Article

Conformational Analysis of N-Alkyl-N-[2-(diphenylphosphoryl)ethyl]amides of Diphenylphosphorylacetic Acid: Dipole Moments, IR Spectroscopy, DFT Study

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Abstract: Experimental and theoretical conformational analysis of N-methyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide, N-butyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide, and N-octyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide was carried out by the methods of dipole moments, IR spectroscopy, and Density Functional Theory (DFT) B3PW91/6-311++G(df,p) calculations. In solution, these N,N-dialkyl substituted bisphosphorylated acetamides exist as a conformational equilibrium of several forms divided into two groups—with Z- or E-configuration of the carbonyl group and alkyl substituent, and syn or anti arrangement of the phosphoryl-containing fragments relative to the amide plane. The substituents at the phosphorus atoms have eclipsed cis- or staggered gauche-orientation relative to the P=O groups, and cis orientation of the substituents is due to the presence of intramolecular H-contacts P=O...H−C phenyl or p,π conjugation between the phosphoryl group and the phenyl ring. Preferred conformers of acetamides molecules are additionally stabilized by various intramolecular hydrogen contacts with the participation of oxygen atoms of the P=O or C=O groups and hydrogen atoms of the methylene and ethylene bridges, alkyl substituents, and phenyl rings. However, steric factors, such as a flat amide fragment, the bulky phenyl groups, and the configuration of alkyl bridges, make a significant contribution to the realization of preferred conformers.

Keywords: carbamoylphosphine oxides; phosphorylated acetamides; dipole moments; conformational analysis; DFT calculations

1. Introduction

Derivatives of carbamoylmethylphosphine oxides (CMPO) are known as effective extractants of rare earth elements and actinides from mineral acid solutions [1–17]. The high coordinating ability of these compounds is due to the presence of polar phosphoryl and amide groups. CMPO are used in industrial processes for reprocessing spent nuclear fuel [18,19] and for the processing of radioactive waste and the separation of transplutonium elements [20,21], as well as for the preparation of composite materials for column chromatography [22–24]. Carbamoylmethylphosphine oxides modified with a dialkylaminomethyl coordination group in the methylene bridge are potential neuroprotective agents [25,26]. Manganese (II) complexes with CMPO ligands exhibit luminescent
properties [27]. Palladium complexes containing phosphoryl-substituted CMPO as ligand demonstrate catalytic activity in Suzuki reaction [28].

Although carbamoylmethylphosphine oxides have found active practical application for a long time, their spatial structure has been poorly studied. The literature contains data on the structure of complexes of CMPO with rare earth elements and actinides [17,29,30], manganese (II) [27], and mercury (II) [31] in the solid state. The crystal structures of (diethylcarbamoyl)methyldiphenylphosphine oxide and its protonated form [32], diphenylmorpholine carbamoylmethyl phosphine oxide [33], diphenyl-\(N,N\)-dimethylcarbamoylmethylphosphine oxide [34], \(N\)-aryl-substituted CMPO modified with a phosphoryl group, and its complexes with Pd (II) and Re (I) [35] were determined by X-ray diffraction. Information on the structure in solution is available only for \(P(X)\)-modified (\(X = O, S\)) \(N\)-aryl-substituted CMPOs: experimental and theoretical conformational analysis by the methods of dipole moments and quantum chemistry DFT B3PW91/6-311++G(df,p).

In the present work, we investigated the spatial structure of \(N\)-methyl-\(N\)-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide 1, \(N\)-butyl-\(N\)-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide 2, and \(N\)-octyl-\(N\)-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide 3 (Scheme 1) by the methods of dipole moments, IR spectroscopy, and quantum chemistry DFT B3PW91/6-311++G(df,p).

![Scheme 1](image-url)  
Scheme 1. Structures of compounds 1–3, selected atom numbers are given.

2. Results and Discussion

2.1. Methodology

The studied amides of phosphorylacetic acid (phosphorylacetamides) 1–3 are polyfunctional polar compounds. Two phosphoryl groups with bulky phenyl substituents at the phosphorus atom are linked through alkyl bridges by an amide group; there is also a flexible alkyl substituent at the nitrogen atom. All fragments of a molecule can freely rotate about single bonds, and functional groups can participate in intra- and intermolecular non-covalent interactions, which leads to the emergence of a large number of possible conformers.

