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THE PREPARATION AND CHARACTERIZATION OF SOME FLUORINATED α -AMINOARYLMETHANEPHOSPHONIC ACIDS

Donovan St. C. Green^a; Ulrike Gruss^b; Gerhard Hägele^b; Harry R. Hudson^a; Lars Lindblom^a; Max Pianka^a

^a School of Applied Chemistry, University of North London, London, United Kingdom ^b Institut für Anorganische Chemie und Strukturchemie I, Heinrich Heine-Universität Düsseldorf, Düsseldorf, Germany

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THE PREPARATION AND CHARACTERIZATION OF SOME FLUORINATED α-AMINOARYLMETHANEPHOSPHONIC ACIDS

DONOVAN ST. C. GREEN,† ULRIKE GRUSS,‡,§ GERHARD HÄGELE,‡,| HARRY R. HUDSON,†, || LARS LINDBLOM† and MAX PIANKA†

†School of Applied Chemistry, University of North London, 166-220 Holloway Road, London N7 8DB, United Kingdom; ‡Institut für Anorganische Chemie und Strukturchemie I, Heinrich Heine-Universität Düsseldorf, Universitätsstraße 1, D-40225 Düsseldorf, Germany

Dedicated to Prof. Dr. M. Baudler, Köln

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 α -Aminoarylmethanephosphonic acids have been prepared with a range of fluoro, fluoroalkyl, or fluoroalkoxy substituents in the benzene ring (4-F, 3-F, 2-F, 3,4-F₂, F₅, 4-CF₃, 3-CF₃, 4-CF₃O, and 3-CF₃O). These compounds have relatively low aqueous solubility and their NMR spectra (¹H, ¹³C, ³¹P and ¹⁹F) were therefore recorded in D₂O in the presence of an excess of alkali. Under these conditions, the ring substituents appear to have little effect on δ_P (15-18 ppm), or on the ¹H and ¹³C parameters for the benzylic group (α -CH), which are mainly in the ranges observed for other types of α -aminoarylmethanephosphonic acids under alkaline conditions ($\delta_{\rm H}$ 3.8-4.0 ppm, $^2J_{\rm PH}$ 15.3-16.5 Hz; $\delta_{\rm C}$ 57-58 ppm, ${}^{1}J_{PC}$ 128-132 Hz). For those examples with fluorine in the ortho position (i.e., the 2-fluoro and pentafluoro derivatives) a slightly higher field chemical shift was observed for the benzylic carbon atom (δ_c 50-51 ppm). In the fast-atom bombardment mass spectra, pseudo-molecular ions, MH⁺, and ions resulting from the elimination of phosphorous acid [MH - H₃PO₃]⁺, provide a further useful means of chararacterization for these compounds.

Key words: Fluorinated aminoarylmethanephosphonic acids, dialkyl N-diphenylmethylaminoarylmethanephosphonates, NMR spectroscopy, FAB mass spectrometry.

INTRODUCTION

Aminophosphonic acids are of widespread interest as biologically active molecules, 1-7 as complexing agents for metal ions, 8 and as building blocks in the synthesis of phosphonopeptides. 9,10 α -Aminoarylmethanephosphonic acids and their esters, the phosphonic analogues of α -C-aryl-substituted glycine, have been reported by a number of workers, 11 although derivatives with fluorine or fluorinated substituents in the aromatic ring are uncommon. Such compounds are of interest because of the modified properties (pKa, lipoidal solubility, etc.), which the introduction of fluorine may engender. The only previous report concerning α -aminofluorophenylmethanephosphonic acids is that of L. Maier et al., 12 who prepared α -amino-4-fluorophenylmethanephosphonic acid and α -amino-3-(4'-fluorophenoxy)phenylmethanephosphonic

[§]Part of forthcoming dissertation from U. Gruss, Heinrich-Heine-Universität Düsseldorf 1996. ||Authors for correspondence.

acid via addition of dialkyl phosphites to N-phenylmethyl protected imines (followed by hydrogenation and hydrolysis). The same method was used for the synthesis of diethyl N-phenylmethylamino-2,3-difluoromethylenedioxyphenylmethanephosphonate, which was converted into the corresponding diethyl aminophosphonate. Other derivatives related to α -aminofluorophenylmethanephosphonic acids are the N-protected diethyl amino-4-fluorophenylmethanephosphonates with chiral N-1-phenylethyl and N-2-methoxy-1-phenylethyl groups, prepared by the imine method; and the hydrobromide salt of diphenyl amino-4-fluorophenylmethanephosphonate form N-phosphoryl- and N-thiophosphorylaminoarylmethanephosphonates synthesized by amidoalkylation of triphenyl phosphite with the corresponding amides). But in all these cases conversion to the free α -aminoarylmethanephosphonic acids was not reported. (Another fluorinated derivative, known in the literature, is α -hydroxy-amino-4-fluorophenylmethanephosphonic acid, prepared by hydroxyamination of diethyl aroylphosphonate.)

In this paper we describe the preparation of a range of fluorinated α -aminoaryl-methanephosphonic acids and we discuss their characterization by NMR spectroscopy (1 H, 13 C, 31 P and 19 F) and FAB mass spectrometry.

RESULTS AND DISCUSSION

Preparations

Preparations were carried out by one or more procedures, involving reduction of the corresponding α -oximinophosphonates¹⁹ by zinc in formic acid²¹ (method A in Scheme I), amidoalkylation of phosphorus trichloride by a one-step process (method B with the variations a) and b) in Scheme II),³⁰ or a two-step process (method B with the variation c) in Scheme III,^{32,33} or the addition of dialkyl phosphite to an N-protected Schiff base derived from diphenylmethylamine, followed by hydrolysis (method C in Scheme IV).^{11,34} These methods have been applied previously in the synthesis of aminoalkanephosphonic acids of various types and were found to be suitable in the present work.

Starting materials for the reaction pathway A, the amination of diethyl aroylphosphonates (Scheme I), are fluorinated benzoic acids 1, which were converted nearly quantitatively to aroyl chlorides 2 by reaction with an excess of thionyl chloride and catalytic amounts of dimethylformamide at 75°C. 18a,b Arbuzov reaction with a slight excess of triethyl phosphite at temperatures below 40°C, followed by distillation at oil pump pressure, led to diethyl aroylphosphonates 3.15,19 These compounds 3 are sensitive to traces of acids, bases and weak nucleophiles (e.g. water, amines); decomposition of the P-C bond occurs easily. They were therefore converted immediately into the more stable diethyl hydroxyiminoarylmethanephosphonates 4 by reaction with hydroxylamine hydrochloride and pyridine in dry ethanol. These compounds 4 were obtained in the form of highly viscous oils or waxy solids (each consisting of a mixture of E and Z isomers, in which the E form predominates) after extractive work-up and drying under vacuum at room temperature. Purification of the diethyl hydroxyiminoarylmethanephosphonates 4 by distillation was not possible, since decomposition occurs at temperatures above 60°C. 15,19,20 Reduction of the oximes 4 to the corresponding amines 5 was carried out by the use of zinc dust in 99%

$$R_{f} = \frac{1.0}{0.00} + \frac{1.0}{0.00$$

SCHEME 1 Synthesis of α -aminoarylmethanephosphonic acids 6. Method A: Amination of diethyl aroylphosphonates. In the text hydrochlorides of 6 are designated 6'.

formic acid at temperatures below 65°C. ²¹ After filtration and evaporation the remaining oils were used for the following hydrolysis step without further purification. They have not been investigated by ¹H NMR spectroscopy, but comparison with similar processes described in the literature ^{21,22,23a,23b} and ³¹P{¹H} NMR spectroscopy showed that in all cases the oils consisted of mixtures 5 of diethyl aminoarylmethanephosphonates, aminoarylmethanephosphonic acids and analogous N-formylated derivatives. Heating of these products with concentrated hydrochloric acid (6–8 h at 100°C) led to hydrolysis of the phosphonate moiety (and cleavage of the N-formyl groups) and yielded α -aminoarylmethanephosphonic acid hydrochlorides 6′. After evaporation of the solvent and repeated dissolving of the remaining oils in ethanol/ water mixtures followed by evaporation, the α -aminoarylmethanephosphonic acids 6 precipitated during heating in boiling water as fine white crystalline solids. ¹⁹

Method A for the synthesis of fluorinated α -aminoarylmethanephosphonic acids 6 produced very clean microcrystalline products, so that additional recrystallization steps were not necessary. This reaction pathway can be applied for α -aminoarylmethanephosphonic acids 6 with fluorinated substituents at any position in the aromatic system. A negative point is that the reaction procedure involved six steps with very long reaction times in some cases. Only the products of the first and the second step have to be isolated and purified (aroyl chlorides 2 were obtained in nearly

No.	R _f	No.	R _f	R'
7, 6 b	3-F	8 ab	3-F	C ₆ H ₅
7, 6 e	4-CF ₃	8 ae	4-CF ₃	C ₆ H ₅
7, 6 f	3-CF ₃	8 af	3-CF ₃	C ₆ H ₅
7, 6 g	4-CF ₃ O	8 ag	4-CF ₃ O	C ₆ H ₅
7, 6 h	3-CF ₃ O	8 ah	3-CF ₃ O	C ₆ H ₅
		8 bb	3-F	СНз
		8 be	4-CF ₃	CH ₃
		8 bf	3-CF ₃	CH ₃
		8 bg	4-CF ₃ O	CH ₃
		8 bh	3-CF ₃ O	CH ₃

SCHEME II Synthesis of α -aminoarylmethanephosphonic acids 6. Method B: Amidoalkylation of phosphorus trichloride in one-pot procedures: variation a): with benzamide; variation b): with acetamide.

quantitative yields, while the preparation of the moisture sensitive and high boiling diethyl aroylphosphonates 3 determined the yield of the whole reaction pathway). For the following steps, each product can be used in its crude form.

Other methods for the synthesis of aminoalkanephosphonic acids and aminophosphonates reported in the literature differ only in the techniques for reduction of the dialkyl hydroxyiminoalkanephosphonates. They include the use of a) aluminum amalgam, 19,24,25 b) Zn/Cu couple, 26 c) B_2H_6 gas in THF²⁴ d) BH_3*THF , 27 e) TiCl₃/NaBH[C(O)CH₃]₃ in acetate buffered solution, 28 and f) catalytic hydrogenation. 20,25,29 In the present work the use of zinc dust in formic acid²¹ was chosen as this required only simple reactants, avoided working with gases (method c)²⁴ or f)^{20,25,29}), the preparation of reduction catalysts, the generation of the reducing species in a separate preceding reaction step (method a)¹⁹ or b)²⁶), or employing expensive complex hydrides (method d)²⁷ or e)²⁸).

A one-pot synthesis for fluorinated α -aminoarylmethanephosphonic acids 6 is the amidoalkylation of phosphorus trichloride in acetic acid solution (method B, variations a) and b) in Scheme II). For this reaction an excess (up to 40%) of a fluorinated benzaldehyde 7 was allowed to react with a mixture of equimolar amounts of benzamide a) or acetamide b) and phosphorus trichloride in acetic acid to give the N-acylaminoarylmethanephosphonic acid 8a,b. After evaporation of the solution the remaining oil was heated with an excess of concentrated hydrochloric acid to remove the N-protection group to form the aminophosphonic acid hydrochloride 6', from which the α -aminoarylmethanephosphonic acid 6 was precipitated by addition of

SCHEME III Synthesis of α -aminoarylmethanephosphonic acids 6. Method B: Amidoalkylation of phosphorus trichloride via a two-step reaction.

pyridine in methanolic solution (pH 4) and cooling to -5 to 0°C. The α -aminoarylmethanephosphonic acids 6 were obtained as amorphous powders which were purified by recrystallization from alcohol/water solutions.³⁰

The application of the different amides in this type of reaction showed different advantages: with benzamide the formation of the N-acylated aminophosphonic acids 8a, and the N-deprotecting reaction were much quicker than with the use of acetamide. The cleavage of the N-benzoyl group was observable by the precipitation of benzoic acid, while the final removal of the N-acetyl group had to be determined by ${}^{31}P\{{}^{1}H\}$ NMR spectroscopy. Nevertheless, the amidoalkylation reaction with acetamide leads in some cases to much better yields for α -aminoarylmethanephosphonic acids 6 (especially for $R_f = 4$ -CF₃ 6e) than the analogous reaction with benzamide.

Method B (Scheme II) makes it possible to synthesize α -aminoarylmethanephosphonic acids 6 in one step without the use of special apparatus and without purification of intermediate products. Disadvantages are the need to use an excess of the expensive fluorinated benzaldehydes 7, long reaction times under drastic acid conditions to split off the N-protection groups, and long crystallization times to precipitate the α -aminoarylmethanephosphonic acids 6 from the complex reaction mixtures. By this type of amidoalkylation reaction in a one-pot procedure only meta- and parasubstituted benzaldehydes 7 can be converted to α -aminophosphonic acids 6 in moderate to good yields; ortho-substituted compounds do not react in this way (as shown with $R_f = 2\text{-CF}_3$, $2\text{-CF}_3\text{O}$).

The third variation of method B for the amidoalkylation of phosphorus trichloride is a two-step reaction (Scheme III). First the fluorinated benzaldehyde 7 was converted into an N,N'-arylidenebis-(acetamide) 9 by heating it with an excess of acetamide, followed by crystallization from methanol.³² After isolation of this compound it was used for the amidoalkylation reaction under moderate conditions at $0-10^{\circ}$ C with equimolar amounts of phosphorus trichloride in acetic acid solution, followed

No.	R _f	N o	Rf	R
7, 10, 6 a	4-F	11 aa	4-F	СНЗ
7, 10, 6 b	3-F	11 ae	4-CF ₃	CH ₃
7, 10, 6 c	2-F	11 ba	4-F	C ₂ H ₅
7, 10, 6 d	3,4-F ₂	11 bb	3-F	C ₂ H ₅
7, 10, 6 e	4-CF ₃	11 bd	3,4-F ₂	C ₂ H ₅
7, 10, 6 f	3-CF ₃	11 be	4-CF ₃	C ₂ H ₅
7, 10, 6 g	4-CF ₃ O	11 bf	3-CF ₃	C ₂ H ₅
7, 10, 6 h	3-CF ₃ O	11 bg	4-CF ₃ O	C ₂ H ₅
7, 10, 6 i	F ₅	11 ca	4-F	C ₆ H ₅
		11 cb	3-F	C ₆ H ₅
		11 cd	3,4-F ₂	C ₆ H ₅
		11 ce	4-CF ₃	C ₆ H ₅
		11 cf	3-CF ₃	C ₆ H ₅
		11 cg	4-CF ₃ O	C ₆ H ₅

SCHEME IV Synthesis of α -aminoarylmethanephosphonic acids 6. Method C: Addition of dialkyl phosphites to N-diphenylmethyl protected imines.

by heating at 100°C for completion of the reaction. Work-up and N-deprotection of the N-acetylaminophosphonic acid **8b** was achieved in the same way as described for the one-pot method. 32,33,42

This two-step procedure is especially suitable for aldehydes sensitive to oxidation, because these compounds were converted first into more stable derivatives, the N,N'-arylidenebis-(acetamides), before the amidoalkylation reaction under drastic acid conditions starts.^{32,33,42} This additional step is not a disadvantage, since N,N'-arylidenebis-(acetamides) 9 were obtained in high yields in a crystalline state; they can be employed without purification for the further synthesis, and only equimolar amounts of the expensive fluorinated benzaldehydes 7 are necessary.

Method B, variation c) (Scheme III) was used for one example, namely 4-fluorobenzaldehyde 7a, and the yield of α -amino-(4-fluorophenyl-)methanephosphonic acid 6a obtained was noticeably higher than yields for fluorinated α -aminoarylmethanephosphonic acids 6 according to the one-pot procedures (method B, variations a) and b) in Scheme II).

