Radical Reactions

α,α-Difluoro-*H*-phosphinates: Useful Intermediates for a Variety of Phosphate Isosteres

Arnaud Gautier,* Goulnara Garipova, Carmen Salcedo, Sébastien Balieu, and Serge R. Piettre*

Dedicated to Professor Clayton Heathcock

Research efforts have long since established *H*-phosphinates **1** as important and valuable intermediates for the preparation of bioactive analogues of natural phosphates (Scheme 1).^[1] Despite various methodologies for their easy preparation, the synthesis of phosphonate **2**, phosphonothioates **3**, and phosphinates **4** remains an area of intense activity.^[2]

Moreover, major progress has been made in the chemistry of α,α -difluorophosphonates **7** since their introduction slightly more than two decades ago.^[3] Indeed, physicochemical studies have provided some rationale for the isosteric behavior of the above functional group to the phosphate group, and numerous applications have flourished.^[4] Among these, analogues targeting phospholipase C (PLC), purine nucleoside phosphorylase (PNP), and protein phosphotyrosine, phosphoserine, or phosphothreonine phosphatases have been reported.^[5] This functional group has also been success-

Angew. Chem. Int. Ed. 2004, 43, 5963-5967

DOI: 10.1002/anie.200460519

^[*] Dr. A. Gautier, G. Garipova, C. Salcedo, S. Balieu, Prof. Dr. S. R. Piettre Laboratoire des Fonctions Azotées et Oxygénées Complexes UMR 6014 CNRS, IRCOF, Université de Rouen 76821 Mont Saint Aignan (France) Fax: (+33) 2-3552-2971 E-mail: Arnaud.Gautier@univ-rouen.fr Serge.Piettre@univ-rouen.fr

Communications



Scheme 1. General structure of *H*-phosphinates **1**, related structures **2–4** obtained therefrom, and target compounds **5–9**.

fully used to mimic the phosphate group in nucleotide monophosphates and triphosphates: analogues of adenosine monophosphate, cyclic adenosine monophosphate, adenosine triphosphate, and adenosyl adenosine triphosphate, as well as structurally related potent inhibitors of the reverse transcriptase of human immunodeficiency virus (HIV), have been described in the literature.^[6]

Despite their potential both as a new class of isosteres of natural phosphates and as important intermediates in the synthesis of numerous α, α -difluorinated organophosphorus compounds, reports on fluorinated H-phosphinates 5 are scarce.^[7] The main reasons behind this situation are the synthetic problems underlying their preparation. Indeed, processes such as nucleophilic substitution of halodifluorinated centers (including the Arbuzov reaction) have long been known to be disfavored,^[8] and we and others have confirmed the sluggishness of phosphorus-centered radicals with respect to addition onto difluoroalkenes, when compared to their nonfluorinated analogues.^[9] In this context, we found that the sodium salt 6 of hypophosphorous acid behaves in a unique way, and have developed an efficient and practical preparation of α, α -difluoro-*H*-phosphinates. As shown below, these compounds can be easily transformed into difluorophosphonates 7, difluorophosphonothioates 8, and difluorophosphinates 9 (Scheme 1).

When a solution of 6 (0.2 M, 1.3 equiv) in methanol was refluxed for four hours with difluoroalkene 10-13 in the presence of tert-butyl peroxypivalate (TBPP) or tert-butyl 2ethylhexyl peroxycarbonate (TBEC)^[10] as a radical initiator, complete consumption of the substrate occurred and led to the formation of a single product in each case. A simple workup led to the isolation of products 14-17 in 75, 85, 83, and 80% yield, respectively, in the case of TBPP (Table 1). Later, we found that the Et₃B/O₂ system also allows a smooth conversion to take place.^[11] The α, α -difluoro-*H*-phosphinates are stable for weeks under standard conditions (room temperature and air). As expected, the ³¹P NMR spectra of compounds 14-17 displayed signals around 20 ppm with a P,H coupling constant of 590 Hz and P,F coupling constants of about 120 Hz. Similarly, the ¹⁹F NMR spectra were characterized by signals with two-bond F,P and three-bond F,H couplings, in accordance with the depicted structures. Additionally, FTIR signals were detected at 1250(s) and around 2370(m) cm⁻¹, which correspond to the P=O and P-H bonds, respectively.^[7]

Table 1: Structures of *gem*-difluoroalkenes and α , α -difluoro-*H*-phosphinates, and yields of the latter.