The method of dipole moments is a sensitive instrument for the determination of structure and the study of fine features of spatial and electronic structure of polar organic and organoelement compounds in solution. To determine the experimental values of the dipole moments, we used the second Debye method based on the measurement of
the dielectric permittivity of the dilute solutions of the polar substance in a nonpolar solvent [38].

To search the possible conformations of isolated molecules of 1–3, we applied the Density Functional Theory (DFT) with B3PW91 hybrid functional. This method has been successfully used to study the polarity and spatial structure of similar organophosphorus compounds with double bond phosphorus-chalcogen and aryl substituents [36, 39]. The choice of the B3PW91 method was also based on the data [40].

2.2. Polarity of Phosphorylacetamides 1–3

We have determined previously unknown polarities of compounds 1–3. The experimental values of the dipole moments were determined in benzene solutions using the second Debye method based on the measurement of the dielectric constant of the dilute solutions of a polar substance in a nonpolar solvent. The experimental dipole moments of 1–3 are listed in Table 1; their values are sufficiently high (Table 1) and are typical for the polarities of the compounds of tetra-coordinated phosphorus (2.5–5.0 Debye) [41].

Table 1. Coefficients of calculated equations (α, γ), orientation polarizabilities (P or.), and experimental dipole moments (µ) of compounds 1–3 in benzene.

| Compound | α   | γ   | P or., cm³ | µ, D |
|----------|-----|-----|------------|------|
| 1        | 6.106 | 0.303 | 547.82     | 5.15 |
| 2        | 5.070 | 0.214 | 496.84     | 4.90 |
| 3        | 4.091 | 0.259 | 432.32     | 4.77 |

2.3. Experimental and Theoretical Conformational Analysis of Phosphorylacetamides 1–3

Conformers for the compounds were built using the Gauss View 6.0 imaging software. All possible conformations for compounds 1–3 were built by sequential rotation of the parts of the molecules relative to single bonds. Conformations with overlapping atoms or with a too close arrangement, which does not correspond to the physical meaning, were immediately discarded. Thus, there were about a thousand conformers. First, the calculations were carried out for them using small basis set 6-31G(d). Based on these results, we have discarded degenerate structures. Then, conformers with high relative energies (more than 15 kJ/mol) were discarded as unlikely. At the next stage, the calculations were carried out using the extended basis set 6-311++G(df,p). We considered the preferred conformers with relative energies less than 6 kJ/mol without taking into account the mirror isomers with identical energies and theoretical dipole moments. For all preferred conformers, their relative energies and theoretical polarities were computed, and the dipole moments were calculated according to the vector-additive scheme (Table 2). Conformational diversity of the compounds 1–3 is due to the presence of a large number of rotation axes and several functional groups in their molecules. The percentage of conformers in the mixture was calculated on the basis of the theoretical values of Gibbs energies (Table 2).

According to the results of theoretical calculations, five energetically preferred conformers were found for N-methylacetamide 1 (Figure 1); their characteristics are listed in Tables 2 and 3. In conformers 1a–1e, the phosphorus atoms have a pyramidal structure, the phenyl substituents have cis- or gauche-orientation, and the methyl or ethyl bridges have gauche-orientation relative to the P=O bonds (α, β, γ, i, κ, and λ angles in Table 3).
Table 2. Relative energies (ΔE), Gibbs free energies (ΔG, 298.15 K) (kJ/mol), theoretical (μ_theor) and calculated by the vector-additive scheme (μ_calc) dipole moments (D), percentage (n) of preferred conformers of 1–3 according to DFT B3PW91/6-311++G(df,p).