The third method C (Scheme IV) for the preparation of α -aminoarylmethane-phosphonic acids 6 uses the addition of substances with P—H bonds to N-protected imines; three steps lead to the desired products. The intermediate compounds, N-

substituted imines and N-substituted dialkyl aminophosphonates, are obtained in high to quantitative yields.

In this study the diphenylmethyl group has been chosen for N-protection.³⁴ Fluorinated benzylidene-1,1-diphenylmethylamines 10 were synthesized in two different ways: In the first route mixing of equimolar amounts of fluorinated benzaldehydes 7 with diphenylmethylamine in dry dichloromethane or dry diethyl ether, and addition of anhydrous potassium carbonate led to the imines 10 under very mild conditions. After filtration and evaporation of the solvents the imines 10 were obtained in the form of waxy solids in nearly quantitative yields.^{11,42} In the second variation the compounds 10 were prepared by heating equimolar amounts of benzaldehydes 7 with diphenylmethylamine. The use of dry solvents, the addition of drying agents and catalysts and/or methods to remove water as formed (e.g. Dean and Stark apparatus) were not necessary, because it separated from the oily reaction mixture, and was removed by dissolving the oil in dichloromethane and the addition of anhydrous sodium sulfate.

These imines 10 were converted into α -aminoarylmethanephosphonic acids 6 by a one-pot method: they were stirred with equimolar amounts of dialkyl phosphite (R = CH₃, C₂H₅) at 120–140°C (30–60 min) to yield *in situ* dialkyl N-diphenylmethylaminoarylmethanephosphonates 11. These oils (without isolation or purification) were heated with concentrated hydrochloric acid at 100°C for N-deprotection and phosphonate hydrolysis at the same time. After extraction of the by-product, diphenylmethyl chloride, with toluene and evaporation of the aqueous solution the remaining oils were dissolved in methanol and the α -aminoarylmethanephosphonic acids 6 were precipitated by the addition of propylene oxide at 40°C.^{11,34,35,42,43}

For isolation of dialkyl or diphenyl N-diphenylmethylaminoarylmethanephosphonates 11 (R = CH₃, C₂H₅, C₆H₅) longer reaction times of the imines 10 with the phosphites were required. The oily reaction mixtures crystallized after dissolving them in alcohols followed by addition of diethyl ether at 0°C, to give waxy white solids (R = CH₃, C_2H_5) or fine, white crystalline powders (R = C_6H_5) in high yields. These aminophosphonates 11 were isolated and investigated by NMR spectroscopy in this work for the first time. They represent an air stable storage form for other aminophosphonic acid derivatives. It is possible to convert them to α -aminoarylmethanephosphonic acids 6 as described above, to N-diphenylmethylaminoaryl methanephosphonic acids or to dialkyl aminoarylmethanephosphonates. N-Diphenylmethylaminoarylmethanephosphonic acids can be obtained by transforming the N-protected dialkyl aminoarylmethanephosphonates 11 first to the corresponding bis-(trimethylsilyl) aminophosphonates by reaction with an excess of bromotrimethylsilane in chlorinated hydrocarbons at room temperature followed by cleavage of the phosphonic ester groups with water at room temperature.¹² Dialkyl aminoarylmethanephosphonates are prepared by cleaving the N-protection group (without hydrolysis of the phosphonate moiety) by catalytic hydrogenation with Pd/charcoal catalyst in alcoholic solution 12,34,36,43 or by reaction in acid solutions e.g. HBr in acetic acid^{16,17,37} or HBr in alcoholic solution^{9d,38} under controlled conditions at room temperature, followed by immediate work-up. These derivatives are especially important for phosphonopeptide synthesis.

The α -aminoarylmethanephosphonic acids 6a-6i were obtained in all cases as anhydrous, microcrystalline solids and were characterized by elemental analysis, NMR spectroscopy and mass spectrometry. Yields and principal NMR parameters are shown in Table I.

NMR Spectroscopy

Because of the low solubility of the α -aminoarylmethanephosphonic acids 6 in water, NMR parameters were determined in the presence of an excess of alkali (NaOD or KOD in D₂O). Under these conditions full deprotonation takes place: the phosphonate is in the bi-negative PO₃⁻ form and the amino group in the neutral NH₂ form. It has been shown elsewhere for simple α -aminoalkanephosphonic acids, that removal of the proton from nitrogen $(NH_1^+ \rightarrow NH_2)$ has the effect of moving the ³¹P chemical shift to lower field than would be observed for the same aminophosphonic acids, in D_2O alone.³⁹ While a similar effect may be expected for the α -aminoarylmethanephosphonic acids, our present results are confined to measurements in alkaline conditions only. Under these conditions the ³¹P chemical shifts for the various fluorinated compounds showed little variation with the substituents present. All were found to lie in the region of δ_P 17-18 ppm (relative to 85% H₄PO₄), except in the case of the pentafluoro derivative 6i for which the chemical shift was at a slightly higher field (δ_P 15.3 ppm). These chemical shifts are, as a group, all slightly upfield from those reported for other non-fluorinated α -aminoarylmethanephosphonic acids 6 (X = H, 4-MeO, 4-Me, 4-Et, 2-HO, 3-MeO, 4-HO), for which δ_P in all cases was found to lie in the range 18-20 ppm in NaOD/D₂O. 11,40 It is interesting that α -amino-4nitrophenylmethanephosphonic acid 6 ($X = 4-NO_2$), with the strongly electron-attracting nitro group present in the para position, also exhibits a chemical shift ($\delta_{\rm p}$ 16.6 ppm) in the same region as the fluorinated compounds.⁴⁰ The reason for this uniformly upfield shift in δ_P when electron-attracting substituents are present in the ring is obscure but the effect is small and it is likely that the shielding of phosphorus is dominated more by the electron density associated with the oxygen atoms of the phosphonate anion, PO₃²⁻, rather than the ring substituents. Conformational and hydrogen-bonding influences may also be involved. Coupling to fluorine broadens the ³¹P{¹H} NMR signals in general but distinct doublets were observed for the monofluorinated derivatives with 4-, 5-, or 6-bond coupling via the aromatic ring (${}^4J_{PF}$ 4.1 Hz, ${}^5J_{PF}$ 1.4 Hz and ${}^6J_{PF}$ 3.9 Hz in 6a, 6b and 6c, respectively).

¹H and ¹³C NMR parameters (see Table I and IX) for the aryl-substituted CH groups of α-aminoarylmethanephosphonic acids 6 also lie, in most cases, within fairly narrow ranges ($\delta_{\rm H}$ 3.8–4.0 ppm, $^2J_{\rm PH}$ 15.3–16.5 Hz; $\delta_{\rm C}$ 57.5–58.3 ppm, $^2J_{\rm PC}$ 128–132 Hz) and are not significantly different from those reported for the non-fluorinated compounds. ^{11,40} The only compounds whose ¹H and ¹³C chemical shifts for the CH group lie outside these ranges are those having one or more substituents ortho to the arylmethane carbon 6c, 6i, where the inductive effect of fluorine might be expected to have the greatest influence. Even in these cases, the ¹H chemical shift is only slightly downfield, at ca. 4.2 ppm although the ¹³C chemical shift, surprisingly, is clearly up-field at 50–51 ppm. There is no significant difference in the magnitude of coupling to phosphorus in either case. It is interesting to note that very similar chemical shifts are shown by α-amino-2-hydroxyphenylmethanephosphonic acid (6, X = 2-HO) ($\delta_{\rm H}$ 4.3 ppm, $\delta_{\rm C}$ 50.23 ppm), suggesting that the effects observed are steric in origin rather than electronic.

¹⁹F NMR chemical shifts for these molecules are similar to those for the starting materials 1 and 7. The proton-decoupled ¹⁹F NMR spectra appear as broad singlets for the trifluoromethyl 6e, 6f, and trifluoromethoxy 6g, 6h compounds and as un-

TABLE I

³¹P{¹H} and ¹H NMR data, yields and melting points of fluorinated α-aminoarylmethanephosphonic acids 6. ³¹P{¹H} NMR: 10% 1m KOH/D₂O, (**D**), or NaOD(xs)/D₂O, (**L**), vs. 85% H₃PO₄ ext. ¹H NMR: 6% 1m KOH/D₂O, (**D**), or NaOD(xs)/D₂O, (**L**), vs. TSP—d₄—Na, δ [ppm], Mult., ^{*}J_{XY} [Hz]

No.	δ _H (L) α-CH	δ _H (D) α-CH	δp (L)	δp (D)	yields [%]	mp. [^O C]
6a	3.86, d,	3.80, d,	17.03, d,	18.50, d,	63.3 ^a , 59.0 ^d ,	278-282
	2 _{JpH} 15.4	2 _{JPH} 15.3	6 _{JPF} 3.0	6 _{JPF} 3.9	55e	
6b	-	3.93, d,	•	18.03, d, 5 _{JPF} 1.4	18.6 ^b , 25.5 ^c	289-290
6c		4.13, d, 2 _{JPH} 15.9			43.3 ^a , 28 ^e	270-273
6d	-	3.79, d, ² J _{PH} 15.7		17.86, dd, 6 _{JPF} 4.2, ⁵ JPF 1.6	56.1 ^a	280-282
6e		3.91, d, 2 _{JPH} 16.5		17.71, q, ⁷ Jp p 2.1	40.9 ^b , 74.8 ^c , 46 ^e	287-289
6f	-	4.01, d, ² J _{PH} 15.8		15.03, s	41.7 ^b , 35.6 ^c	253-255
6g	-	3.85, d, ² J _{PH} 15.8	•	18.12, s	31.1 ^b , 45.1 ^c ,	305-307
6h				15.99, s	25.9 ^b , 35.0 ^c , .43 ^e	277-280
6i		-	15.31, m	-	18 ^e	268-270

^a Preparation by method A (Amination of diethyl aroylphosphonates), SCHEME I.

resolved multiplets in other cases due to coupling with phosphorus and other fluorine atoms if present (in 6a, 6b, 6c, 6d and 6i).

Detailed NMR data are quoted under the experimental section.

Mass Spectroscopy

As reported for other classes of aminophosphonic acids, ^{6,10b,41} FAB MS (with a primary beam of xenon atoms) or LSIMS (using fast caesium ions) provide additional

b Preparation by method B, variation a) (Amidoalkylation of phosphorus trichloride with benzamide, one-pot procedure), SCHEME II.

^c Preparation by method B, variation b) (Amidoalkylation of phosphorus trichloride with acetamide, one-pot procedure), SCHEME II.

d Preparation by method B, variation c) (Amidoalkylation of phosphorus trichloride with arylmethylidenebisacetamide, two-step method), SCHEME III.

^e Preparation by method C (addition of dialkyl phosphites to N-protected imines, followed by hydrolysis), SCHEME IV.

TABLE II
Characteristic ions in the FAB mass spectra of α -aminoarylmethanephosphonic
acids 6a—6i

No.	Methoda	Matrix		m/z [%]		
		(Z)	[2M + H] ⁺	[MH + Z] ⁺	[MH] ⁺	[MH - H ₃ PO ₃]+
6a	Cs+	glycerol	411 (17.6)	298 (8.9)	206 (33.7)	124 (100)
6b	Cs+	glycerol	411 (3.8)	298 (5.7)	206 (20.1)	124 (100)
6c	Cs+	glycerol	-	298 (5.0)	206 (13.5)	124 (71.2)
6d	Cs+	glycerol	-	-	224 (7.4)	142 (98.7)
6e	Cs+	glycerol	511 (6.3)	348 (2.2)	256 (14.1)	174 (100)
6e	Xe	thioglycerol	511 (20.6)	-	256 (52.5)	174 (100) ^b
6f	Cs+	glycerol	-	-	256 (2.9)	174 (100)
6g	Cs+	glycerol	-	364 (4.2)	272 (10.2)	190 (100)
6h	Cs+	glycerol	-	-	272 (13.7)	190 (100)
6h	Xe	3-NOBA	-	-	272 (24.6)	190 (6.4) ^C
6i	Cs ⁺	glycerol	555 (11.4)	370 (10.2)	278 (34.8)	196 (100)
6i	Xe	thioglycerol	555 (40.2)	386 (2.7)	278 (74.6)	196 (100)

^a Bombardment by primary beam of Cs⁺ ions or Xe atoms as indicated.

$$Ar - CH = P OH OH OH OH OH OH$$

$$Ar - CH = NH$$

$$OH OH OH OH$$

$$OH OH$$

$$O$$

SCHEME V Elimination of phosphorous acid from the pseudo-molecular ions of α -aminoarylmethane-phosphonic acids 6.

useful confirmation of identity (Table II). The spectra in most cases show prominent pseudomolecular ions, MH^+ , although these are generally weaker in intensity than those observed for aminoalkanephosphonic acids and in one case **6f** this ion was not distinguishable from the background. In those cases in which comparison was made (for **6e**, **6h**, **6i**), more intense pseudomolecular ions were obtained using xenon bombardment with thioglycerol or 3-nitrobenzyl alcohol (3-NOBA) as matrix, than with glycerol and Cs^+ ion bombardment. The base peak in all spectra was generally formed by the elimination of phosphorous acid from the pseudomolecular ion (Scheme V), the so-formed iminium ion being stabilized in these aromatic systems by conjugation with the benzene ring. Cluster ions such as $[MH + Z]^+$ and $[2M + H]^+$ were also observed in certain cases. In the case of the 4-CF₃ compound **6e** there was evidence for the elimination of metaphosphoric acid from the pseudomolecular ion, ^{6.10b.41} as shown by the appearance of a peak at m/z = 176 (12.1%), and of the loss of fluorine from the iminium ion to give m/z = 155 (29.2%).

b Also m/z [%] = 176, ([MH - HPO₃]⁺, 12.1), 155 ([MH - H₃PO₃ - F]⁺, 29.2).

^c Peak at m/z [%] = 189 (15.8) corresponds to unexpected [M - H_3PO_3]⁺; also m/z [%] = 254 ([MH - H_2O]⁺, 28.5).

EXPERIMENTAL

Fluorinated benzaldehydes, fluorinated benzoic acids and other reagents were obtained commercially; benzaldehydes and phosphites were distilled before use and solvents were dried according to standard procedures. Melting points were determined using a Reichert Thermopan Kofler Hot Stage or on an Electrothermal Digital Melting Point Apparatus (mp. >250°C); lower melting points were determined on a Büchi Melting Point Apparatus SMP 200; they are all uncorrected. Preparative studies were performed in London and Düsseldorf.

NMR Spectroscopy

Results from ³¹P-NMR for compounds 6 and 11 are given in Table I and VI-VIII. Compounds 6, 10 and 11 were characterized by extensive ¹³C-NMR investigations, corresponding data are listed in Tables IX to XIII. Some ¹H-data for 6 and 10 are given in Tables I and V. In general the ¹H and ¹⁹F-NMR spectra of the aromatic ring systems in all compounds described in this study are of second order character. Labourious analyses and iterations of those high resolution spectra were not performed, but a few, significant, directly accessible parameters are given under the preparative section.

The following spectrometers and referencing techniques were used by the London (L) and Düsseldorf (D) teams: 'H NMR: Perkin Elmer R12B (L), 60 MHz, Bruker AM 250 (L), 250.13 MHz, Bruker AM 200 SY (D), 200.13 MHz. 'C NMR: Bruker WP 80 (L), 20.12 MHz, Bruker AM 250 (L), 62.90 MHz, Bruker AM 200 SY (D), 50.32 MHz. Chemical shifts were recorded relative to TSP-d₄ (Me₃Si—CD₂—CO₂—COONa) for aqueous solutions of aminophosphonic acids or TMS for solutions of the Schiff bases or dialkyl N-protected aminoarylmethanephosphonates in CDCl₃. ³¹P NMR: WP 80 (L), 32.39 MHz, AM 200 SY (D), 81.02 MHz with 85% H₃PO₄ as an external reference. ¹⁹F NMR: Varian VXR 400 (L), 376 MHz with CFCl₃ as an external reference, AM 200 SY (D), 188.28 MHz with C₆F₆ as an internal reference for solutions in CDCl₃ and C₆D₆ or as an external reference for D₂O solutions. The NMR parameters (δ, given in [ppm] and "J_{XY}, given in [Hz]) were taken directly from the spectra.

