1,5-Phosphonium betaines from N-triflylpropiolamides, triphenylphosphane, and active methylene compounds

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Abstract

\(N\)-Phenyl-\(N\)-(trifluoromethylsulfonyl)propiolamides react with triphenylphosphane in the presence of various active methylene compounds \(\text{CH}_2\text{XY}\) in a 1:1:1 molar ratio to furnish 1-phosphonium-5-oxabetaines, \(\text{Ph}_3\text{P}^+\text{–C} (\text{R}) = \text{CH} = \text{C} (\text{O}^−) = \text{CXY}\). These betaines are formed preferentially, but not exclusively, as \(E\)-diastereoisomers with respect to the vinylic double bond. In some cases, separation of the two diastereoisomers was achieved by fractionating crystallization. Structure determination by X-ray diffraction analysis revealed marked conformational differences around the \(\text{CH} – \text{C} (\text{O}^−)\) single bond of \(E\) and \(Z\)-isomers and extended charge delocalization in the anionic part.

Introduction

Beside the well-known phosphonium ylides (Wittig ylides, methyleneephosphoranes), various other types of zwitterions containing a tetravalent phosphonium moiety (phosphonium betaines) exist. They are often considered as reaction intermediates, but reports on their isolation and structural characterization are still less common. A very convenient access to such betaines is provided by the nucleophilic addition of tertiary phosphanes to electron-deficient alkenes, alkynes and allenes (phospha-Michael addition), which generates the betaines as reactive intermediates that can be transformed intra- or intermolecularly into a wide array of acyclic, carbocyclic and heterocyclic structures. In these processes, the tertiary phosphanes can act as nucleophilic organocatalysts (for reviews, see [1-4] or be incorporated in the (pre-)final products, typically as alkylidene phosphoranes which in turn may undergo Wittig olefination with elimination of \(\text{Ph}_3\text{P}=\text{O}\) [1,5]. Recently, a study of the tributylphosphane-catalyzed reaction of ethyl 2-butyroate and ethanol by in-situ ESIMS and NMR techniques has been published [6], and mechanistic aspects of trialkylphosphane-catalyzed reactions of acetylenic ketones and esters with pinacolborane have been discussed [7,8].

The Michael addition of \(\text{PPH}_3\) at acetylenic carbonyl compounds generates phosphonium/vinyl anion intermediates which
Scheme 1: Stable betaines I–IV.

Results and Discussion

Synthesis of betaines 3

The combination of equimolar amounts of triphenylphosphane and N-triflyl-propiolanilide 1a in dichloromethane yielded a product mixture, from which only N-phenyltriflylamide (HN(Ph)Tf; $^{19}$F NMR: $\delta$ 76.5 ppm rel. to C$_6$F$_6$) could be extracted, the remainder being undefined oligomeric/polymeric material. However, when an N-triflylpropiolamide 1a–e (1.03 molar equivalents) was added to a solution of PPh$_3$ and an active methylene compound 2a–c,e,f, a 1:1:1 three-component reaction occurred smoothly within 0.5–6 hours and yielded the phosphonium-1,5-betaines 3a–o in mostly high yields (Scheme 3 and Figure 1). Meldrum’s acid (2a), $N,N’$-dimethyl-barbituric acid (2b), 1,3-indanedione (2c), malononitrile (2e), 4-cyanomethyl-2,3,5,6-tetrafluorobenzonitrile (2f) were successfully applied as active methylene compounds; all of them have been trapped with CH-, NH-, OH- and SH-acids [5]. In this manner, the 1:1:1 reaction of triphenylphosphane with alkyl propiolates [9,10] or dialkyl acetylenedicarboxylates [11] and strong CH-acids, such as Meldrum’s acid, 1,3-dimethylbarbituric acid, dimedone and indane-1,3-dione, leads to stable 1,4-dionic phosphonium betaines I or II (Scheme 1) in high yields. An analogous reaction has been achieved with 1,1,1,5,5,5-hexafluoropentane-2,4-dione as the active methylene component [12]. Some of these three-component reactions were even performed in aqueous media [13]. In mechanistic terms, the initially formed 1,3-betaine is C-protonated by the CH-acid, the anion of which then combines with the vinylphosphonium ion, and a proton transfer finally yields the 1,4-betaine. A kinetic study of the PPh$_3$/DMAD/Meldrum’s acid reaction by spectrophotometric and stopped-flow methods has been published [14]. The initial 1,3-betaine has also been trapped with other electrophiles. The stable 1,4-betaines betaines III (PR$_3$ = alkyl-substituted phosphane, but not PPh$_3$) have been obtained from isonicotinaldehyde [15], and betaines IV from aryl tosylimines [16]. For the stability of betaines III, the nature of the aryl substituent is critical; with the combination P$_3$H$_2$/DMAD/EWG-substituted benzaldehydes, cyclization leading to butenolides occurs easily [17].
Scheme 3: Synthesis of betaines by a three-component reaction. N-triflylpropiolamides 1: R = Ph (a), 4-Cl-C₆H₄ (b), t-Bu (c), n-Bu (d), cyclopropyl (e), SiMe₃ (f). See Figure 1 for products.