| Conformer | ΔE  | ΔG  | μ_theor | μ_calc | n, % |
|-----------|-----|-----|---------|--------|------|
| 1a        | Z, syn | 0.0 | 0.0     | 5.67   | 4.75 | 41.3 |
| 1b        | E, syn | 1.5 | 0.4     | 5.44   | 5.15 | 35.3 |
| 1c        | Z, syn | 2.9 | 8.2     | 1.72   | 1.77 | 1.5  |
| 1d        | E, anti | 3.7 | 1.6     | 5.53   | 5.84 | 21.5 |
| 1e        | Z, anti | 5.6 | 11.5    | 3.74   | 3.40 | 0.4  |
| 2a        | Z, syn | 0.0 | 2.1     | 5.48   | 4.74 | 23   |
| 2b        | E, syn | 1.6 | 0.0     | 5.22   | 5.33 | 55   |
| 2c        | Z, syn | 3.0 | 9.8     | 2.35   | 2.05 | 1    |
| 2d        | Z, syn | 3.4 | 5.7     | 5.71   | 4.77 | 6    |
| 2e        | E, syn | 5.1 | 6.0     | 5.54   | 5.51 | 5    |
| 2f        | E, anti | 5.4 | 4.3     | 5.30   | 5.80 | 10   |
| 3a        | Z, syn | 0.0 | 1.6     | 5.44   | 4.75 | 21   |
| 3b        | E, syn | 1.5 | 0.0     | 5.18   | 5.43 | 43   |
| 3c        | Z, syn | 2.9 | 9.3     | 2.34   | 2.28 | 1    |
| 3d        | Z, syn | 3.4 | 5.0     | 5.69   | 4.32 | 6    |
| 3e        | E, syn | 5.0 | 3.8     | 5.55   | 5.44 | 5    |
| 3f        | E, anti | 5.3 | 2.0     | 5.27   | 5.80 | 20   |

Figure 1. Preferred conformers of 1 according to DFT calculations (some hydrogen atoms are omitted). Distances are given in Å.
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Table 3. Selected dihedral angles (degree) for the preferred conformers of 1–3 according to DFT B3PW91/6-311+G(df,p): α O1 = P1–C1–C2, β O1 = P1–C3–C4, γ O1 = P1–C6–C7, δ O1–C7–N–C15, ε O2 = P2–C10–C11, η N–C8–C9–P2, θ C8–C9–P2 = O2, κ O2 = P2–C10–C11, λ O2 = P2–C13–C14.

|     | α    | β    | γ    | δ    | ε    | η    | Θ    | τ    | κ    | λ    |
|-----|------|------|------|------|------|------|------|------|------|------|
| 1a  | −1   | 22   | 82   | 93   | 0    | −84  | 168  | −65  | −14  | 7    |
| 1b  | 6    | 24   | 75   | 99   | −178 | −100 | −179 | 60   | 8    | 31   |
| 1c  | 59   | 12   | 47   | 89   | −2   | −73  | −166 | 41   | 72   | 5    |
| 1d  | 7    | 25   | 77   | 104  | 173  | −94  | 178  | −61  | −30  | −9   |
| 1e  | 61   | 15   | 71   | 100  | 3    | 80   | 134  | −58  | 36   | 17   |
| 2a  | −1   | 22   | 82   | 92   | 0    | 87   | −84  | 167  | −65  | −15  | 7    |
| 2b  | 5    | 25   | 74   | 97   | −179 | −97  | 100  | −178 | 60   | 8    | 31   |
| 2c  | 59   | 12   | 47   | 89   | −2   | −73  | −166 | 41   | 72   | 5    |
| 2d  | −2   | 22   | 79   | 95   | −1   | −77  | −79  | 168  | −67  | −15  | 7    |
| 2e  | 7    | 26   | 74   | 98   | −179 | −97  | 100  | −179 | 60   | 9    | 31   |
| 2f  | 6    | 24   | 77   | 100  | −179 | −102 | −109 | 179  | −60  | −30  | −8   |
| 3a  | −1   | 22   | 81   | 92   | −1   | 87   | −84  | 168  | −65  | −14  | 7    |
| 3b  | 6    | 25   | 74   | 97   | −179 | −96  | 100  | −178 | 60   | 8    | 31   |
| 3c  | 58   | 13   | 60   | 76   | −3   | −84  | −91  | −153 | 49   | 70   | 5    |
| 3d  | −1   | 23   | 79   | 95   | −1   | −77  | −79  | 167  | −68  | −15  | 6    |
| 3e  | 8    | 26   | 75   | 103  | 179  | 121  | 108  | −179 | 60   | 8    | 31   |
| 3f  | 6    | 25   | 77   | 100  | −179 | −101 | −108 | 179  | −60  | −29  | −7   |

The amide fragment is flat, the carbonyl and methyl groups have Z-orientation relative to the C(sp2)=N bond in conformers 1a, 1c, 1e, and E-orientation in 1b and 1d; the bulky diphenylphosphoryl fragments are syn-located relative to amide plane in conformers 1a–1c and anti-located in 1d and 1e (Figure 1).