Mass Spectrometry

Low resolution electron impact mass spectra were obtained on a Kratos Profile mass spectrometer (L) operating at 70 eV. FAB mass spectra were obtained with a VG Micromass ZAB-E mass spectrometer, with a primary beam of xenon atoms operating at 8 kV. LSIMS mass spectra were obtained using the Kratos Profile instrument, equipped with a Cs⁺ ion gun operating at 10 kV.

Preparation of Fluorinated α -Aminoarylmethanephosphonic Acids by Method A (Amination of Diethyl Aroylphosphonates) (Scheme I)

Fluorinated Benzoyl Chlorides $2a-d^{44}$: 0.50 mol fluorinated benzoic acid 1, 0.75 mol thionyl chloride and 2 drops of dimethylformamide were mixed under an atmosphere of nitrogen and heated (2-3 h), with stirring at 70-80°C (oil bath temperature). Gas evolution occurred with the formation of a greenish-brown solution. The excess of thionyl chloride was removed by distillation at atmospheric pressure and the fluorinated benzoyl chloride 2 was purified by fractional distillation in vacuo (oil pump).

The fluorinated benzoyl chlorides 2 were obtained as colourless, strongly-smelling liquids; they were stored under nitrogen on account of their sensitivity to hydrolysis.

```
4-Fluorobenzoyl Chloride 2a
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M.Wt. = 158.56, yield: 98.8%, bp. 69-71^{\circ}C/15 mmHg, n_{20} = 1.5296.
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¹H NMR (6% CDCl₃, TMS), δ : 7.12-7.23 (m, $2H_{arom.}$, 4-F— C_6H_4 —C), 8.09-8.19 (m, $2H_{arom.}$, 4-F— C_6H_4 —C).

¹⁹F NMR (6% CDCl₃, C_6F_6 int.), δ : 60.92–61.07 (m, 1F, 4-F— C_6H_4 —C).

2-Fluorobenzoyl Chloride 2c

```
M.Wt. = 158.56, yield: 98.3%, bp. 48-50^{\circ}C/1.5 mmHg, n_{20} = 1.5369.
```

¹H NMR (6% CDCl₃, TMS), δ : 7.14–7.36 (m, 2H_{arom.}, 2-F—C₆H₄—C), 7.62–7.73 (m, 1H_{arom.}, 2-F—C₆H₄—C), 8.08–8.17 (m, 1H_{arom.}, 2-F—C₆H₄—C).

¹⁹F NMR (6% CDCl₃, C₆F₆ int.), δ: 53.57-53.70 (m, 1F, 2-F--C₆H₄---C).

3,4-Difluorobenzoyl Chloride 2d

M.Wt. = 176.55, yield: 96.3%, bp. 72-73°C/13 mmHg.

¹H NMR (6% CDCl₃, TMS), δ: 7.27-7.40 (m, $1H_{arom}$, $3.4-F_2-C_6H_3-C$), 7.92-8.02 (m, $2H_{arom}$, $3.4-F_2-C_6H_3-C$).

¹⁹F NMR (6% CDCl₃, C_6F_6 int.), δ : 27.21 – 27.44 (m, 1F, 3.4-F₂— C_6H_3 —C), C 36.60 – 36.83 (m, 1F, 3.4-F₂— C_6H_3 —C).

Fluorinated Diethyl Benzoylphosphonates 3a-d: 0.20 mol fluorinated benzoyl chloride 2 was mixed with 0.22 mol triethyl phosphite by dropwise addition (during 2 h), with vigorous stirring under nitrogen. A strongly exothermic reaction occurred with the evolution of ethyl chloride and the reaction mixture became yellow. The temperature of the mixture was maintained at 40°C with the aid of an ice/water bath. Stirring was then continued at room temperature under a gentle stream of nitrogen (4-8 h). After removal of the excess of triethyl phosphite, the fluorinated diethyl benzoylphosphonate 3 was purified by fractional distillation in vacuo (oil pump).

The fluorinated diethyl benzoylphosphonates 3 are clear, oily, yellow liquids. As they undergo decomposition in the presence of traces of acids or bases, or weak nucleophiles (like water or amines), they were stored under argon.

```
Diethyl 4-Fluorobenzoylphosphonate 3a
M.Wt. = 260.20, yield: 95.1%, bp. 97-99°C/0.1 mmHg.
<sup>31</sup>P{<sup>1</sup>H} NMR (10% CDCl<sub>3</sub>, 85% H<sub>3</sub>PO<sub>4</sub> ext.), \delta: -0.88 (d, <sup>6</sup>J<sub>PF</sub> 0.6).
<sup>1</sup>H NMR (6%, C_6D_6, TMS), \delta: 1.00 (dt, {}^3J_{HH} 7.1, {}^4J_{PH} 0.5, 6H, 2 × CH<sub>3</sub>), 3.99 (qd, {}^3J_{HH} 7.1, {}^3J_{PH} 8.1,
4H, 2 \times CH_2), 6.56-6.60 (m, 2H_{arom.}, 4-F-C_6H_4-C), 8.33-8.43 (m, 2H_{arom.}, 4-F-C_6H_4-C).
<sup>19</sup>F NMR (6% C_6D_6, C_6F_6 int.), \delta: 60.65–60.80 (m, 1F, 4-F—C_6H_4—C).
Diethyl 2-Fluorobenzoylphosphonate 3c
M.Wt. = 260.20, yield: 68.8\%, bp. 108-111°C/0.05 mmHg.
<sup>31</sup>P{^{1}H} \delta: -1.90 (d, ^{4}J_{PF} 1.6).
<sup>1</sup>H \delta: 1.02 (dt, <sup>3</sup>J_{HH} 7.1, <sup>4</sup>J_{PH} 0.5, 6H, 2 × CH<sub>3</sub>), 4.03 (qd, <sup>3</sup>J_{HH} 7.1, <sup>3</sup>J_{PH} 8.2, 4H, 2 × CH<sub>2</sub>), 6.60–6.76
(m, 2H_{arom}, 2-F-C_6H_4-C), 6.81-6.91 (m, 1H_{arom}, 2-F-C_6H_4-C), 8.35-8.45 (m, 1H_{arom}, 2-F-C_6H_4-C)
C_6H_4--C).
<sup>19</sup>F δ: 51.78–51.91 (m, 1F, 2-F—C_6H_4—C).
Diethyl 3,4-Difluorobenzoylphosphonate 3d
M.Wt. = 278.19, yield: 83.4\%, bp. 116-117°C/0.1 mmHg.
<sup>31</sup>P{<sup>1</sup>H} \delta: -1.68 (dd, <sup>6</sup>J_{PF} 3.3, <sup>5</sup>\tilde{J}_{PF} 0.6).
<sup>1</sup>H \delta: 1.41 (dt, <sup>3</sup>J_{HH} 7.1, <sup>4</sup>J_{PH} 0.6, 6H, 2 × CH<sub>3</sub>), 4.31 (qd, <sup>3</sup>J_{HH} 7.1, <sup>3</sup>J_{PH} 8.1, 4H, 2 × CH<sub>2</sub>), 7.27–7.42
(m, 1H_{arom.}, 3,4-F_2—C_6H_3—C), 8.05-8.23 (m, 2H_{arom.}, 3,4-F_2—C_6H_3—C).
^{19}F δ: 26.65–26.89 (m, 1F, 3,4-F<sub>2</sub>—C<sub>6</sub>H<sub>3</sub>—C), 35.55–35.78 (m, 1F, 3,4-F<sub>2</sub>—C<sub>6</sub>H<sub>3</sub>—C).
```

Fluorinated Diethyl Benzoyloximinophosphonates 4a-d: 0.20 mol hydroxylamine hydrochloride was suspended in 130 ml of absolute ethanol, with the exclusion of moisture, and 0.23 mol dry pyridine was added. A solution of 0.15 mol fluorinated diethyl benzoylphosphonate 3 in 50 ml of absolute ethanol was added dropwise (during 2 h). Slight warming occurred to give a clear, yellow solution and the reaction temperature was maintained below 30°C with the aid of an ice/water bath. The mixture was finally stirred at room temperature (55-75 h) and became colourless (the reaction was monitored by $^{31}P\{^{1}H\}$ NMR). The solvent was then removed on a rotary evaporator under water pump pressure to leave an oil which was taken up in 200 ml of water and extracted five times with dichloromethane. The organic phase was dried over Na₂SO₄ and filtered, and after removal of solvent the yellowish-brown oil was dried in vacuo (oil pump) at room temperature.

The oil consists of a mixture of the cis- and trans-isomer of the fluorinated diethyl benzoyloximinophosphonate 4 (with the E-form in excess), contaminated with phosphates. Purification by vacuum distillation is impossible since decomposition occurs above 60°C.

```
Diethyl 4-Fluorobenzoyloximinophosphonate 4a M.Wt = 275.22, yield of the crude oil: 94.8%. 

^{31}P\{^{1}H\} NMR (reaction mixture in C_{2}H_{3}OH, D_{2}O ext., 85% H_{3}PO_{4} ext.), \delta: 10.06 (s), (E), 4.93 (s), (Z), (relation E:Z = 1.5:1). 

Diethyl 2-Fluorobenzoyloximinophosphonate 4c
```

M.Wt. = 275.22, yield of the crude oil: 96.0%. $^{31}P\{^{1}H\}$ NMR (10% CDCl₃, 85% H₃PO₄ ext.), δ : 8.60 (d, $^{4}J_{PF}$ 1.9), (E), 3.77 (d, $^{4}J_{PF}$ 1.1), (Z), (relation E:Z = 1.5:1).

```
Diethyl 3,4-Difluorobenzoyloximinophosphonate 4d M.Wt. = 293.21, yield of the crude oil: 93.2%. ^{31}P\{^{1}H\} \delta: 9.26 (s), (E), 4.64 (s), (Z) (relation E:Z = 2:1).
```

Fluorinated Diethyl Aminoarylmethanephosphonates and N-Formylated Derivatives 5a-d: 0.10 mol of the oily fluorinated diethyl benzoyloximinophosphonate 4 was added dropwise (during 1 h), under nitrogen, to a suspension of 0.40 mol zinc dust in 100 ml of 99% formic acid. An exothermic reaction occurred with the vigorous evolution of gas; the reaction temperature was maintained below 65°C by cooling. After addition was complete the suspension was stirred at room temperature (12 h) and the reaction was monitored by ³¹P{¹H} NMR. Solids were filtered using a G3-sintered glass filter and washed

repeatedly with formic acid to give a yellow filtrate which was evaporated on a rotary evaporator under water pump pressure to give a brownish oil.

From the ³¹P{¹H} NMR spectrum it could be seen that in addition to reduction to give diethyl aminoarylmethanephosphonate 5 partial hydrolysis had occurred in the acid solution to give the aminoarylmethanephosphonic acid (in form of its formic acid salt) 6'.

Diethyl Amino-(4-fluorophenyl-)methanephosphonate 5a

M.Wt. = 261.23.

³¹P{¹H} NMR (reaction mixture in HCOOH, D₂O ext., 85% H₃PO₄ ext.), δ: 21.27 (s), diethyl aminoarylmethanephosphonate, 16.92 (s), aminoarylmethanephosphonic acid.

Diethyl Amino-(2-fluorophenyl-)methanephosphonate 5c

M.Wt. = 261.23.

³¹P(¹H) NMR (reaction mixture in HCOOH, D₂O ext., 85% H₃PO₄ ext.), δ: 19.19 (s), diethyl aminoarylmethanephosphonate, 14.72(s), aminoarylmethanephosphonic acid.

Diethyl Amino-(3,4-difluorophenyl-)methanephosphonate 5d

M.Wt. = 279.22.

³¹P{¹H} NMR (reaction mixture in HCOOH, D₂O ext., 85% H₃PO₄ ext.), δ: 18.43 (s), diethyl aminoarylmethanephosphonate, 11.14 (s), aminoarylmethanephosphonic acid.

Fluorinated α -Aminoarylmethanephosphonic Acids **6a-d**: The brownish oil was heated under reflux (6-8 h) with 50 ml of concentrated hydrochloric acid (oil bath temperature $100-110^{\circ}$ C). The hydrolysis gave a clear yellow solution (monitored by ${}^{31}P\{{}^{1}H\}$ NMR), which was evaporated at water pump pressure on a rotary evaporator to leave a yellowish oil of the aminoarylmethanephosphonic acid hydrochloride **6'**. Water was added and the solution was evaporated repeatedly until a white solid was formed. The residue was heated under reflux (2 h) with 50-100 ml of water, and the slightly soluble α -aminoarylmethanephosphonic acid **6** separated as a white crystalline solid. The solid was filtered off (by a Büchner funnel), washed with ethanol, dried in a vacuum desiccator over silicagel, and recrystallized from water/ ethanol mixtures.

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α-Amino-(4-fluorophenyl-)methanephosphonic Acid 6a
```

M.Wt. = 205.13, yield: 63.3% (based on diethyl benzoylphosphonate), mp. 278-282°C.

³¹P{¹H} NMR (10% 1m KOD/D₂O, 85% H₃PO₄ ext.), δ : 18.50 (d, ${}^{6}J_{PF}$ 3.9).

¹H NMR (10% 1m KOD/D₂O, TSP—d₄—Na), δ : 3.80 (d, ² J_{PH} 15.3, 1H, PCH), 7.03 – 7.15 (m, 2H_{arom.}, 4-F—C₆H₄—C), 7.33 – 7.45 (m, 2H_{arom.}, 4-F—C₆H₄—C).

¹⁹F NMR (10% 1m KOD/D₂O, C_6F_6 ext.), δ : 45.00 – 45.18 (m, 1F, 4-F— C_6H_4 —C).

α-Amino-(2-fluorophenyl-)methanephosphonic Acid 6c

M.Wt. = 205.13, yield: 43.3% (based on diethyl benzoylphosphonate), mp. 270-273°C.

³¹P{¹H} δ : 18.14 (d, ⁴ J_{PF} 4.1).

¹H δ : 4.13 (d, ² J_{PH} 15.9, 1H, PCH), 7.06–7.35 (m, 3H_{arom}, 2-F—C₆H₄—C), 7.42–7.52 (m, 1H_{arom}, 2-F—C₆H₄—C).

¹⁹F δ: 45.01-45.21 (m, 1F, 2-F—-C₆H₄—-C).

α-Amino-(3,4-difluorophenyl-)methanephosphonic Acid 6d

M.Wt. = 223.12, yield: 56.1% (based on diethyl benzoylphosphonate), mp. 280-282°C.

 $^{31}P\{^{1}H\}$ δ : 17.86 (dd, $^{6}J_{PF}$ 4.2, $^{5}J_{PF}$ 1.6).

¹H δ : 3.79 (d, ² J_{PH} 15.7, 1H, CH), 7.09–7.34 (m, 3H_{aron.}, 3,4-F₂—C₆H₃—C).

¹⁹F δ : 23.95–24.18 (m, 1F, 3,4-F₂—C₆H₃—C), 27.37–27.61 (m, 1F, 3,4-F₂—C₆H₃—C).