Entry	Substrate		Product		Yield [%]
1	₩ B CF2	10		14	75 ^[a]
2		11	СF ₂ -СF ₂ -Н ОН	15	85 ^[a,b]
3	tBu-CF2	12	<i>t</i> Bu—CF ₂ —P ^O ₄ H OH	16	83 ^[a,c]
4		13	OH CF2-P, H OH	17	80 ^[a]

[a] Using TBPP. [b] A complete conversion also occurred with $B(Et)_3/O_2$. [c] Isolated as a 4:1 mixture of diastereoisomers.

The weak phosphorus–hydrogen bond present in adducts **14–17** renders it prone to homolytic cleavage and highlights the possibility of generating yet another phosphorus-centered radical by treating these compounds with a radical initiator, and of a second radical addition on an alkene. Thus, interaction of a 1.6 m methanolic solution of **15** with methylenecyclohexane in the presence of a catalytic amount of radical initiator (TBPP) led to the formation of the expected α,α -difluorophosphinic acid, which was isolated in the form of its methyl ester **18** (diazomethane) in 53 % yield (Scheme 2). A similar two-step process involving sequential addition of 4-phenylbut-1-ene and diazomethane gave methyl difluorophosphinate **19** in 80 % yield.

The documented tautomeric equilibrium between an *H*-phosphinate and the corresponding phosphite led us to envision the possible transformation of α , α -difluoro-*H*-phosphinates into the corresponding bis-*O*,*O*-silylated difluoroal-



Scheme 2. Transformation of α,α-difluoro-*H*-phosphinates into difluoro-phosphonates, difluorophosphonothioates, and difluorophosphinates. a) 1. Alkene, **6** (0.3 equiv), C₆H₆, 55 °C, 18 h; 2. CH₂N₂, Et₂O, 12 h; b) bis(trimethylsilyl)acetamide (BSA, 3 equiv), CH₂Cl₂, 25 °C, 1 h; c) O₂, CH₂Cl₂, 25 °C, 0.5 h; d) S₈, CH₂Cl₂, 25 °C, 0.5 h; e) 1. but-3-en-2-one, BSA (3 equiv), CH₂Cl₂, 25 °C, 18 h, 2. CH₂N₂, Et₂O, 12 h; f) 1. pivalalde-hyde, BSA (5.5 equiv), CH₂Cl₂, 25 °C, 3 h, 2. CH₂N₂, Et₂O, 12 h.



kylphosphites, and the use of the nucleophilic P^{III} species.^[12] Accordingly, treatment of **16** with a threefold excess of trimethylsilyl chloride (TMSCl) and pyridine led to quantitative formation of the corresponding air-sensitive phosphites **20** and **21**, as demonstrated by a shift of the ³¹P NMR signals to about 130 ppm, and the lack of any onebond P,H coupling. Exposure of the bis-*O*,*O*-trimethylsilyl phosphites to oxygen or elemental sulfur quickly transformed these products into difluorophosphonates **22**.

Interactions between bissilylated difluoroalkylphosphites and electrophilic carbon atoms were exemplified through the 1,2- and 1,4-addition reactions (Abramov and Pudovic reactions, respectively).^[13] Thus, a degassed solution of 15, methyl vinyl ketone, BSA, and TMSCl in dichloromethane was stirred at room temperature, and the crude sample was sequentially subjected to a classical workup and treatment with diazomethane, which delivered difluorophosphinate 24 in 57% yield. Similarly, trimethylacetaldehyde (pivalaldehyde) reacted with 21 to furnish α,α -difluoro- α' hydroxyphosphinate 25 (64% yield). Few

examples of difluorophosphinates have so far been reported,^[2b,9a,14] and the present methodology constitutes the first general preparation of these compounds. It was particularly gratifying to note that the presence of the fluorine atoms did not prevent the P^{III} species from interacting with electrophiles.