The conformers 1a–1e differ in the arrangement of the following bonds relative to each other: the P1–C(sp3) and C=O bonds have synclinal orientation in 1c and anticlinal orientation in all other conformers; the N–C(sp3) and C(sp3)–P2 bonds are practically coplanar in conformers 1a–1d (antiperiplanar orientation), and they have anticlinal orientation in 1e. The C(sp3)–C(sp3) and N–C(sp3)(methyl) bonds have synclinal orientation in conformers 1a, 1c, 1e, and anticlinal orientation in 1b and 1d; that is, the ethyl fragment is practically perpendicular to the amide plane (Table 3). The N–C(sp3) and C(sp3)–P2 bonds lie in the same plane in 1a–1e (θ angle, Table 3).

All conformers 1a–e are additionally stabilized by intramolecular hydrogen contacts (Figure 1, Table 4). In conformers 1b and 1d, there is an interaction between the oxygen atom of the phosphoryl group P1=O and the hydrogen atom of the methyl substituent, resulting in the formation of seven-membered intramolecular heterocycles. The bifurcate hydrogen bond between the oxygen atom of the P1=O group and the hydrogen atoms of the ethylene bridge and one of the phenyl substituents at the second diphenylphosphoryl group is observed in 1a, 1c, and 1e, resulting in the formation of seven- or eight-membered intramolecular cycles. In addition, in conformer 1c, H-contacts between the oxygen atom of the second phosphoryl group P2=O and the hydrogen atom of one of the phenyl substituents at the P1 atom (C2–H1…O2=P 2.391 Å, C2–H1…O2 156°), as well as between the oxygen atom of the carbonyl group and the hydrogen atom of the other phenyl substituent at the P1 atom (C5–H2…O3–C7 2.300 Å, C5–H2…O3 141°) are observed. The second bifurcate bond is formed by the oxygen atom of the P2 = O group and hydrogen atoms of the phenyl ring at the first phosphorus atom and the methylene bridge (C2–H1…O2 = P2 2.304 Å, C2–H1…O2 167° and C6–H3…O2=P2 2.283 Å, C6–H3…O2 164° respectively) in conformer 1e.
Table 4. Selected geometry characteristics of hydrogen bonds (distance, Å; angle, degrees) according to DFT B3PW91/6-311+G(df,p).

|          | H5···O1 | C9–H5···O1 | H6···O1 | C12–H6···O1 |
|----------|---------|------------|---------|-------------|
| 1a       | 2.194   | 153        | 2.339   | 164         |
| 1b       | 2.247*  | 151*       |         |             |
| 1c       | 2.160   | 144        | 2.305   | 157         |
| 1d       | 2.236   | 152        | 2.341   | 164         |
| 1e       | 2.175** | 152**      | 2.422   | 154         |
| 2a       | 2.189   | 154        | 2.341   | 164         |
| 2b       | 2.271*  | 153*       | 2.350   | 163         |
| 2c       | 2.152   | 162        | 2.255   | 172         |
| 2d       | 2.248   | 147        | 2.323   | 163         |
| 2e       | 2.259*  | 153*       |         |             |
| 2f       | 2.189   | 154        | 2.248   | 172         |
| 3a       | 2.272*  | 153*       |         |             |
| 3b       | 2.162   | 161        | 2.250   | 163         |
| 3c       | 2.250   | 147        | 2.323   |             |
| 3d       | 2.260*  | 153*       |         |             |
| 3f       | 2.194   | 153        | 2.339   | 164         |

*H7···O1, C15–H7···O1; **H4···O1, C8–H4···O1.

Elongation of the alkyl substituent at the nitrogen atom led to an increase in the number of energetically preferred conformers of N-butyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide 2 to 6 (Figure 2); their characteristics are listed in Tables 2 and 3. The common features of conformers 2a–f are a pyramidal structure of phosphorus atoms and a planar amide fragment, which are also characteristic for conformers 1a–e. In conformers 2a–f, the phenyl substituents are predominantly cis-oriented, and the methyl or ethyl bridges are gauche-orientated relative to the P = O bonds ($\alpha$, $\beta$, $\gamma$, $\iota$, $\kappa$, and $\lambda$ angles in Table 3).

The carbonyl group and butyl substituent at the nitrogen atom have mutual Z-orientation in conformers 2a, 2c, 2d, and E-orientation in 2b, 2e, and 2f (e angle, Table 3): the bulky diphenylphosphoryl fragments are syn-located relative to amide plane in conformers 2a–e and anti-located in 2f (Figure 2).