Preparation of Fluorinated α -Aminoarylmethanephosphonic Acids by Method B (Amidoalkylation of Phosphorus Trichloride)

One-Pot Procedure: Variation a) with Benzamide; $R = C_6H_5$ (Scheme II)

Fluorinated N-Benzoylaminoarylmethanephosphonic Acids 8a/b-h: Under a gentle stream of nitrogen, 0.05 mol benzamide was suspended in 10 ml of acetic acid, and 0.05 mol phosphorus trichloride was added with vigorous stirring. An exothermic reaction occurred with the evolution of hydrogen chloride, and the mixture was stirred until it had cooled to room temperature. 0.07 mol Fluorinated benzaldehyde 7 was added dropwise (during 30 min), so that the reaction temperature did not exceed 40°C. A clear, light yellow solution was formed. In order to complete the conversion, the mixture was heated on a boiling water bath (1-2 h) until gas evolution ceased. Rotary evaporation at water pump pressure gave the N-benzoylaminoarylmethanephosphonic acid 8a as a yellow oil.

Fluorinated α -Aminoarylmethanephosphonic Acids **6b-h**: In order to remove the benzoyl protecting group from nitrogen, the oil was heated under reflux (6-24 h) with 30 ml of concentrated hydrochloric

TABLE III

Fluorinated α-aminoarylmethanephosphonic acids 6b-h
(Scheme II, variation a)

no.	M.Wt.	yields	mp.	31P{1H} NMR
	· · · · · · · · · · · · · · · · · · ·	[%]	[oC]	δ [ppm], Mult., ⁿ JpF [Hz]
6b	205.13	18.6	289-290	18.03, d, ⁵ J _{PF} 1.4
6e	255.13	40.9	287-289	17.71, q, ⁷ JpF 2.1
6f	255.13	41.7	253-255	15.08, bs
6g	271.13	31.1	305-307	18.12, s
6h	271.13	25.9	277-280	15.99, s

acid. After cooling, two phases were formed, the lower being a yellowish oil of the aminoarylmethanephosphonic acid hydrochloride 6' and the upper being an aqueous phase with needle-like crystals of the resulting benzoic acid. The aqueous phase was separated, filtered to remove benzoic acid and, together with the oily layer, was evaporated at water pump pressure on a rotary evaporator. The residual oil was repeatedly taken up in water and evaporated until a white solid was formed. The residue was dissolved in hot methanol and, after cooling, pyridine was added dropwise until the pH was 4. After several days in a refrigerator the α -aminoarylmethanephosphonic acid 6 separated as a fine, white solid, which was filtered off (on a G3-sintered glass filter), washed with methanol, and dried in a vacuum desiccator over silica gel (Table III).

```
α-Amino-(3-fluorophenyl-)methanephosphonic Acid 6b ^{31}P\{^{1}H\} NMR (10% 1m KOD/D<sub>2</sub>O, 85% H<sub>3</sub>PO<sub>4</sub> ext), δ: 18.03 (d, ^{5}J_{PF} 1.4). ^{1}H NMR (10% 1m KOD/D<sub>2</sub>O, TSP—d<sub>4</sub>—Na), δ: 3.93 (d, ^{2}J_{PH} 15.6, 1H, PCH), 6.99–7.10 (m, 1H<sub>arom.</sub>, 3-F—C<sub>6</sub>H<sub>4</sub>—C), 7.14–7.23 (m, 2H<sub>arom.</sub>, 3-F—C<sub>6</sub>H<sub>4</sub>—C), 7.32–7.43 (m, 1H<sub>arom.</sub>, 3-F—C<sub>6</sub>H<sub>4</sub>—C). ^{19}F NMR (10% 1m KOD/D<sub>2</sub>O, C<sub>6</sub>F<sub>6</sub> ext.), δ: 48.42–48.61 (m, 1F, 3-F—C<sub>6</sub>H<sub>4</sub>—C).
```

α-Amino-(4-trifluoromethylphenyl-)methanephosphonic Acid **6e**

³¹P{¹H} δ : 17.71 (q, ⁷ J_{PF} 2.1).

 1 H δ: 3.91 (d, $^{2}J_{PH}$ 16.5, 1H, PCH), 7.51–7.58 (m, 2H_{arom.}, 4-F—C₆H₄—C), 7.63–7.72 (m, 2H_{arom.}, 4-F—C₆H₄—C).

¹⁹F δ: 101.16 (bs, 3F, 4-CF₃— C_6H_4 —C).

 $\alpha\text{-}Amino\text{-}(3\text{-}trifluoromethylphenyl-)methanephosphonic Acid 6f <math display="inline">^{31}P\{^{1}H\}$ 8: 15.03 (s).

 1 H δ: 4.01 (d, $^{2}J_{PH}$ 15.8, 1H, PCH), 7.49 – 7.68 (m, 3H_{arom.}, 3-CF₃—C₆H₄—C), 7.73 – 7.76 (m, 1H_{arom.}, 3-CF₃—C₆H₄—C).

¹⁹F δ: 100.97 (bs, 3F, 3-CF₃— C_6H_4 —C).

α-Amino-(4-trifluoromethoxyphenyl-)methanephosphonic Acid 6g

 $^{31}P\{^{1}H\}$ δ : 18.12 (s).

¹H δ: 3.85 (d, $^2J_{PH}$ 15.8, 1H, PCH), 7.25–7.32 (m, 2H_{arom.}, 4-CF₃O—C₆H₄—C), 7.42–7.51 (m, 2H_{arom.}, 4-CF₃O—C₆H₄—C).

¹⁹F δ: 104.94 (bs, 3F, 4-CF₃O—C₆H₄—C).

α-Amino-(3-trifluoromethoxyphenyl-)methanephosphonic Acid 6h

³¹P{¹H} δ : 15.99 (s).

¹H δ: 3.91 (d, ${}^{2}J_{PH}$ 16.0, 1H, PCH), 7.20–7.26 (m, 1H_{arom.}, 3-CF₃O—C₆H₄—C), 7.32–7.49 (m, 3H_{arom.}, 3-CF₃O—C₆H₄—C).

¹⁹F δ: 105.54 (bs, 3F, 3-CF₃O—C₆H₄—C).

One-Pot Procedure: Variation b) with Acetamide; $R = CH_3$ (Scheme II)

Fluorinated N-Acetylaminoarylmethanephosphonic Acids 8b/b-h: Under a gentle stream of nitrogen, 0.05 mol acetamide was dissolved in 10 ml of acetic acid and 0.05 mol phosphorus trichloride was added with vigorous stirring. Slight warming occurred, with gas evolution, and a white suspension was formed. The mixture was allowed to cool to room temperature and 0.07 mol fluorinated benzaldehyde 7 was added (during 30 min) with the temperature below 40° C. A weakly exothermic reaction occurred with gas evolution and the solid dissolved. In order to complete the reaction, the mixture was heated (2-3 h) on a boiling water bath. Rotary evaporation of the solution at water pump pressure finally gave the N-acetylaminoarylmethanephosphonic acid 8b as a light yellow oil.

TABLE IV
Fluorinated α-aminoarylmethanephosphonic acids 6b-h
(Scheme II, variation b)

no.	M.Wt.	mp.	yields [%]	31 _P {1 _H }-NMR δ [ppm], Mult., ⁿ J _{PF} [Hz]
_	005.40	888 888	oc c	18.03, d, ⁵ Jp p 1.4
6b	205.13	289-290	25.5	
6e	255.13	287-289	74.8	17.71, q, ⁷ JpF 2.1
6f	255.13	253-255	35.6	15.08, bs
6g	271.13	305-307	45.1	18.12, s
6h	271.13	277-280	35.0	15.99, s

Fluorinated α -Aminoarylmethanephosphonic Acids **6b-h**: The acetyl group was removed from nitrogen by heating the yellowish oil under reflux (12-30 h) with 30 ml of concentrated hydrochloric acid, to give a clear, light yellow solution. After cooling, insoluble components were filtered off, and the aqueous phase was evaporated on a rotary evaporator at water pump pressure to give the aminoarylmethanephosphonic acid hydrochloride **6'** as a yellow oil; this was then repeatedly taken up in water and evaporated until a solid separated in the oil. The residue was dissolved in hot methanol, and after cooling, pyridine was added dropwise until the pH was 4. After several days in a refrigerator the α -aminoarylmethanephosphonic acid separated as a fine, white solid, which was filtered off (on a G3-sintered glass filter), washed with methanol, and dried in a vacuum desiccator over silica gel (Table IV).

Preparation of Fluorinated α -Aminoarylmethanephosphonic Acids by Method B (Amidoalkylation of Phosphorus Trichloride)

Two-Step Procedure (Scheme III)

N,N'-(4-Fluorophenyl-)methylidenebisacetamide 9a: 0.20 mol 4-Fluorophenyl-)methylidenebisacetamide 9a: 0.20 mol 4-Fluorophenyl-)methyliden

