Phosphorus-nitrogen compounds- (Part 50): correlations between structural parameters for cyclophosphazene derivatives containing ferrocenylic pendant arm(s)

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Abstract: The results of a systematic study of spiro-cyclotri/tetraphosphazenes with ferrocenylic pendant arm on the basis of correlation between structural parameters were presented. The main parameters were obtained from X-ray crystallography and 31P NMR results in order to investigate the relationship between the δPspiro shift values and endocyclic and exocyclic NPN bond angles, and electron density transfer parameters. Structural parameters derived from 11 types of the ferrocenylic cyclophosphazene derivatives with 5- to 7-membered spiro-rings introduced to the literature from our research group were studied and compared with each other.

Key words: spirocyclic Ferrocenylic phosphazene, 31P NMR, X-ray crystallography, endocyclic bond angle, exocyclic bond angle, electron density transfer parameter

1. Introduction
The phosphazene chemistry has attracted much attention since 1960 [1,2]. Especially, hexachlorocyclotriphosphazene (N3P3Cl6, trimer) and octachlorocyclotetraphosphazene (N4P4Cl8, tetramer) are of particular interest to both theoretical and experimental researchers concerning phosphazene-based chemistry. Because of their tendency to react with the nucleophilic mono-, di-, or multi-functional groups [3–6], both of the cyclophosphazenes were used in the syntheses of a considerable range of organocyclotri/tetraphosphazene derivatives with diverse applications [7,8]. The substantial efforts have been performed on the nucleophilic substitution reactions, in which the 1- to 6-Cl-atoms on trimer and 1- to 8-Cl atoms on tetramer have been replaced by the NH and/or OH functioned reagents, forming isomeric products e.g., structural (spiro-, ansa- and bino-architectures or a mixed of the same or different architectures), geometrical (geminal, non-geminal cis/trans-), and optical isomers [9,10]. The nature of the products strongly depends on the various chemical factors which control the replacement reaction mechanisms such as chain lengths of nucleophilic groups, the polarity of solvents, and the reaction temperature [11]. So far only a limited number of published studies on cyclophosphazene derivatives with ferrocenylic pendant arm is present in the literature [12–16].

The organocyclophosphazene derivatives have several potential applications in different fields of science as flame-retardant additives for organic polymers [17], liquid crystals [18,19], antibacterial [20] and anti-cancer [21] agents, fluorescence chemosensors [22], ion-transferring agents for rechargeable lithium batteries [23,24] and light-emitting diodes (LEDs) [25]. Besides, ferrocene-containing compounds have been used for molecular sen-
sors, biosensors, electron-transfer mediators, non-linear optical materials, liquid crystals, and redox-active probe materials [26,27]. In this context, we were therefore interested in synthesizing of ferrocenyl cyclophosphazenes and thought that the presence of both ferrocene moiety as a substituent and a trimeric/tetrameric phosphazene as a skeleton in a molecule could give rise to a novel kind of cyclophosphazene derivatives and bring together many biological and physicochemical properties of the molecule. Furthermore, cyclotri/tetraphosphazene ring systems are electrochemically inert, and ferrocenyl group is an excellent redox-active precursor. Hence, ferrocenyl cyclophosphazenes were synthesized to investigate the electrochemical behavior of the phosphazenes [28–30]. Furthermore, substituted spiro-monoferroenyl cyclotri/tetraphosphazenes were prepared by our group to evaluate in terms of their antituberculosis, anticancer, and antimicrobial activities. According to these studies, it was observed that geminal vanillinato (Van)-substituted spiro-monoferroenyl cyclotriphosphazenes [31], tetra pyrrolidine (Pyr)-substituted spiro-mono [32,33] and bisferrocenyl [33] cyclotriphosphazenes and hexa Pyr-substituted spiro-monoferroenyl cyclotetraphosphazenes [34] inhibited the growth of Mycobacterium tuberculosis H37Rv. While 1,4-dioxa-8-azaspiro[4,5]decane (DASD)-substituted spiro-bisferrocenyl [35] and partly substituted dispiro-bisferrocenyl [36] cyclotriphosphazenes, the fully and nongeminal (cis) [37] Van-substituted spiro-monoferroenyl cyclotriphosphazenes, were effective against the human cervical cancer cell line (HeLa), bis(diamino) substituted dispiro-bisferrocenyl cyclotetraphosphazene was found to be more active against colon cancer DLD-1 cells than Doxorubicin [38]. It was also found that the DASD and Pyr-substituted ferrocenyl cyclotriphosphazenes were active against some gram-positive and gram-negative bacteria [32,33,35] and Pyr-substituted ferrocenyl cyclotetraphosphazenes were more effective than the commercial antifungal drug Ketoconazole against fungi [34].

