FULL PAPER

The eccentric connectivity index of polycyclic aromatic hydrocarbons (PAHs)

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Mathematical chemistry is the area of research engaged in new application of Mathematics in Chemistry. Major areas of research in mathematical chemistry include chemical graph theory. Chemical graph theory applies graph theory to mathematical modeling of chemical phenomena. If $G=(V(G),E(G))$ is a connected graph, where $V(G)$ is a non-empty set of vertices and $E(G)$ is a set of edges, then the eccentric connectivity index of $G$ (denoted by $\xi(G)$) was defined as $\xi(G) = \sum_{v \in V(G)} d_v \cdot \varepsilon(v)$, where $d_v$ is the degree of a vertex $v$ and $\varepsilon(v)$ is its eccentricity. In this study, we investigated the eccentric connectivity index of polycyclic aromatic hydrocarbons (PAHs).

KEYWORDS
Connected Molecular Graph; eccentric connectivity index; distance; polycyclic aromatic hydrocarbons (PAHs).

Introduction

Let $G=(V,E)$ be a simple connected graph with vertex set $V(G)$ and edge set $E(G)$. A molecular graph is a simple connected graph such that its vertices correspond to the atoms and the edges to the bonds. The degree of a vertex/atom $v$ in $G$ is the number of edges incident to it and is denoted by $d_v$. The distance between the vertices $u$ and $v$, denoted by $d(u,v)$, is the length of the shortest path joining them. The eccentricity $\varepsilon(v)$ of a vertex $v$ is the distance to a vertex farthest from $v$, that is

$$\varepsilon(v) = \text{Max}\{d(u,v)\mid \forall u \in V(G)\}$$

The radius $R(G)$ of a graph $G$ is the minimum eccentricity of the vertices and the diameter $D(G)$ of $G$ is the maximum eccentricity.

$$D(G) = \text{Max}_{v \in V(G)}\{d(u,v)\mid \forall u \in V(G)\}$$

$$R(G) = \text{Min}_{v \in V(G)}\{\text{Max}\{d(u,v)\mid \forall u \in V(G)\}\}$$

An eccentric vertex of a vertex $v$ is a vertex farthest away from $v$. An eccentric path of a vertex $v$ denoted by $P(v)$ is a path of length $e(v)$ joining $v$ and its eccentric vertex. There may exist more than one eccentric path for a given vertex.

A topological index of a graph is a number related to a graph which is invariant under graph automorphisms and is a numeric quantity from the structural graph of a molecule.

The eccentric connectivity index $\zeta(G)$ of a graph $G$ is defined by V. Sharma, R. Goswami, A.K. Madan, in 1997 [1], as

$$\zeta(G) = \sum_{v \in V(G)} d_v \cdot \varepsilon(v)$$

Recently, we know the eccentric connectivity polynomial of a graph $G$ as

$$ECP(G,x) = \sum_{v \in V(G)} d_v \cdot x^{\varepsilon(v)}$$

Obviously, the eccentric connectivity index is the first derivative of $ECP(G,x)$ evaluated at $x=1$. Also, the total eccentricity index of $G$ is defined as

$$\Theta(G) = \sum_{v \in V(G)} \varepsilon(v)$$
M.J. Morgan et al. [2] investigated the eccentric connectivity index of trees. For further study and more details see references [3-6].

In this paper, we investigate the eccentric connectivity index of a family of hydrocarbon molecules, which are called Polycyclic Aromatic Hydrocarbons (PAHs).

**Results and discussion**

Mathematical chemistry is the area of research engaged in new application of mathematics in chemistry. Major areas of research in mathematical chemistry include chemical graph theory. Chemical graph theory applies graph theory to mathematical modeling of chemical phenomena.

Polycyclic aromatic hydrocarbons (PAHs) are a group of over 200 different chemicals formed when wood, coal, oil, gasoline or other organic materials are burned. They can also be formed in food when fish or meats are charbroiled. "Polycyclic aromatic hydrocarbon" is just a mouthful for an organic molecule in space—not just any organic molecule, but one with a structure sturdy enough to survive the harsh environment between stars. Usually called "PAHs," this class of molecule is almost certainly the culprit responsible for the unidentified infrared emission (UIR) bands discovered in the early 1970s. The UIR bands are seen in many different kinds of objects from starburst galaxies to comets, so understanding their origin is very important. Because the UIR bands are so common, the PAHs which produce them must account for a sizeable fraction of all interstellar carbon in our galaxy and others. Estimates run as high as 15% or more.

Polycyclic Aromatic Hydrocarbons (PAHs) consist several copies of benzene C6 on circumference and is ubiquitous combustion products. They have been implicated as carcinogens and play a role in graphitization of organic materials [7]. In addition, they are of interest as molecular analogues of graphite [8] as candidates for interstellar species [9] and as building blocks of functional materials for device applications [8-10]. Synthetic routes to PAHs are available [11] and a detailed knowledge of all these features would therefore be necessary for the tuning of molecular properties towards specific applications (see the references [7-12]).