The difference between conformers is due to the different mutual arrangement of the following bonds: P1–C$_{sp3}$ and C$_{sp2}$=O bonds have synclinal orientation in 2c and anticlinal orientation in all other forms ($\delta$ angle, Table 3). The C$_{sp3}$–C$_{sp3}$ and N–C$_{sp3}$(butyl) bonds have synclinal orientation in 2a and 2d, and anticlinal orientation in 2b, 2c, 2e, and 2f (n angle, Table 3); the N–C$_{sp3}$ and C$_{sp3}$–P$_{2}$ bonds are antiperiplanar in 2a–2f ($\theta$ angle, Table 3).

An additional variety of conformers 2a–f compared to compound 1 is due to the presence of a longer alkyl substituent. However, the difference between preferred conformers is only due to the rotation relative to the N–C$_{sp3}$(R) bond, because conformers with a zigzag alkyl radical possess low energies. The C$_{sp2}$–N and C$_{sp3}$(R)–C$_{sp3}$(R) bonds have synclinal orientation in 2a, 2c, and 2d and anticlinal orientation in 2b, 2e, and 2f ($\zeta$ angle, Table 3).

For conformers 2a–d and 2f, an additional stabilization effect is observed due to the formation of intramolecular hydrogen contacts (Table 4). In 2b and 2f, the interaction occurs between the oxygen atom of the group P1=O and one of the hydrogen atoms of the butyl substituent, resulting in the formation of seven-membered intramolecular rings. In conformers 2a, 2c, 2d, and 2f, a bifurcate hydrogen bond is observed between the oxygen atom of the group P1=O and the hydrogen atoms of the ethylene bridge and one of the phenyl rings at the P2 atom, resulting in the formation of seven- or eight-membered intramolecular cycles. In addition to the bifurcate bond, a contact between the oxygen atom of the second phosphoryl group and the hydrogen atom of one of the phenyl substituents at the P1 atom is formed in conformer 2c (C2–H1···O2=P2 2.257 Å, C2–H1···O2 176°).

According to the quantum chemical calculations, for N-octyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylacetamide 3, six energetically preferred conformers were found (Figure 3); their characteristics are listed in Tables 2 and 3. As for compounds 1 and 2,
conformers 3a–f are characterized by a pyramidal structure of the phosphorus atoms and a flat structure of the nitrogen atom. In conformers 3a–f, the phenyl substituents are predominantly cis-oriented and the methyl or ethyl bridges are gauche-orientated relative to the P=O groups ($\alpha$, $\beta$, $\gamma$, $\iota$, $\kappa$, and $\lambda$ angles in Table 3).

Figure 2. Preferred conformers of 2 according to DFT calculations (some hydrogen atoms are omitted). Distances are given in Å.
As in the case of butyl substituted amide 2, the carbonyl group and octyl substituent at the nitrogen atom have mutual Z-configuration in conformers 3a, 3c, 3d, and E-configuration in 3b, 3e, and 3f (dihedral angle $\epsilon$, Table 3); the diphenylphosphoryl fragments are syn-located relative to amide plane in conformers 3a–e and anti-located in 3f (Figure 2).

Conformers 3a–e differ in the structure of the carbamoylmethyl fragment and the ethylene bridge; the P1–C$_{sp3}$ and C=O bonds are synclinal in 3c, while in the rest of the
conformers, these bonds are anticlinal (dihedral angle δ, Table 3). The C\textsubscript{sp3}–C\textsubscript{sp3} and N–C\textsubscript{sp3}(R) bonds are synclinal in 3\textsubscript{a} and 3\textsubscript{d}, and anticlinal in 3\textsubscript{b}, 3\textsubscript{c}, 3\textsubscript{e}, and 3\textsubscript{f} (dihedral angle η, Table 3). The N–C\textsubscript{sp3} and C\textsubscript{sp3}–P2 bonds have antiperiplanar mutual arrangement in conformers 3\textsubscript{a–f} (dihedral angle ζ, Table 3).

An additional difference between conformers is due to the rotation of the octyl substituent relative to the N–C\textsubscript{sp3}(R) bond. The C\textsubscript{sp2}–N and C\textsubscript{sp3}(R)–C\textsubscript{sp3}(R) bonds have mutual synclinal arrangement in 3\textsubscript{a, 3c, and 3d} and anticlinal arrangement in 3\textsubscript{b, 3e, and 3f} (dihedral angle η, Table 3).