Figure 1: Phosphonium betaines prepared. An E/Z ratio of 100:0 means that only the E-isomer was observed in the NMR spectra.
have $pK_a$ values between 4.70 (2b, in dioxane/H$_2$O 3:1 [20]) and $\approx$15.80, reported for 2-(pentafluorophenyl)acetonitrile in DMSO [21], the value for 2f should be similar). On the other hand, the combination of $N,3$-diphenyl-$N$-triflylpropiolamide (1a) with PPh$_3$ and dibenzoylmethane (2d) was far from clean and furnished in low yield betaine $E$-$3d$, which could not be purified by repeated recrystallization. An undefined polymeric material rather than betaines was obtained from 1,1,1,5,5,5-hexafluoropropane-2,4-dione ($pK_a$ 2.30 in DMSO) and dimedone ($pK_a$ 11.42 in DMSO).

Figure 1 also shows that 3-aryl- as well as various 3-alkyl-substituted propiolamides 1 furnish betaines 3 in high yields. A separate case was observed for 3-trimethylsilylpropiolamide 1f, which reacted with PPh$_3$ and malononitrile to furnish the desilylated betaine $E$-$3o$; the required proton probably stems from the adventitious presence of water (Scheme 4). Notably, 3o could not be obtained directly from the 3-H substituted $N$-triflylpropiolamide 1g.

Some betaines 3 were isolated solely as $E$-diastereoisomers, others as $E$/Z mixtures. In some cases, the two diastereoisomers were separated by fractionating crystallization from a mixed-solvent solution. In the case of 3a, a trace amount ($\approx$2%) of the Z-isomer was detected in the reaction mixture by $^1$H NMR (the chemical shift of the olefinic proton being characteristic), but got lost during work-up. An inspection of Figure 1 reveals no obvious reason for the obtained $E$/Z ratios. Beside characteristic spectroscopic data (vide infra), the diastereoisomeric betaines can also be distinguished by their colors: the isolated $E$-isomers are usually colorless, while the Z-isomers have a yellow shade. As an exception, $E$-$3f$ was obtained as dark red needle-shaped crystals, in contrast to pale yellow blocks of Z-$3f$. All betaines could be stored unchanged at least for a year under a dry atmosphere. They appear to be somewhat hygroscopic, as the broad O–H absorption bands in their IR spectra ($\approx$3400–3500 cm$^{-1}$) and the presence of hydrogen-bonded water molecules in the crystal structure of $E$-$3e$ (vide infra) indicate.

The formation of betaines 3 includes the elimination of the $N$-phenyltriflamide anion, which is known as an excellent leaving group in transamidation reactions with nucleophilic amines [18,22]. Therefore, it was interesting to know the performance of propiolic acid chlorides in the same betaine syntheses. We found that acid chlorides 4 indeed underwent the three-component reaction with $N,N'$-dimethylbarbituric acid and furnished betaines 3b, 3j and 3k, respectively, with about the same $E$/Z ratios as in Figure 1 (Scheme 5). However, the products were obtained in low yield, because unidentified side-reactions occurred and purification was cumbersome. Thus, propiolic acid chlorides are not well suited for these three-component reactions.

