product, with regioselectivity (79–90%) not as high as that observed in the reaction of TFP. The Ru catalyst, Ru$_4$(CO)$_{12}$, showed rather low catalytic activity (49% conversion), giving isoaldehyde as the major isomer (22% yield, b/n = 74/26), accompanied by a substantial amount of hydrogenated product, C$_5$F$_7$CH$_2$CH$_3$ (25%). The Pt catalyst, PtCl$_2$(DIOP)/SnCl$_2$, showed a high catalytic activity (100% conversion in 4 h, 76% aldehyde yield), but virtually no regioselectivity was observed and the hydrogenation of PFS took place as a severe side reaction (20%). Thus, the dependency of regioselectivity on the metal species was similar to that for TFP, and the observed regioselectivity was also remarkably high compared with that reported for styrene.32–34

A kinetic study was performed for the Rh$_4$(CO)$_{12}$- and Co$_2$(CO)$_8$-catalyzed reactions of PFS.30 At 100 °C and 82 atm (CO/H$_2$ = 1) with $1.0 \times 10^{-5}$ M catalyst concentration, the Rh-catalyzed reaction was first order in PFS concentration, and the apparent rate constant was calculated to be $6.2 \times 10^4$ s$^{-1}$; i.e., the turnover number was estimated to be 55 800 h$^{-1}$ per Rh metal. The Co-catalyzed reaction with $1.0 \times 10^{-2}$ M catalyst concentration at 100 °C and 82 atm (CO/H$_2$ = 1) was also first order in PFS concentration, and the apparent rate constant was calculated to be $1.6 \times 10^3$ s$^{-1}$; i.e., the turnover number per cobalt metal was 2.88 h$^{-1}$. Thus, the Rh catalyst was ca. 20000 times more active than the Co catalyst per metal provided that all metal species participate in the catalysis.

On the basis of the fact that the addition or the introduction of tertiary phosphines to the catalyst caused only a slight change in regioselectivity, in sharp contrast to the hydroformylation of propene or styrene using the same catalysts,
stabilization of the formal negative charge developing on Cα. Thus, the formation of isoaoyl-[M] species (IIiso) should be much more favorable than that of the n-acyl-[M] species (IIs) regardless of the group VIII transition-metal species. In fact, the results of the reactions of vinyl fluoride (VF) provide strong supporting evidence for this hypothesis.

The iso/n ratio of aldehydes should reflect the ratio of the intermediate iso- and n-acyl-[M] species (IIiso and IIs) (Scheme 2) under sufficient pressure of hydrogen. Thus, it is deduced that in the Rh-catalyzed reaction, $k_{\text{iso}} \gg k_{\text{i}}$ and $k_{\text{CO}} \gg k_{\text{n}}$, and thus the initially formed isoaoyl-[Rh] species (IIiso, M = Rh) generates the isoacyl-[Rh] species (IIIiso, M = Rh) and gives the isoaalkyldehyde with high regioselectivity. In sharp contrast, the rate constants in the Co-catalyzed reaction are $k_{\text{iso}} \gg k_{\text{i}}$, $k_{\text{n}}$, and $k_{\text{CO}} \gg k_{\text{n}}$, and thus the initially formed isoaoyl-[Rh] species (IIiso, M = Rh) generates the isoacyl-[Rh] species (IIIiso, M = Rh) and gives the isoaalkyldiene with high regioselectivity. The Rh- and Co-catalyzed reactions are extremely selective cases, and the Pt- and Ru-catalyzed reactions are in between the two extreme cases.

In addition to these kinetic aspects, we should take into account the fundamental difference between each isoaoyl-[M] intermediate, i.e., the size of the metal and the polarizability of the metal–carbon bond. Thus, the relative stability of IIiso can be estimated to increase in the order Rf(Me)CH-CoLM < Rf(Me)CH-PtLM < Rf(Me)CH-RuLM < Rf(Me)CH-RhLM.30

It is worth noting that these mechanistic details were revealed because of the use of fluoro-olefins as unique substrates for the hydroformylation reaction. In this case, organometallic chemistry and catalysis research greatly benefited from fluorne compounds. In turn, fluorne chemistry also benefited from the discovery of highly regioselective hydroformylation processes, which provided versatile fluorine-containing aldehydes. In fact, immediately after the discovery of highly regioselective hydroformylation of TFP by Rh-catalyzed process, I envisioned that we should be able to produce a series of “CF3-chemicals” from TFMA.

1.2. Hydroesterification and Hydrocarboxylation of Fluoro-olefins. Hydrocarboxylations of olefins serves as a convenient method for the synthesis of the corresponding esters or carboxylic acids.32,36 Despite extensive mechanistic studies as well as applications of the reactions to organic syntheses, little attention had been paid to the reactions of fluoro-olefins before we started the investigation on this subject.

The screening of typical transition-metal complexes in the hydrocarboxylations of TFP and PFS revealed that only Pd complexes with phosphine ligands showed sufficient catalytic activity to promote the reaction under the given reaction conditions.37 As Scheme 3 shows, the Pd-complex-catalyzed hydroesterification of TFP and PFS gave branched esters, Rf(Me)CHCOOR, in good to excellent regioselectivity, while the corresponding hydrocarboxylation afforded linear acids, RfCH2CH2COOH, in excellent yield and regioselectivity. Plausible mechanisms were proposed to accommodate the observed marked difference in regioselectivity for these two reactions.

2. SYNTHESIS OF FLUORO ANALOGUES OF ALIPHATIC AND AROMATIC α-AMINO ACIDS

By the mid 1980s, it was shown that fluorinated analogues of naturally occurring biologically active compounds often exhibited unique physiological activities.25,28,38 For example, fluorinated pyrimidines acted as anticancer/antiviral agents, and some fluorinated aromatics as well as CF3-aromatic compounds were used as nonsteroidal anti-inflammatory drugs, antifungal agents, human antiparasitic agents, central nervous system agents for psycho-pharmacology, diuretics agents, and anthypertensive agents. Some fluoro-amino acids acted as “suicide substrate enzyme inactivators”, showing strong antibacterial activities and some of them also acted as antihypertensive agents.28 In the mid to late 1980s, there was an increasing interest in the incorporation of fluoro-amino acids into peptides.39–43 Accordingly, it was timely and important to develop new and efficient methods for the synthesis of fluoroo-amino acids at that time (and even now in the 2010s!).

We found that the fluoro-aldehydes obtained in the hydroformylation of TFP and PFS, described above, served as excellent intermediates for the synthesis of fluoro-amino acids. Thus, we developed efficient synthetic routes to 4,4,4-trifluorovaline (TFV), 5,5,5-trifluoronvaline (TFNV), 5,5,5-trifluoroleucine (TFL), 6,6,6-trifluoronleucine (TFNL), 4,5,5,6-tetrafluorotryptophan (tryptophan-f4), and related compounds from the fluoro-aldehydes by means of transition-metal-catalyzed transformations as well as enzymatic reactions.

2.1. 4,4,4-Trifluorovaline (TFV) and 5,5,5-Trifluoronvaline (TFNV). TFV and TFNV were synthesized using Co-catalyzed amidoformylation of 2-TFMPA and 3-TFMPA, respectively. The amidoformylation of 2-TFMPA and 3-TFMPA with acetyamide catalyzed by Co2(CO)8 (CO/H2 (1/1) 100 atm, 120 °C) gave Ac-TFV and N-Ac-TFNV, respectively, in good yields, which were further hydrolyzed to the corresponding free amino acids (eqs 4 and 5).44

We also successfully carried out the kinetic optical resolution of Ac-TFV, using porcine kidney acylase I (25 °C, pH 7.0) to give (S)-TFV and (R)-Ac-TFV with high enantiopurities, which were readily separated (eq 6).44 (R)-Ac-TFV was further hydrolyzed to (R)-TFV with 3 M HCl. The enantiopurities of (S)-TFV and (R)-TFV were determined by Mosher’s MTPA method (H and 19F NMR).
It has been shown that TFNV inhibits the growth of *E. coli* and may be used as a growth regulatory factor in microbiology.\(^4^6\) TFV serves as a modifier of biologically active peptides in protein engineering and chemical biology. In fact, both TFNV and TFV are commercially available from more than several vendors now, and indeed, (2S)-TFV has been extensively used in protein design and chemical biology, as we envisioned in late 1980s.\(^4^7\)

### 2.2. 5,5,5-Trifluoroleucine (TFL) and 6,6,6-Trifluoronorleucine (TFNL)

TFL and TFNL were synthesized via azlactones prepared from 2-TFMPA and 3-TFMPA, respectively (Scheme 4, (a) and (b)).\(^4^4\) The azlactones were subjected to alcoholysis to give the corresponding dehydroamino acid esters 3 and 4. (Z)-Dehydroamino acid 3c was obtained by hydrolysis of 1-Z. Then, the (Z)-dehydroamino acid and esters (3-Z and 4-Z) were hydrogenated over Pd/C followed by hydrolysis to give the corresponding amino acids TFL and TFNL. The azlactones (1-Z and 2-Z) were also treated with hydriodic acid/red phosphorus to give TFL and TFNL directly.

It is worthy of note that the chiral 2,2,2-trifluoroisopropyl group acted as an effective stereogenic center in the hydrogenation of 3-Z over Pd/C, yielding Bz-TFL-OR (Scheme 4, (a)).\(^4^4\) Thus, the hydrogenation of 3a-Z, 3b-Z, and 3c-Z in THF at ambient temperature and pressure of H\(_2\) gave Bz-TFL-Ome (85:15 dr), Bz-TFL-OEt (90:10 dr), and Bz-TFL-Oh (82:18 dr), respectively, in quantitative yield. It was rather surprising that the "chiral isopropyl group" was able to induce a high degree of stereoselectivity. It was strongly suggested that the trifluoromethyl group was not only a bulkier substituent than methyl but also imposed a unique electronic effect on the Pd—metal surface.