The intramolecular hydrogen contacts are possible in conformers 3\textsubscript{a–d} and 3\textsubscript{f} (Table 4). In conformers 3\textsubscript{b} and 3\textsubscript{f}, an interaction between the oxygen atom of the P1=O group and one of the hydrogen atoms of the octyl substituent is observed, resulting in the formation of seven-membered intramolecular cycles. The bifurcate hydrogen bonds between the oxygen atom of the first phosphoryl group and the hydrogen atoms of the ethylene bridge and one of the phenyl rings at the P2 atom arise in conformers 3\textsubscript{a, 3c, 3d, and 3f}. Moreover, a contact between the oxygen atom of the second phosphoryl group and the hydrogen atom of one of the phenyl substituents at the P1 atom is formed in conformer 3\textsubscript{c} (C2–H1···O2 = P2 2.273 Å, C2–H1···O2 176°). It should be noted that in conformers 1\textsubscript{a, 2a, and 2d, 3a, and 3d}, the phosphoryl group P1=O and one of the phenyl substituents at the P1 atom have a completely eclipsed cis-orientation (α angle, Table 3); their coplanar arrangement promotes p,π-conjugation in the molecules and additionally stabilizes these conformers (Figure 4 and Figure S1). A similar fact was previously described for (arylcarbamoylmethyl)diphenylphosphine oxides and sulfides [38].

Conformers 1\textsubscript{c, 1e, 2c, and 3c} have the smallest dipole moments, both theoretical and calculated by the vector-additive scheme (exaltation between the experimental and calculated values—Δµ is 3.38, 1.75, 2.85, and 2.49 D, respectively—see Tables 1 and 2), which can presumably be explained by the largest number of intramolecular hydrogen contacts in these forms, causing a decrease in their dipole moments. However, these low-polarity forms have slightly higher Gibbs energies, and their content in conformational equilibrium is insignificant (Table 2).

We have registered the IR spectra of compounds 1–3 in the solid state, in the melt, and in solution of the trichloromethane (Figure 5, Figures S2 and S3). The comparison of IR spectra of 1–3 showed a change in the number of bands in the range of 700–800 cm\textsuperscript{-1}, corresponding to the out-of-plane deformation vibrations of the C–H bonds in the phenyl substituents. For compound 1, three absorption bands are observed in the melt, whereas seven bands are observed in the solid sample: the 717 cm\textsuperscript{-1} band splits in two (717 cm\textsuperscript{-1} and 724 cm\textsuperscript{-1}), and new bands at 737 cm\textsuperscript{-1}, 758 cm\textsuperscript{-1}, and 770 cm\textsuperscript{-1} appear. In the case of compound 2, there are also three absorption bands in the melt and seven bands in the solid state: the 717 cm\textsuperscript{-1} band splits in two (715 cm\textsuperscript{-1} and 723 cm\textsuperscript{-1}), the 793 cm\textsuperscript{-1} band splits in two (790 cm\textsuperscript{-1} and 795 cm\textsuperscript{-1}), and new bands at 740 cm\textsuperscript{-1} and 774 cm\textsuperscript{-1} appear. The IR spectra of 3 also contain three vibrational bands in the melt and six bands in the solid state. New bands at 760 cm\textsuperscript{-1}, 777 cm\textsuperscript{-1}, and 809 cm\textsuperscript{-1} appear. These IR spectral data indicate the presence of conformational equilibrium in acetamides 1–3.

It should be noted that there is no noticeable shift and no change in the number of vibration bands of the phosphoryl and carbonyl groups when changing the state from solid to liquid (Table 5). A split peak is observed for the stretching vibrations of the P=O bond in the spectrum of the solid sample of 1 as well as in simulated spectra of conformers 1\textsubscript{a–e, 2a–f, and 3a–f}. The theoretical frequencies of the stretching vibrations of the C=O and P=O groups were calculated using the scaling factor 0.96 [42].
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Figure 4. Visualization of the molecular orbitals № 102 for 1a, № 112 for 2a, № 112 for 2d, № 124 for 3a, and № 123 for 3d corresponding to the interaction of the P=O bond and phenyl substituents. The positive and negative wave functions are shown as blue and red areas, respectively. The isocontour value is 0.02 a.u. The frontier orbitals are not given, since we do not consider the reactivity of the compounds in this context.