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N,N'-(4-Fluorophenyl-)methylidenebisacetamide 9a
```

M.Wt. = 224.23, yield: 70.0%, white needles, mp. 165-167°C.

¹H NMR (6% CDCl₃, TMS), δ: 1.94 (bs, 6H, $2 \times$ CH₃), 6.20 (m, 3H, CH + $2 \times$ NH), 6.98-7.09 (m, 2H_{arom.}, 4-F—C₆H₄), 7.95-8.05 (m, 2H_{arom.}, 4-F—C₆H₄).

¹⁹F NMR (10% C_6D_6 , C_6F_6 int.) δ : 56.02 (m, 1F, 4-F— C_6H_4).

N-Acetylamino-(4'-fluorophenyl-)methanephosphonic Acid 8ba: Under a gentle stream of nitrogen, 0.10 mol N,N'-(4-fluorophenyl-)methylidenebisacetamide 9a was suspended in 30 ml of acetic acid and 0.10 mol phosphorus trichloride was added dropwise (during 60 min) at a reaction temperature of 10°C. A strongly exothermic reaction occurred with the evolution of hydrogen chloride and the formation of a white gel. After the addition was complete, the mixture was heated further (2-3 h) on a boiling water bath to give a clear, light yellow solution. Cooling, followed by rotary evaporation at water pump pressure, gave N-acetylamino-(4'-fluorophenyl-)methanephosphonic acid 8ba, as a light yellow oil.

 α -Amino-(4-fluorophenyl-)methanephosphonic Acid 6a: In order to remove the acetyl group from nitrogen, the oil of N-acetylamino-(4'-fluorophenyl-)methanephosphonic acid 8ba was heated under reflux (24 h) with 100 ml of concentrated hydrochloric acid. A yellowish solution was formed, with fine needles of acetamide hydrochloride which were filtered off after cooling, and washed with water. The aqueous filtrate was evaporated at water pump pressure on a rotary evaporator to give amino-(4-fluorophenyl-)methanephosphonic acid hydrochloride as a yellow oil. This was repeatedly taken up in water, followed by evaporation, until a solid formed in the oil. The residue was dissolved in a little hot methanol and, after cooling, pyridine was added dropwise until the pH was 4. The solution was allowed to stand for several days in an ice/water bath, when α -amino-(4-fluorophenyl-)methanephosphonic acid 6a was obtained as a fine white solid which was filtered off (using a G3-sintered glass filter), washed with methanol, and dried in a vacuum desiccator over silica gel.

TABLE V	
Fluorinated N-benzylidene-1,1-diphenylmethylamines 10a	-g

no.	M.Wt.	yields	reacting	¹ H NMR
		[%]	time [h]	δ (=CH) [ppm]
10a	289.35	99.5	4	8.35
10b	289.35	97.8	6	8.38
10d	307.34	93.5	6	8.31
10e	339.36	98.2	4	8.43
10f	339.36	93.1	10	8.44
10g	355.36	95.5	8	8.39

TABLE VI Fluorinated dimethyl N-diphenylmethylaminoarylmethanephosphonates 11aa-ae

no.	M.Wt.	yields [%]	reacting time [h]	³¹ P{ ¹ H} NMR δ [ppm], Mult., ⁿ Jp _F [Hz]
11aa	399.40	79.7	4	26.49, d, ⁶ J _{PF} 4.8
11ae	449.41	89.4	5	25.72, q, ⁷ JpF 2.3

α-Amino-(4-fluorophenyl-)methanephosphonic Acid 6a

M.Wt. = 205.13, yield: 59.0%, fine, white solid, mp. 278-282°C.

Preparation of Fluorinated α-Aminoarylmethanephosphonic Acids by the Method C (Addition of Substances with P-H Bonds to N-Protected Imines, Variation a)) (Scheme IV)

Fluorinated N-Benzylidene-1,1-diphenylmethylamines 10a-h: The fluorinated benzaldehyde 7 was added dropwise with stirring to a solution of an equimolar quantity of diphenylmethylamine in anhydrous ether (5% w/w) or dichloromethane (10-30% w/w) in the presence of anhydrous potassium carbonate (1-2 mol equiv) at 0°C. The solution was stirred (1-1.5 h) at room temperature and then filtered. Solvent was removed by rotary evaporation, followed by vigorous shaking at 0.1 mmHg (2 h), to leave the Schiff base as follows:

N-4-Fluorobenzylidene-I, I-diphenylmethylamine 10a

Yield: 97%, bright yellow crystalline solid, C, H, N analysis: C₂₀H₁₆FN, M.Wt. = 289.35, requires: C 83.02, H 5.57, N 4.84, found: C 82.91, H 5.60, N 4.89%.

¹H NMR (CDCl₃), δ 5.57 (s, 1H, NCH), 7.06 (m, 2H_{arom.}, 4-F—C₆H₄), 7.18–7.40 (m, 10H_{arom.}, 2 \times C_6H_5), 7.81 (m, $2H_{arom.}$, $4\text{-F}—C_6H_4$), 8.35 (bs, 1H, =CH). ¹⁹F NMR (CDCl₃), δ : -109.63--109.71 (m, $4\text{-F}—C_6H_4$).

EI ms, m/z [%]: 289 (M^+ , 15), 167 ((C_6H_5)₂ CH^+ , 100).

N-2-Fluorobenzylidene-1,1-diphenylmethylamine 10c

Yield: 99%, sticky, orange-yellow solid, C, H, N analysis: C₂₀H₁₆FN, M.Wt. = 289.35, requires: C 83.02, H 5.57, N 4.84, found: C 83.01, H 5.53, N 4.86%.

 1 H NMR (CDCl₃), δ : 5.60 (s, 1H, NCH), 7.02 (m, 1H_{aron.}, 2-F—C₆H₄), 7.11–7.41 (m, 10H_{aron.}, 2 × $C_6H_5 + 2H_{arom.}$, 2-F— C_6H_4), 8.18 (m, $1H_{arom.}$, 2-F— C_6H_4), 8.74 (bs, $1H_5$ —CH).

¹⁹F NMR (CDCl₃), δ : - 124.57 (m, 2-F—C₆H₄).

EI ms, m/z [%]: 290 ([M+1]⁺, 20.6), 289 ([M]⁺, 49.8), 270 ([M - F]⁺, 6.3), 212 ([M - C₆H₅]⁺, 25.9), 182 ([M - F— C_6H_4C]⁺, 23.2), 181 ([M - F— C_6H_4CH]⁺, 18.6), 167 ([M - F— C_6H_4 — $CH = N^{+}, 100, 108 ([M - (C_6H_5)_2CH - N]^{+}, 35.1).$

N-4-Trifluoromethylbenzylidene-1,1-diphenylmethylamine 10e

Yield: 99%, orange solid, C, H, N analysis: C₂₁H₁₆F₃N, M.Wt. = 339.36, requires: C 74.33, H 4.75, N 4.13, found: C 74.30, H 4.82, N 4.07%.

N-3-Trifluoromethoxybenzylidene-1,1-diphenylmethylamine 10h

Yield: 88%, light brown viscous oil, C, H, N analysis: C₂₁H₁₆F₃NO, M.Wt. = 355.36, requires: C 70.98, H 4.54, N 3.94, found: C 70.89, H 4.54, N 3.84%.

TABLE VII

Fluorinated diethyl N-diphenylmethylaminoarylmethanephosphonates

11ba - bg

no.	M.Wt.	yields	reacting	31P{1H} NMR
		[%]	time [h]	δ [ppm], Mult., ⁿ JpF [Hz]
11ba	427.45	90.4	10	24.13, d, ⁶ JpF 4.8
11bb	427.45	81.8	11	23.68, d, ⁵ J _{PF} 1.7
11bd	445.44	87.5	12	23.43, dd, 6JpF 5.5, 5JpF 1.8
11be	477.46	89.3	8	23.39, q, ⁷ JpF 2.3
11bf	477.46	78.6	12	23.30, q, ⁷ J _{PF} 0.6
11bg	493.46	91.6	14	23.79, s

TABLE VIII
Fluorinated diphenyl N-diphenylmethylaminoarylmethanephosphonates
11ca - cg

no.	M.Wt.	yields [%]	reacting time [h]	³¹ P{ ¹ H} NMR: δ [ppm], Mult., ⁿ Jp _F [Hz]
11ca	523.54	73.3	4	16.78, d, ⁶ Jp ; 5.1
11cb	523.54	86.1	8	16.33, d, ⁵ JpF 1.8
11cd	541.53	68.2	6	15.97, dd, ⁶ J _{PF} 5.4, ⁵ J _{PF} 1.8
11ce	573.55	78.6	5	15.95, q, ⁷ JpF 2.3
11cf	573.55	70.3	5	15.87, q, ⁷ JpF 0.7
11cg	589.55	77.8	4	16.37, s

N-Pentafluorobenzylidene-1,1-diphenylmethylamine 10i

Yield: 99%, bright yellow solid, C, H, N analysis: $C_{20}H_{12}F_5N$, M.Wt. = 361.31, requires: C 66.49, H 3.35, N 3.88, found: C 66.67, H 3.39, N 3.91%.

Fluorinated α -Aminoarylmethanephosphonic Acids 6a-i: A mixture of the fluorinated benzylidenediphenylamine 10 and an equimolar amount of dimethyl phosphite or diethyl phosphite was heated at 120-140°C (30 min). Concentrated hydrochloric acid (80 ml) was added to the product 11 and the resultant solution was heated under reflux for 3 h. After the removal of by-products by extraction with toluene the aqueous layer was concentrated under reduced pressure, dissolved in methanol, and treated with a moderate excess of propylene oxide at 40-50°C. The crystalline product which separated was filtered off, washed with acetone and dry ether, and dried in vacuo at 80°C to give the following products:

α-Amino-(4'-fluorophenyl-)methanephosphonic Acid 6a

Yield: 55%, m.p. 278-282°C, C, H, N analysis: $C_7H_9FNO_3P$, M.Wt. = 205.13, requires: C 40.99, H 4.42, N 6.83, found: C 40.98, H 4.93, N 6.80%.

 $^{31}P\{^{1}H\}$ δ : 17.03 (d, $^{6}J_{PF}$ 3.0).

¹H (NaOD/D₂O), δ : 3.86 (d, ² J_{PH} 15.4, 1H, PCH), 7.11 (m, 2H_{arom.}, 4-F—C₆H₄), 7.40 (m, 2H_{arom.}, 4-F—C₆H₄).

 $\alpha\text{-}Amino\text{-}(2'\text{-}fluorophenyl\text{-})methanephosphonic\ Acid\ \textbf{6c}$

Yield: 28%, m.p. 270–273°C, C, H, N analysis: $C_7H_9FNO_3P$, M.Wt. = 205.13, requires: C 40.99, H 4.42, N 6.83, found: C 41.05, H 4.41, N 6.61%.

³¹P{ 1 H} δ : 17.59 (d, $^{4}J_{PF}$ 4.0).

¹H (NaOD/D₂O), δ: 4.41 (d, $^2J_{PH}$ 16.1, 1H, PCH), 7.07 – 7.34 (m, 3H_{aron.}, 2-F—C₆H₄), 7.46 (m, 1H_{aron.}, 2-F—C₆H₄).

¹⁹F (NaOD/D₂O), δ: 117.97 (s).

 $\alpha\text{-}Amino\text{-}(4'\text{-}trifluoromethylphenyl\text{-})methanephosphonic\ Acid\ \textbf{6e}$

Yield: 46%, m.p. 287-289°C, C, H, N analysis: $C_8H_9F_3NO_3P$, M.Wt. = 255.13, requires: C 37.66, H 3.56,

TABLE IX

 $^{13}C\{^{1}H\}$ NMR data of the fluorinated α -aminoarylmethanephosphonic acids 6. $^{13}C\{^{1}H\}$ NMR (10% KOH/D₂O, TSP—d₄—Na), δ [ppm], Mult., " J_{XY} [Hz]. ov.: overlapping

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No.	α-СН	C1	C2	C3	C4	C5	C6	R
6a	57.52, d,	139.82, s	131.90, dd,	117.41, d,	164.12, d,	117.41, d,	131.90, dd,	
			$^{3}J_{FC}$ 7.8,	² J _{FC} 21.6	¹ J _{FC} 243.5	² J _{FC} 21.6	$^{3}J_{FC}$ 7.8,	
	¹ J _{PC} 131.1		³ J _{PC} 5.1	10			$^{3}J_{PC}$ 5.1	
6b		147.85, dd,		165.16, dd,	115.57, dd,	132.29, dd,	126.33, dd,	_
	⁴ J _{FC} 1.6,	³ J _{FC} 7.3,	2 _{JFC} 21.8,	1 _{JFC} 241.1,		$^{3}J_{FC}$ 8.5,	⁴ J _{FC} 2.6,	
	1 _{JPC} 129.5		³ J _{PC} 4.7	⁴ J _{PC} 2.5	5J _{PC} 2.6	⁴ J _{PC} 2.1	3 _{JPC} 5.0	
6c		131.82, dd,			131.55, dd,	126.69, s	130.53, dd,	-
	³ J _{FC} 1.2,	ov.	¹ J _{FC} 242.6	2 _{JFC} 22.8	³ J _{FC} 8.1,		³ J _{FC} 8.3,	
	¹ J _{PC} 132.1	$2_{J_{PC}}$ 1.9	ov		ov		3 _{JPC} 2.4	
6d		140.98, dd,	119.05, dd,	151.49, ddd,	152.30, ddd,	119.38, dd,	126.63, ddd,	-
	⁴ J _{FC} 0.9,	$^{3}J_{FC}^{3.0}$	² J _{FC} 16.9,	1 _{JFC} 242.8,	¹ J _{FC} 243.2,	$^{2}J_{FC}$ 15.6,	$^{3}J_{FC}$ 6.4,	
			³ J _{FC} 4.5,	² J _{FC} 12.6,	² J _{FC} 12.6,		⁴ J _{FC} 3.4,	
	¹ J _{PC} 129.5	$^{2}J_{PC}$ 2.9		⁴ J _{PC} 2.7	⁵ J _{PC} 2.3	$^{4}J_{PC}$ 1.7,	³ J _{PC} 5.2	
			OV.			ov.		
6e	58.27, d,	149.09, s	127.54, s	130.62, d,	130.05, q,	130.62, d,	127.54, s	127.34, q,
				$^{3}J_{FC}$ 4.7	² J _{FC} 63.8	$^{3}J_{FC}$ 4.7		¹ J _{FC} 271.2
	¹ J _{PC} 128.1							
6f	58.28, d,	146.15, d,	126.93, dd,	131.96, dq,	134.04, dd,	131.96, d,	125.63, dq,	127.32, q,
			³ J _{FC} 4.4,	$^{2}J_{FC}$ 31.5,	$^{3}J_{FC}$ 4.8,		⁵ J _{FC} 2.6,	¹ J _{FC} 271.7
	¹ J _{PC} 129.3	$^{2}J_{PC}$ 2.6	³ J _{PC} 3.9	⁴ J _{PC} 2.1	⁵ J _{PC} 1.2	⁴ J _{PC} 2.0	³ J _{PC} 3.9	
6g	57.91, d,	144.21, d,	131.75, d,	123.28, d,	150.02, q,	123.28, d,	131.75, d,	123.20, dq,
	, ,		, ,		³ J _{FC} 2.6		, ,	1 _{JFC} 255.6,
	¹ J _{PC} 130.1	$^{2}J_{PC}$ 2.3	$^{3}J_{PC}$ 4.9	⁴ J _{PC} 1.0	••	⁴ J _{PC} 1.0	³ J _{PC} 4.9	8 _{JPC} 1.1
6h		147.46, s	128.90, d,	151.24, s	132.00, s	121.13, s	122.69, d,	123.03, q,
	, ,	,	, ,	,	,	,	, ,	l _{JFC} 255.8