Optically active (S)-N-Bz-TFNL-Ome (87—89% ee) was obtained quantitatively by asymmetric hydrogenation of 4a-Z (R = Me) using a cationic Rh catalyst with diPAMP\(^4^8\) at 40 °C and 1 atm of H\(_2\) in ethanol.\(^4^4\) (S)- and (R)-TFNL with excellent enantiopurities (>98% ee) were obtained through enantiomeric resolution of racemic Ac-TFNL using porcine kidney acylase I in a manner similar to that for Ac-TFNV (eq 7).\(^4^4\)

We envisioned that TFL and TFNL would serve as unique modifiers of biologically active peptides in protein engineering and chemical biology. As we anticipated in the late 1980s, both TFL and TFNL are commercially available from more than several vendors now, and indeed, (2S)-TFL has been extensively used in protein design and chemical biology.\(^4^7\)

### 2.3. 4,5,6,7-Tetrafluorotryptophan and Related Compounds

As the importance of biologically active compounds bearing the indole skeleton such as tryptophan, tryptamine, indoleacetic acid, and alkaloids was well recognized, tetrafluoro analogues of indoles were synthesized from 2-PFPPA.\(^4^4\) The reaction of 2-PFPPA with allylamine followed by cyclization using lithium diisopropylamide (LDA) as a base and protection of the indole nitrogen gave 3-Me-indole-f4 in 72% from 2-PFPPA (eq 8).

The SeO\(_2\) oxidation of N-Ac-3-Me-indole-f4 \(5\) gave indole-f4 3-CHO in 86% yield (eq 9). The SeO\(_2\) oxidation of \(5\) in the
presence of acetic anhydride gave 1-Ac-3-(AcO-methyl)indole-f₄ 6 in 60% yield (eq 9). 3-(AcO-methyl)indole-f₄ 6 and indole-f₄-3-CHO were very useful intermediates for the synthesis of tetrafluoro analogues of tryptophan, tryptamine, and indoleacetic acid.44

Thus, tryptamine-f₄ was synthesized from indole-f₄-3-CHO in 83% overall yield through condensation with nitromethane, followed by LiAlH₄ reduction, while tryptophan-f₄ was obtained in four steps in 51% overall yield through Erlenmeyer’s azlactone method (eq 10). The reaction of 6 with piperidine gave 3-(piperidinomethyl)indole-f₄ in 97% yield, which was a known key intermediate for tryptophan-f₄ in 2 steps (eq 11). Since it was shown that tryptophan-f₄ strongly inhibits both the tryptophanyl hydroxamate and aminoacyl t-RNA formation,50,51 the tetrafluoro analogues of tryptamine, indoleacetic acid, and other indole derivatives were expected to possess unique physiological activities. Tryptophan-f₄ has been used in enzymology, protein engineering and chemical biology.52,53

3. CARBONYLATIONS OF α-(TRIFLUOROMETHYL)VINYL BROMIDE

3.1. Synthesis of 2-Trifluoromethylacrylic Acid through Pd-Catalyzed Carboxylation. The bromination of TFP by photoirradiation followed by dehydrobromination on KOH gave 2-Br-TFP in high yield.44 The carboxylation of 2-Br-TFP catalyzed by a Pd catalyst, e.g., PdCl₂(PPh₃)₂ or PdCl₂(dppf), in the presence of Et₃N in DMF or THF afforded 2-(trifluoromethyl)acrylic acid (2-TFMAA) in 65−78% yield (eq 12).55

A variety of trifluoromethacrylates, CH₂=C(CF₃)COOR, were readily prepared from 2-TFMAA, which were potentially very useful monomers for fluorine-containing polymethacrylates (Figure 1). Copolymerization with other olefins, e.g., methyl methacrylate (MMA) and styrenes, were also possible. In fact, the homo- and copolymerization of methyl trifluoromethacrylate (MTFMA) were reported56 with regard to the development of new radiation-sensitive polymers for resists in microelectronic fabrication processes, wherein MTFMA was prepared from trifluoroaceton. Copolymers of TFMA esters with styrene and substituted styrenes were also prepared.57 We synthesized a variety of new trifluoromethacrylates bearing polyfluoroalkyl ester moieties,58,59 which would serve as monomers for potential photoresists and as a component of optical fibers. Although we did not follow up this line of research in my laboratory, TFMA esters have been extensively studied and are still under active investigation for the development of photoresists for lithography using short wavelength light, e.g., 157/193 nm,60−62 as we envisioned in the 1980s.24,58

3.2. α-Trifluoromethyl-β-alanine (α-TFM-β-Ala). We found that the addition of gaseous ammonia to 2-TFMAA at 0−5 °C in CH₃Cl₂ gave a novel β-amino acid, α-trifluoromethyl-β-alanine (α-TFM-β-Ala), in excellent yield.44 However, the reaction sometimes gave double and triple Michael addition products, depending on the reaction conditions. The use of hexamethyldisilazane (HMDS) in an attempt to protect the C-terminus of TFMAA with a TMS group resulted in an addition of H₂NTMS, generated in situ, to O-TMS-TFMAA, giving N,O-bis-TMS-α-TFM-β-Ala (7) in quantitative yield. No trace of N,N,O-tris-TMS-α-TFM-β-Ala was detected. Also, Michael addition of HMDS to the methyl and benzyl esters of TFMAA did not proceed at all. The disilylated α-TFM-β-Ala (7), thus obtained, was treated with MeOH to give α-TFM-β-Ala in nearly quantitative yield (eq 13).44 α-TFM-β-Ala did not exhibit any antibacterial activity in our preliminary screening. However, an enkephalin analogue bearing α-TFM-β-Ala, Tyr-β-Ala-(α-TFM-β-Ala)-Phe-Met, has shown fairly strong analgesic effects.53,64 This suggested that the novel fluoro-β-amino acid would serve as a modifier for a variety of peptide hormones and other physiologically active peptides, although a method to obtain enantiopure material was needed. It should be noted that α-TFM-β-Ala is now commercially available from more than several vendors.

3.3. “Ureidocarboxylation” of 2-Bromotrifluoropropene (2-Br-TFP) Catalyzed by a Pd−Phosphine Complex. The Pd-complex-catalyzed amidation of vinyl halides was shown to be a convenient method for the synthesis of αβ-unaturated amides.65 However, nothing was known for the Pd-catalyzed reaction of vinyl halides with ureas instead of amines. Accordingly, we investigated the Pd-catalyzed carboxylation reaction of 2-Br-TFP with a urea. Our hypothesis was that both nitrogen termini of a urea would possess sufficient nucleophilicity so that the reaction should give the dihydroxuracil skeleton.
in one step. Actually, the reaction proceeded as anticipated to give 5-CF₃-5,6-dihydrouracil 8 in good yield, and this novel carbonylation process was termed “ureidocarbonylation”. A general scheme for this novel process is shown in eq 14.⁶⁶ However, when unsubstituted urea was employed, the yield of 8d (R₁ = R₂ = H) was low. 5-CF₃-dihydrouracils 8, thus obtained, were readily converted to the corresponding 5-CF₃-uracils 16 by treating with bromine⁶⁷ in nearly quantitative yields (eq 9).⁶⁶ It should be noted that 8 exhibited substantial antitumor activity against ascitic mastocarcinoma MM2 cells.⁵⁵ A proposed mechanism for the “ureidocarbonylation” is illustrated in Scheme 5.⁶⁶

We also found a simple method for the synthesis of 8 as well as its thio analogues just by heating a mixture of 2-TFMAA and a urea or thiourea in the presence of acetic anhydride at 80−100 °C, which afforded the corresponding 5-CF₃-uracils 16 by treating with bromine⁶⁷ in nearly quantitative yields (eq 9).⁶⁶ It should be noted that 8 exhibited substantial antitumor activity against ascitic mastocarcinoma MM2 cells.⁵⁵ A proposed mechanism for the “ureidocarbonylation” is illustrated in Scheme 5.⁶⁶

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It is worthy of note that the processes for producing 2-TFMAA from 2-Br-TFP as well as 9d from 2-TFMAA and urea followed by dehydrogenation were developed as commercial processes by Japan Halon (now Tosoh F-Tech). Furthermore, 9d was successfully applied to the commercial synthesis of trifluridine (trifluorothymidine), an antiviral drug, primarily used on the eye topically, such as “Viroptic”. This commercial process was developed by Japan Halon and Tokyo Yuki Gosei Kogyo in Japan in the early 1990s and is still operating at present.

4. HYDROFORMYLATION−AMIDOCARBONYLATION OF FLUORO-OLEFINS: HIGHLY REGIOSELECTIVE DIRECT SYNTHESIS OF FLUOROAMINO ACIDS FROM FLUORO-OLEFIN

4.1. Hydroformylation−Amidocarbonylation of Tri-fluoropropene (TFP). The hydroformylation−amidocarbonylation (HF-AC) of TFP was investigated since the reaction should give the corresponding normal or iso N-acetylamino acid, Ac-TFNV or Ac-TFV, directly from TFP if the extremely regioselective hydroformylation was successfully combined with amidocarbonylation. As Scheme 6 shows, the Co-catalyzed reaction gave Ac-TFNV with 96% selectivity, while the reaction catalyzed by the Rh−Co binary system (Co₂(CO)₈/Rh₆(CO)₁₆ = 50) under the same conditions gave Ac-TFV with 94% selectivity.⁷⁰ The latter result clearly indicated that the Rh-catalyzed hydroformylation took place exclusively in the first step to give 2-TFMPA with high selectivity, which was effectively incorporated into the subsequent Co-catalyzed amidocarbonylation.

4.2. Hydroformylation−Amidocarbonylation of Pentfluorostyrene (PFS). In contrast to the results obtained for the reactions of TFP, the attempted regioselective HF-AC of PFS catalyzed by the Co−Rh binary catalyst system as well as Co₂(CO)₈ under similar conditions gave unexpected results. The detailed study of the reaction revealed interesting mechanistic aspects of Co−Rh mixed-metal catalyst systems, including a novel CoRh(CO)₇-catalyzed process.⁷⁰ The HF-AC of PFS with acetamide catalyzed by Co₂(CO)₈ gave N-Ac-4-C₆F₅-homoalanine (10) with 90−92% regioselectivity (Scheme 7). This regioselectivity was much higher than that (79%) of the simple hydroformylation in benzene.³⁰ The reaction catalyzed by Co₂(CO)₈/Rh₆(CO)₁₆ gave N-acetyl-3-C₆F₅-homoalanine (11) with only ca. 80% regioselectivity, which was much lower than the excellent regioselectivity (98%).
of the simple hydroformylation in benzene. To accommodate these unexpected results, a detailed mechanistic study was performed to clarify these anomalies.