We propose two mechanistic pathways for the synthesis of betaines 3 from $N$-triflylpropiolamides 1 (Scheme 6). Both of them begin with the conjugate addition of PPh$_3$ at the C–C-triple bond, leading to the vinyl anion intermediate 5. On pathway A, 5 is protonated by the active methylene compound to furnish the vinylphosphonium ion 6. These two steps have been proposed earlier for the formation of related betaines from acetylenic ketones and esters (see Introduction). A replacement of

\[
\begin{align*}
R-\equiv\equiv N-P\text{H} & \quad + \quad \text{PPh}_3 \quad + \quad \text{CH}_2(\text{CN})_2 \\
& \quad \xrightarrow{\text{CH}_2\text{Cl}_2, \ 0 \ ^\circ \text{C, } 2 \ h} \quad \text{71%} \\
& \quad \xrightarrow{\text{R = SiMe}_3} \text{(1f)} \\
& \quad \xrightarrow{\text{R = H}} \text{(1g)} \\
& \quad \text{no defined products}
\end{align*}
\]

**Scheme 4:** Unexpected synthesis of $E$-$3o$.

\[
\begin{align*}
\text{R-} & \equiv\equiv \text{Cl} \quad + \quad \text{O} \quad + \quad \text{O} \quad + \quad \text{N} \quad + \quad \text{O} \\
& \quad \text{PPh}_3 \quad \xrightarrow{\text{CH}_2\text{Cl}_2, \ rt} \quad \text{3} \\
& \quad \text{3a: } \text{R = Ph; 23%} \\
& \quad \text{3j: } \text{R = n-Bu; 18%} \\
& \quad \text{3k: } \text{R = c-C}_3\text{H}_6; 27%
\end{align*}
\]

**Scheme 5:** Betaines 3 from propiolic acid chlorides.
the N-phenyltriflamide group by the conjugate base of the active methylene compound followed by deprotonation of the CHXY group converts 6 into the final betaine 3.

On an alternative reaction pathway (path B in Scheme 6), the initial vinyl anion 5 could be split into a 3-phosphonio-substituted propadienone 7 and the N-phenyltriflamide anion. The latter is able to deprotonate the active methylene compound, the anion of which finally would add to the propadienone. This pathway is notable insofar as propadienones (methylene ketenes) are short-lived compounds which have been generated by flash vapor pyrolysis at high temperatures and characterized spectroscopically or by follow-up products [23]. Under synthetically more convenient conditions, some propadienones have been generated at subambient temperature from 2-bromoacryloyl chlorides by 1,2-elimination induced by the [Mn(CO)5]− anion [24]. For both strategies, the intermediate formation of the cumulene was confirmed in favorable cases by the isolation of [2 + 2] cyclodimerization products (e.g., 3,3-diphenylpropadienone → 2,4-diphenylmethylenecyclobutane-1,3-dione [24]). As we have not obtained so far an experimental evidence for the intermediate formation of cumulenes 7, pathway B is currently speculative but may suggest a novel approach to propadienones.

The presentation of 1,5-zwitterions 3 with the negative charge residing on a carbon atom is more or less a formal one. The structural and spectroscopic data (vide infra) support the view that the mesomeric enolate structure 3’ (see Scheme 3) significantly contributes to the bonding state and that an even wider delocalization of the negative charge, represented by formula 3”, does occur. Therefore, these zwitterions are better described as 1-phosphonium-5-oxabetaines than -5-carbabetaines. Furthermore, they represent a novel type of phosphonium enolate betaines and may be considered as vinylogues of recently reported acylphosphonium zwitterions [25].

**Structural studies**
The solid-state molecular structures of betaines E-3a, E-3b, E-3e and Z-3e were determined by single-crystal X-ray diffraction analysis and are shown in Figures 2–5. Some data charac-

**Scheme 6**: Two mechanistic scenarios for the formation of betaines 3.
Figure 3: Solid-state structure of E-3b·CH₂Cl₂ (ORTEP plot); CH₂Cl₂ solvate molecule not shown.

Figure 4: Solid-state structure of E-3e·H₂O·CH₂Cl₂ (ORTEP plot). The CH₂Cl₂ solvate molecule is disordered. Hydrogen-bonded H₂O molecule: O1···O2 2.907(2) Å, angle O1···H–O2 169(3)°.

Figure 5: Solid-state structure of Z-3e (ORTEP plot).

terizing the bonding geometry of the P⁺–C1–C2–C3(–O)–C4 backbone are compiled in Table 1. For all molecules, the bond distances P–C1, C1–C2 and C2–C3 are in the typical ranges of the respective bond type and the values are more or less identical within the estimated standard deviations; exceptions can be seen, however, for E-3e vs Z-3e. On the other hand, the bond lengths C3–C4, C4–C5 and C3–O1 clearly indicate a partial double bond character, as can be expected when the negative charge is dispersed over the almost planar O–C3–C4–(X,Y) moiety (see formulae 3’ and 3” in Scheme 3). The NMR observation of magnetic non-equivalence of chemically equivalent atoms in the CXY fragment of some betaines (mainly those with X = Y = CN) also indicates that free rotation around the C3–C4 bond is slow on the NMR time scale around 295 K.