Besides, the chiral properties of mono Van-substituted dispiro-bis ferrocenyl cyclophosphazenes were investigated by $^{31}$P NMR spectroscopy upon the addition of the chiral solvating agent [39].

On the other hand, we also succeeded in the preparation of ultrathin and highly ordered Langmuir-Blodgett films of tetrachloro-, and mono and gem DASD-substituted mono-ferrocenyl cyclotriphosphazenes [40,41]. These compounds are the first phosphazene derivatives prepared as thin films in the literature.

Shaw described the first systematic study of the relationship between the bond angles around the phosphorus atoms and $^{31}$P NMR spectral data in phosphazene derivatives [42]. The changes in structural parameters for different kinds of structurally analogous cyclotriphosphazenes (cyclotriphosphazenes possessing 6-membered spiro ring/rings [43], monospiro-, dispiro-, spiro-ansa-spiro- and spiro-bino-spiro-cyclotriphosphazenes [44–46], spiro-cyclotriphosphazenic lariat (PNP-pivot) ether derivatives [47,48], monotopic and ditopic spiro-crypta cyclotriphosphazenes [49–51]) were investigated previously. It was found that the systematic variations in the $^{31}$P NMR chemical shifts depend fundamentally on some electronic (electron-releasing and electron-withdrawing capacities of substituent groups), steric (the steric hindrance between the exocyclic groups) and conformational factors, and on the changes in bond lengths and bond angles around the phosphorus atoms [especially endocyclic ($\alpha$) and exocyclic ($\alpha'$) bond angles] in cyclotriphosphazene derivatives. The current study deals with a number of correlations between structural parameters [e.g., $^{31}$P NMR spectral data and X-ray crystallographic data (endocyclic and exocyclic NPN bond angles, and bond lengths)] in spirocyclic ferrocenyl cyclophosphazenes introduced to the literature from our research group (Table 1) [33–41,52]. Therefore the content of this report includes: (i) a brief description of the synthesis methods of 11 different structural types and a total of 28 spirocyclic ferrocenyl phosphazenes with 5- to 7-membered spiro-rings used for the graph construction, (ii)
the relationship between the $\delta P_{\text{spiro}}$ shifts and the values of electron density transfer parameters $\Delta(P-N)$, and (iii) the correlation of $\delta P_{\text{spiro}}$ shifts and endocyclic ($\alpha$) and exocyclic ($\alpha'$) NPN bond angles of the compounds.

**Table 1.** The endocyclic ($\alpha$) and exocyclic ($\alpha'$) NPN bond angles and bond lengths (a, a', b, and b') on the formulae of cyclophosphazenes.
2. Results and discussion
2.1. Syntheses