The scope of the methodology was investigated by developing an application in the field of ribofuranosyl and cyclitol phosphates. Thus, reaction between 6 and ribofuranose derivatives 26 or 27 resulted in complete conversions, and the desired adducts 30 and 31 could be isolated in yields of 78 and 63%, respectively (Table 2). Similarly, cyclitol derivatives 28 and 29 reacted smoothly with 6 to deliver compounds 32 and 33 in good yields. These results are significant in light of the complete lack of reactivity of both diethyl phosphite and diethyl thiophosphite under similar conditions.^[15]

Table 2 clearly indicates the complete failure of any addition process in the case of phosphonyl and phosphonothioyl radicals, despite the demonstrated higher reactivity of the latter.^[9a-b,16] The involvement of a tautomeric P^{III} species of the radical generated from **6** has been ruled out by Beckwith,^[17] and the nature of this radical and those generated from phosphites and thiophosphites should therefore be similar. Additional physicochemical studies will be needed to explain the peculiar, but synthetically useful, behavior of **6** and shed light on the steric and electronic factors at play in this reaction.^[18] α,α -Difluoro-*H*-phosphinates **30** and **31** are useful intermediates to various compounds with potential applications in the fields of modified nucleotides and oligonucleotides (hence the antisense and

Table 2: Addition of hypophosphorous acid sodium salt, diethyl phosphite, and diethyl thiophosphite to difluoroalkenes **26–29**.

Entry	Substrate		Product		Phosphorus precursor	Conversion [%] (yield [%])
1 2 3	Pivo F ₂ C O F	26		30	NaOP(O)H $_2$ (EtO) $_2$ P(O)H (EtO) $_2$ P(S)H	100 (78 ^[a,b]) 0 0
4 5 5		27	4-CIBzO O H~p'-CF2 O (Et) ₃ NHO	31	NaOP(O)H ₂ (EtO) ₂ P(O)H (EtO) ₂ P(S)H	100 (63 ^[b,c]) 0 0
7 3 9	F ₂ C MeO ¹ , O O-C ₆ H ₁₀ , O O-C ₆ H ₁₀	28	$\begin{array}{c} H_{-} \overset{O}{P} = CF_{2} \\ NaO' \\ MeO'' \\ O = C_{6}H_{10} \end{array}$	32	NaOP(O)H ₂ (EtO) ₂ P(O)H (EtO) ₂ P(S)H	100 (76 ^[a]) 0 0
10 11 12	TBSO ¹¹ MeO	29	HO NaO ^C F ² CF ² TBSO ^{VI} OOMe	33	NaOP(O)H ₂ (EtO) ₂ P(O)H (EtO) ₂ P(S)H	100 (70 ^[a,d]) 0 0

[a] Using TBPP. [b] The addition proceeded with a diastereoselectivity greater than 95%. [c] Using $B(Et)_3/O_2$. [d] Isolated as a 3:7 mixture of diastereoisomers.

antigene strategies). For example, *H*-phosphinate **30** was easily transformed into difluorophosphonate **34** and difluorophosphonothioate **35** by treatment with TMSCl, pyridine, and the requisite O_2 or S_8 (Scheme 3).^[2a,15,19] In addition, *H*phosphinate **31** could easily be esterified by reaction with ribofuranose **36** in the presence of dicyclohexylcarbodiimide (DCC) and trifluoroacetic acid (TFA). The resultant *H*phosphinate was treated with sulfur and TMSCl in pyridine to



Scheme 3. Transformation of α , α -difluoro-*H*-phosphinates 30 and 31 into difluorophosphonate 34 and difluorophosphonothioates 35 and 37. a) Pyridine/TMSCl/O₂ or pyridine/TMSCl/S₈; b) DCC/TFA; c) pyridine/TMSCl/S₈. Bz = benzoyl, Piv = pivaloyl.

Angew. Chem. Int. Ed. 2004, 43, 5963-5967

www.angewandte.org

Communications

afford difluorophosphonothioic acid monoester **37** in 50% yield (a 1:1 mixture of two diastereoisomers at the phosphorus center).^[2a] This last result demonstrated the efficacy of the methodology in providing precursors of modified dinucleotides with new phosphorus-centered linkers.

In summary, radical addition of **6** on *gem*-difluoroalkenes constitutes a powerful method of constructing the previously unreported α, α -difluoro-*H*-phosphinates. This new functional group is easily and efficiently transformed into difluorophosphonates, difluorophosphonothioates, and difluorophosphinates. The methodology can be expected to have a major impact on the preparation of difluorophosphonyl, difluorophosphonothioyl, and difluorophosphinyl analogues of natural phosphates.