It was found that the hydroformylation was the rate- and regioselectivity-determining step and the presence of acetamide substantially increased the normal selectivity, probably by forming an active species HCo(CO)₅(CH₃CONH₂). This catalyst species, however, substantially increased the formation of hydrogenation product, C₆F₅Et. If the Co and Rh catalysts worked independently, the ratio of normal aldehyde formation should increase at higher Co/Rh ratios and eventually the normal aldehyde should become the major product. However, contrary to this assumption, an interesting leveling phenomenon was observed, i.e., the iso/n ratio decreases from 94/6 at Co/Rh = 5 to 88/12 at Co/Rh = 25, but the ratio was unchanged (87/13) even at a Co/Rh ratio of 100! This leveling phenomenon was best interpreted by taking into account the formation of and the catalysis by a Co–Rh mixed-metal complex. When we reached this conclusion, Horváth, Bor, and Pino at ETH reported synthesis, characterization, and some reactions of an interesting coordinatively unsaturated Co–Rh mixed metal complex, CoRh(CO)₇ (eq 16), which was eventually identified as the catalyst species in our HF-ADS reaction.

\[
\text{Rh}_4(\text{CO})_{12} + 2\text{Co}_2(\text{CO})_8 \overset{K}{\rightleftharpoons} 4\text{CoRh}(\text{CO})_7
\]  

In order to directly confirm the catalyst species in dioxane, we performed a high pressure FT-IR study on the Co–Rh mixed metal complex system, which provided strong supporting evidence for the CoRh(CO)₇ catalysis.

Next, the relative activities of catalyst species were evaluated on the basis of the equilibrium constant \(K\) reported for eq 14. The iso/n ratio vs Co/Rh ratios were calculated and plotted with several given relative catalytic activities. In dioxane, the relative catalytic activity of 54/mol (9/Rh) for \(\text{Rh}_6(\text{CO})_{16}/\text{Co}_2(\text{CO})_8\) showed a very good agreement with the experimental results. The relative catalytic activity for \(\text{Rh}_6(\text{CO})_{16}/\text{Co}_2(\text{CO})_8\) was calculated to be ca. 400000, i.e., 133000 per metal. Finally, kinetic studies were performed to compare the results with those predicted by calculations based on the regioselectivity. The kinetic measurements in dioxane provided the relative activity values as follows: \(\text{Rh}_6(\text{CO})_{16}/\text{Co}_2(\text{CO})_8\) = 3.8/Rh, \(\text{Rh}_6(\text{CO})_{16}/\text{Co}_2(\text{CO})_8\) = 398000. Consequently, the results of these two independent evaluation methods were in very good agreement in spite of various assumptions and simplification for calculations.

Overall, this study provided a rare and successful example of the elucidation of mixed-metal catalysis, in which actual active catalyst species and their direct precursors were detected spectroscopically and the observation corresponded almost perfectly to the mechanism proposed based on the regioselectivity analysis. It should be emphasized that the discovery of CoRh(CO)₇ catalysis as well as the successful mechanistic studies in organometallic chemistry and catalysis were only possible by the use of unique fluoro-olefin, PFS.

From the synthetic viewpoint, a highly regioselective formation of fluoro-amino acid 12, directly from PFS is noteworthy. The reaction of PFS using a Co₂(CO)₈ (5.0 mol %)–Rh₄(CO)₁₂ (0.05 mol %) catalyst system gave 12 in 80% yield and 98.2% regioselectivity at 60 °C for 6 h and then 125 °C for 5 h under 75 atm of CO and 48 atm of H₂. Base-promoted cyclization of 12 gave N-Ac-2-hydroxycarbonyl-3-methyl-2,3-dihydro-4,5,6,7-tetrafluorindole (13) in 92% yield, which could be converted to a variety of fluoro-indoles and fluoro-alkaloids (Scheme 8).

Scheme 8. Tetrafluorindole Synthesis from PFS via Highly Regioselective Hydroformylation–Amidocarbonylation

5. APPLICATIONS OF TRIFLUOROMETHYL-CONTAINING AMINO ACIDS TO ENZYME INHIBITORS

5.1. Trifluoromethyl Analogues of Captopril as Inhibitors of Angiotensin-Converting Enzyme. It has been shown that inhibitors of angiotensin converting enzyme (ACE) play key roles in the control of blood pressure as therapeutic agents. Since the development of potent ACE inhibitors captopril74 and enalaprilat,75 various analogues of these drugs have been designed and synthesized, and their ACE inhibitory activity has been examined. However, little attention had been paid to the synthesis and activity of fluorinated analogues until we started our investigation in the late 1980s.76 Thus, we designed and synthesized CF₃ analogues of captopril and evaluated their ACE inhibitory activity to examine the effect of fluorine incorporation on the potency.

Chiral Michael acceptor 2-CF₃-acyrloyl-(S)-Pro-OBu-f (14) was prepared through the facile coupling of (S)-Pro-OBu-f and \(\alpha\)-CF₃-acyrloyl chloride,55 derived from 2-TMFAA in 85% yield (Scheme 9).78 Conjugate addition of thiolacetic acid to 14 gave a diastereomeric mixture of adducts, (R,S)-15 and (S,S)-15 (RS/SS = 1/2) in 70% yield, which were separated by MPLC (Scheme 9).78,79 The stereochemical assignments were performed to clarify these anomalies, a detailed mechanistic study was performed to clarify these anomalies.
unambiguously made on the basis of the X-ray crystal structure of (S,S)-15. Syntheses of (R,S)-captopril-3 and (S,S)-captopril-3 are illustrated in Scheme 9.78,79

The CF3-analogues of captopril were subjected to in vitro enzyme inhibitory assay against ACE based on the method of Holmquist et al. using the tripeptide, [3-((2-furyl)acryloyl)]-Phe-Gly-Gly, as the substrate.80 The results are listed in Table 1.78,79 (S,S)-Captopril was also used as the reference in this assay.

Table 1. ACE Inhibitory Activity of CF3-Analogues of Captopril

<table>
<thead>
<tr>
<th>ACE inhibitor</th>
<th>IC50 (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(R,S)-captopril-3</td>
<td>2.9 × 10⁻¹⁰</td>
</tr>
<tr>
<td>(S,S)-captopril-3</td>
<td>4.8 × 10⁻⁷</td>
</tr>
<tr>
<td>(S,S)-captopril</td>
<td>3.6 × 10⁻⁹</td>
</tr>
</tbody>
</table>

As Table 1 shows, (R,S)-captopril-3 was found to be extremely potent (IC50, 10⁻¹⁰ M level), while the corresponding diastereomer, (S,S)-captopril-3, was much less potent by 3 orders of magnitude. A similar difference in potency was reported for (S,S) and (R,S)-captopril, i.e., the (S,S)-isomer was more potent than the (R,S)-isomer by a factor of 100.74 It is worthy of note that (R,S)-captopril-3 is at least 1 order of magnitude more potent than (S,S)-captopril.

The considerably higher potency of (R,S)-captopril-3 than (S,S)-captopril may be ascribed to the hydrophobicity and the stereoelectronic effects of the CF3 group. Namely, the incorporation of the (2R)-CF3 group may significantly contribute to an increase in attractive interaction with the hydrophobic binding site of ACE. Also, the stereospecific incorporation of (2R)-CF3 may cause strong restriction of rotation around the amide bond because of the stereoelectronic effect of the CF3 group. This would fix the inhibitor in the favorable conformation such that strong binding with the active site is achieved without sacrificing energy for conformational change.

In fact, the energy minima for hypothetical binding conformations based on molecular mechanics calculations of (S,S)-captopril and (R,S)-captopril-3 indicated the latter compound to be more favorable by 1.3 kcal/mol, which corresponds to 10-fold difference in activity.78,79 Also, the 8.33 kcal/mol difference between (R,S)-captopril-3 and (S,S)-captopril-3 calculated corresponds to the 1,700 times difference in the activity. In addition, a semiempirical approach was employed to further examine the inhibitor-enzyme interaction by using the n-SCF-molecular mechanics program (PIMM).81 This calculation indicated that the binding energy of (R,S)-captopril-3 was 2.4 kcal/mol more favorable than (S,S)-captopril.

5.2. Potent Enkephalin Analogues with Trifluoromethyl-Containing Amino Acid Residues. Enkephalins, “opioid peptides” in the brain, are known to play important roles as analgesics, regulators of blood pressure, and neurotransmitters. In the hope of developing nontoxic, nonaddictive, and effective analgesics, replacing morphine, structural modifications of enkephalins have been extensively studied, and a variety of analogues have been developed.82 Neurobiological studies have revealed the presence of the major opiate receptors μ, δ, and κ that mediate the analgesic effects of opiates and opioid peptides including enkephalins.83 Thus, the discovery and development of opiate receptor specific ligands has been of active interest in medicinal chemistry as well as neurobiology. However, in spite of extensive studies on enkephalin analogues, little attention had been paid to the fluoro-analogues of enkephalins84 when we started our investigation in the early 1990s.

The major nemeses of these opioid peptides is a group of degrading enzymes that cleave the peptide into inactive fragments.85,86 These enzymes include (i) aminopeptidase M and membrane-bound aminopeptidase(s), cleaving enkephalins at the Tyr1-Gly2 bond, which is responsible for 80% of the degradation pathway,85,87 (ii) enkephalinase, cleaving the Gly3-Phe4 bond,86 and (iii) dipetidylaminopeptidase, cleaving the Gly2-Gly3 bond.85 Accordingly, the development of inhibitors for these degrading enzymes is an important approach to the enhancement of analgesic activity by increasing the duration of the effect.

In the course of our study on the synthetic and medicinal chemistry of CF3-containing amino acids, enzyme inhibitors and peptide hormones by exploring unique effects of the CF3 group,24,35,44,79 we designed and synthesized a series of novel enkephalin analogues bearing TFNV and TFNL. Then, their in vivo analgesic activity as well as in vitro receptor binding ability was examined.88

Synthesis of Novel Fluoro-enkephalin Analogues. A series of fluoro-enkephalin analogues was synthesized through solid-phase peptide synthesis by replacing either Gly2 or Gly3 by (R)-TFNV, (S)-TFNL, or (R)-TFNL, as shown in Scheme 10.88 An analogue bearing (N-Me)Phe in place of Phe was also synthesized.