Figure 5. IR spectra of compound 1 in different aggregate states. D is the absorbance.
Molecules exist as two sets of conformers: in the first set, the C=O and alkyl groups have
N-configuration in rotamers 1a, 1c, and 1e; 2a, 2c, and 2d; 3a, 3c, and 3d, while in the second set, these groups have E-configuration in rotamers 1b and 1d; 2b, 2e, and 2d; 3b, 3e, and 3d, and in all forms except 1e, 2f, and 3f, the phosphoryl-containing fragments are syn-located relative to the plane of the amide fragment. We believe that the first set of conformers corresponds to the minor conformer, and the second set corresponds to the

Table 5. Selected vibration frequencies of the phosphoryl and carbonyl groups of 1–3; theoretical values are listed for conformers a–f.

|         | P=O       | C=O       |
|---------|-----------|-----------|
|         | 1         | 2         | 3         | 1         | 2         | 3         |
| ν_{exp} solid | 1179, 1188 | 1187      | 1181      | 1648      | 1642      | 1632      |
| ν_{exp} melt | 1194      | 1194      | 1195      | 1643      | 1640      | 1641      |
| ν_{exp} solution | 1184      | 1184      | 1182      | 1639      | 1639      | 1634      |
| ν_{theor} a | 1178, 1187 | 1177, 1182, 1194 | 1179, 1184, 1187 | 1650      | 1644      | 1644      |
| ν_{theor} b | 1187, 1192 | 1190, 1193 | 1188, 1192, 1193 | 1642      | 1639      | 1639      |
| ν_{theor} c | 1193      | 1162, 1175 | 1162, 1175 | 1641      | 1652      | 1652      |
| ν_{theor} d | 1171, 1181 | 1179, 1181, 1191 | 1179, 1184 | 1647      | 1643      | 1643      |
| ν_{theor} e | 1180, 1183 | 1190, 1194 | 1191, 1194 | 1659      | 1632      | 1632      |
| ν_{theor} f | 1190, 1193 | 1192, 1193 | -         | 1635      | 1635      |           |

A comparison of experimental and calculated according to vector scheme dipole moments, IR data, and theoretical results showed that in solution, compounds 1–3 exist as a conformational equilibrium of several forms, among which highly polar ones prevail.

Preferred conformers 1a–e, 2a–f, and 3a–f can be divided into two groups, with Z- or E-configuration of the carbonyl and alkyl groups relative to the C_{sp2}=N bond in the planar amide fragment; in addition, the phosphoryl-containing fragments can be syn- or anti-oriented relative to the amide plane (Table 2, Figure 6).

According to NMR spectroscopy data (1H, 13C{1H}, 31P{1H}, in CDCl₃ solutions), diphenylphosphinylacetic acid amides 1–3 exist as two conformers in a ratio of ≈2.5–3.0:1, depending on the structure of the alkyl substituent at the nitrogen atom [37]. However, it was not possible to establish the structure of these two conformers.

A comparative analysis of the obtained experimental and theoretical results and the data of NMR spectroscopy [37] allowed us to conclude that in solution, compounds 1–3 exist as two sets of conformers: in the first set, the C=O and alkyl groups have Z-configuration in rotamers 1a, 1c, and 1e; 2a, 2c, and 2d; 3a, 3c, and 3d, while in the second set, these groups have E-configuration in rotamers 1b and 1d; 2b, 2e, and 2d; 3b, 3e, and 3d, and in all forms except 1e, 2f, and 3f, the phosphoryl-containing fragments are syn-located relative to the plane of the amide fragment. We believe that the first set of conformers corresponds to the minor conformer, and the second set corresponds to the
majority conformer, found by NMR spectroscopy data, which is confirmed by the total conformer ratio (Table 2). Elongation of the alkyl substituent at the nitrogen atom (methyl–butyl–octyl) in a series of 1–2–3 led to an increase in the number of preferred conformers from five for 1 to six for 2 and 3. However, the structures of the corresponding preferred conformers differ slightly, and steric factors—a flat amide fragment, the size of bulky phenyl groups, and the configuration of alkyl bridges—are important.

3. Materials and Methods

3.1. Materials

N-methyl-N-[2-(diphenylphosphoryl)ethyl]diphenylphosphorylamide 1, N-butyl-N-[2-(diphenylphosphoryl)ethyl] diphenylphosphorylacetamide 2, and N-octyl-N-[2-(diphenylphosphoryl)ethyl] diphenylphosphorylacetamide 3 were synthesized according to the following procedure [37].