Experimental Section

General procedure for the synthesis of α,α -difluoro-*H*-phosphinates using *tert*-butyl peroxypivalate or *tert*-butyl 2-ethylhexyl peroxycarbonate as initiator: The requisite 1,1-difluoroalkene^[20-22] (1.0 equiv) and initiator (0.3 equiv) were added to a solution of hypophosphorous acid sodium salt monohydrate (0.2 M, 1.3 equiv) in degassed methanol. The solution was refluxed for 4 h under a nitrogen atmosphere and cooled to room temperature. The solution was poured into water and the aqueous phase was extracted with diethyl ether and lyophilized after separation of the layers. The solid was dissolved in aqueous NaHSO₄ (2.0 M) and extracted with dichloromethane. The organic layer was dried and evaporated to give a viscous oil.

Gram-scale synthesis of 31: Hypophosphorous acid sodium salt monohydrate (1.24 g, 1.7 mmol, 4.0 equiv) was added to a solution of 27 (1.0 g, 2.8 mmol, 1.0 equiv) in nondegassed methanol (25 mL) at room temperature in an open flask. Triethylborane (5 mL, 1M solution in hexane, 5.0 mmol, 5.0 equiv) was added under vigorous stirring, and the solution was stirred for 10 min. This operation was repeated twice. The fast addition of Et₃B solution was crucial. After the third addition of Et₃B, the solution was stirred for 1 h and then the solvent removed by evaporation. Water (50 mL) was added and the aqueous layer was extracted with ethyl acetate (20 mL). The aqueous layer was lyophilized and the crude solid was dissolved in aqueous triethylammonium carbonate (1.0 M, 30 mL). The solution was extracted twice with dichloromethane (30 mL); the organic layer was dried and evaporated to give a viscous oil (930 mg, 63 %). $[a]_{20}^{D} =$ + 33.9 (c = 1.08 in CHCl₃); ¹H NMR (200 MHz, CDCl₃): $\delta = 7.10$ (dd, J = 556 Hz, J = 6 Hz, 1 H), 7.92 (d, J = 8.7 Hz, 2 H), 7.33 (d, J = 8.7 Hz, 2H), 5.80 (d, J=3.9 Hz, 1H), 5.04 (t, J=3.9 Hz, 1H), 4.8 (m, 1H), 4.73 (d, J = 12 Hz, 1 H), 4.32 (dd, J = 12 Hz, J = 5.5 Hz, 1 H), 3.00 (q, J = 7.3 Hz, 6H), 2.70 (m, 1H), 1.50 (s, 3H), 1.26 (s, 3H), 1.26 ppm (t, J = 7.3 Hz, 9H); ¹³C NMR (50 MHz, CDCl₃): $\delta = 165.1$, 139.1, 130.9, 128.4, 128.1, 120.2 (td, J = 257 Hz, J = 121 Hz, CF₂), 112.5, 104.7, 79.4 (d, J = 7.6 Hz), 74.4, 64.6, 49.0 (td, J = 22.7 Hz, J = 13.6 Hz), 45.3, 26.3 (d, J = 6.1 Hz), 8.26 ppm; ³¹P NMR (81 MHz, CDCl₃): $\delta = 9.7$ ppm (dd, J = 92 Hz, J = 86 Hz); ¹⁹F NMR (188 MHz, CDCl₃, C₆F₆): $\delta =$ 51.5 (ddt, J = 301 Hz, J = 85 Hz, J = 10 Hz), 47.4 ppm (ddd, J =301 Hz, J = 92 Hz, J = 23 Hz); IR (NaCl): $\tilde{\nu} = 2986$, 1721, 1455, 1275, 1091 cm⁻¹. MS (MALDI, matrix: 2,4,6-trihydroxyacetophenone) m/z = 425.1 [M - 102.1].