Scheme 10. Schematic Illustration of Modification Strategy of Methionine—Enkephalin with Trifluoronorvaline (TFNV) and Trifluoronorleucine (TFNL)

Analytical Activity Assay in Vivo.88 The in vivo analgesic activity of fluoro-enkephalin analogues was evaluated by the standard writhing test using ddY male mice via intracerebroventricular (i.c.v.) administration. As Table 2 shows, the substitution of Gly2 by (R)-TFNV exhibited a remarkable increase in potency, i.e., 100000 times stronger than methionine—enkephalin, and even 1 order of magnitude stronger than morphine (entry 8). The absolute configuration of TFNV at this position is very important. Thus, ([S]-TFNV2, Met3-NH2)enkephalin showed only ca. 30 times increase in potency (entry 7). The substitution of Gly3 with TFNL improved potency by the factor of 30–60, in which the (R)-TFNL analogue was twice as effective as the (S)-TFNL analogue (entries 10, 11). The modification of carboxyl terminus to amide showed 5–6 times improvement in potency.
As Table 2 shows, 

\[
\text{entry} \quad \text{compound} \quad \text{ED}_{50} (10^{-6} \text{ mol/mouse})
\]

1 methionine-enkephalin 700
2 morphine-HCl 0.07
3 Tyr-(R)-Ala-Gly-Phe-Met-NH₂ 0.05
4 sedapain (morphine analogue) 0.05
5 Tyr-(S)-TFNV-Gly-Phe-Met 120
6 Tyr-Gly-(S)-TFNV-Phe-Met 140
7 Tyr-(S)-TFNV-Gly-Phe-Met-NH₂ 25
8 Tyr-(R)-TFNV-Gly-Phe-Met-NH₂ 0.007
9 Tyr-(R)-Nval-Gly-Phe-Met-NH₂ 0.04
10 Tyr-Gly-(S)-TFNV-Phe-Met-NH₂ 22
11 Tyr-Gly-(R)-TFNV-Phe-Met-NH₂ 12
12 Tyr-(R)-TFNL-Gly-Phe-Met-NH₂ 0.07
13 Tyr-(R)-TFNV-Gly-(N-Me)Phe-Met-NH₂ 0.002

(entries 5, 6, 7, and 10). The substitution of Gly² with (R)-TFNL, a homologue of (R)-TFNV, exhibited 10000 times increase in potency, but it was 1 order of magnitude lower than the corresponding (R)-TFNV analogue (entry 12). The observed remarkable increase in potency for [(R)-TFNV², Met³-NH₂]enkephalin was not entirely exceptional since a known analogue, [(R)-Ala², Met³-NH₂]enkephalin, showed a 10000 fold increase in potency in the same in vivo assay (entry 3). In order to assess a "fluorine effect" on potency, (R)-norvaline (R)-Nval) was synthesized and assayed. As entry 10 shows, this analogue exhibited almost equivalent potency to that of the (R)-Ala² analogue, which was 1 order of magnitude weaker than the (R)-TFNV analogue. Accordingly, it is clear that there was a "fluorine effect", which improved the potency by 1 order of magnitude even after a major enhancement factor, i.e., (R)-amino acid residue at Gly² position, was introduced. The most potent analogue in this series was [(R)-TFNV², (N-Me)Phe², Met³-NH₂]enkephalin (entry 13), which was 3.5 times more potent than [(R)-TFNV², Met³-NH₂]enkephalin.

Receptor Binding Assay in Vitro. In order to investigate the origin of the remarkable enhancement in potency by the introduction of (R)-TFNV at the Gly² position, the in vitro receptor binding assays for [(R)-TFNV², Met³-NH₂]enkephalin were carried out against µ, δ, and κ receptors using tritium-labeled standard ligands. As Table 2 shows, [(R)-TFNV², Met³-NH₂]enkephalin exhibited a 10⁻¹⁰ M level IC₅₀ against µ-receptor, but it was only a half order of magnitude enhancement in the binding ability compared with methionine-enkephalin. For δ-receptor, [(R)-TFNV², Met³-NH₂]enkephalin showed almost the same level binding ability as methionine–enkephalin. Interestingly, [(R)-TFNV², Met³-NH₂]enkephalin bound to the κ-receptor at the 10⁻⁷ M level IC₅₀ whereas methionine–enkephalin did not show any appreciable binding.

The results clearly indicate that the observed remarkable enhancement in the in vivo potency of [(R)-TFNV², Met³-NH₂]enkephalin was not based on much stronger binding to receptor sites, but mainly due to the extremely efficient inhibition of degradation by aminopeptidase(s). Possible enhancement of the rates of absorption and transport, arising from the lipophilicity of CF₃ group should also be taken into account as the secondary effect. It is worth noting that [(R)-TFNV², Met³-NH₂]enkephalin was found to cross the blood–brain barrier.

The studies described in this section clearly demonstrated the uniqueness and usefulness of fluoroamino acids as modifiers of peptides and enzyme inhibitors of medicinal interest. Thus, we envisioned that further research in this direction would explore newer and exciting aspects of organofluorine chemistry in medicinal chemistry and chemical biology.
troscopy along with large $^{19}$F--$^1$H coupling constants and the virtual absence of $^{19}$F in the living tissue render fluorine incorporation a particularly powerful tool for the investigation of biological processes.\textsuperscript{8,10,16,19} Therefore, fluorine-containing $\alpha$-hydroxy-$\beta$-amino acids are expected to serve as useful bioactive compounds with a wide range of potential applications in medicinal chemistry and chemical biology. However, only a few methods had been reported for the synthesis of fluorine-containing $\alpha$-hydroxy-$\beta$-amino acids when we started our investigation in the mid-late 1990s.\textsuperscript{19,110--114}

Accordingly, newer and efficient approaches to enantiopure fluorine-containing $\alpha$-hydroxy-$\beta$-amino acids needed to be developed. To this end, the “$\beta$-lactam synthon method” invented in my laboratory and developed by us and others\textsuperscript{115--120} offered an attractive protocol for the synthesis of enantiopure CF$_2$H- and CF$_3$-containing $\alpha$-hydroxy-$\beta$-amino acids and their congeners.

### 6.1. Synthesis of Enantiopure 3-Hydroxy-4-CF$_2$H-$\beta$-lactams

Racemic cis-$\beta$-lactam ($\pm$)-16 was prepared through $[2 + 2]$ ketene lactams. The enzymatic optical resolution of racemic cis-3-BnO-4-CF$_2$-$\beta$-lactam ($\pm$)-23 in moderate yield, as reported by us and others.\textsuperscript{111,124} Hydrogenolysis of $\beta$-lactam ($\pm$)-23, followed by acetylation, gave the corresponding racemic cis-3-AcO-CF$_2$-$\beta$-lactam ($\pm$)-25 in good overall yield (Scheme 13).\textsuperscript{122,123}

### 6.2. Synthesis of Enantiopure 3-Hydroxy-4-CF$_3$-$\beta$-lactams

A different strategy was employed for the synthesis of enantiopure 4-CF$_3$-$\beta$-lactams. Namely, the CF$_3$ moiety was introduced, from the very beginning, to the imine to be used for the $[2 + 2]$ ketene--imine cycloaddition. N-PMP-trifluoroacetaldimine (22) was reacted with the ketene generated in situ from benzoylxyacetyl chloride to afford racemic cis-3-BnO-4-CF$_2$-$\beta$-lactam ($\pm$)-23 with 99.9% ee as well as (3S,4S)-3-hydroxy-4-CF$_3$-$\beta$-lactam ($\pm$)-26 with 97% ee in good to high yield.\textsuperscript{122}

### 6.3. Synthesis of Enantiopure 1-Acyl-3-hydroxy-4-R$_F$-$\beta$-lactams

The $\beta$-lactam synthon method has been developed by exploiting the unique nature of this strained four-membered skeleton for its facile ring-opening reactions with a variety of nucleophiles.\textsuperscript{115,116} When the nitrogen of this strained cyclic amide is acylated (including carbalkoxy, carbamoyl, thiocarboxyl, and sulfonyl groups besides the standard acyl groups), the resulting $N$-acyl-$\beta$-lactam becomes exceptionally reactive for
nucleophilic attacks, leading to facile ring-opening coupling. This unique feature of N-acyl-β-lactams has been successfully utilized in organic synthesis and medicinal chemistry.\textsuperscript{115,117–120}

We prepared a series of N-acyl-3-TIPSO-3-Rf-β-lactams\textsuperscript{29} (Rf = CF$_2$H or CF$_3$), as examples, in good to high yield through acylation (including carbalkoxylation and sulfonylation) of 3-TIPSO-4-CF$_2$H-β-lactam\textsuperscript{21} and 3-TIPSO-4-CF$_3$-β-lactam\textsuperscript{28}, using (Boc)$_2$O, ClCO$_2$Bn, 4-F-C$_6$H$_4$COCl and p-TolSO$_2$Cl in the presence of an appropriate base such as DMAP/Et$_3$N in CH$_2$Cl$_2$ (eq 17: only (3$\text{R}$,4$\text{R}$) series is shown for simplicity).\textsuperscript{122,123}

6.4. Synthesis of Enantiopure β-CF$_2$H- and β-CF$_3$-α-hydroxy-β-amino Acids and Esters via Facile Hydrolysis/Alcoholysis of N-Acetyl-β-lactams. Enantiopure β-R$_f$-α-hydroxy-β-amino acids (R$_f$ = CF$_2$H or CF$_3$) were readily obtained through facile ring-opening hydrolysis of (+)- or (−)-29. For example, the reaction of 29 with KOH in aqueous THF at ambient temperature gave the corresponding O-protected amino acid 30, in high yield (eq 18: only (2S,3S) series is shown for simplicity).\textsuperscript{122,123} The O-TIPS protecting group could be easily removed by HF/pyridine as needed to give 31.