Routes for the synthesis of spirocyclic ferrocenyl cyclophosphazenes clarified their solid-state structures using X-ray crystallography by our research group and investigated in this study are summarized in Scheme. The syntheses of mono and bisferrocenyl diamines, as the starting compounds, were carried out according to the published procedures, in which ferrocenecarboxaldehyde reacted with appropriate diamines and followed by reduction of the azomethine bonds in the intermediate products [53,54]. The reactions of trimer with mono and bisferrocenyl diamines gave partly substituted spiro-mono (I) [33] and spiro-bis (V) [33,52] ferrocenyl cyclotriphosphazenes, respectively. The substituted phosphazene derivatives were synthesized by stepwise substitutions of partly substituted spiro-mono (I) and spiro-bis (V) ferrocenyl cyclotriphosphazenes which consist of 4 reactive P-Cl units. The reactions of 1 equimolar amount of partly substituted spiro-bis (V) and spiro-mono (I) ferrocenyl cyclotriphosphazenes with 1 and 2 equimolar amounts of heterocyclic amines (DASD and Pyr) produced corresponding mono heterocyclic amine (DASD) substituted spiro-bis (VI) [35] and spiro-mono (II) [40] and geminal heterocyclic amine (DASD and Pyr) substituted spiro-bis (VII) [35] and spiro-mono (III) [35,40,41] ferrocenyl cyclotriphosphazenes in the presence of NEt₃ in refluxing dry THF. The fully heterocyclic amine [DASD, Pyr, and morpholine (Morp)] substituted spiro-bis (V) [33] and spiro-mono (I) [33,35,52] ferrocenyl cyclotriphosphazenes were prepared by replacing 4 Cl-atoms on partly substituted derivatives (I) and (V), respectively, with excess heterocyclic amines in boiling THF. The reactions of equimolar amounts of partly substituted spiro-mono ferrocenyl cyclotriphosphaze (I) and potassium vanillinate were found to yield the corresponding mono Van-substituted spiro-mono ferrocenyl cyclotriphosphaze (II) as a major product and geminal (III) [37] and nongeminal (cis) (IV) substituted spiro-mono ferrocenyl cyclotriphosphazenes as minor products. Fully Van-substituted spiro-bisferrocenyl cyclotriphosphazene (V) was synthesized from the reaction carried out with excess potassium vanillinate [37]. The Cl-replacement reactions of trimer with 2 equimolar amounts of mono-ferrocenylkdiamines resulted in the formation of the corresponding partly substituted cis- (meso) and trans- (racem) dispiro-bisferrocenyl cyclotriphosphazenes (VIII) as the major products and spiro-mono (I) ferrocenyl cyclotriphosphazenes as minor products [36]. Three products were separated performing column chromatography. The reactions of 1 equimolar amount of cis- and trans-dispiro-bisferrocenyl cyclotriphosphazenes (VIII) having 2 reactive Cl-atoms with 2 equimolar amounts of potassium vanillinate in refluxing THF afforded the mono (IX) and fully (VIII) Van-substituted cis- and trans-dispiro-bisferrocenyl cyclotriphosphazenes (IX) and (VIII) [39]. The mono and fully substituted derivatives were separated using column chromatography. On the other hand, the partly substituted spiro-mono (X) [34] and cis- and trans-dispiro-bis (XI) [38] ferrocenyl cyclotetraphosphazenes were obtained from the reactions of tetramer with 1 and 2 equimolar amounts of mono-ferrocenyl diamines in dry THF. The fully Pyr-substituted (X) and trans-(XI) were prepared by the reaction of partly substituted ones with excess Pyr in dry THF at ambient temperature.
Scheme. Routes for the synthesis of spirocyclic ferrocenyldicyclopophazene derivatives investigated in this study.
2.2. Correlations between structural parameters

The endocyclic (α) and exocyclic (α’) NPN bond angles, and the bond lengths (a, a’, b, and b’) were defined in the generalized structures for the 11 types of cyclotri/tetraphosphazenes containing ferrocenyl pendant arm/arms and 5-, 6- and 7-membered spiro-ring/rings shown in Table 1. δP_s piro shifts, α, and α’ bond angles, and \( \Delta(P-N) \) values that are needed to be used for graph construction are listed in Table 2. The corresponding values of the standard compounds trimer \((N_3P_3Cl_6)\) [55,56] and tetramer \((N_4P_4Cl_8)\) [57,58] were taken from the literature. Types \( I \) and \( V \) members are partly and fully substituted spiro-mono and spiro-bisferocenyl cyclotriphosphazenes, respectively. Mono and geminal substituted spiro-mono/bisferocenyl cyclotriphosphazenes are the types \( II \) and \( VI \), and the types \( III \) and \( VII \) group members, respectively. Nongeminal (cis) substituted spiro-monoferocenyl cyclotriphosphazene constitutes the type \( IV \). Members of types \( VIII \) and \( IX \) derivatives comprise partly and fully substituted and monosubstituted cis/trans-dispiro-bisferocenyl cyclotriphosphazenes, respectively. spiro-Mono and trans-dispiro-bisferocenyl cyclotetrahosphazenes constitute the types \( X \) and \( XI \) compounds.