37: Compound **24** (58 mg, 0.28 mmol, 1.5 equiv), DCC (118 mg, 0.76 mmol, 4.0 equiv), and trifluoroacetic acid (72 μ L, 0.95 mmol, 5.0 equiv) were added to a solution of **31** (100 mg, 0.19 mmol, 1.0 equiv) in degassed dichloromethane (3 mL). A white precipitate immediately formed and the slurry was stirred for 10 min. Powdered S₈ (200 mg, 6.2 mmol, 32.6 equiv), pyridine (1 mL, 12.2 mmol, 43.8 mmol), and TMSCl (1 mL, 7.9 mmol, 28.1 equiv) were then added sequentially. The reaction was stirred for an additional 15 min, then water (1 mL) was added, and the solution was filtered. The

filtrate was extracted with ice-cold aqueous HCl (0.1m, 10 mL). washed with aqueous triethylammonium carbonate (1.0 M), and evaporated. The residue was purified by flash chromatography over silica gel (AcOEt/MeOH/triethylamine 90:8:2) to give a viscous oil (70 mg, 50%) as a 1:1 mixture of two diastereoisomers (stereogenic phosphorus atom). A second chromatography procedure was performed which allowed partial separation of one of the diastereoisomers from the mixture. $[\alpha]_{25}^{D} = +17.2$ (c = 0.5 in CHCl₃); ¹H NMR (200 MHz, CDCl₃): $\delta = 8.00$ (d, J = 8.4 Hz, 2 H), 7.30 (d, J = 8.4 Hz, 2H), 5.80 (d, J=3.7 Hz, 1H), 5.10 (t, J=3.7 Hz, 1H), 4.90 (s, 1H), 4.82 (d, J = 5.8 Hz, 1 H), 4.70 (dd, J = 10.2 Hz, J = 3.6 Hz, 1 H), 4.54 (d, J = 5.8 Hz, 1 H), 4.40–3.90 (m, 4 H), 3.30 (s, 3 H), 3.10 (q, J = 7.3 Hz, 6H), 1.50 (s, 3H), 1.4 (s, 3H), 1.3-1.2 ppm (m, 16H); ¹³C NMR $(50 \text{ MHz}, \text{ CDCl}_3): \delta = 165.4, 139.2, 131.0, 128.6, 128.4, 112.5, 111.9,$ 109.0, 104.5, 85.4, 84.9, 81.6, 80.4(m), 75.0, 66.7, 65.1, 54.6, 47.4(m), 45.4, 26.8, 26.3, 26.3, 24.8, 8.3 ppm (CF₂ not observed); 31 P NMR (81 MHz, CDCl₃): $\delta = 61.5$ ppm (dd, J = 93 Hz, J = 90 Hz); ¹⁹F NMR (188 MHz, CDCl₃, C₆F₆): $\delta = 56.4$ (ddd, J = 306 Hz, J = 97 Hz, J =14 Hz), 50.0 ppm (ddd, J = 306 Hz, J = 97 Hz, J = 17 Hz); IR (NaCl): $\tilde{\nu} = 2987$, 2935, 1722, 1274, 1108, 1093, 1013 cm⁻¹; MS (MALDI, matrix: 2,4,6-trihydroxyacetophenone) m/z = 643.1[M-103.1].