3.2. IR Spectroscopy

The infrared spectra of crystals were collected on a FTIR Bruker Vertex 70 spectrometer (Bruker, Ettlingen, Germany) (600–4000 cm\(^{-1}\)) with a single reflection, germanium crystal ATR accessory (MIRacle, PIKE Technologies) purged under dry air to remove atmospheric water vapor. The interferograms were recorded with a resolution of 2 cm\(^{-1}\), 128 scans, and Fourier transformed using a Blackman–Harris apodization function. The thin films of molten compounds were produced by the heating of crystal between KBr plates in home-made electrical oven. Temperature was measured by the PT100 sensor and was kept constant using a PID controller to ensure a standard deviation smaller than 1 K. The solid phase of samples was produced by slow cooling of thin film of molten compound (liquid) between KBr plates. The crystallization of the films was observed visually between crossed polarizers. The comparison spectra of the solid phases after melting and followed crystallization and the crystals of the same compounds showed their identity. Thus, there was no decomposition of the samples studied upon melting. KBr cells were used with a spacer (0.2 mm) to achieve the best signal/noise ratio. Concentrations of compounds were varied from 0.05 to 0.1 mol/L. Chloroform purified using molecular sieves was used as a solvent.

3.3. Dipole Moments

The experimental values of the dipole moments were determined using the second Debye method [38]. Physical parameters of 1–3 were measured for series consisting of 4–6 solutions in benzene at 25 °C. The dielectric permittivities of solutions of 1–3 were determined on a BI-870 instrument (Brookhaven Instruments Corporation, New York, USA); the accuracy is ±0.01. The refractive indices of solutions were measured on a RA-500 refractometer (Kyoto Electronics, Kyoto, Japan); the accuracy is ±0.0001.

The experimental dipole moments were calculated by the Debye Equation (1) [38]:

\[
\mu = 0.01283 \sqrt{P_{or} \cdot T}.
\]  

The orientation polarizabilities \(P_{or}\) were calculated by the Guggenheim-Smith Equation (2) [43,44]:

\[
P_{or} = \frac{M}{d} \left[ \frac{3a}{(\varepsilon_0 + 2)^2} - \frac{3\gamma}{(n_0^2 + 2)^2} \right]
\]  

where \(M\) is the molecular weight of a substance, \(d\) is the solvent density, \(a\) and \(\gamma\) are slopes of the \(\varepsilon_i - \omega_i\) and \(n_i^2 - \omega_i\) plots; \(\varepsilon_i\), \(n_i\), and \(\omega_i\) are the dielectric constant, refractive index, and mass fraction of the solute of the \(i\)th solution, respectively. Equations for \(a\) and \(\gamma\) and the \(\varepsilon_i - \omega_i\) and \(n_i^2 - \omega_i\) plots (Figures S4–S6) are given in Supplementary Materials.

In the calculations of dipole moments according to the vector-additive scheme, we used the theoretical geometry parameters and following dipole moments of bonds and groups: \(m(C_\text{Ph} = \text{P})\) 1.09 D, calculated from \(\mu_{\text{exp}} (\text{C}_6\text{H}_5)\text{P}\) [41]; \(m(C = \text{O})\) 1.94 D [45];
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1a with a predominance of highly polar rotamers. Preferred conformers were characterized by Hessians containing only positive frequencies.

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Supplementary Materials: Figure S1: IR spectra of compound 2 in different aggregate states, Figure S2: IR spectra of compound 3 in different aggregate states, Figure S3. Visualization of the molecular orbitals for 1e (the interaction between the P = O bond and phenyl substituents is absent). The positive and negative potentials are shown as blue and red areas, respectively. Equations for α and γ calculations (Guggenheim–Smith equation) and the ε−μ and n2−μ plots, Figure S4. The ε−μ and n2−μ plots for compound 1. Figure S5. The ε−μ and n2−μ plots for compound 2. Figure S6. The ε−μ and n2−μ plots for compound 3.

Author Contributions: Conceptualization, N.B. and Y.V.; investigation, A.K., D.C. and Y.V.; synthesis, N.B. and O.A.; methodology, Y.V.; software, D.C.; project administration, Y.V.; supervision, Y.V.; visualization, A.K. and Y.V.; writing—original draft, A.K., D.C. and Y.V.; writing—review and editing, D.C. and Y.V. All authors have read and agreed to the published version of the manuscript.

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