The concept of the double-bond character of the P-N linkage in the cyclophosphazene derivatives is still not clearly understood. Negative hyperconjugation and ionic bonding alternatives are exclusive [59]. The natural bond orbital and topological electron-density analyses of phosphazenes have proved the crucial role of negative hyperconjugation in the description of the P-N bond.

2.2.1. The correlation of \( \delta P_{\text{spiro}} \) shifts and values of electron density transfer parameters \( \Delta(P-N) \)

The electron density transfer parameter \( \Delta(P-N) \) is the difference between the bond lengths of 2 adjacent endocyclic P-N bonds as defined in Table 2 for spirocyclic ferrocenyl phosphazenes. It shows a measure of the electron releasing and withdrawing capacities of the substituent groups on cyclophosphazene ring. The relationship between the \( \delta P_{\text{spiro}} \) shifts and \( \Delta(P-N) \) values is illustrated in Figure 1 for partly and heterocyclic amine [Pyr, piperidine (Pip), Morp and DASD] (i) and Van (ii) substituted spirocyclic ferrocenyl phosphazenes, respectively. A cluster of points rather than the linear trend was observed between the \( \Delta(P-N) \) and \( \delta P_{\text{spiro}} \)-shifts. In Figure 1i, all types of triphosphazene structures were accumulated in 6 regions A, B, C, D, E, and F. The points of partly substituted types \( I \) and \( V \) and type \( VIII \) phosphazenes accumulate in regions A and B, respectively. The points of mono (types \( II \) and \( VI \)), geminal (types \( III \) and \( VII \)) and fully heterocyclic amine substituted cyclotr (types \( I \) and \( V \)) and cyclotetra (type \( X \)) phosphazenes accumulate in regions C, D, E, and F, respectively.

According to Figure 1i, some comparisons can be made on the electron-releasing power of the substituent depending on whether the substituent is a chloro or heterocyclic amine group of the compounds with the same membered spiro-rings. For example, the \( \Delta(P-N) \) values of fully heterocyclic amine substituted \( I_d, I_c, \) and \( t-XIb \) are respectively; 0.0055, 0.0035, and –0.005 and –0.0055, indicating that the electron releasing power of the nitrogen atoms of heterocyclic amine groups is greater than that of the chloro groups in \( I_a \) (0.087), \( V_b \) (0.0675), and \( t-XIa \) (0.046) with the larger \( \Delta(P-N) \) values. A similar situation is observed for fully Van (\( V_d \) and \( c/t-VIIIId \)) and partly \([V_a \text{ and } (t-VIIId \text{ and } c/t-VIIIId)]\) substituted ferrocenyl cyclophosphazenes (Figure 1ii), showing the oxygen atoms of Van groups bonded to phosphorus atoms release electrons to the cyclophosphazene ring. It is not possible to say whether heterocyclic amines or vanniline release more electrons to the phosphazene ring since we do not have crystallographic data of the heterocyclic amines and Van substituted derivatives of any type are not available.
Table 2. Endocyclic (α) and exocyclic (α') NPN bond angles, bond lengths (a, a', b, and b'), δP\text{spiro} shifts and Δ(P-N) values for the compounds [δP\text{spiro} shifts in ppm, α and α' angles in °, a, a', b, and b' lengths in Å].