	1 _{JPC} 129.8		$^{3}J_{PC}$ 4.7				$^{3}J_{PC}$ 4.5	10
6i	-	118.45, dd,		140.14, ddd.	142.09, dm,	140.14, ddd,		
	, ,	² J _{FC} 16.2,			¹ J _{FC} 249.4,			
	1 _{JPC} 130.9		ov.	ov.	ov.	ov.	re ·	
	, r C == 4.0	- * *			•	•		

N 5.49, found: C 37.50, H 3.55, N 5.40%.

Yield: 43%, m.p. 277-280°C, C, H, N analysis: $C_8H_9F_3NO_4P$, M.Wt. = 271.13, requires: C 35.44, H 3.35,

 $^{^{31}}P\{^{1}H\}$ δ : 16.96 (s).

¹H (NaOD/D₂O), δ: 3.95 (d, ²J_{PH} 16.41, 1H, PCH), 7.57 (m, 2H_{arom.}, 4-CF₃—C₆H₄), 7.67 (m, 2H_{arom.}, 4-CF₃—C₆H₄), 7.67 (m, 2H_{arom.}, 4-CF₃—C₆H₄), 7.67 (m, 2H_{arom.}), 7.67 (CF_3 — C_6H_4). ¹⁹F (NaOD/D₂O), δ : 61.94 (s).

FAB ms (thioglycerol), m/z [%]: 511 (2M + H, 20.6), 256 (M + H, 52.5), 186 (14.6), 176 (M + H - H, 20.6) HPO_3 , 12.1), 174 (M + H - H_3PO_3 , 100), 155 (M + H - F, 29.2), 130 (72.7).

 $[\]alpha$ -Amino-(3'-trifluoromethoxyphenyl-)methanephosphonic Acid **6h**

TABLE X

Part I: ¹³C{¹H} NMR data of the fluorinated N-benzylidene-1,1-diphenylmethylamines 10. ¹³C{¹H} NMR (10% CDCl₃, TMS), δ [ppm], Mult., ⁿJ_{XY} [Hz]

No.	δ _C C1	C2	C3	C4	C5	C6	R
10a	132.55, d,	130.32, d ,	115.58, d,	164.33, d,	115.58, d,	130.32, d,	_
(L)	⁴ J _{FC} 2.8	³ J _{FC} 8.6	² J _{FC} ^{21.9}	¹ J _{FC} 250.8	² J _{FC} ^{21.9}	³ J _{FC} 8.6	
10a	133.28, d,	131.00, d,	116.25, d,	165.02, d,	116.25, d,	131.00, d,	-
(D)	⁴ J _{FC} 3.1	3 _{JFC} 8.6	² J _{FC} ^{21.9}	¹ J _{FC} 250.8	² J _{FC} ^{21.9}	³ J _{FC} 8.6	
10b	139.32, d,	115.08, d,	163.70, d,	118.37, d,	130.71, d,	125.27, d,	-
(D)	$^{3}J_{FC}$ 7.3	² J _{FC} 22.2	¹ J _{FC} 246.4	² J _{FC} 21.7	³ J _{FC} 8.0	⁴ J _{FC} 2.9	
10c	123.86, d,	162.33, d,	115.63, d,	124.24, d,	128.07, d,	132.31, d,	-
(L)	² J _{FC} 9.4	1 _{JFC} 252.5	² J _{FC} 21.1	³ J _{FC} 3.6	⁴ J _{FC} 2.8	³ J _{FC} 8.6	
10 d	134.24, dd,	117.95, dd,	151.36, dd,	152.72, dd,	117.11, dd,	125.86, dd,	-
(D)	$^{3}J_{FC}$ 5.5,	$^{2}J_{FC}$ 17.8,	¹ J _{FC} 249.1,	¹ J _{FC} 252.9,	$^{2}J_{FC}$ 18.0,	$^{3}J_{FC}$ 6.8,	
	⁴ J _{FC} 3.6	$^{3}J_{FC}$ 0.7	$^{2}J_{FC}$ 13.1	$^{2}J_{FC}$ 13.1	$^{3}J_{FC}$ 1.1	⁴ J _{FC} 3.5	
10e	139.35, s	128.65, s	125.49, q,	*	125.49, q,	128.65, s	132.29, q,
(L)			$^{3}J_{FC}$ 3.8		³ J _{FC} ^{3.8}		¹ J _{FC} 259.3
10e	140.07, q,	129.32, s	126.16, q,	133.00, q,	126.16, q,	129.32, s	124.66, q,
(D)	⁵ J _{FC} 1.3		$^{3}J_{FC}$ 3.8	2 _{JFC} 32.4	$^{3}J_{FC}$ 3.8		¹ J _{FC} 272.6
10f	137.68, s	125.78, q,	131.79, q,	127.89, q,	132.26, q,	129.73, s	124.66, q,
(D)		$^{3}J_{FC}$ 3.8	$^{2}J_{FC}$ 32.6	³ J _{FC} 3.8	$^{4}J_{FC}$ 1.2		¹ J _{FC} 272.4
10g	151.61, q,	130.59, s	121.51, q,	129.83, q,	121.51, q,	130.59, s	121.11, q,
(D)	⁶ Ј _{FC} 1.8		$^{4}J_{FC}$ 1.0	$^{3}J_{FC}$ 89.4	$^{4}J_{FC}$ 1.0		¹ J _{FC} 257.8
10h	138.37, s	129.92, s	149.58, q,	126.94, s	123.02, s	120.43, s	120.49, q,
(L)			$^{3}J_{FC}$ 1.9				¹ J _{FC} 257.5
10i	111.34, dd,	146.05, dm,	137.77, dm,	141.89, dm,	137.77, dm,	146.05, dm,	-
(L)	² J _{FC} 11.6,	$^{1}J_{FC}$ 255.4,	¹ J _{FC} 249.8,	$^{1}J_{FC}$ 268.1,	¹ J _{FC} 249.8,	¹ J _{FC} 255.4,	
	overlapping	overlapping	overlapping	overlapping		overlapping	

N 5.17, found: C 36.62, H 3.54, N 5.40%.

α-Amino-(pentafluorophenyl-)methanephosphonic Acid 6i

Yield: 18%, m.p. $268-270^{\circ}$ C, C, H, N analysis: $C_7H_5F_5NO_3P$, M.Wt. = 277.09, requires: C 30.34, H 1.82, N 5.06, found: C 31.04, H 2.15, N 5.38%.

FAB ms (thioglycerol), m/z [%]: 555 (2M + H, 40.2), 386 (M + H + T, 2.7), 278 (M + H, 74.6), 196 (M + H - H₃PO₃, 100), 178 (67.9), 130 (72.7).

Preparation of Fluorinated Dialkyl N-Diphenylmethylaminoarylmethanephosphonates by Method C (Addition of Substances with P—H Bonds to N-Protected Imines, Variation b)) (Scheme IV)

Fluorinated N-Benzylidene-1,1-diphenylmethylamines 10a-g: 0.05 mol of the fluorinated benzaldehyde 7 was mixed under nitrogen with 0.05 mol diphenylmethylamine. A strongly exothermic reaction oc-

 $^{^{31}}P\{^{1}H\}$ δ : 17.29 (s).

 $^{^{1}}$ H (NaOD/D₂O), δ: 3.87 (d, $^{2}J_{PH}$ 16.2, 1H, PCH), 7.21 (m, 1H_{aroen}, 3-CF₃O—C₆H₄), 7.33–7.47 (m, 3H_{aroen}, 3-CF₃O—C₆H₄).

¹⁹F (NaOD/D₂O), δ : -57.62 (s).

FAB ms (3-NOBA), m/z [%]: 308 (100), 290 (70.8), 272 (M + H, 24.6), 254 (M + H - H_2O , 28.5), 189 (15.8), 166 (51.7).

 $^{^{31}}P\{^{1}H\}$ δ : 15.32 (m).

¹H (NaOD/D₂O), δ: 4.20 (d, ²J_{PH} 17.5, 1H, PCH).

¹⁹F (NaOD/D₂O), δ : -9.47 (m), -25.96 (m), -31.22 (m).

TABLE X Part II: $^{13}C\{^{1}H\}$ NMR data of the fluorinated N-benzylidene-1,1-diphenylmethylamines 10

No.	δC	N-CH	α-СН	C7/ 7'	C8/ 8'	C9/ 9'	C10/ 10'	C11/11'	C12/ 12'
10a		77.80, s	159.30, s	143.79, s	*	128.90, s	126.66, s	128.90, s	*
(L) 10a (D)		78.47, s	159.96, d, ⁵ J _{FC} 0.6	144.55, s	128.30, s	129,10, s	127.69, s	129,10, s	128.30, s
10b (D)		78.47, s		144.33, s	128.29, s	129.16, s	127.76, s	129.16, s	128.29, s
10c (L)		78.38, s	154.02, d, ³ J _{FC} 4.8	144.19, s	127.59, s	128.45, s	127.03, s	128.45, s	127.59, s
10d (D)		78.36, s	158.95, dd, ⁴ J _{FC} 2.5,	144.25, s	128.25, s	129.18, s	127.80, s	129.18, s	128.25, s
10e (L)		77.98, s	⁵ J _{FC} 1.7 159.37, s	143.49, s	127.61, s	128.55, s	127.19, s	128.55, s	127.61, s
10e (D)		78,65, s	160.02, d, 6 _{JFC} 0.4	144.19, s	128.31, s	129.22, s	127.86, s	129.22, s	128.31, s
10f (D)		78.60, s	159.94, s	144.18, s	128.33, s	129.22, s	127.85, s	129.22, s	128.33, s
10g (D)		78.54, s	159.76, s	144.36, s	128.31, s	129.17, s	127.78, s	129.17, s	128.31, s
10h (L)		77.86, s	159.18, s	143.54, s	127.61, s	128.52, s	127.14, s	128.52, s	127.61, s
10i (L)		79.78, s	149.10, d, ³ J _{FC} 2.5	143.02, s	127.46, s	128.63, s	127.37, s	128.63, s	127.46, s

TABLE XI
Part I: $^{13}\text{C}\{^{1}\text{H}\}$ NMR data of the fluorinated dimethyl N-diphenylmethylaminoarylmethanephosphonates 11a. $^{13}\text{C}\{^{1}\text{H}\}$ NMR (10% CDCl₃, TMS), δ [ppm], Mult., $^{1}J_{XY}$ [Hz]

No.	δC	CH ₃ (1)	CH ₃ (2)	СНР	CHN
		50.01.1	54.50 I		
llaa		53.91, d,	54.60, d,	57.64, d,	64.30, d,
(D)		2 J $_{PC}$ 7.0	² J _{PC} 6.9	1 J $_{PC}$ 156.2	³ J _{PC} 16.8
11ae		54.02, d,	54.76, d,	58.17, d,	64.60, d,
(D)		$^{2}J_{PC}$ 7.0	$^{2}J_{PC}$ 7.0	¹ J _{PC} 154.5	$^{3}J_{PC}$ 16.5

TABLE XI
Part II: 13C(1H) NMR data of the fluorinated dimethy
N-diphenylmethylaminoarylmethanephosphonates 11a

No.	δC	C1	C2	C3	C4	C5	C6	R
llaa		132.07. dd.	130.85. dd.	116.26. dd.	163.19, dd,	116.26. dd.	130.85. dd.	-
(D)		⁴ J _{FC} 2.0,		, ,	¹ J _{FC} 246.7,	, ,	, ,	
		2 _{JPC} 3.2	3 _{JPC} 6.4	⁴ J _{PC} 2.2	5 _{JPC} 3.2	⁴ J _{PC} 2.2	³ J _{PC} 6.4	
llae		141.07, dq,	129.62, d,	126.08, dq,	130.77, dq,	126.08, dq,	129.62, d,	124.78, dq,
(D)			⁴ J _{FC} 3.7,	$^{3}J_{FC}$ 32.4,	$^{2}J_{FC}$ 3.7,	$^{3}J_{FC}$ 32.4,	$^{4}J_{FC}$ 3.7,	¹ J _{FC} 272.1,
		² J _{PC} 1.3, overlapping	³ J _{PC} 6.0	⁴ J _{PC} 2.5	⁵ J _{PC} 3.1	⁴ J _{PC} 2.5	³ J _{PC} 6.0	⁶ J _{PC} 1.2

TABLE XI

Part III: ¹³C{¹H} NMR data of the fluorinated dimethyl
N-diphenylmethylaminoarylmethanephosphonates 11a

No.	δ _C	C7/ 7'	C8/ 8', C	9/ 9'	C 10/ 10'	C 11/11',	C 12/ 12'
llaa		142.36, s,	127.83, s,	129.12, s,	127.91, s,	127.83, s,	129.12, s,
(D)		144.05, s	128.38, s,	129.36, s	128.17, s	128.38, s,	129.36, s
11ae		142.19, s,	127.84, s,	129.20, s,	128.04, s,	127.84, s,	129.20, s,
(D)		143.81, s	128.37, s,	129.47, s	128.26, s	128.37, s,	129.47, s

curred, with the development of turbidity, and the solution became light yellow. The condensation reaction was completed by heating on a boiling water bath (4-10 h) to give a clear, yellow solution, with water droplets separating on the upper part of the flask. After cooling, the oily reaction mixture was dissolved in dichloromethane and dried over Na₂SO₄. Filtration, followed by rotary evaporation of the filtrate gave a yellow oil which solidified after 1-3 days at room temperature to give a waxy, light yellow solid (Table V).

N-4-Fluorobenzylidene-1,1-diphenylmethylamine 10a

 1 H NMR (6% CDCl₃, TMS), δ: 5.57 (bs, 1H, CHPh₂), 7.0–7.11 (m, 2H_{arom.}, 4-F—C₆H₄—C), 7.16–7.41 (m, 10H_{arom.}, 2 × C₆H₃), 7.71–7.88 (m, 2H_{arom.}, 4-F—C₆H₄—C), 8.35 (bs, 1H, —CH). 19 F NMR (6% CDCl₃, C₆F₆ int.), δ: 52.26–52.51 (m, 1F, 4-F—C₆H₄—C).

N-3-Fluorobenzylidene-1,1-diphenylmethylamine 10b

C, H, N analysis: $C_{20}H_{10}FN$, M.Wt. = 289.35, requires: C 83.02, H 5.57, N 4.84, found: C 82.71, H 5.41, N 4.70.%.

 1 H δ: 5.60 (bs, 1H, CHPh₂), 7.05–7.15 (m, 1H_{arom.}, 3-F—C₆H₄—C), 7.18–7.42 (m, 10H_{arom.}, 2 × C₆H₅ + 1H_{arom.}, 3-F—C₆H₄—C), 7.49–7.55 (m, 1H_{arom.}, 3-F—C₆H₄—C), 7.59–7.66 (m, 1H_{arom.}, 3-F—C₆H₄—C), 8.38 (bs, 1H, —CH).

¹⁹F δ: 48.69-48.82 (m, 1F, $3-F-C_6H_4-C$).

N-3,4-Difluor obenzylidene-1,1-diphenylmethylamine 10d

C, H, N analysis: $C_{20}H_{15}F_2N$, M.Wt. = 307.34, requires: C 78.16, H 4.92, N 4.56, found: C 78.12, H 4.84, N 4.44.%.

TABLE XII

Part I: $^{13}C\{^{1}H\}$ NMR data of the fluorinated diethyl N-diphenylmethylaminoarylmethanephosphonates 11b. $^{13}C\{^{1}H\}$ NMR (10% CDCl₃, TMS), δ [ppm], Mult., $^{17}J_{XY}$ [Hz]

No.	δC	CH ₃ (1)	CH ₃ (2)	CH ₂ (3)	CH ₂ (4)	СНР	CHN
11ba		16.83, d,	17.16, d,	63.30, d,	63.72, d,	57.92, d,	64.35, d,
(D)		$^{3}J_{PC}$ 5.9	3 _{JPC} 6.1	² J _{PC} 6.9	$^{2}J_{PC}$ 7.0	¹ J _{PC} 155.9	$^{3}J_{PC}$ 16.7
11bb		16.80, d,	17.13, d,	63.36, d,	63.77, d,	58.35, dd,	64.48, d,
(D)		³ J _{PC} 5.9	³ J _{PC} 6.0	² J _{PC} 6.9	² J _{PC} 7.0	¹ J _{PC} 154.8, ⁴ J _{FC} 1.7	³ J _{PC} 16.5
11bd		16.90, d,	17.18, d,	63.42, d,	63.87, d,	57.86, dd,	64.53, d,
(D)		³ J _{PC} 5.8	³ J _{PC} 6.0	² J _{PC} 6.9	² J _{PC} 7.0	¹ J _{PC} 155.4, ⁴ J _{FC} 1.2	³ J _{PC} 16.2
11be		16.82, d,	17.15, d,	63.44, d,	63.93, d,	58.48, d,	64.61, d,
(D)		$^{3}J_{PC}$ 5.9	3 JPC $^{6.0}$	$^{2}J_{PC}$ 7.0	$^{2}J_{PC}$ 7.0	¹ J _{PC} 153.9	$^{3}J_{PC}$ 16.3
11bf		16.76, d,	17.09, d,	63.50, d,	63.90, d,	58.52, d,	64.70, d,
(D)		$^{3}J_{PC}$ 6.0	$^{3}J_{PC}$ 6.0	$^{2}J_{PC}$ 7.0	$^{2}J_{PC}$ 7.0	$^{1}J_{PC}$ 154.3	³ J _{PC} 16.1
11bg		16.79, d,	17.15, d,	63.38, d,	63.84, d,	58.05, d,	64.52, d,
(D)		³ J _{PC} 5.9	$^{3}J_{PC}$ 6.0	$^{2}J_{PC}$ 7.0	$^{2}J_{PC}$ 7.0	$^{1}J_{PC}$ 155.1	³ J _{PC} 16.4

 1 H δ: 5.58 (bs, 1H, CHPh₂), 7.09–7.51 (m, 10H_{arom.}, 2 × C₆H₅ + 2H_{arom.}, 3,4-F₂—C₆H₃—C), 7.69–7.82 (m, 1H_{arom.}, 3,4-F₂—C₆H₃—C), 8.31 (bs, 1H, —CH). 19 F δ: 24.52–24.75 (m, 1F, 3,4-F₂—C₆H₃—C), 27.75–27.98 (m, 1F, 3,4-F₂—C₆H₃—C).

