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# Stereochemistry of phosphorus compounds: A short communication 

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#### Abstract

The phosphorus element having a polymeric nature and formed the macro-compounds or polymers. The phosphorus compounds such as the poly-(phosphates and phosphazenes) were shown the stereochemistry as optical-geometrical isomerism with chair-boat conformation and molecular symmetry (C3v, C2h and D3h) in their novel structure. The structure of these polymeric compounds have been characterized by IR, X-ray, Raman spectra and by 31P NMR analyses as well.


Keywords: Stereochemistry, conformation, symmetry, polyphosphate, polyphosphazene

## 1. Introduction

In periodic table, the Phosphorus is a nonmetallic chemical element of group 15 or nitrogen family with symbol P and having atomic number as 15 . The phosphorus is a colorless, soft and waxy solid at room temperature that glow in dark conditions. Recently, a large number of compounds of phosphorus-containing P-C, P-N and P-O bonds have been reported from natural and synthetic resources ${ }^{[1-3]}$. Many of them have shown interesting in biological process as play a vital role in metabolic intermediates, coenzyme and in nucleic acid ${ }^{[4]}$. They are part of ADP, ATP, RNA and DNA as a number of important biological elements of living organisms with object of intensive scientific research. These area of research on phosphorus compounds is still little studies ${ }^{[5,6]}$, that prompted us to discuss and analyze it in our review research work. In this article, we have reported about a stereochemistry as conformation and symmetry in P - compounds such as in poly- (phosphates and phosphazenes) molecules ${ }^{[7-9]}$, (figure-1 a, b).


Fig 1: The linear structure of (a) Polyphosphate and, (b) Polyphosphazene

## 2. Poly-(Phosphates and phosphazene)

Polyphosphates are a salt or ester of polymeric oxy-anions and formed from tetrahedral phosphate $\left(\mathrm{PO}_{4}\right)$ units, which is linked together by sharing phosphorus-oxygen atoms ${ }^{[10]}$. The polyphosphates can have a linear, branched or cross-linked and cyclic ring structures with electrolytic properties ${ }^{[11]}$. In biology, the phosphates ester are involved in nucleotides, energy storage, protein synthesis and in carbohydrate metabolite action with great attention. The polyphosphazenes (Phosphonitrilic chloride) are included a wide range of hybrid organic-inorganic polymers with $-\mathrm{P}=\mathrm{N}$ - backbone and having the different skeletal or molecular architectures with different side groups (R) ${ }^{[12,13]}$. These phosphorus-containing polymeric polyphosphates and polyphosphazenes have been finding in a variety of applications.

## 3. Results and Discussion



Linear polyphosphate
(b)


Cyclic tripolyphosphate (metaphosphate)
(c)

(A)


Cyclophosphazenes

Linear phosphazenes | $\begin{array}{l}\text { Replace chlorine } \\ \text { atoms by organic } \\ \text { groups, } \mathrm{R}\end{array}$ |
| :--- |



## Linear polyphosphazenes high polymers

Fig 2: The various type of (A) Polyphosphates and, (B) Polyphosphazenes


Fig 3: The possible imaginary (a) chair and, (b) boat shape for poly- (phosphates and phosphazenes)