A major difference between the E and Z-betaines 3 lies in the conformation at the C2–C3 single bond. The torsion angle C1–C2–C3–C4 amounts to 78.7–82.3° in E-3a and E-3e, 120.0° in E-3b, but 178.5° in Z-3e. Thus, the olefinic π-bond and the delocalized anionic π-system are electronically decoupled in the E-isomers, but coplanar in Z-3e. These conformational differences can explain why the E-isomers are colorless and Z-3e is yellow [UV–vis, acetonitrile, 4 × 10⁻⁴ mol L⁻¹: λmax (lg ε) = 368 nm (2.42) for E-3e, 390 nm (2.94) for Z-3e].

Notably, Z-3e does not undergo a 1,5-cyclization leading to a 5-methylene-2,2,2-triphenyl-2,5-dihydro-1,2λ₅-oxaphosphole. Other studies have come to the conclusion, that the bonding situation in annelated oxaphospholes 8 (Scheme 7) is best described by the contribution of two canonical structures: an oxaphosphole 8A with a pentacovalent phosphorus atom and a P–O bond, and a betaine structure 8B with a phosphonium group and a P···O distance that varies with the electronic properties of the substituent R (Scheme 7) [26-28]. In Z-3e, the intramolecular P···O distance is 2.691 Å, significantly longer than in compounds 8 (2.00–2.36 Å), but still much shorter than the sum of the van der Waals radii (3.32 Å). A distance similar to that in Z-3e has been reported for a betaine III (Scheme 1) (PR₄ = PMe₂Ph, R’ = Ph, 2.597 and 2.620 Å in two symmetry-independent molecules [15]). Furthermore, the coordination around the phosphorus atom comes close to a trigonal-pyramidal geometry with the oxygen and the carbon atom C13 in apical positions: the angle O···P–C13 is 177.1° and the sum of bond angles involving atoms C1, C19 and C25 in the equatorial plane is 346.8° (compared to 328.4° for an ideal tetrahedral coordina-
Table 1: Bond distances and torsion angles in betaines E-3a, E-3b, E-3e, and Z-3e based on XRD data.a

|                  | E-3a molecule 1 | E-3a molecule 2b | E-3b     | E-3e     | Z-3e     |
|------------------|-----------------|------------------|----------|----------|----------|
| Distances [Å]    |                 |                  |          |          |          |
| P–C1             | 1.794(6)        | 1.801(6)         | 1.799(2) | 1.8076(16)| 1.8237(13)|
| C1–C2            | 1.342(9)        | 1.341(9)         | 1.332(3) | 1.335(2) | 1.406(2) |
| C2–C3            | 1.507(4)        | 1.504(9)         | 1.512(3) | 1.508(2) | 1.406(2) |
| C3–C4            | 1.428(9)        | 1.416(9)         | 1.437(3) | 1.406(2) | 1.411(2) |
| C4–C5            | 1.432(9)        | 1.426(9)         | 1.435(3) | 1.406(2) | 1.411(2) |
| C3–O1            | 1.248(8)        | 1.241(8)         | 1.238(3) | 1.246(3) | 1.249(2) |
| Torsion angles [deg] |                  |                  |          |          |          |
| P–C1–C2–C3      | −171.4(5)       | −174.2(5)        | −173.84(15) | −178.0(3) | −2.5(2) |
| C1–C2–C3–O1     | −100.0(7)       | −98.4(7)         | 63.5(3)  | 99.5(2)  | 0.8(2)  |
| C1–C2–C3–C4     | 78.7(8)         | 81.3(8)          | −120.0(2)| −82.3(2) | −178.5(1)|
| C2–C3–C4–C5     | 12.9(9)         | −176.0(6)        | −170.9(2)| 0.8(2)   | 175.7(1)|
| C2–C3–C4–C9     |                 |                  | 6.2(3)   |          |          |
| C2–C3–C4–C6     |                 | −177.7(1)        |          |          | −5.3(2) |

aNote that the atom numbering refers to the molecule plots shown in Figures 2–5 and is different from the systematic nomenclature. bBond geometries involving the same atoms as in molecule 1; compare Figure 2 and Figure 3.

All these data point to the description of Z-3e as a 1,5-betaine with an attractive electrostatic interaction of phosphorus and oxygen. The 31P NMR chemical shift (δ = 23.53 ppm) further confirms the presence of a tetravalent phosphorus atom (see [28]).