6.5. Synthesis of R$_f$-Containing Isoserine Dipeptides through Efficient Ring-Opening Coupling of N-Acyl-β-lactams with α- and β-Amino Esters. The ring-opening coupling of N-acyl-β-lactam, (+)- or (−)-36, with various α- and β-amino acid esters provided a very easy access to a library of R$_f$-containing isoserine dipeptides. Since these coupling reactions do not need any peptide coupling reagents such as DCC, DIC and EDC, the “atom economy”\textsuperscript{125} is extremely high. The coupling reactions gave the corresponding R$_f$-containing isoserine dipeptides, 34 and 35, in good to quantitative yields (Scheme 16: Only (S,S,S) or (S,S) series is shown for simplicity).\textsuperscript{122,123}

6.6. Synthesis of R$_f$-Containing Isoserine Dipeptides through Feasible Transformations of 1-Carbalkoxy-4-R$_f$-β-lactams to Peptides and Peptidomimetics. The results described above as well as our previous works\textsuperscript{126–128} allow us to envision the versatile utility of the N-carbalkoxy-3-TIPSO-4-R$_f$-β-lactams for the synthesis of R$_f$-containing isoserine dipeptides, depsipeptides, peptidomimetics, and key synthetic building blocks for R$_f$-containing hydroxyethylene, dihydroxyethylene, and hydroxyethylene dipeptide isosteres. Possible transformations of N-carbalkoxy-3-TIPSO-4-R$_f$-β-lactams are illustrated in Scheme 17.
most important chemotherapeutic drugs, currently used for the
treatment of advanced ovarian cancer, metastatic breast cancer,
melanoma, non-small cell lung cancer, and Kaposi’s sarcoma.129–131
More recently, these drugs have been used for the
treatment of neck, prostate, and cervical cancers.130,131
The mechanism of action of paclitaxel involves its binding to the β-
subunit of αβ-tubulin dimer, accelerating the formation of
microtubules. The resulting paclitaxel-bound microtubules are
much more stable and less dynamic than the natural GTP-
bound microtubules. The resulting paclitaxel-bound microtubules are
frozen out. Accordingly, F2-paclitaxel SB-T-31031 and F-
docetaxel SB-T-3001 were selected as probes (Figure 5) for
the study of the solution structures and dynamic behavior of
paclitaxel and docetaxel, respectively, in protic and aprotic
solvent systems.134

7.2. The Fluorine Probe Approach: Solution-Phase
Structure and Dynamics of Taxoids. The rational design of
new generation taxoid anticancer agents would be greatly
facilitated by the development of reasonable models for the
biologically relevant conformations of paclitaxel. In this regard,
we recognized that the design and synthesis of fluorine-
containing taxoids would have a very useful offshoot of
providing us with the capability of studying bioactive
conformations of taxoids using a combination of 19F/1H
NMR techniques and molecular modeling.134

The early conformational analysis of paclitaxel and docetaxel
in solution largely identified two major conformations, with
minor variations between studies.135 Structure I, characterized by
gault conformation with a H2–C2′–C3′–H3′ dihedral angle of ca. 60°, was based on the X-ray crystal structure of
docetaxel136 and was believed to be commonly observed in
aprotic solvents (Figure 4, I).135 Structure B, characterized by the antti conformation with a H2′–C2′–C3′–H3′ dihedral angle of ca. 180°, was observed in theoretical conformational
analysis137,138 as well as 2D NMR analyses,139 and found in the
X-ray structure of the xylene extracted from a dioxane/H2O/
xylene solution (Figure 4, II).140 Despite extensive structural
studies, no systematic study on the dynamics of these two as
well as other possible bioactive conformations of paclitaxel had
been reported when we started our study on this problem. The
relevance of the “fluorine probe” approach to study dynamic
properties prompted us to conduct a detailed investigation into
the solution dynamics of fluorine-containing paclitaxel and
docetaxel analogues.

The use of 19F NMR for a variable temperature (VT) NMR
study of fluorine-containing taxoids was obviously advantageous
over the use of 1H NMR because of the wide dispersion of the
19F chemical shifts that allows fast dynamic processes to be
frozen out. Accordingly, F2-paclitaxel SB-T-31031 and F-
docetaxel SB-T-3001 were selected as probes (Figure 5) for
the study of the solution structures and dynamic behavior of
paclitaxel and docetaxel, respectively, in protic and aprotic
solvent systems.134

Analysis of the low temperature VT-NMR (19F and 1H) and
19F–1H heteronuclear NOE spectra of SB-T-31031 and SB-T-
3001 in conjunction with molecular modeling revealed the
presence of equilibrium between two conformers in protic solvent systems. Interpretation of the temperature dependence
of the coupling constants between H2′ and H3′ for SB-T3013 indicated that one of these conformers (conformer C,
Figure 6 and Figure 7) possessed an unusual near-eclipsed
arrangement around the H2′–C2′–C3′–H3′ dihedral angle
(JH2′-C2′-H3′ = 5.2 Hz, corresponding to the H2′–C2′–C3′–H3′
torsion angle of 124° based on the MM2 calculation) and was
found to be more prevalent at ambient temperatures.134 The
other one corresponded to the anti conformer (conformer B,
JH2′-H3′ = 10.1 Hz, corresponding to the H2′–C2′–C3′–H3′
torsion angle of 178° based on the MM2 calculation) and was
quite closely related to the structure I in Figure 4. These
conformers were different from the one observed in aprotic
solvents (conformer A, H2′–C2′–C3′–H3′ torsion angle of 54°) that was related to the X-ray crystal structure of docetaxel
represented by the structure I in Figure 4.136 Figure 6 shows the Newman projections for these three conformers and Figure
7 the structures of the conformers A, B, and C.134
Restrained molecular dynamics (RMD) studies presented evidence for the hydrophobic clustering of the 3′-phenyl and 2- benzoate (Ph moiety) for both conformers B and C. Although the conformer C possessed a rather unusual semieclipsed arrangement around the C2′−C3′ bond, the unfavorable interaction associated with such a conformation was apparently offset by significant solvation stabilization observed in the comparative RMD study in a simulated aqueous environment for the three conformers. The solvation stabilization term for the conformer C was estimated to be about 10 kcal/mol greater than those for the conformers A and B. Accordingly, the “fluorine probe” approach succeeded in finding a new conformer that had never been predicted by the previous NMR and molecular modeling studies.

Strong support for the conformer C was found in its close resemblance to a proposed solution structure of a water-soluble paclitaxel analog, paclitaxel-7-MA (MPA = N-methylpyridinium acetate), wherein the H2′−C2′−C3′−H3′ torsion angle of the N-phenylisoserine moiety is 127°, which is only a few degrees different from the value for the conformer C.

Thus, the “fluorine probe” approach has proved highly useful for the conformational analysis of paclitaxel and taxoids in conjunction with the determination of possible bioactive conformations. The previously unrecognized conformer C might be the molecular structure first recognized by the β-tubulin binding site on microtubules.

**Determination of the Binding Conformation of Taxoids in Microtubules Using Fluorine Probes.** The knowledge of the solution structures and dynamics of paclitaxel and its analogues is necessary for a good understanding of the recognition and binding processes between paclitaxel and its binding site on the microtubules. Such knowledge would also provide crucial information for the design of future generation anticancer agents. However, the elucidation of the microtubule-bound conformation of paclitaxel was critical for the rational design of efficient inhibitors of microtubule disassembly. The lack of information about the three-dimensional tubulin binding site by protein X-ray crystallography prompted us to apply our fluorine probe approach to the determination of the F−F distances in the microtubule-bound F2-taxoids. The results of such a study should provide the relevant distance map for the identification of the bioactive (binding) conformation of paclitaxel.

We successfully applied the fluorine probe approach to the estimation of the F−F distance in the microtubule-bound F2-10-Ac-docetaxel (Figure 5) using the solid-state magic angle spinning (SS MAS) 19F NMR coupled with the radio frequency driven dipolar recoupling (RFDR) protocol in collaboration with L. Gilchrist, A. E. McDermott, K. Nakanishi (Columbia University), S. B. Horwitz, and M. Orr (A. Einstein College of Medicine). 107

F2-10-Ac-docetaxel was first studied in a polycrystalline form by the RFDR protocol. Based on the standard simulation curves derived from molecules with known F−F distances (distance markers), the F−F distance of two fluorine atoms in F2-10-Ac-docetaxel was estimated to be 5.0 ± 0.5 Å (Figure 8). This value corresponded quite closely to the estimated F−F distances for the conformers B and C (F−F distance was ca. 4.5 Å for both conformers) based on our RMD studies for F2- paclitaxel (SB-T-31031). This means that the microcrystalline structure of F2-10-Ac-docetaxel was consistent with the hydrophobic clustering conformer B or C, but not with the conformer A in which the F−F distance was ca. 9.0 Å.

The microtubule-bound complex of F2-10-Ac-docetaxel revealed the F−F distance to be 6.5 ± 0.5 Å (Figure 8), which was larger than that observed in the polycrystalline form by ca. 1 Å. It is very likely that the microtubule-bound conformation of F2-10-Ac-docetaxel was achieved by a small distortion of the solution conformation, i.e., either conformer B or C. 107

The above account demonstrated the power of the fluorine probe approach that is evident from its ability to supply extremely valuable and precise information about both bound and dynamic conformations of biologically active molecules. Such information is especially useful in the absence of knowledge about the three-dimensional crystal structure of their binding site. It is worthy of note that our preliminary study on the structure of the protein-bound fluorine-labeled

**Figure 6.** Newman projections of the isoserine moieties of the three conformers of F2-paclitaxel identified.

**Figure 7.** Three conformers of F2-paclitaxel identified.

**Figure 8.** Determination of the F−F distance in F2-10-Ac-paclitaxel using the RFDR protocol by solid state MAS 19F NMR spectroscopy.
docetaxel, by means of the Solid State MAS$^{19}$F NMR using the REDOR protocol, was the very first to tackle such an important and challenging problem in the structural and chemical biology of paclitaxel and tubulin/microtubules, which made a solid foundation for further investigations along this line (see section 7.6).

7.3. Second-Generation Taxoids. It has been shown that a primary mechanism of drug resistance is the overexpression of ABC transporters, e.g., P-glycoprotein, an integral membrane glycoprotein that acts as a drug-efflux pump to maintain the intracellular concentration of drugs below therapeutically active level.$^{142}$ In the course of our extensive studies on the design, synthesis and the structure–activity relationship (SAR) of taxoid anticancer agents, we discovered second-generation taxoids that possess 1 order of magnitude better activity against drug-sensitive cell lines and more than 2 orders of magnitude better activity against drug resistant cell lines.$^{143,144}$ Several examples are shown in Table 3 (see Figure 9 for structures). It was found that meta-substitution of the benzoyl group at the C2 position substantially increased the cytotoxicity of taxoids against drug-resistant cell lines.$^{144,145}$ This class of the second-generation taxoids is 3 orders of magnitude more potent than paclitaxel and docetaxel against drug-resistant cancer cell lines, expressing MDR phenotype (Table 3).$^{144,145}$

Because of the aforementioned advantages of introducing fluorine into biologically active molecules, we synthesized fluorine-containing paclitaxel and docetaxel analogues to investigate the effects of fluorine—incorporation on the cytotoxicity and the blockage of known metabolic pathways.