| Compound | a       | a'      | b       | b'      | Δ(P-N)  | α       | α'      | δP\text{NPN} |
|----------|---------|---------|---------|---------|---------|---------|---------|--------------|
| Ia\textsuperscript{33} | 1.613(3) | 1.614(3) | 1.548(3) | 1.548(3) | 0.0070  | 113.14(15) | 102.68(16) | 13.62        |
| Ib\textsuperscript{33} | 1.587(3) | 1.598(3) | 1.603(3) | 1.588(2) | -0.003  | 117.66(13) | 102.16(13) | 20.76        |
| Ic\textsuperscript{35} | 1.586(1) | 1.600(1) | 1.594(1) | -0.005  | 117.67(17) | 103.02(7) | 19.38        |
| Id\textsuperscript{52} | 1.604(1) | 1.597(1) | 1.597(1) | 0.0055  | 115.47(7) | 104.06(7) | 22.10        |
| IIa\textsuperscript{40} | 1.630(8) | 1.596(8) | 1.547(9) | 1.566(8) | 0.0565  | 113.7(4)  | 101.7(4)  | 17.00        |
| IIIa\textsuperscript{35} | 1.592(1) | 1.628(1) | 1.592(1) | 1.560(1) | 0.034   | 114.62(6) | 104.68(6) | 17.32        |
| IIIb\textsuperscript{41} | 1.579(3) | 1.616(3) | 1.598(3) | 1.562(3) | 0.0175  | 113.18(15) | 93.24(15)  | 22.06        |
| IIIc\textsuperscript{40} | 1.6208(10) | 1.5903(10) | 1.5634(10) | 1.5921(10) | 0.0278  | 113.32(5) | 95.00(5)  | 21.06        |
| IVa\textsuperscript{57} | 1.612(2) | 1.612(2) | 1.555(2) | 1.556(2) | 0.0565  | 113.50(10) | 100.13(10) | 15.41        |
| Va\textsuperscript{52} | 1.620(3) | 1.620(3) | 1.544(3) | 1.544(3) | 0.076   | 110.0(2)  | 105.0(2)  | 6.20         |
| Vb\textsuperscript{33} | 1.613(3) | 1.619(3) | 1.549(3) | 1.548(3) | 0.0675  | 113.09(13) | 103.14(13) | 14.56        |
| Vc\textsuperscript{33} | 1.599(4) | 1.591(3) | 1.592(3) | 1.591(4) | 0.0035  | 114.96(18) | 98.94(17)  | 22.08        |
| Vd\textsuperscript{57} | 1.613(3) | 1.613(3) | 1.563(3) | 1.563(3) | 0.05    | 113.3(2)  | 100.6(2)  | 21.60        |
| Vla\textsuperscript{35} | 1.592(2) | 1.626(2) | 1.574(2) | 1.554(2) | 0.045   | 114.42(13) | 102.85(11) | 14.41        |
| Vlla\textsuperscript{35} | 1.558(2) | 1.600(2) | 1.603(3) | 1.593(2) | -0.019  | 113.26(9) | 101.45(9) | 18.32        |
| t-VIIib\textsuperscript{36} | 1.620(4) | 1.587(3) | 1.555(4) | -        | 0.0485  | 113.31(17) | 94.49(18)  | 24.22        |
| c-VIIib\textsuperscript{36} | 1.618(3) | 1.593(3) | 1.552(3) | -        | 0.0535  | 112.98(16) | 94.33(18)  | 22.57        |
| t-VIIib\textsuperscript{36} | 1.614(3) | 1.589(3) | 1.558(3) | -        | 0.0435  | 113.24(16) | 94.61(16)  | 22.63        |
| c-VIIIc\textsuperscript{36} | 1.634(2) | 1.583(2) | 1.549(3) | -        | 0.0595  | 113.84(13) | 102.10(13) | 19.63        |
| t-VIIic\textsuperscript{36} | 1.6055(19) | 1.5951(17) | 1.5602(19) | -        | 0.0401  | 115.57(9)  | 104.35(10) | 19.65        |
| c-VIIId\textsuperscript{39} | 1.6014(19) | 1.588(2) | 1.5716(18) | -        | 0.0231  | 114.03(10) | 95.22(9)  | 27.32        |
| t-VIIId\textsuperscript{39} | 1.608(3) | 1.592(4) | 1.566(3) | -        | 0.034   | 113.43(17) | 93.75(19)  | 27.52        |

\[ \Delta (P - N) = \frac{a + a'}{2} - \frac{b + b'}{2} \]

for (I-VII), (X) and (XI)

\[ \Delta (P - N) = \frac{a + a'}{2} - b \]

for (VIII) and (IX)
Table 2. (Continued).