NMR data of betaines 3

The 1H, 13C and 31P data of the P–C3–C2–C1 chain in betaines 3 are listed in Table 2.

Based on the spectra of those betaines, for which the molecular geometry has been established by XRD analysis (vide supra), the stereochemistry at the P-substituted olefinic bond could be determined in all cases from characteristic chemical shifts of the olefinic proton and from 3J(P,H) and 3J(P,C) coupling constants. The olefinic proton signal of the E-isomers was found in the δ range 6.8–7.7 ppm, the signal of the Z-isomers at 7.9–8.4 ppm. In all cases where both diastereoisomers were available, the known relationship 3J(P,H)trans < 3J(P,H)trans [29,30] was valid; the values were in the range 20.3–30.8 Hz for the cis coupling and 38.3–46.1 Hz for the trans coupling. The magnitude of the coupling between the tetravalent phosphorus nucleus and the carbonyl carbon atom C-1 also follows the known trend 3J(P,C)trans < 3J(P,C)trans [29]. The ranges –3.7–7.4 Hz vs 15.3–21.4 Hz – are sufficiently different to distinguish between E and Z-isomers even if only one of them is available.

The 31P NMR chemical shifts were found between 19.26 (for Z-3f) and 30.23 ppm (for E-3l), the typical range being 23–27 ppm. As in other alkenes bearing a tetravalent phosphorus atom [29], the values of the E-isomers are always somewhat higher than those of the Z-isomers. Considering the question of an intramolecular P···O coordination in the Z-isomers (vide supra), the chemical shift values by themselves (compare:...
δₚ = 18.7 ppm for ([E]-Ph₃P⁺−CH=CHMe Br⁻ [31]) and the similarity of the chemical shifts for E/Z pairs indicate a tetracovalent rather than a pentacovalent phosphorus atom [28].

**Conclusion**

We have observed that the reaction of N-trityl-N-phenyl-propiolamides with triphenylphosphane results in undefined polymerization and formation of N-phenyltriflimide. In the presence of several active methylene compounds, however, a clean three-component reaction leads to a novel type of phosphonium betaines, which can formally be named 1,5-phosphonium-carbabetaines, but because of the extended delocalization of the negative charge are better described as 1-phenonium-5-oxabetaines. The excellent leaving group character of the N-phenyltriflimide anion ([N(Ph)SO₂]⁻) is a key factor in the reaction course, because analogous three-component reactions using propionic acid esters as Michael acceptors are known to furnish another type of betaines in which the ester group (COOR) is retained. Questions on some mechanistic details, in particular at which stage of the reaction sequence the triflimide anion departs, remain to be answered.

**Supporting Information**

**Supporting Information File 1**

Experimental procedures, characterization data, NMR spectra (¹H, ¹³C, ³¹P, ¹⁹F) and IR spectra for the synthesized compounds, and data for the X-ray crystal structure determinations.