<table>
<thead>
<tr>
<th>Taxoid</th>
<th>MCF7 IC50 (nM)</th>
<th>NCI/ADR IC50 (nM)</th>
<th>LCC6-WT IC50 (nM)</th>
<th>LCC6-MDR IC50 (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paclitaxel</td>
<td>1.7</td>
<td>300</td>
<td>3.1</td>
<td>346</td>
</tr>
<tr>
<td>Docetaxel</td>
<td>1.0</td>
<td>235</td>
<td>1.0</td>
<td>120</td>
</tr>
<tr>
<td>SB-T-1103</td>
<td>0.36</td>
<td>0.33</td>
<td>1.0</td>
<td>0.9</td>
</tr>
<tr>
<td>SB-T-1103</td>
<td>0.36</td>
<td>0.43</td>
<td>0.9</td>
<td>0.8</td>
</tr>
</tbody>
</table>

7.4. 3′-Difluoromethyltaxoids and 3′-Trifluoromethyltaxoids. In the course of our extensive SAR studies of the taxoid anticancer agents, we have synthesized a good number of fluorotaxoids by means of the “β-lactam synthon method” to investigate the effects of fluorine on cytotoxicity and metabolic stability.$^{106,107,146}$ Along this line, second-generation fluorotaxoids possessing CF$_2$H and CF$_3$ groups at the 3′-position were synthesized and their biological activity evaluated.$^{122,147}$ The synthesis of these fluorotaxoids is shown in Scheme 18.

![Scheme 18. Synthesis of 3′-CF$_2$H- and 3′-CF$_3$-taxoids](image)

7.5. 3′-Difluoroxytaxoids. As described above, the introduction of isobutyl, isobutenyl, CF$_2$H, and CF$_3$ groups to the 3′-position of taxoids, replacing the phenyl group of paclitaxel and docetaxel, has led to the development of highly potent second-generation taxoids, especially against drug-resistant cancer cell lines, expressing MDR phenotype. Our...
metabolism studies on 3′-isobutyl- and 3′-isobutenyl-taxoids disclosed that the metabolism of second-generation taxoids (SB-T-1214 and SB-T-1103) was markedly different from that of docetaxel and paclitaxel. These taxoids were metabolized (via hydroxylation) by CYP 3A4 of the cytochrome P450 family enzymes primarily at the two allylic methyl groups of the 3′-isobutyl group and the methyne moiety of the 3′-isobutyl group (see Figure 10). This finding was in sharp contrast to the known result that the tert-butyl group of the C3′N-t-Boc moiety was the single predominant metabolic site for docetaxel. These unique metabolic profiles prompted us to design and synthesize 3′-difluorovinyltaxoids in order to block the allylic oxidation by CYP 3A4 mentioned above, which should enhance the metabolic stability and activity in vivo.

For the synthesis of a series of 3′-difluorovinyltaxoids 48, novel (3R,4S)-1-t-Boc-3-TIPSO-4-difluorovinyl-β-lactam (+)-47 was the key component for the coupling with baccatins 43 (Scheme 19). We prepared β-lactam (+)-47 in three steps from 4-formyl-β-lactam (+)-26 (see Scheme 1) using the Wittig reaction of the formyl moiety with difluoromethylphosphorus ylide generated in situ from (Me₂N)₃P/CF₂Br₂/Zn (Scheme 19). The ring-opening coupling reaction of β-lactam (+)-47 with baccatins 43 (X = H, MeO, N₃) and the subsequent removal of the silyl protecting groups gave the corresponding 3′-difluorovinyltaxoids 41 in good to excellent yields (Scheme 19). Cytotoxicities of the 3′-difluorovinyltaxoids 41 were evaluated against 4 human cancer cell lines. Results for selected taxoids are summarized in Table 5.

As Table 5 shows, all difluorovinyltaxoids 41 are exceedingly potent as compared to paclitaxel. A clear effect of C2-benzoate modification at the meta position (X = H vs X = MeO or N₃) was observed on the increase in potency against MCF7 (Pgp-) and NCI/ADR (Pgp+) cell lines (entries 2–5 vs entries 6–11). Difluorovinyltaxoids with 2,10-modifications (entries 6–11) exhibited impotent potency, exhibiting IC₅₀ values in the >100 pM range (78–92 pM) except one case against MCF7 (entry 7) and in the subnanomolar range (0.34–0.57 nM) against NCI/ADR, which was 3 orders of magnitude more potent than paclitaxel. The resistance factor for these taxoids was 1.7–6.4, while that for paclitaxel was 250. Difluorovinyltaxoids with an unmodified C2-benzoate moiety (entries 2–5) also showed highly enhanced potency against MCF7 and NCI/ADR as compared to paclitaxel. These taxoids exhibited impressive potency against HT-29 (human colon) and PANCl-1 (human pancreatic) cancer cell lines as well. SB-T-12853 appeared particularly promising against these gastrointestinal (GI) cancer cell lines.

7.6. Possible Bioactive Conformations of Fluorotaxoids. As described in section 7.2, we have successfully used fluorine-containing taxoids as probes for NMR analysis of the
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Table 5. In Vitro Cytotoxicity (IC_{50}, nM) of 3′-Difluorovinyltaxoids 41

<table>
<thead>
<tr>
<th>entry</th>
<th>taxoid</th>
<th>R</th>
<th>X</th>
<th>MCF_{78} (breast)</th>
<th>NCI/ADR* (ovarian)</th>
<th>R/S</th>
<th>HT-29d (colon)</th>
<th>PANC-1* (pancreatic)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>paclitaxel</td>
<td></td>
<td></td>
<td>1.2</td>
<td>300</td>
<td>250</td>
<td>3.6</td>
<td>25.7</td>
</tr>
<tr>
<td>2</td>
<td>SB-T-12851</td>
<td>Ac</td>
<td>H</td>
<td>0.099</td>
<td>0.95</td>
<td>9.6</td>
<td>0.41</td>
<td>1.19</td>
</tr>
<tr>
<td>3</td>
<td>SB-T-12852</td>
<td>c-Pr-CO</td>
<td>H</td>
<td>0.12</td>
<td>6.0</td>
<td>50</td>
<td>0.85</td>
<td>5.85</td>
</tr>
<tr>
<td>4</td>
<td>SB-T-12853</td>
<td>Et-CO</td>
<td>H</td>
<td>0.12</td>
<td>1.2</td>
<td>10</td>
<td>0.34</td>
<td>0.65</td>
</tr>
<tr>
<td>5</td>
<td>SB-T-12854</td>
<td>Me,N-CO</td>
<td>H</td>
<td>0.13</td>
<td>4.3</td>
<td>33</td>
<td>0.46</td>
<td>1.58</td>
</tr>
<tr>
<td>6</td>
<td>SB-T-12852-1</td>
<td>c-Pr-CO</td>
<td>MeO</td>
<td>0.092</td>
<td>0.48</td>
<td>5.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>SB-T-12853-1</td>
<td>Et-CO</td>
<td>MeO</td>
<td>0.34</td>
<td>0.57</td>
<td>1.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>SB-T-12851-1</td>
<td>MeO-CO</td>
<td>MeO</td>
<td>0.078</td>
<td>0.50</td>
<td>6.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>SB-T-12851-3</td>
<td>Ac</td>
<td>N_3</td>
<td>0.092</td>
<td>0.34</td>
<td>3.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>SB-T-12853-3</td>
<td>MeO-CO</td>
<td>N_3</td>
<td>0.078</td>
<td>0.40</td>
<td>5.3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3. See footnotes of Table 3. *Human pancreatic carcinoma.

In 2005, we proposed a new bioactive conformation of paclitaxel, "REDOR-Taxol", based on (i) the 19F→13C distances obtained by the REDOR experiment, (ii) the photoaffinity labeling of microtubules, (iii) the crystal structure (pdb code: 1TUB) of Zn^{2+}-stabilized paclitaxel and docetaxel. Then, Schaefer and co-workers used the rotational echo double resonance (REDOR) to investigate the structure of the microtubule-bound paclitaxel by determining the 19F→13C distances of an fluorine-probe of paclitaxel (Figure 11). These SS MAS 19F NMR studies have provided critical information on the bioactive conformation of paclitaxel and docetaxel.

Figure 11. Solid-state NMR studies on microtubule-bound fluorotaxoid probes.

In 2007, three additional intramolecular distances of the key atoms in the microtubule-bound 19F/H-labeled paclitaxel were determined by the REDOR method (Figure 11). Also, it has been shown that the optimized cryo-EM crystal structure of tubulin-bound paclitaxel (pdb code: 1JFF) serves better for the computational structure analysis. Accordingly, we have optimized our REDOR-Taxol structure, using the 1JFF coordinates as the starting point, by means of molecular dynamics simulations (Macromodel, MMFF94) and energy minimization (InsightII 2000, CVFF).

We applied the same computational protocol to investigate the microtubule-bound structures of the 3′-CF_2H, 3′-CF_3, and 3′-CF_2C≡CH-taxoids, using the updated REDOR-Taxol as the starting structure. Three fluorotaxoids, SB-T-1284, SB-T-1282, and SB-T-12853 (Figure 12), were docked into the binding pocket of paclitaxel in the β-tubulin subunit by superimposing the baccatin moiety with that of the REDOR-Taxol and their energies minimized (InsightII 2000, CVFF). The resulting computer-generated binding structures of these fluorotaxoids are shown in Figure 13 (A, B, and C).