| Compound | a   | a'  | b   | b'  | Δ(P-N) | α      | α'     | δP<sub>NP</sub>N |
|----------|-----|-----|-----|-----|--------|--------|--------|-----------------|
| c-IXa<sup>39</sup> | 1.605(2) | 1.5922(19) | 1.562(2) | -   | 0.036  | 113.64(10) | 94.54(10) | 26.68           |
|          | 1.612(2) | 1.586(2)    | 1.563(2) | -   | 0.036  | 112.50(10)| 93.93(11) |                |
| t-IXa<sup>39</sup> | 1.610(3) | 1.593(3)    | 1.560(4) | -   | 0.0415 | 113.77(19)| 94.88(18) | 25.91           |
|          | 1.622(3) | 1.584(3)    | 1.550(4) | -   | 0.053  | 113.34(18)| 93.89(19) |                |
| Xa<sup>34</sup>   | 1.587(3) | 1.583(3)    | 1.589(3) | 1.586(3) | -0.0025 | 118.19(15)| 94.31(13) | 12.61           |
| Xb<sup>34</sup>   | 1.584(2) | 1.581(2)    | 1.592(2) | 1.588(2) | -0.0075 | 117.68(9) | 94.94(8)  | 12.27           |
| XIa<sup>38</sup>  | 1.5940(13)| 1.5853(14) | 1.5301(13)| 1.5570(13)| 0.046 | 114.91(7) | 102.01(7) | 1.19            |
| XIb<sup>38</sup>  | 1.584(3) | 1.573(3)    | 1.585(3) | 1.582(3) | -0.005  | 121.07(14)| 102.14(13) | 3.46            |
Moreover, there is no significant difference between the $\Delta (P-N)$ values of cis- and trans-structures of the same compound for types VIII and IX phosphazenes (0.00825 for VIIIb, 0.002475 for VIIIc, and 0.01095 for IXa). However, the difference between the $\Delta (P-N)$ values of cis- and trans-structures of the phosphazenes with 5-membered spiro-rings (VIIIb and IXa) is slightly larger than that of the phosphazene with 6-membered spiro-rings (VIIIc). That could be significantly attributed to the fact that 5-membered spiro-rings of c-VIIIb, t-VIIIb, c-IX and t-IX are in envelope conformation and 6-membered spiro-rings of c-VIIIc and t-VIIIc are in the chair conformation [36,39].

![Figure 1](image.png)

**Figure 1.** The relationship between $\delta P_{spiro}$ shifts and $\Delta (P-N)$ values for partly and heterocyclic amine (Pyr, Pip, Morp, DASD) (i) and Van (ii) substituted spirocyclic ferrocenyl phosphazenes. $\delta P_{ClPCl}$ shift values of $N_3P_3Cl_6$ and $N_4P_4Cl_8$ are 19.60 [56] and $-5.45$ [58] ppm, respectively.

For fully heterocyclic amine substituted phosphazenes (cycle E), the $\Delta (P-N)$ and $\delta P_{spiro}$ values of cyclotriphosphazenes having the 7-membered spiro-ring (Id and Vc) are similar, regardless of whether the compounds are mono (type I) and bis (type V) ferrocenyl phosphazenes.