N-4-Trifluoromethylbenzylidene-1,1-diphenylmethylamine 10e

 1 H δ: 5.62 (bs, 1H, CHPh₂), 7.18–7.42 (m, 10H_{arom.}, 2 × C₆H₅), 7.60–7.67 (m, 2H_{arom.}, 4-CF₃—C₆H₄—C), 7.88–7.95 (m, 2H_{arom.}, 4-CF₃—C₆H₄—C), 8.43 (bs, 1H, ==CH). 19 F δ: 98.97 (bs, 3F, 4-CF₃—C₆H₄—C).

N-3-Trifluoromethylbenzylidene-1,1-diphenylmethylamine 10f

C, H, N analysis: $C_{21}H_{16}F_3N$, M.Wt. = 339.36, requires: C 74.33, H 4.75, N 4.13, found: C 74.16, H 4.67, N 4.11.%.

 1 H δ: 5.63 (bs, 1H, CHPh₂), 7.19 – 7.43 (m, 10H_{arom.}, 2 × C₆H₅), 7.46 – 7.56 (m, 1H_{arom.}, 3-CF₃—C₆H₄—C), 7.62 – 7.69 (m, 1H_{arom.}, 3-CF₃—C₆H₄—C), 7.97 – 8.02 (m, 1H_{arom.}, 3-CF₃—C₆H₄—C), 8.09 – 8.11 (m, 1H_{arom.}, 3-CF₃—C₆H₄—C), 8.44 (bs, 1H, —CH). 19 F δ: 99.09 (bs, 3F, 3-CF₃—C₆H₄—C).

N-4-Trifluoromethoxybenzylidene-1, I-diphenylmethylamine 10g

C, H, N analysis: $C_{21}H_{16}F_3NO$, M.Wt. = 355.36, requires: C 70.98, H 4.54, N 3.94, found: C 70.73, H 4.49, N 3.92.%.

¹H δ: 5.60 (bs, 1H, CHPh₂), 7.17–7.42 (m, $10H_{arom.}$, $2 \times C_6H_5 + 2H_{arom.}$, 4-CF₃O—C₆H₄—C), 7.82–7.89 (m, $2H_{arom.}$, 4-CF₃O—C₆H₄—C), 8.39 (bs, 1H, —CH). ¹⁹F δ: 104.05 (bs, 3F, 4-CF₃O—C₆H₄—C).

Fluorinated Dimethyl N-Diphenylmethylaminoarylmethanephosphonates 11aa-ae: 0.02 mol of the fluorinated N-benzylidene-1,1-diphenylmethylamine 10 and 0.03 mol dimethyl phosphite were mixed together under an atmosphere of nitrogen. An endothermic reaction occurred and a yellowish suspension was formed. In order to complete the phosphite-addition reaction the mixture was heated at 100°C (4-5 h) to give a clear, yellowish, oily liquid. Cooling in an ice/water bath gave a highly viscous oil with a white solid. The suspension was dissolved in hot methanol. After cooling the solution was placed in a refrigerator: a white precipitate of the nitrogen protected dimethyl aminophosphonate 11a formed after a few days, which was filtered off (using a G3-sintered glass filter), washed with ether, dried in a vacuum desiccator over silica gel, and purified by recrystallization from methanol/ether mixtures (Table VI).

Dimethyl N-Diphenylmethylamino-(4'-fluorophenyl-)methanephosphonate 11aa C, H, N analysis: C₂₂H₂₃FNO₃P, M.Wt. = 399.40, requires: C 66.16, H 5.80, N 3.51, found: C 65.93, H 5.94, N 3.62%.

TABLE XII Part II: 13C{1H} NMR data of the fluorinated diethyl N-diphenylmethylaminoarylmethanephosphonates 11b

No.	δ _C	C1	C2	C3	C4	C5	C6	R
11ba		132.36, dd,	130.88, dd,	116.10, dd ,	162.70, dd,	116.10, dd ,	130.88, dd,	-
(D)		$^{4}J_{FC}$ 2.0,	$^{3}J_{FC}$ 8.1,	$^{2}J_{FC}$ 21.5,	¹ J _{FC} 246.4,	$^{2}J_{FC}$ 21.5,	$^{3}J_{FC}$ 8.1,	
		$^{2}J_{PC}$ 3.2	$^{3}J_{PC}$ 6.3	⁴ J _{PC} 2.2	⁵ J _{PC} 3.6	⁴ J _{PC} 2.2	$^{3}J_{PC}$ 6.3	
11bb		139.48, dd,	116.11, dd,	163.57, dd,	115.51, dd,	130.59, dd,	124.94, dd,	-
(D)		$^{3}J_{FC}$ 6.8,	² J _{FC} 21.9,	1 _{JFC} 246.6,	$^{2}J_{FC}$ 21.2,	$^{3}J_{FC}$ 8.1,	⁴ J _{FC} 2.9,	
		$^{2}J_{PC}$ 2.0	³ J _{PC} 6.1	$^{4}J_{PC}$ 2.5	⁵ J _{PC} 3.0	$^{4}J_{PC}$ 2.4	³ J _{PC} 6.3	
11bd		133.97, ddd,	118.12, ddd,				125.31, ddd,	-
(D)		⁴ J _{FC} 3.8,		$^{2}J_{FC}$ 12.4,	$^{2}J_{FC}$ 12.4,		⁴ J _{FC} 2.8,	
			$^{2}J_{FC}$ 17.8,	l _{JFC} 248.3,	1 _{JFC} 248.5,	$^{2}J_{FC}$ 17.3,	$^{3}J_{FC}$ 6.4,	
		² J _{PC} 2.3	³ J _{PC} 5.9, overlapping	⁴ J _{PC} 3.3	⁵ J _{PC} 2.6	⁴ J _{PC} 2.3, overlapping	³ J _{PC} 3.6	
11be		140.72, dq,	129.59, d,	126.27, dq,	130.96, dq,	126.27, dq,	129.59, d,	124.74, dq,
(D)				$^{3}J_{FC}$ 3.8,	$^{2}J_{FC}$ 32.5,	$^{3}J_{FC}$ 3.8,		¹ J _{FC} 272.1,
		$2J_{PC} = 1.3$, overlapping	³ J _{PC} 6.1		⁵ J _{PC} 3.1	⁴ J _{PC} 2.4	³ J _{PC} 6.1	6J _{PC} 1.3
11bf		138.01, d,	126.24, dq,	131.48, dq,	125.37, dq,	132.46, dq,	129.66, d,	124.72, dq,
(D)			$^{3}J_{FC}$ 3.8,	$^{2}J_{FC}$ 32.3,	$^{3}J_{FC}$ 3.7,	⁴ J _{FC} 5.7,		$^{1}J_{FC}$ 272.3,
		$^{2}J_{PC}$ 2.3	$^{3}J_{PC}$ 6.4	$^{4}J_{PC}$ 2.4	⁵ J _{PC} 3.2	$^{4}J_{PC}$ 1.2	$^{3}J_{PC}$ 2.4	$^{5}J_{PC}$ 0.5
11bg		135.47, d,	130.67, d,	121.54, dq,	149.56, dq,	121.54, dq,	130.67, d,	121.14, dq,
(D)					$^{3}J_{FC}$ 3.5,			¹ J _{FC} 257.2,
		² J _{PC} 2.1	³ J _{PC} 6.2	⁴ J _{PC} 1.1, overlapping		⁴ J _{PC} 1.1, overlapping	³ J _{PC} 6.2	⁷ J _{PC} 1.1

 $^{31}P\{^{4}H\}$ NMR (6% CDCl₃, 85% H₃PO₄ ext.), δ : 26.49 (d, $^{4}J_{PF}$ 4.8). ^{1}H NMR (6% CDCl₃, TMS), δ : 2.50 (bs, 1H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, CH₃), 3.86 (d, $^{3}J_{PH}$ 10.6, 3H, NH), 3.51 (d, $^{3}J_{PH}$ 10.5, 3H, NH), 3.51 (d, $^{3}J_$ CH₃), 3.95 (d, ${}^{2}J_{PH}$ 21.6, 1H, PCH), 4.64 (d, ${}^{3}J_{HH}$ 1.2, 1H, CHPh₂), 7.00 – 7.13 (m, 2H_{wom.}, 4-F— C_6H_4 —C), 7.13–7.37 (m, 10 H_{arom} , 2 × C_6H_5 + 2 H_{arom} , 4-F— $\overline{C_6H_4}$ —C). ¹⁹F NMR (6% CDCl₃, C_6F_6 int.), 8: 47.71–47.89 (m, 1F, 4-F— C_0H_4 —C).

Dimethyl N-Diphenylmethylamino-(4'-trifluoromethylphenyl-)methanephosphonate 11ae ³¹P{¹H} δ : 25.72 (q, ⁷ J_{PF} 2.3).

¹H δ : 3.55 (d, ³ J_{PH} 10.6, 3H, CH₃), 3.86 (d, ³ J_{PH} 10.6, 3H, CH₃), 4.05 (d, ² J_{PH} 22.7, 1H, PCH), 4.63 (d, $^{3}J_{HH}$ 1.1, 1H, CHPh₂), 7.14–7.36 (m, 10H_{arom.}, 2 × C₆H₅), 7.43–7.49 (m, 2H_{arom.}, 4-CF₃—C₆H₄—C), 7.61 – 7.67 (m, $\overline{2H}_{arom}$, 4-CF₃—C₆H₄—C). ¹⁹F δ: 99.24 (bs, 3F, 4-CF₃— C_6H_4 —C).

Fluorinated Diethyl N-Diphenylmethylaminoarylmethanephosphonates 11ba - bg: 0.02 mol of the fluorinated N-benzylidene-1,1-diphenylmethylamine 10 was mixed with 0.02 mol diethyl phosphite under nitrogen; with slight cooling a pale yellow suspension resulted. The reaction was completed by heating (8-14 h) at 100°C when an intensively yellow solution was formed. Cooling gave a highly viscous oil, which was dissolved in methanol. After concentration on a rotary evaporator at water pump pressure, the solution was placed in a refrigerator to give the nitrogen protected diethyl aminophosphonate 11b as a waxy, pale yellow, solid precipitate, which was filtered off (using a G3-sintered glass filter), washed with cooled ether, dried in a vacuum desiccator over silica gel, and purified by recrystallization from methanol/ether mixtures (Table VII).

Diethyl N-Diphenylmethylamino-(4'-fluorophenyl-)methanephosphonate 11ba C, H, N analysis: $C_{24}H_{27}FNO_3P$, M.Wt. = 427.45, requires: C 67.44, H 6.37, N 3.28, found: C 67.14, H 6.51, N 3.38%.

³¹P{¹H} NMR (6% CDCl₃, 85% H₃PO₄ ext.), δ : 24.13 (d, ${}^6J_{PF}$ 4.8).

TABLE XII Part III: 13C{1H} NMR data of the fluorinated diethyl N-diphenylmethylaminoarylmethanephosphonates 11b

No.	δС	C7/ 7'	C8/ 8', C9/ 9'	C10/ 10'	C11/11', C12/12'
			10001		
11ba		142.52, s,	127.84, s, 129.07, s,	127.84, s,	127.84, s, 129.07, s,
(D)		144.15, s	128.42, s, 129.28, s	128.03, s	128.42, s, 129.28, s
11bb		142.45, s,	127.83, s, 129.07, s,	127.82, s,	127.83, s, 129.07, s,
(D)		144.05, s	128.40, s, 129.28, s	128.05, s	128.40, s, 129.28, s
11bd		142.35, s,	127.85, s, 129.15, s,	127.96, s,	127.85, s, 129.15, s,
(D)		143.92, s	128.39, s, 129.40, s	128.18, s	128.39, s, 129.40, s
11be		142.37, s,	127.84, s, 129.13, s,	127.94, s,	127.84, s, 129.13, s,
(D)		143.92, s	128.39, s, 129.37, s	128.16, s	128.39, s, 129.37, s
11bf		142.31, s,	127.90, s, 129.13, s,	127.94, s,	127.90, s, 129.13, s,
(D)		143.88, s	128.39, s, 129.38, s	128.18, s	128.39, s, 129.38, s
11bg		142.50, s,	127.87, s, 129.12, s,	127.90, s,	127.87, s, 129.12, s,
(D)		144.05, s	128.43, s, 129.34, s	128.12, s	128.43, s, 129.34, s

 1 H NMR (6% CDCl₃, TMS), δ: 1.20 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5, 3H, CH₃), 1.35 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5, 3H, CH₃), 3.76–4.04 (m, 2H, CH₂), 4.07–4.34 (m, 2H, CH₂), 3.92 (d, $^{2}J_{PH}$ 22.2, 1H, PCH), 4.65 (d, $^{3}J_{HH}$ 1.1, 1H, CHPh₂), 6.99-7.13 (m, 2H_{arom.}, 4-F—C₆H₄—C), 7.14-7.36 (m, 10H_{arom.}, $2 \times C_6H_5 + 2H_{arom.}$ $4-F-C_6H_4-C$). ¹⁹F NMR (6% CDCl₃, C_6F_6 int.), δ : 47.37–47.55 (m, 1F, 4-F— C_6H_4 —C).

Diethyl N-Diphenylmethylamino-(3'-fluorophenyl-)methanephosphonate 11bb

C, H, N analysis: C₂₄H₂₇FNO₃P, M.Wt. = 427.45, requires: C 67.44, H 6.37, N 3.28, found: C 67.37, H 6.32, N 3.28%.

 $^{31}P\{^{1}H\}$ δ : 23.68 (d, $^{5}J_{PF}$ 1.7).

 ^{1}H 8: 1.11 (dt, $^{3}J_{\text{HH}}$ 7.1, $^{4}J_{\text{PH}}$ 0.5, 3H, CH₃), 1.35 (dt, $^{3}J_{\text{HH}}$ 7.1, $^{4}J_{\text{PH}}$ 0.5, 3H, CH₃), 3.69 – 4.06 (m, 2H, CH₂), 4.07-4.34 (m, 2H, CH₂), 3.93 (d, ²J_{PH} 22.4, 1H, PCH), 4.67 (d, ³J_{HH} 1.0, 1H, CHPh₂), 7.04-7.42 $(m, 10H_{arom.}, 2 \times C_6H_5 + 4H_{arom.}, 3-F-C_6H_4-C)$ ¹⁹F δ: 48.70–48.84 (m, 1F, 3-F— C_6H_4 —C).

Diethyl N-Diphenylmethylamino-(3',4'-difluorophenyl-)methanephosphonate 11bd C, H, N analysis: $C_{24}H_{26}F_2NO_3P$, M.Wt. = 445.44, requires: C 64.71, H 5.88, N 3.14, found: C 64.55, H 5.91, N 3.18%.

 $^{31}P\{^{1}H\}$ δ : 23.43 (dd, $^{6}J_{PF}$ 5.1, $^{5}J_{PF}$ 1.8).

 1 H δ : 1.15 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5, 3 H, CH₃), 1.35 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5 Hz, 3H, CH₃), 3.74–4.08 (m, 2H, CH₂), 4.07-4.33 (m, 2H, CH₂), 3.90 (d, ²J_{PH} 22.2, 1H, PCH), 4.65 (d, ³J_{HH} 0.9, 1H, CHPh₂), 7.00-7.56 (m, $10H_{aron.}$, $2 \times C_6H_5 + 3H_{aron.}$, $3,4 \cdot F_2 - C_6H_3 - C$). ^{19}F δ : 23.02 - 23.28 (m, 1F, $3,4 \cdot F_2 - C_6H_3 - C$), 24.57 - 24.80 (m, 1F, $3,4 \cdot F_2 - C_6H_3 - C$).

Diethyl N-Diphenylmethylamino-(4'-trifluoromethylphenyl-)methanephosphonate 11be

 $^{31}P\{^{1}H\} \delta: 23.39 \text{ (q, } ^{7}J_{PF} 2.3).$ $^{1}H \delta: 1.12 \text{ (dt, } ^{3}J_{HH} 7.1, ^{4}J_{PH} 0.5, 3H, CH_{3}), 1.35 \text{ (dt, } ^{3}J_{HH} 7.1, ^{4}J_{PH} 0.5, 3H, CH_{3}), 3.72-4.07 \text{ (m, 2H, CH₂), 4.07-4.34 (m, 2H, CH₂), 4.02 (d, ^{2}J_{PH} 22.6, 1H, PCH), 4.64 (bs, 1H, CHPh₂), 7.20-7.56 (m, 10H_{arom.}, 2 × C₆H₅ + 2H_{arom.}, 4-CF₃—C₆H₄—C) 7.59-7.67 (m, 2H_{arom.}, 4-CF₃—C₆H₄—C).$

TABLE XIII Part I: ¹³C(¹H) NMR data of the fluorinated diphenyl N-diphenylmethylaminoarylmethanephosphonates 11c. $^{13}C\{^{1}H\}$ NMR (10% CDCl₃, TMS), δ [ppm], Mult., $^{13}J_{XY}$ [Hz]

No.	δ _C	CHP	CHN	C13/ 13'	C14/ 14'	C 15/ 15'	C16/ 16'	C17/ 17'	C 18/ 18'
11ca		57.64, d,	64.30, d,	150.99, d,	120.89, d,	130.18, d,	125.63, d,	130.18, d,	