As shown in Figure 13 (A, B, and C), the baccatin moiety occupies virtually the same space in all cases, as expected. Each fluorotaxoid fits comfortably in the binding pocket without any high-energy contacts with the protein. There is a very strong hydrogen bond between the C2′-OH of a fluorotaxoid and His^219 of β-tubulin in all cases, which shares the same key feature with the REDOR-Taxol structure. It should be noted that our preliminary study on the tubulin-bound structures of these three fluorotaxoids using the 1TUB coordinates led to different structures in which the C2′-OH had a hydrogen bonding to Arg359 of β-tubulin. However, the use of the updated REDOR-Taxol structure based on the 1JFF coordinates unambiguously led to the fluorotaxoids structures bearing a strong hydrogen bond between the C2′-OH and His^219.

The CF_2H and CF_3 moieties fill essentially the same space, as anticipated. However, the CF_2C≡CH moiety occupies more extended hydrophobic space than the CF_2H and CF_3 moieties. It is likely that this additional hydrophobic interaction is substantially contributing to the exceptional cytotoxicity of 3′-
representative second-generation taxoid, SB-T-1213 shows systems for anticancer agents, di ff from cancer cells or tissues, is an urgent need to improve the tri modi component in medicinal chemistry and may serve as a versatile Anticancer Agents. Traditional chemotherapy depends on to be destroyed by cytotoxic agents than normal cells. In reality, di ff groups rather than two methyl groups electronically. Accord-ing, di ff groups in size, and two fluo-ine atoms may mimic two hydroxyl groups rather than two methyl groups electronically. Accord-ingly, the di ff group is a very unique structural component in medicinal chemistry and may serve as a versatile modi fier of pharmacological properties in a manner similar to the trifluromethyl group.

7.7. Use of Fluorine in Taxoid-Based Tumor-Targeting Anticancer Agents. Traditional chemotherapy depends on the premise that rapidly proliferating tumor cells are more likely to be destroyed by cytotoxic agents than normal cells. In reality, however, these cytotoxic agents have little or no specificity, which leads to systemic toxicity causing undesirable side effects. Accordingly, the development of tumor-specific drug delivery systems for anticancer agents, differentiating the normal tissues from cancer cells or tissues, is an urgent need to improve the efficacy of cancer chemotherapy. Various drug delivery systems have been studied over the past few decades to address this problem.161 Rapidly growing cancer cells overexpress tumor-specific receptors to enhance the uptake of nutrients and vitamins. These receptors can be used as targets for cancer cell-specific delivery of cytotoxic agents through receptor-mediated endocytosis. Furthermore, the characteristic physiology of tumor and cancer cells can be exploited to selectively accumulate and release a cytotoxic agent inside these cells. Monoclonal antibodies, polyunsaturated fatty acids, folic acid, biotin, aptamers, transferrin, oligopeptides, and hyaluronic acid, for example, have been employed as tumor-specific “guiding modules” to construct tumor-targeting drug conjugates.161–166

As a general structure, tumor-targeted drug delivery systems consist of a tumor-targeting module (TTM) conjugated to a cytotoxic warhead directly or through a suitable “smart” linker (Figure 14). These drug conjugates should be stable in blood circulation to minimize systemic toxicity and should be effectively internalized inside the target tumor cells. Upon internalization, the drug conjugate should efficiently release the cytotoxic agent without loss of potency. Thus, the “smart” linkers should possess proper characteristics to provide suitable stability and reactivity.

Fluorine-containing anticancer agents can certainly serve as potent “warheads” for the tumor-targeting drug conjugates.104 In addition, fluorine can be strategically incorporated to create useful biochemical tools that facilitate the development of such tumor-targeting drug conjugates. For example, 18F NMR can provide techniques to directly observe time-dependent processes in complex biological systems since the fluorine nucleus is virtually nonexistent in biological systems.167 As 18F is a common radioisotope used for positron emission tomography (PET) imaging, selective fluorination for biodistribution studies has found extensive use in diagnostic applications.168–170

This section concisely describes our recent approaches to the strategic incorporation of fluorine(s) into taxoid-based tumor-targeted drug delivery systems and ongoing investigations with future perspective.

7.7.1. Polyunsaturated Fatty Acid (PUFA)—Taxoid Conjugates. Omega-3 polyunsaturated fatty acids (PUFA), such as linolenic acid (LNA), eicosapentaenoic acid (EPA), and docosahexaenoic acid (DHA), are naturally occurring compounds found in vegetable oils, cold-water fish, and meat.171 Perfusion studies have shown that some PUFAs are taken up more rapidly by tumor cells than normal cells.172 It has also been shown that PUFAs are readily incorporated into cellular membranes, catabolized as an energy resource, and produce metabolites that act as signaling molecules, modulating various intracellular processes ranging from inflammatory response to cellular proliferation.173 It has been shown that the conjugation of a drug to a PUFA greatly alters the pharmacokinetics (PK) profile of the parent compound, leading to tumor-selective accumulation of the drug conjugate.174 Thus, a number of PUFA–drug conjugates are currently undergoing preclinical and clinical evaluations.175 For example, DHA—paclitaxel (Taxoprexin), currently in phase III clinical trials, has exhibited better efficacy than paclitaxel in some studies174,176 but does not show efficac against multidrug-resistant (MDR) tumors that overexpress P-glycoprotein. As mentioned above, second-generation taxoids and fluorotaxoids exhibited 2–3 orders of magnitude higher potency than paclitaxel against MDR cancer cell lines. Thus, PUFA conjugates, bearing a second-generation taxoid or fluorotaxoid should be more efficacious than DHA-paclitaxel against drug-resistant tumors.

Accordingly, novel DHA— and LNA—taxoid conjugates were synthesized and assayed in vivo for their efficacy against...
different drug-resistant and drug-sensitive human tumor xenografts in severe combined immune deficiency (SCID) mice. Several of these conjugates led to a complete regression of the tumor in all surviving mice with minimal systemic toxicity. For example, DHA-SB-T-1214 led to a complete regression of the highly drug-resistant DLD-1 colon tumor xenograft in mice without appreciable systemic toxicity, wherein no recurrence of tumor growth was observed for more than 190 days after treatment. DHA-SB-T-1214 is currently undergoing extensive late-stage preclinical evaluation, and an Investigational New Drug (IND) application will be going forward in the near future. Since 3′-difluorovinytaxoids exhibit equivalent or even higher potency than SB-T-1214 with excellent metabolic stability, DHA and LNA conjugates of SB-T-12854 (Figure 15) have been synthesized and their efficacy in vivo is currently under active investigation. A preliminary in vivo efficacy study on LNA-SB-T-12854 against highly metastatic MX-1 human breast tumor xenograft in nude mice has shown promising results, wherein the complete eradication of tumor was achieved using weekly regimen for drug administration.

7.7.2. Self-Immobilative Disulfide Linkers for Tumor-Targeted Drug Delivery. Monoclonal Antibody–Taxoid Conjugates with First-Generation Disulfide Linker. Cancer cells overexpress certain antigens on the cell surface and these tumor-specific antigens can be used as biomarkers to differentiate tumor tissues from normal tissues. Certain monoclonal antibodies (mAb) have high binding specificity to tumor-specific antigens and can be used as drug delivery vehicles to carry a payload of cytotoxic agents specifically to the tumor site. The mAb–drug conjugate is internalized upon binding to the tumor antigen via receptor-mediated endocytosis (RME) and the payload is released inside the cancer cell. For example, brentuximab vedotin (Adcetris) is an mAb–drug conjugate targeting CD30 and recently approved by the FDA for the treatment of Hodgkin’s lymphoma, and several other mAb–drug conjugates are currently in human clinical trials.

The efficacy of mAb–drug immunoconjugates depends not only on the specificity of the mAb and the potency of the cytotoxic drug, but also on the linker which connects the mAb to the drug. We successfully conjugated a highly cytotoxic C-10 methylsulfonylpropanoyl taxoid to immunoglobulin G class mAbs, recognizing the epidermal growth factor receptor (EGFR), through a disulfide-containing linker (Figure 16). These conjugates showed excellent selectivity in vitro and remarkable antitumor activity in vivo against A431 human squamous tumor xenografts in SCID mice, resulting in eradication of the tumor without appreciable systemic toxicity. However, the modification at the 10 position of the taxoid resulted in 8–10 times loss of potency relative to the parent taxoid. Accordingly, the mechanism-based second-generation linker system was designed and developed to allow the release of the unmodified taxoid with uncompromised potency.

Second-Generation Self-Immobilative Disulfide Linkers. Second-generation mechanism-based bifunctional disulfide linkers can be generally used to connect a warhead to one end and a tumor-targeting module (TTM) to the other end. This self-immobilative disulfide linker module can release a taxoid warhead efficiently inside cancer cells by taking advantage of 1000 times higher concentration of glutathione in tumor as compared to that in blood plasma. When the TTM navigates the drug-conjugate to the target receptors on the tumor surface, the whole conjugate is internalized via RME. Then, an intracellular thiol-triggered cascade drug-release takes place through thiolactonization (Figure 17) and the released potent anticancer drug attacks its target protein, i.e., microtubules for taxoids. To promote the thiolactonization process, a phenyl moiety was strategically placed to direct the cleavage of the disulfide bond by intracellular thiol (e.g., glutathione), generating a thiophenolate or sulfhydrylphenyl species which attacks the ester linkage to the drug molecule (Figure 18).

The validity of this self-immobilative drug-release mechanism has been proven in a model system using fluorine-labeling and monitoring by 19F NMR spectroscopy (Figure 18) as well as in a real system with cancer cells using fluorescence-labeling and confocal fluorescence microscopy (CFM). These self-immobilative disulfide linkers have been successfully incorporated into various tumor-targeting drug conjugates and their efficacy evaluated in cancer cells.
and vitamin B12, are overexpressed on the cancer cell surface, providing useful targets for tumor-targeted drug delivery.187 Receptors for vitamins, including those for folic acid (vitamin M, vitamin B9), biotin (vitamin H, vitamin B7, coenzyme R), and vitamin B12, are overexpressed on the cancer cell surface, providing useful targets for tumor-targeted drug delivery.188 Biotin serves as a coenzyme for dependent carboxylases and plays a significant role in epigenetic regulation, fatty acid synthesis, energy production and the metabolism of fats and amino acids.189,190 Biotin receptors had not been studied until 2004 when vitamin receptor-targeted rhodamine polymers indicated that receptors for biotin were even more overexpressed on the surface of cancer cells than those for folic acid and vitamin B12. Thus, the biotin receptor has emerged as a novel target for tumor-targeted drug delivery in addition to the well-established folate receptors.