[https://www.beilstein-journals.org/bjc/content/ supplementary/1860-5397-15-253-S1.pdf]
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References
1. Guo, H.; Fan, Y. C.; Sun, Z.; Wu, Y.; Kwon, O. Chem. Rev. 2016, 118, 10049–10293. doi:10.1021/acs.chemrev.6b00081
2. Enders, D.; Saint-Dizier, A.; Lannou, M.-I.; Lenzen, A. Eur. J. Org. Chem. 2006, 29–49. doi:10.1002/ejoc.200500593
3. Ye, L.-W.; Zhou, J.; Tang, Y. Chem. Soc. Rev. 2008, 37, 1140–1152. doi:10.1039/b717758e
4. Methot, J. L.; Roush, W. R. Adv. Synth. Catal. 2004, 346, 1035–1050. doi:10.1002/adsc.200404087
5. Ramazani, A.; Kazemizadeh, A. R.; Ahmadi, E.; Noshiranzadeh, N.; Soudlozi, A. Curr. Org. Chem. 2008, 12, 59–82. doi:10.2174/13852720878333055
6. Stoddard, R. L.; Luo, J.; van der Wal, N.; O’Rourke, N. F.; Wulff, J. E.; McInerney, J. S. New J. Chem. 2014, 38, 5382–5390. doi:10.1039/c4nj01070a
7. Nagao, K.; Yamaazaki, A.; Ohmiya, H.; Sawamura, M. Org. Lett. 2018, 20, 1861–1865. doi:10.1021/acs.orglett.8b00390
8. Zi, Y.; Schömberg, F.; Seifert, F.; Görß, H.; Viljojvic, I. Org. Biomol. Chem. 2016, 16, 6341–6349. doi:10.1039/c6ob01343h
9. Yavari, I., Magsoudioli, M. T. Tetrahedron Lett. 1996, 39, 4579–4580. doi:10.1016/0040-4039(96)00811-9
10. Yavari, I.; Anary-Abbasinejad, M.; Alizadeh, A. Phosphorus, Sulfur Silicon Relat. Elem. 2002, 177, 93–103. doi:10.1080/10426500878333005
11. Yavari, I.; Islami, M. R.; Bijanzadeh, H. R. Tetrahedron 1999, 55, 5547–5554. doi:10.1016/s0040-4020(99)00220-3
12. Shabani, A.; Temouri, M. B.; Yavari, I.; Arasi, H. N.; Bijanzadeh, H. R. J. Fluorine Chem. 2000, 103, 155–157. doi:10.1016/s0022-1139(99)00305-x
13. Heydari, R.; Magsoudioli, M. T.; Yami, R. N. Chin. Chem. Lett. 2009, 20, 1175–1176. doi:10.1016/j.ccl.2009.05.019
14. Ghodsi, F.; Habibi-Khorasani, S. M.; Shahrazi, M. Molecules 2016, 21, 1514. doi:10.3390/21111514
15. Zhu, X.-F.; Henry, C. E.; Kwon, G. J. Am. Chem. Soc. 2007, 129, 6722–6723. doi:10.1021/ja071990s
16. Liu, H.; Zhang, Q.; Wang, L.; Tong, X. Chem. – Eur. J. 2010, 16, 1968–1972. doi:10.1002/chem.200902105
17. Deng, J.-C.; Chen, F.-W.; Kuo, C.-W.; Cheng, C.-A.; Huang, C.-Y.; Chuang, S.-C. Eur. J. Org. Chem. 2012, 5738–5747. doi:10.1002/ejoc.201200400
18. Fiore, V. A.; Maas, G. Tetrahedron 2019, 75, 3586–3595. doi:10.1016/j.tet.2019.05.027
19. Fiore, V. A.; Maas, G. Z. Naturforsch., B: J. Chem. Sci. 2019, 74, 671–676. doi:10.1515/znb-2019-0100
20. Hennig, L.; Alva-Astutillo, M.; Mann, G.; Kappe, T. Monatsh. Chem. 1992, 123, 571–579. doi:10.1007/bf00816851
21. Bordwell, F. G.; Branca, J. C.; Bares, J. E.; Filler, R. J. Org. Chem. 1988, 53, 780–782. doi:10.1021/jo00039a016
22. Hendrickson, J. B.; Bergeron, R. Tetrahedron Lett. 1973, 14, 4607–4610. doi:10.1016/s0040-4020(01)87289-0
23. Brown, R. F. C.; Eastwood, F. W. Synlett 1993, 9–19. doi:10.1055/s-1993-22330
24. Masters, A. P.; Sorensen, T. S.; Tran, P. M. Can. J. Chem. 1987, 65, 1499–1502. doi:10.1139/v87-256
25. Saijo, R.; Uno, H.; Mori, S.; Kawase, M. Chem. Commun. 2016, 52, 8006–8009. doi:10.1039/c6cc01627h
26. Kawamoto, I.; Hata, T.; Kishida, Y.; Tamura, C. Tetrahedron Lett. 1971, 12, 2417–2420. doi:10.1016/s0040-4020(01)96878-9
27. Kawamoto, I.; Hata, T.; Kishida, Y.; Tamura, C. Tetrahedron Lett. 1972, 13, 1611–1614. doi:10.1016/s0040-4020(01)84699-2
28. Naya, S.-I.; Nitta, M. J. Chem. Soc., Perkin Trans. 2 2002, 1017–1023. doi:10.1039/b109076n
29. Duncan, M.; Gallagher, M. J. Org. Magn. Reson. 1981, 15, 37–42. doi:10.1002/mrc.1270150110
30. Espenlaub, S.; Gerster, H.; Maas, G. ARKIVOC 2007, No. iii, 114–131.
31. Albright, T. A.; Freeman, W. J.; Schweizer, E. E. J. Am. Chem. Soc. 1975, 97, 2946–2950. doi:10.1021/ja00844a003

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2611