It can be seen from Figure 1i that there are greater changes in $\Delta (P-N)$ values for types II and VI with 1 heterocyclic amine substituent per P atom, types III and VII with 2 heterocyclic amine substituents per P atom and types I and V with 4 heterocyclic amine substituents. Therefore, the $\Delta (P-N)$ values of these types phosphazenes can be compared with each other according to the number of heterocyclic amine substituents. As expected, the $\Delta (P-N)$ value of mono substituted compounds is between the $\Delta (P-N)$ value of partly (cycle A) and fully (cycle E) substituted phosphazenes, while geminal substituted derivatives except for VIIa (cycle D) have the $\Delta (P-N)$ value between those of mono (cycle C) and fully (cycle E) substituted ones. The $\Delta (P-N)$ value of VIIa appears to the left more than other geminal substituted derivatives (IIIa-IIIc) (cycle D) or is greater than those of the fully substituted derivatives (cycle E). This situation may be related to the higher basicity of the DASD substituent in VIIa. A similar relationship was observed between the $\Delta (P-N)$ values of nongeminal cis- (IVa) and fully (Vd) Van substituted cyclophosphazenes and partly substituted Ia and Va, respectively (Figure 1ii). Furthermore, the $\Delta (P-N)$ values of fully heterocyclic amine substituted types X (cycle F) and XI cyclotetraphosphazenes and types I and V cyclotriphosphazenes, respectively, are quite close together.
Although the compounds IIIa and VIIa both have geminal structure and 7-membered spiro-ring, and are close in δPspiro shifts, the major difference in their Δ(P−N) values and basicities is that the phosphazenes contain mono and bis ferrocenyl pendant arms, respectively. On the other hand, based on the electron-releasing capacity of the ferrocenyl pendant group for partly substituted phosphazenes (cycles A and B), it has been made the following order: Type VIII > type V > type I. Type I (Ia), and type V compounds (Va and Vb) are mono-spiro mono and bis structures, while type VIII (t-VIIIa, e/t-VIIlb, and c/t-VIIIc) phosphazenes are di-spiro bis structures. As expected, the electron releasing powers of 2 ferrocenyl pendant groups are greater than those of 1 ferrocenyl pendant group. Moreover, in partly substituted phosphazenes (cycle A), the δPspiro shifts of 7-membered Ia and Vb are close to each other while 6-membered Va has a lower δPspiro shift.

In the case of 5-membered spiro-ring geminal (IIIb and IIIc) and 6-membered spiro-ring fully ( Ib and Ic) substituted phosphazenes, the electron releasing capacity of DASD group is much larger than that of Pip and Pyr, respectively.

Besides, when the number of atoms increases in the spiro-ring, the electron releasing capacity of the phosphazene decreases. In general, the electron releasing power of the rings is in the following order: spiro-rings with 5-membered > spiro-rings with 6-membered > spiro-rings with 7-membered.

As a result, electron − withdrawing substituents, like chlorine group, increase Δ(P−N) values, pulling away electrons from spiro-ring/rings to the phosphorus atom bonded to the electron− withdrawing groups. Whereas the electron-releasing substituents, like heterocyclic amines, decrease Δ(P−N) values, resulting in decreased the bond lengths a and a’ and increased the bond lengths b and b’ when compared bond lengths of partly substituted derivatives. Hence, the shortening of the endocyclic P−N bonds and decreased electron charge density at the exocyclic P-N bonds is likely to be a measure of the electron-releasing power of the substituent and the increase in negative hyperconjugation.

The relationship between the Δ(P−N) and δPspiro shifts makes sense in the basicity of the ring nitrogen atoms in phosphazenes. The basicity of the chlorocyclophosphazene ring nitrogen atoms is quite low, and it may be improved by replacing Cl-atoms with electron-releasing substituents on phosphorus. Thus, the basicity of the phosphazene ring nitrogen atoms (N1−PX2 and N2−Pspiro) in fully substituted cyclotriphosphazenes with those in partly substituted ones can be compared. The basicity of N1 atom/atoms in fully substituted phosphazenes appears to have increased due to electron-releasing power of the heterocyclic amine groups, while N2 atom/atoms in partly substituted phosphazenes due to electron-withdrawing power of the chloro groups. As a result, an increase in the electron-releasing power of heterocyclic amine substituents seems to bring about an increase in the basicity of the nitrogen atom (N1) and the negative hyperconjugation.