A general mechanism is illustrated in Figure 19 for the internalization of a tumor-targeting drug conjugate through receptor-mediated endocytosis (RME), drug release, and the binding of the released drug to the target protein (taxoid binds microtubules).185 The internalization of the fluorescent probes via RME, drug release via disulfide bond cleavage and binding of the released taxoid “warhead” to microtubules were validated by CFM analysis of three fluorescent probes, i.e., (a) biotin–FITC conjugate, (b) biotin–linker–coumarin conjugate and (c) biotin–linker–taxoid–fluorescein.166,184,186

In addition, biotin–linker–taxoid conjugate was synthesized and assayed in vitro against L1210 (mouse lymphocytic leukemia), L1210FR (folic and biotin receptors overexpressed L1210 leukemia) and WI38 (normal human lung ﬁbroblastoma) cells to examine the efﬁcacy of the biomarker-speciﬁc targeting of the conjugate.166,184,186 The IC50 values of the conjugate (taxoid = SB-T-1214) against L1210FR, L1210, and WI38 cell lines were 8.80 nM, 522 nM and 570 nM, respectively. The results clearly indicate the high target-speciﬁcity of the drug conjugate, which is consistent with the RME-based internalization and drug release observed by CFM and flow cytometry using ﬂuorescent probes. Accordingly, this tumor-targeted drug delivery system bearing a vitamin as the tumor-targeting module is ready to utilize highly potent fluorotaxoids as “warheads”.

### 7.7.3. Vitamins as Tumor-Targeting Modules.

Cancer cells critically require certain vitamins, such as those essential for cell division, to sustain their rapid growth although vitamins are essential for the growth and development of all living cells. Vitamins are not functionalized carbon nanotubes (CNTs) have emerged as a unique “warhead” are linked via a 1,3,5-triazine splitter with a watersolubilizing triethylene glycol (PEG3) spacer and a self-immolative disulfide linker. The imaging arm is attached to the triazine splitter using click chemistry. The 15F-(PEG3)-moiety can be introduced either through the click reaction of 15F-(PEG3)-N3 with the propargylaminotriazine conjugate or 15F fluorination of the MsO-(PEG3)-triazole linked to the triazine linker in the drug conjugate, with BuN18F. For a feasibility study, we have successfully synthesized a prototype conjugate with a 15F-(PEG3)-arm, using biotin as TTM and SB-T-1214 as “warhead”. The corresponding “hot” chemistry experiment will be performed in collaboration with the Fowler laboratory at the Brookhaven National Laboratory, shortly. Results of this radio-synthesis and biodistribution analysis of the tumor-targeting drug conjugate in real-time will be reported elsewhere in due course. Again, fluorotaxoids would serve as powerful “warheads” for these “theranostics” as well.


In the past decade, carbon nanotubes (CNTs) have emerged as a unique and efficient vehicle for transport and delivery of drugs.191,192 Since functionalized carbon nanotubes (f-CNTs) were found
Asymmetric Bowtie Dendrimers As Vehicles for Tumor-Targeted Drug Delivery. Dendrimers are monodisperse, starburst polymers with a defined architecture originating from a central core. The high density of surface functional groups allows for polyvalent conjugation of TTM, drugs, and tracers, which has made dendrimers highly versatile vehicles for tumor-targeted drug delivery, and extensive studies have been done. Recently, an orthogonal diblock strategy for the generation of asymmetric bowtie dendrimers with a polyamido-amine (PAMAM) backbone and a cystamine core has been developed in our laboratory as well as by others. First, two dendrimers are fully functionalized with TTM, drugs or imaging agents. Second, each dendrimer is reductively cleaved to produce two half dendrons that can be asymmetrically joined via a bis-maleimido spacer. This synthetic approach offers unique advantages of selective modification and quantification of TTM, drugs, and tracers on each half-dendron prior to coupling. Therefore, this strategy should produce tumor-targeting nanoscale drug conjugates with well-defined drug loading and enhanced tumor specificity.

We have designed a novel asymmetric bowtie dendrimer scaffold bearing a 1,3,5-triazine splitter in the bis-maleimide linker module to introduce an imaging module (Figure 23). We set out to construct a prototype dendrimer conjugate wherein the vitamin TTM is biotin, the "warhead" is fluorotaxoid SB-T-12854 or SB-T-1214, and the imaging module is fluorine (19F for cold material and 18F for PET). In this prototype, an asymmetric bow-tie dendrimer, consisting of the G3 and G1 PAMAM dendrons, is used. The G3 dendron allows the conjugation of 16 biotin molecules, while the G1 dendron holds 4 taxoids. We have successfully synthesized key components and the final assembly to the designed conjugate is in progress. Synthesis and biological evaluation of this dendrimer-based tumor-targeted drug delivery system will be reported in due course.

Figure 21. Novel SWNT-based tumor-targeting "Trojan Horse" drug conjugate 42 and the CFM images of L1210FR cells treated with 42 incubated (A) before and (B) after the addition of GSH-ethyl ester. Image B clearly highlights the presence of fluorescent microtubule networks in the living cells generated by the binding of taxoid-fluorescein upon cleavage of the disulfide bond in the linker by either GSH or GSH-ethyl ester. (CFM images adapted from ref 184. Copyright 2008 American Chemical Society).

Figure 22. Novel SWNT-based tumor-targeting "Trojan Horse" drug conjugate 43 bearing a fluorine probe.

CONCLUDING REMARKS

This Perspective has covered the evolution of my research endeavor on fluorine chemistry not as a specialist in this field, but as an explorer of its interfaces with multidisciplinary fields...
in chemistry and biology. As mentioned, I was a synthetic chemist specializing in organometallic chemistry and homogeneous catalysis when I was brought into a very unique world of “fluorine chemistry” at the end of the 1970s. My laboratory started exploring the interface of fluorine chemistry and transition-metal catalysis, especially hydrocarbonylations and amidocarbonylation, which opened highly efficient synthetic routes to a variety of organofluorine compounds. Among them, CF₃-containing enantiopure amino acids were successfully applied to the enzyme inhibitor design, leading to the discovery of highly potent ACE inhibitor and enkephalin analogues. This line of research brought us into the field of medicinal chemistry. Also, trifluoromethacrylic acid was the key compound for the synthesis of trifluorothymine and this process was incorporated into the commercial process for the production of antiviral drug, trifluridine. We also introduced fluorine chemistry to the “β-lactam synthon method” and demonstrated the versatile and robust utility of fluorine-containing N-acyl-β-lactams as key intermediates to a library of fluorine-containing α-hydroxyl-β-amino acids and their peptides. Through expansion of this chemistry, we synthesized novel fluorine-containing taxoids and used them as excellent probes for the identification of bioactive conformations of paclitaxel and taxoids by means of ¹⁹F NMR in solution and solid phase. We also designed and developed a series of fluorine-containing taxoids, which are highly potent in vitro and in vivo, especially against multidrug resistant tumors through strategic incorporation of fluorine for potency and metabolic stability. In the past decade, my laboratory has been developing new and efficacious tumor-targeted drug delivery systems for new generation cancer chemotherapy. These novel drug delivery systems consist of tumor-targeting modules, mechanism-based self-immolative disulfide linkers and warheads. Fluorotaxoids are obviously highly potent “warheads” for these tumor-targeting drug conjugates. In order to monitor the tumor-specific drug delivery, internalization, drug release, and drug binding to the target protein, we have been exploring the potential of the “fluorine probe” approach based on ¹⁹F NMR as well as ¹⁸F-tracer for PET analysis in addition to fluorescence-based probes. The use of multifunctionalized SWNTs as a drug delivery system bearing multiple-warheads and multiple-targeting modules has shown highly promising results on the benefit of mass-delivery (“Trojan Horse” strategy) of anticancer drug molecules to cancer cells with high specificity. We have also been constructing another nanoscale drug delivery system based on asymmetric bowtie PAMAM dendrimers, bearing an imaging module, including ¹⁸F tracer for PET. Our future efforts will be concentrated on the design and synthesis of “tailor made nano-medicines” bearing imaging modules (for MRI, PET, fluorescence imaging and ¹⁹F NMR), which would enable us to perform diagnostics and therapy in the real time.

Thus, my laboratory has been exploring the interfaces of fluorine chemistry and multidisciplinary field of research involving medicinal chemistry, chemical biology, cancer biology, and molecular imaging. When we explore new areas of research, naturally we enjoy and struggle with many “first” findings, successes, and problems. This is, to my opinion, the most exciting thing as a researcher.

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Notes

The authors declare no competing financial interest.

Biography

Iwao Ojima received his Ph.D. in organic chemistry from the University of Tokyo. He is currently a University Distinguished Professor and the Director of the Institute of Chemical Biology & Drug Discovery at Stony Brook University. His broad research program can be summarized as synthetic organic chemistry at the biomedical interface, including medicinal chemistry, organofluorine chemistry, and chemical biology.

ACKNOWLEDGMENTS

The author acknowledges numerous research grants from the National Institutes of Health (National Institute of General Medical Sciences and National Cancer Institute), National Science Foundation, and ACS Petroleum Research Fund, which have been supporting his research for the last three decades. Generous support from the New York State Science and Technology Foundation, Author C. Cope Fund, New York State Office of Science, Technology and Academic Research (NYSTAR), Rhone-Poulenc Rorer (now Sanofi), Indena SpA,
Mitsubishi Chemical Corp., Ajinomoto Co., Inc, Central Glass Co. Ltd., Japan Halon, Inc. (now Toho F-Tech, Inc.), Tokyo Yuki Gosei Kogyo Co. Ltd., E. Merck, Darmstadt, and Fuji Chemical Ind. Co. Ltd. is acknowledged. The author also thanks his research group members at the Sagami Institute, who participated in the pioneering work on fluorine chemistry. He is grateful to all of his former and current graduate students, postdoctoral research associates, senior staff assistants, and staff of the Institute of Chemical Biology & Drug Discovery (ICB&DD), especially those who were engaged in the “fluorine chemistry” projects at Stony Brook, for their cooperation, support, and hard work, which made my endeavor fruitful. For the cover art, the author thanks Dr. Raphaël Geney for designing the background graphics, showing a tubulin-bound REDOR–Taxol molecule and a macrocyclic analogue, as well as Alexandra A. Athan for technical assistance to put together the images.

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