Received: April 30, 2004

Keywords: phosphinates · phosphorus · radical reactions · synthesis design

- a) W. H. Parsons, A. A. Patchett, H. G. Bull, W. R. Schoen, D. Taub, J. Davidson, P. L. Combs, J. P. Springer, H. Gadebusch, B. Weissberger, M. E. Valiant, T. N. Mellin, R. D. Busch, J. Med. Chem. 1988, 31, 1772–1778; b) D. S. Karanewsky, M. C. Badia, D. W. Cushman, J. M. DeForrest, T. Dejneka, M. J. Loots, M. G. Perri, E. W. Petrillo, Jr., J. R. Powell, J. Med. Chem. 1988, 31, 204–212; c) N. S. Sampson, P. A. Bartlett, J. Org. Chem. 1988, 53, 4500–4503; d) M. T. Martin, T. S. Angeles, R. Sugasawara, N. I. Aman, A. D. Napper, M. J. Darsley, R. I. Sanchez, P. Booth, R. C. Titmas, J. Am. Chem. Soc. 1994, 116, 6508–6512; e) F. Tian, J.-L. Montchamp, J. W. Frost, J. Org. Chem. 1996, 61, 7373–7381.
- [2] a) J. Stawinski, A. Kraszewski, Acc. Chem. Res. 2002, 35, 952–960; b) W. Froestl, S. J. Mickel, G. Sprecher, P. J. Diel, R. G. Hall, L. Maier, S. Dietrich, V. Melillo, P. A. Baumann, R. Bernasconi, C. Gentsch, K. Hauser, J. Jaekel, G. Karlsson, K. Klebs, L. Maitre, C. Marescaux, M. F. Pozza, M. Schmutz, M. W. Steinmann, H. Riezen, A. Vassout, C. Mondadori, H. R. Olpe, P. C. Waldmeier, H. Bittiger, J. Med. Chem. 1995, 38, 3313–3331.
- [3] a) C. E. McKenna, P.-D. Shen, J. Org. Chem. 1981, 46, 4773–4776; b) G. M. Blackburn, D. A. England, F. Kolmann, J. Chem. Soc. Chem. Commun. 1981, 930–932; c) G. M. Blackburn, D. E. Kent, F. Kolkmann, J. Chem. Soc. Chem. Commun. 1981, 1188–1190.
- [4] a) R. D. Chambers, R. Jaouhari, D. O'Hagan, *Tetrahedron* 1989, 45, 5101-5108; b) R. D. Chambers, D. O'Hagan, R. B. Lamont, S. C. Jain, J. Chem. Soc. Chem. Commun. 1990, 1053-1054.
- [5] a) T. R. Burke, Jr., M. S. Smyth, M. Nomizu, A. Otaka, P. P. Roller, J. Org. Chem. 1993, 58, 1336–1340; b) S. F. Martin, Y.-L. Wong, A. S. Wagman, J. Org. Chem. 1994, 59, 4821–4831;
 c) D. B. Berkowitz, Q. Shen, J.-H. Maeng, Tetrahedron Lett. 1994, 35, 6445–6448; d) T. R. Burke, Jr., B. Ye, M. Akamatsu, H. Ford, Jr., X. Yan, H. K. Kole, G. Wolf, S. E. Shoelson, P. P. Roller, J. Med. Chem. 1996, 39, 1021–1027; e) D. B. Berkowitz, M. Eggen, Q. Shen, R. K. Shoemaker, J. Org. Chem. 1996, 61, 4666–4675; f) B. Ye, T. R. Burke, Jr., Tetrahedron 1996, 52, 9963–9970; g) M. N. Qabar, J. Urban, M. Khan, Tetrahedron 1997, 53, 11171–11178; h) T. Yokomatsu, H. Abe, T. Yamagishi,



K. Suemune, S. Shibuya, J. Org. Chem. **1999**, 64, 8413-8418; i) T. R. Burke, Jr., K. Lee, Acc. Chem. Res. **2003**, 36, 426-433.