2.2.2. The correlation of the δPspiro shifts, endocyclic (α), and exocyclic (α’) NPN bond angles
A cluster of points between the δPspiro shifts and the endocyclic NPN bond angles (α) [A, B, C, D, E, and F given in Figure 2i)] and a trend of approximate linearity between the δPspiro shifts and the exocyclic NPN bond angles (α’) [(a), (b), (c), and (d) given in Figure 2ii] were observed.

The changes in α and α’ bond angles show parallelism except for a contrasting trend observed for partly substituted types I and V cyclotriphosphazenes (cycle A) and fully substitute type X cyclotetraphosphazenes (cycle F). Small changes in α’ bond angles lead to significant changes in δPspiro shifts. The number of members in the spiro-ring seems to be effective on α’ bond angles. In fact, the α’ bond angles of cyclotriphosphazenes with 5-membered spiro-ring are narrower than those with larger 6- and 7-membered ones and even narrower
The relationship between δP spiro shifts and endocyclic (α) (i) and exocyclic α’ (ii) NPN bond angles for partly and heterocyclic amine (Pyr, Pip, Morp, DASD) substituted spirocyclic ferrocenyl phosphazenes. δP ClP Cl shift values of N$_3$P$_3$Cl$_6$ and N$_4$P$_4$Cl$_8$ are 19.60 [56] and –5.45 [58] ppm, respectively. The α and α’ values are 118.3(2) and 101.2(1)° for N$_3$P$_3$Cl$_6$ [55] and 121.2 and 102.8° for N$_4$P$_4$Cl$_8$ [57] respectively.

Figure 2. The relationship between δP spiro shifts and endocyclic (α) (i) and exocyclic α’ (ii) NPN bond angles for partly and heterocyclic amine (Pyr, Pip, Morp, DASD) substituted spirocyclic ferrocenyl phosphazenes. δP ClP Cl shift values of N$_3$P$_3$Cl$_6$ and N$_4$P$_4$Cl$_8$ are 19.60 [56] and –5.45 [58] ppm, respectively. The α and α’ values are 118.3(2) and 101.2(1)° for N$_3$P$_3$Cl$_6$ [55] and 121.2 and 102.8° for N$_4$P$_4$Cl$_8$ [57] respectively.
substituted type XI cyclotetraphosphazenes, $\alpha$ angle is much affected by the substitution, but, $\alpha'$ angle is less affected. Moreover, the correlations between the $\delta P_{\text{spiro}}$ shifts and $\alpha$ (Figure 3i) and $\alpha'$ (Figure 3ii) NPN bond angles show contrasting trends. For example, the $\alpha$ and $\alpha'$ angles of 6-(Va) and 4-(Ia) membered partly substituted phosphazenes are smaller than 6-membered fully Van-(Vd) and 4-membered nongeminal cis- (IVA) substituted phosphazenes, respectively.

![Figure 3](image_url)

**Figure 3.** The relationship between $\delta P_{\text{spiro}}$ shifts and endocyclic ($\alpha$) (i) and exocyclic $\alpha'$ (ii) NPN bond angles for partly and Van substituted spirocyclic ferrocenyl phosphazenes.

Although there are few examples of spiro-ferrocenyl substituted cyclotetraphosphazenes, the structural parameters of these compounds are given in the figures with the aim of comparison purposes. More values are necessary to learn more about the correlations for cyclotetraphosphazenes.

3. **Conclusions**

A systematic study concerning the correlations between structural parameters [e.g., $^{31}$P NMR spectral data and X-ray crystallographic data (endocyclic and exocyclic NPN bond angles, and bond lengths)] displayed some characteristic results for mono- and di-spirocylic phosphazene derivatives bearing ferrocenyl pendant arm/arms. Naturally, these results become more reliable when more cyclic phosphazenes from this series are synthesized and the $^{31}$P NMR spectroscopic and X-ray crystallographic data of these molecules are taken into account. It is necessary to extend the study for other members of the spirocyclic ferrocenyl cyclophosphazene family to get a more general and including views about the correlations between structural parameters of these molecules. Research along these lines is actually under development in our laboratory and results will be presented elsewhere in the forthcoming future.

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