Indeed, the polyphosphates and polyphosphazenes have the linear, branched or cross-linked and cyclic ring structures (Figure-2.1). These are may be cyclic -trimer, -tetramer or other higher polymers type. These can have possible stereochemistry with chair and boat conformation, as in figure-2.2, (source: from internet), which are well described earlier in our previous paper ${ }^{[14]}$. There is an ample evidence that the trimeta-phosphate $\left(\mathrm{P}_{3} \mathrm{O}_{9}\right)^{3-}$ ion rings have chair conformation with $\mathrm{C}_{3 \mathrm{v}}$ point-group ${ }^{[15]}$ in crystalline solid state and in solution consistent with $\mathrm{D}_{3 \mathrm{~h}}$ molecular symmetry, whereas the tetra-metaphosphate $\left(\mathrm{P}_{4} \mathrm{O}_{12}\right)^{4-}$ ion rings are puckered but conformations vary from compound to compound. The vibrational spectroscopic data of $\left(\mathrm{P}_{3} \mathrm{O}_{9}\right)^{3-}$ ion having the $\mathrm{P}-\mathrm{O}$ bond length of 0.1615 nm (endocyclic) and 0.1484 nm (exocyclic) even the larger one is about 0.01 nm shorter than would be anticipated for ion a single bond supporting the presence of pi- ( $\mathrm{p} \pi-\mathrm{d} \pi$ ) bonding in all P-O bonds. Where the crystal structure of the chair form of $\left(\mathrm{NH}_{4}\right) \quad \mathrm{P}_{4} \mathrm{O}_{12}$ which gives equal ring ( 0.1607 nm ) and terminal ( 0.1479 nm ) P-O bonds with $\mathrm{C}_{2 \mathrm{~h}}$ molecular symmetry. The structure of hexametaphosphate $\left(\mathrm{P}_{6} \mathrm{O}_{18}\right)^{6-}$ ion also shows equivalence in the two kinds of P-O bond, both in close agreement with these observed for the tri- and tetrahomologues of phosphates.

The extensive interest in the phosphazenes, the hydrolysis of trimeric chloride to the acid, takes place slowly with a tautomeric change as shown for the derivatives of alcohol and amines. The chelated trimer spiro phosphazenes such as 1, 1-dichloro-trans-3, 5-bis (p-tolyl)-3, 5-diphenyl cyclotriphosphazene have optical isomers ${ }^{[16]}$. However, some cyclophosphazene derivatives and aryl-halogen ophosphazenes also have geometrical (Cis-trans) isomerism ${ }^{[17]}$. Consequently, the alkoxy- and aryloxy phosphazenes show tautomerism and the aziridinyl phosphazene's derivatives, with partial replacement not only give-rise to positional but also cis-trans isomerism ${ }^{14}$. The molecular geometry of P-N polymers has been assigned by Raman, IR and X-Ray analysis resembles that of the $\left(\mathrm{NPF}_{2}\right)_{x}$, where $\mathrm{x}=$ 3 or 4; are planar, but larger rings are not planar. For other $\left(\mathrm{NPX}_{2}\right)_{\mathrm{n}}$ compounds the six rings are planar with $\mathrm{D}_{3 \mathrm{~h}}$ symmetry, but larger rings are generally non-planar and NPN angles of $\sim 120^{\circ}$ and PNP angles of $\sim 132^{\circ} \mathrm{C}$. The P-N bond distances which are generally equal in these ring systems, lie in the range $1.55-1.61 \AA$; there considerable attention to $\mathrm{P}-\mathrm{N}$ stretching is at an appreciable higher frequency ( $1220 \mathrm{~cm}^{-1}$ ) than that expected for a P-N single (about $750 \mathrm{~cm}^{-1}$ ) bond of as $1.75-1.80 \AA$ and paid to the nature of $\mathrm{P}-\mathrm{N} \pi$ bonding ${ }^{[14]}$.
Although, these rings are conformationally flexible and $\pi$ bonding is only one of many factors that influence the conformation, but the matter is still subject to controversy. The main question concerns the extent of delocalization all around the rings show a kind of aromatic character or whether these are more localized 'islands' within the NPN segments. Of course, there may be considerable differences between the essentially planar rings and those that are puckered. The problem is a complicated one owing to the large number of orbitals potentially involved and to the general lack of ring planarity which means that rigorous
assignment of $\sigma$ and $\pi$ character to individual orbitals is impossible respectively.

## 4. Conclusions

The various spectroscopic data and chemical analyses confirm that, the polymer of phosphorus such as poly(Phosphates and phosphazenes) have been shown the stereoand geometrical isomerism with chair-boat conformation and molecular symmetry (mostly $\mathrm{C}_{3 \mathrm{v}}, \mathrm{C}_{2 \mathrm{~h}}$ and $\mathrm{D}_{3 \mathrm{~h}}$ ), in their structure.

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