(D)		¹ J _{PC} 156.2	$^{3}J_{PC}$ 16.8	² J _{PC} 9.6,	$^{3}J_{PC}$ 4.4,	⁴ J _{PC} 0.9,	⁵ J _{PC} 1.1,	⁴ J _{PC} 0.9,	$^{3}J_{PC}$ 4.4,
				151.53, d,	121.10, d,	130.32, d,	125.73, d,	130.32, d,	121.10, d,
				$^{2}J_{PC}$ 9.6	$^{3}J_{PC}$ 4.4	$^{4}J_{PC}$ 0.8	$^{5}J_{PC}$ 1.1	⁴ J _{PC} 0.8	³ J _{PC} 4.4
11cb		58.53, dd,	64.68, d,	150.97, d,	120.88, d,	130.20, d,	125.68, d,	130.20, d,	120.88, d,
(D)		¹ J _{PC} 158.3,	$^{3}J_{PC}$ 17.3	² J _{PC} 9.7,	3 JPC 4.4,	⁴ J _{PC} 0.9,	⁵ J _{PC} 1.2,	⁴ J _{PC} 0.9,	$^{3}J_{PC}$ 4.4,
		⁴ J _{FC} 1.8		151.49, d,	121.08, d,	130.33, d,	125.76, d,	130.33, d,	
				$^{2}J_{PC}$ 9.7	$^{3}J_{PC}$ 4.5	⁴ J _{PC} 0.9	$^{5}J_{PC}$ 1.1	4J _{PC} 0.9	³ J _{PC} 4.5
11cd		58.23, dd,	64.69, d,	150.92, d,	120.84, d,	130.30, d,	125.81, d,	130.30, d,	120.84, d,
(D)		¹ J _{PC} 158.6,	$^{3}J_{PC}$ 17.1	$^{2}J_{PC}$ 9.6,	$^{3}J_{PC}$ 4.4,	⁴ J _{PC} 0.9,	$^{5}J_{PC}$ 1.2,	⁴ J _{PC} 0.9,	$^{3}J_{PC}$ 4.4,
		⁴ J _{FC} 1.3		151.42, d,	121.04, d,	130.40, d,	125.88, d,	130.40, d,	121.04, d,
				$^{2}J_{PC}$ 9.6	$^{3}J_{PC}$ 4.5	⁴ J _{PC} 0.9	⁵ J _{PC} 1.1	⁴ J _{PC} 0.9	³ J _{PC} 4.5
11ce		58.66, d,	64.79, d,	150.93, d,	120.83, d,	130.26, d,	125.79, d,	130.26, d,	120.83, d,
(D)		$^{1}J_{PC}$ 157.0	³ J _{PC} 17.1	² J _{PC} 9.6,	3 JPC 4.4,	⁴ J _{PC} 0.9,	⁵ J _{PC} 1.2,	⁴ J _{PC} 0.9,	$^{3}J_{PC}$ 4.4,
				151.43, d,	121.07, d,	130.40, d,	125.88, d,	130.40, d,	121.07, d,
				$^{2}J_{PC}$ 9.7	$^{3}J_{PC}$ 4.5	4 _{JPC} 0.9	⁵ J _{PC} 1.1	⁴ J _{PC} 0.9	³ J _{PC} 4.5
11cf		58.61, d,	64.79, d,	150.93, d,	120.78, d,	130.28, d,	125.76, d,	130.28, d,	120.78, d,
(D)		¹ J _{PC} 157.3	$^{3}J_{PC}$ 17.0	² J _{PC} 9.6,	$^{3}J_{PC}$ 4.5,	⁴ J _{PC} 0.8,	⁵ J _{PC} 1.1,	⁴ J _{PC} 0.8,	$^{3}J_{PC}$ 4.5,
				151.40, d,	121.04, d,	130.41, d,	125.89, d,	130.41, d,	121.04, d,
				$^{2}J_{PC}$ 9.6	$^{3}J_{PC}$ 4.5	$^{4}J_{PC}$ 0.8	⁵ J _{PC} 1.1	⁴ J _{PC} 0.8	³ J _{PC} 4.5
11cg		58.25, d,	64.68, d,	150.97, d,	120.82, d,	130.19, d,	125.68, d,	130.19, d,	120.82, d,
(D)		¹ J _{PC} 158.0	$^{3}J_{PC}$ 17.2	² J _{PC} 9.6,	$^{3}J_{PC}$ 4.4,	⁴ J _{PC} 0.9,	$^{5}J_{PC}$ 1.2,	⁴ J _{PC} 0.9,	³ J _{PC} 4.4,
				151.47, d,	121.08, d,	130.35, d,	125.80, d,	130.35, d,	121.08, d,
				² J _{PC} 9.7	³ J _{PC} 4.5	4 _{JPC} 0.9	⁵ J _{PC} 1.1	⁴ J _{PC} 0.9	³ J _{PC} 4.5

¹⁹F δ: 99.27 (bs, 3F, 4-CF₃— C_6H_4 —C).

Diethyl N-Diphenylmethylamino-(3'-trifluoromethylphenyl-)methanephosphonate 11bf

 $^{31}P\{^{1}H\}$ δ : 23.30 (q, $^{6}J_{PF}$ 0.6). ^{1}H δ : 1.12 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5, 3H, CH₃), 1.34 (dt, $^{3}J_{HH}$ 7.1, $^{4}J_{PH}$ 0.5, 3H, CH₃), 3.73–4.07 (m, 2H, CH_2), 4.05-4.32 (m, 2H, CH_2), 4.02 (d, ${}^2J_{PH}$ 22.4, 1H, PCH), 4.65 (bs, 1H, $CHPh_2$), 7.18-7.64 (m, $10H_{arom.}$, $2 \times C_6H_5 + 4H_{arom.}$, $3-CF_3-C_6H_4-C$). ¹⁹F δ: 99.14 (bs, 3F, 3-CF₃—C₆H₄—C).

Diethyl N-Diphenylmethylamino-(4'-trifluoromethoxyphenyl-)methanephosphonate 11bg C, H, N analysis: C₂₅H₂₇F₃NO₄P, M.Wt. = 493.46, requires: C 60.85, H 5.51, N 2.84, found: C 60.70, H 5.65, N 2.87%.

 $^{31}P\{^{1}H\}$ δ : 23.79 (s).

¹H δ : 1.10 (dt, ³ J_{BH} 7.1, ⁴ J_{PH} 0.5, 3H, CH₃), 1.35 (dt, ³ J_{HH} 7.1, ⁴ J_{PH} 0.6, 3H, CH₃), 3.69–4.05 (m, 2H, CH₂), 4.07-4.33 (m, 2H, CH₂), 3.96 (d, ²J_{PH} 22.3, 1H, PCH), 4.66 (d, ³J_{HH} 0.9, 1H, CHPh₂), 7.13-7.40 $(m, 10H_{arom.}, 2 \times C_6H_5 + 4H_{arom.}, 4-CF_3O-C_6H_4-C).$ ¹⁹F δ: 103.96 (bs, 3F, 4-CF₃— C_6H_4 —C).

Fluorinated Diphenyl N-Diphenylmethylaminoarylmethanephosphonates 11ca-cg: 0.02 mol of the fluorinated N-benzylidene-1,1-diphenylmethylamine 10 and 0.02 mol diphenyl phosphite were mixed together under an atmosphere of nitrogen, when a weakly exothermic reaction occurred and a yellowish suspension was formed. In order to complete the phosphite-addition reaction the mixture was heated at 100°C (4-8 h), to give a clear, yellow, oily liquid. Solidification occurred on cooling to give a waxy,

TABLE XIII Part II: 13C(1H) NMR data of the fluorinated diphenyl N-diphenylmethylaminoarylmethanephosphonates 11c

No.	δC	<u>C1</u>	C2	C3	C4	C5	C6	
11ca (D)		131.28, dd,	131.24, dd, ³ J _{FC} 8.2,	116.43, dd, ² J _{FC} 21.6,	163.41, dd, 1 _{JFC} 247.3,	2	131.24, dd, ³ J _{FC} 8.2,	-
(D)		² J _{PC} 1.4, overlapping	³ J _{PC} 6.9	⁴ J _{PC} 2.2	⁵ J _{PC} 3.3	⁴ J _{PC} 2.2	³ J _{PC} 6.9	
11cb		138.31, dd,	116.49, dd,	163.68, dd,	116.09, dd,	130.94, dd,	125.23, dd,	-
(D)		$^{3}J_{FC}$ 6.8,	² J _{FC} 22.2,				⁴ J _{FC} 3.0,	
. ,		$^{2}J_{PC}$ 1.3	³ J _{PC} 6.6	⁴ J _{PC} 2.5				
11cd			118.50, ddd,			, 118.25, ddd		l, -
(D)		$^{3}J_{FC}$ 5.0,	$^{2}J_{FC}$ 17.4,		¹ J _{FC} 249.0,	$^{2}J_{FC}$ 17.0,	$^{3}J_{FC}$ 6.7,	
		$^{4}J_{FC}$ 3.8,		$^{2}J_{FC}$ 12.0,			⁴ J _{FC} 3.1,	
		$^{2}J_{PC}$ 1.5	$^{3}J_{PC}$ 7.0,	⁴ J _{PC} 2.5	⁵ J _{PC} 3.4	⁴ J _{PC} 0.6,	$^{3}J_{PC}$ 3.6,	
			overlapping				overlapping	
11ce		139.94, dq,	129.96, d,	126.38, dq,	131.31, dq,		129.96, d,	124.71, dq,
(D)		⁵ J _{FC} 1.3,		$^{3}J_{FC}$ 3.8,	$^{2}J_{FC}$ 32.5,	$^{3}J_{FC}$ 3.8,		$^{1}J_{FC}$ 272.2,
		$^{2}J_{PC}$ 1.3	3 J $_{PC}$ 6.6	⁴ J _{PC} 2.4	⁵ J _{PC} 3.2	4 J $_{PC}$ 2.4	$^{3}J_{PC}=6.6$	6 _{JPС} 3.2
11cf		136.91, d,	126.56, dq,	131.76, dq,	125.94, dq,		130.00, d,	124.60, dq,
(D)			$^{3}J_{FC}$ 3.8,	² J _{FC} 32.5,		⁴ J _{FC} 6.1,		$^{1}J_{FC}$ 272.5,
		$^{2}J_{PC}$ 1.4	$^{3}J_{PC}$ 7.2	$^{4}J_{PC}$ 2.3		⁴ J _{PC} 1.2	$^{3}J_{PC}$ 2.4	⁵ J _{PC} 0.6
					overlapping			
11cg		134.34, d,	131.03, d,	121.84, dq,	149.90, q,	121.84, dq,	131.03, d ,	121.13, dq,
(D)				⁴ J _{FC} 1.1,	$^{3}J_{FC}$ 1.8	⁴ J _{FC} 1.1,		l _{JFC} 257.3,
		$^{2}J_{PC}$ 1.3	$^{3}J_{PC}$ 6.7					7 _{JPC} 1.0
				overlapping		overlapping		

yellowish solid, which was dissolved by boiling under reflux in a mixture of ethanol and methanol. On cooling in an ice/water bath, the nitrogen protected diphenyl aminophosphonate 11c separated as a fine white powder which was filtered off, washed free of oil with cooled ether, and finally dried in a vacuum desiccator over silica gel. The product was purified by recrystallization form THF/ether mixtures (Table VIII).

Diphenyl N-Diphenylmethylamino-(4'-fluorophenyl-)methanephosphonate 11ca

C, H, N analysis: $C_{32}H_{27}FNO_3P$, M.Wt. = 523.54, requires: C 73.41, H 5.20, N 2.68, found: C 73.34, H 5.36, N 2.70%.

³¹P{¹H} NMR (6% CDCl₃, 85% H₃PO₄ ext.), δ : 16.78 (d, ⁶ J_{PF} 5.1).

¹H NMR (6% CDCl₃, TMS), δ: 4.21 (d, ²J_{PH} 22.5, 1H, PCH), 4.64 (d, ³J_{HH} 1.2, 1H, CHPh₂), 6.72–6.80 + 6.92-7.35 (m, $10H_{arom.}$, $CH(C_6H_5)_2$ + $10H_{arom.}$, $P(OC_6H_5)_2$ + $4H_{arom.}$, 4-F— C_6H_4 —C). ¹⁹F NMR (6% CDCl₃, C_6F_6 int.), $\overline{\delta}$: 48.24-48.42 (m, 1F, 4-F— C_6H_4 —C).

Diphenyl N-Diphenylmethylamino-(3'-fluorophenyl-)methanephosphonate 11cb C, H, N analysis: $C_{32}H_{27}FNO_3P$, M.Wi. = 523.54, requires: C 73.41, H 5.20, N 2.68, found: C 73.30, H 5.30, N 2.76%.

³¹P{¹H} δ : 16.33 (d, ⁵ J_{PF} 1.8).

 1 H δ : 2.75 (bs, 1H, NH), 4.31 (d, $^{2}J_{PH}$ 22.8, 1H, PCH), 4.75 (d, $^{3}J_{HH}$ 1.1, 1H, CHPh₂), 6.83-6.91 + 6.99–7.40 (m, $10H_{arom}$, $CH(C_6H_5)_2 + 10H_{arom}$, $P(OC_6H_5)_2 + 4H_{arom}$, $3-F-C_6H_4-C$). ¹⁹F δ : 49.46–49.59 (m, 1F 3-F- C_6H_4-C).

Diphenyl N-Diphenylmethylamino-(3',4'-difluorophenyl-)methanephosphonate 11cd C, H, N analysis: $C_{32}H_{26}F_2NO_3P$, M.Wt. = 541.53, requires: C 70.97, H 4.84, N 2.59, found: C 70.77, H 5.00, N 2.65%.

TABLE XIII Part III: 13C{1H} NMR data of the fluorinated diphenyl N-diphenylmethylaminoarylmethanephosphonates 11c

No.	δ _C C7/7'	C8/ 8', C9/ 9'	C10/ 10'	C11/11', C12/12'
11ca	142.36, s,	127.83, s, 129.12, s,	127.91, s,	127.83, s, 129.12, s,
(D)	144.05, s	128.38, s, 129.36, s	128.17, s	128.38, s, 129.36, s
11cb	142.09, s,	127.81, s, 128.50, s,	128.03, s,	127.81, s, 128.50, s,
(D)	143.82, s	129.19, s, 129.41, s	128.23, s	129.19, s, 129.41, s
11cd	141.96, s,	127.80, s, 129.24, s,	128.12, s,	127.80, s, 129.24, s,
(D)	143.66, s	128.46, s, 129.50, s	128.34, s	128.46, s, 129.50, s
11ce	142.01, s,	127.82, s, 129.24, s,	128.12, s,	127.82, s, 129.24, s,
(D)	143.69, s	128.48, s, 129.49, s	128.33, s	128.48, s, 129.49, s
11cf	141.94, s,	127.85, s, 129.25, s,	128.12, s,	127.85, s, 129.25, s,
(D)	143.65, s	128.47, s, 129.50, s	128.35, s	128.47, s, 129.50, s
11cg	142.12, s,	127.82, s, 128.48, s,	128.05, s,	127.82, s, 128.48, s,
(D)	143.78, s	129.19, s, 129.43, s	128.25, s	129.19, s, 129.43, s

 $^{31}P\{^{1}H\} \ \delta: \ 15.91 \ (dd, \, ^{6}J_{PF} \ 5.4, \, ^{5}J_{PF} \ 1.8).$ $^{1}H \ \delta: \ 2.59 \ (bs, \ 1H, \ NH), \ 4.19 \ (d, \, ^{3}J_{PH} \ 22.4, \ 1H, \ PCH), \ 4.65 \ (d, \, ^{3}J_{HH} \ 1.1, \ 1H, \ C\underline{H}Ph_{2}), \ 6.79-7.31 \ (m, \, 1.1)$ $10H_{arom.}$, $CH(C_6H_5)_2 + 10H_{arom.}$, $P(OC_6H_5)_2 + 3H_{arom.}$, 3,4- F_2 — C_6H_3 —C). ¹⁹F δ : 23.90–24.17 (m, 1F 3,4-F₂—C₆H₃—C), 25.14–25.39 (m, 1F, 3,4-F₂—C₆H₃—C).

Diphenyl N-Diphenylmethylamino-(4'-trifluoromethylphenyl-)methanephosphonate 11ce ³¹P{¹H} δ : 15.95 (q, ⁷ J_{PF} 2.3).

 1 H δ : 4.39 (d, $^{2}J_{PH}$ 22.9, 1H, PCH), 4.72 (d, $^{3}J_{HH}$ 0.7, 1H, CHPh₂), 6.81–6.89 + 7.03–7.39 (m, 1 0H_{arom.}) $CH(C_6H_5)_2 + 10H_{arom.}$, $P(OC_6H_5)_2$, 7.51-7.67 (m, $4H_{arom.}$, $4-CF_3-C_6H_4-C$). ¹⁹F δ: $\overline{99}.15$ (bs, 3F 4-CF₃— \overline{C}_6H_4 —C)

Diphenyl N-Diphenylmethylamino-(3'-trifluoromethylphenyl-)methanephosphonate 11cf ³¹P{¹H} δ : 15.87 (q, ⁶ J_{PF} 0.7).

¹H δ : 4.29 (d, ² J_{PH} 22.8, 1H, PCH), 4.63 (bs, 1H, CHPh₂), 6.74-6.82 + 6.91-7.23 (m, 10H_{arom.}, $CH(C_6H_5)_2 + 10H_{arom.}$, $P(OC_6H_5)_2$, 7.37-7.57 (m, $4H_{arom.}$, $3-CF_3-C_6H_4-C$). ¹⁹F δ : $\overline{99}$.16 (bs, 3F, 3-CF₃— \overline{C}_6H_4 —C)

Diphenyl N-Diphenylmethylamino-(4'-trifluoromethoxyphenyl-)methanephosphonate 11cg C, H, N analysis: C₃₃H₂₇F₃NO₄P, M.Wt. = 589.55, requires: C 67.23, H 4.62, N 2.38, found: C 67.30, H 4.69, N 2.38%. ³¹P{¹H} δ : 16.37 (s).

¹H δ : 4.24 (d, ² J_{PH} 22.6, 1H, PCH), 4.65 (d, ³ J_{HH} 0.7, 1H, CHPh₂), 6.67–6.75 + 6.90–7.38 (m, 10H_{arom.}, $CH(C_6H_5)_2 + 10H_{arom.}, P(OC_6H_5)_2 + 4H_{arom.}, 4-CF_3-C_6H_4-C).$ ¹⁹F δ: $\overline{104.01}$ (bs, 3F, 4-CF₃O—C₆H₄—C).

¹³C-NMR-data of compounds 6, 10, and 11 are listed in Tables IX-XIII.

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