- [6] a) G. M. Blackburn, D. E. Kent, J. Chem. Soc. Chem. Commun. 1981, 511-513; b) G. M. Blackburn, D. E. Kent, F. Kolkmann, J. Chem. Soc. Perkin Trans. 1 1984, 1119-1125; c) G. M. Blackburn, M.-J. Guo, S. P. Langston, G. E. Taylor Tetrahedron Lett. 1990, 31, 5637-5640; d) G. M. Blackburn, S. P. Langston, Tetrahedron Lett. 1991, 32, 6425-6428; e) J. Matulic-Adamic, N. Usman, Tetrahedron Lett. 1994, 35, 3227-3230; f) J. Matulic-Adamic, P. Haeberli, N. Usman, J. Org. Chem. 1995, 60, 2563-2569; g) S. G. Levvy, B. Wasson, D. A. Carson, H. B. Cottam, Synthesis 1996, 843-846; h) C. J. Hamilton, S. M. Roberts, A. Shipitsin, Chem. Commun. 1998, 1087-1088; i) C. J. Hamilton, S. M. Roberts, J. Chem. Soc. Perkin Trans. 1 1999, 1051-1056.
- [7] a) H. J. Emeleus, R. N. Haszeldinze, R. C. Paul, J. Chem. Soc. 1955, 563-574; b) A. B. Burg, J. E. Griffiths, J. Am. Chem. Soc. 1961, 83, 4333-4337; c) A. Golovanov, I. G. Maslennikov, A. N. Lavrent'ev, Zh. Obshch. Khim. 1988, 58, 1525-1529.
- [8] a) L. Z. Soborovskii, N. F. Baina, *Zh. Obshch. Khim.* 1959, 29, 1142–1143; b) D. J. Burton, R. M. Flynn, *J. Fluorine Chem.* 1977, 10, 329–332; c) D. J. Burton, R. M. Flynn, *Synthesis* 1979, 615.
- [9] a) S. R. Piettre, *Tetrahedron Lett.* **1996**, *37*, 2233–2236; b) T. F. Herpin, W. B. Motherwell, B. P. Roberts, S. Roland, J.-M. Weibel, *Tetrahedron* **1997**, *53*, 15085–15100; c) J. Kovensky, M. McNeil, P. Sinay, J. Org. Chem. **1999**, *64*, 6202–6205.
- [10] For the synthesis of *tert*-butyl peroxypivalate (TBPP), see: P. D. Bartlett, E. P. Benzing, R. E. Pincock, J. Am. Chem. Soc. 1960, 82, 1762–1768; TBPP has a half-life of 5 h at 60 °C. Tert-butyl 2-ethylhexyl peroxycarbonate (TBEC) is commercially available and has a half-life of 10 h at 77 °C and 1 h at 95 °C.
- [11] a) S. Deprele, J.-L. Montchamps, J. Org. Chem. 2001, 66, 6745–6755; b) A. Gautier, G. Garipova, O. Dubert, H. Oulyadi, S. R. Piettre, *Tetrahedron Lett.* 2001, 42, 5673–5676.
- [12] a) J. Jankowska, A. Sobkowska, J. Cieslak, M. Sobkowski, A. Kraszewski, J. Stawinski, D. Shugar, *J. Org. Chem.* **1998**, *63*, 8150–8156; b) M. A. Maier, A. P. Guzaev, M. Manoharan, *Org. Lett.* **2000**, *2*, 1819–1822.
- [13] a) A. N. Pudovik, I. V. Konovalova, Zh. Obshch. Khim. 1959, 29, 3342–3346; b) V. S. Abramov, Dokl. Akad. Nauk SSSR 1950, 73, 487–489; c) R. Engel, Handbook of Organophosphorus Chemistry, Marcel Dekker, New York, 1992.
- [14] J. Ong, D. I. B. Kerr, W. Froestl, Eur. J. Pharmacol. 1999, 374, 351–354.
- [15] A considerable amount of time and effort was spent attempting the addition of phosphonyl and phosphonothioyl radicals onto difluoroalkenes related to 26 and 27, without any success; see: C. Lopin, A. Gautier, G. Gouhier, S. Piettre, J. Am. Chem. Soc. 2002, 124, 14668–14675.
- [16] A. N. Pudovik, I. V. Konovalova, Zh. Obshch. Khim. 1959, 29, 3342–3346.
- [17] A. L. Beckwith, Aust. J. Chem. 1972, 25, 1887-1905.
- [18] J. M. Tedder, J. C. Walton, *Acc. Chem. Res.* **1976**, *9*, 183–191, and references therein.
- [19] This type of furanose derivative has recently been investigated:
 a) A. H. Butt, J. M. Percy, N. S. Spencer, *Chem. Commun.* 2000, 1691–1692;
 b) T. Murano, S. Muroyama, T. Yokomatsu, S. Shibuya, *Synlett* 2002, 1657–1660;
 c) T. Murano, Y. Yuasa, S. Muroyama, T. Yokomatsu, S. Shibuya, *Tetrahedron* 2003, *59*, 9059–9073.
- [20] The 1,1-difluoroalkenes 10–13, 26, 27, and 29 described in this paper were obtained from the corresponding ketone in yields ranging from 55 to 65% by either the method of Wheaton and Burton or the modification of the Motherwell procedure by Serafinowski and Barnes^[21] For instance, a 6.0-g batch of compound 27 was prepared according to reference 21a. Sub-

Angew. Chem. Int. Ed. 2004, 43, 5963-5967

- [21] a) G. A. Wheaton, D. J. Burton, J. Org. Chem. 1983, 48, 917– 927; b) P. J. Serafinowski, C. L. Barnes, Tetrahedron 1996, 52, 7929–7938.
- [22] J. S. Sabol, J. R. McCarthy, *Tetrahedron Lett.* **1992**, *33*, 3101–3104.