![FIGURE 1](https://example.com/figure1.png)

**FIGURE 1** [20] Some example of polycyclic aromatic hydrocarbon PAHs.

The PAHs can be thought as small pieces of graphene sheets with the free valences of the dangling bonds saturated by H. Vice versa, a graphene sheet can be interpreted as an infinite PAH molecule. Successful utilization of PAH molecules in modeling graphite surfaces has been reported earlier [13-34] and references therein.

The first famous members of this hydrocarbon family (PAH family) are denoted and shown as follows (See Figures 1 and 2):
- **PAH₁** be the Benzene with six carbon (C) and six hydrogen (H) atoms,
- **PAH₂** be the Coronene with 24 carbon and twelve hydrogen atoms,
- **PAH₃** be the Circumcoronene with 54 carbon and eighteen hydrogen atoms.
Now, by reference to the eccentric connectivity index \( \zeta(H) \) of Circumcoronene series of Benzenoid \( H_k \) [31-34] and figures therein (see Figure 3), we can compute the eccentric connectivity index \( \zeta(PAH_n) \) easily as:

At first, we denote all vertices as degree three (carbon \( C \) atoms) of polycyclic aromatic hydrocarbon \( PAH_n \) by \( \beta \) and \( \gamma \) (similar to Figure 4) and all vertices as degree two (hydrogen \( H \) atoms) by \( \alpha \). Alternatively, the vertex/atom set and edge/bond set of \( PAH_n \) are equal to:

\[
V(PAH_n)=\{ \alpha_{z,j}, \beta_{z,j}, \gamma_{z,j} \mid i=1, ..., k; \ z \in \mathbb{Z} \ & I=1, ..., l; \ j \in \mathbb{Z} \} \ & E(PAH_n)=\{
\alpha_{z,j}, \gamma_{z,j}^k, \beta_{z,j}^l, \gamma_{z,j}^{-1}, \beta_{z,j}^{-1}, \gamma_{z,j}^{i+1}, \beta_{z,j}^{i+1}, \gamma_{z,j}^{-i} & j \in \mathbb{Z} \ \& z \in \mathbb{Z} \}
\]

where \( \mathbb{Z} = \{1, 2, ..., l\} \) is the cycle finite group of order \( i \).

So, we divide all vertices in some partitions "ring-cuts", such that a \( i \)th ring-cut consists of vertices \( \beta_{z,j}^l, \gamma_{z,j}^{-i} \ (\forall i=1, k; \ z \in \mathbb{Z}, \ j \in \mathbb{Z}) \) and the size of this ring-cut is equal to \( 6i+6(i-1) \). Also, one common property of members of a ring-cut is their farthest vertices. And another properties of them is \( d(\gamma_{z,j}, \gamma_{z,j}^{-i})=d(\beta_{z,j}, \gamma_{z,j}^l)\)

\[=2(k-i), \] in others words the distance of these vertices are equal to two times of difference between order of their ring-cuts. By attention to ring cuts of polycyclic aromatic hydrocarbon \( PAH_n \) and using above mentions and results from [29], we have following formula:

- For all vertices \( \beta_{z,j}^i \) of \( PAH_k \) (\( \forall i=1, ..., k; \ z \in \mathbb{Z} \), \( j \in \mathbb{Z}, i \) 1)

\[
e^{-\frac{e(\beta_{z,j}^i)}{4l-3}}+d(\beta_{z,j}^i, \gamma_{z,j}^l)+d(\gamma_{z,j}^l, \alpha_{z,j}^k)
\]

\[=2(k-i-1) \]

- For all vertices \( \gamma_{z,j}^{-i} \) of \( PAH_k \) (\( \forall i=1, ..., k; \ z \in \mathbb{Z} \), \( j \in \mathbb{Z} \))

\[
e^{-\frac{e(\gamma_{z,j}^{-i})}{4l-1}}+d(\gamma_{z,j}^{-i}, \gamma_{z,j}^l)+d(\gamma_{z,j}^l, \alpha_{z,j}^k)
\]

\[=2(k-i-1) \]
\[ \xi(\text{PAH}_n) = 2(k + i) \]

- For all vertices \( \alpha_{ij} \) (hydrogen (H) atoms) of \( \text{PAH}_n \) (\( \forall j = 1, \ldots, k; \ z = 1, \ldots, 6 \))

\[ e(\alpha_{ij}) = d(\alpha_{ij}, \gamma_{ij}) + d(\gamma_{ij}, \gamma_{ij}') + d(\gamma_{ij}', \alpha_{ij}) \]

\[ = 4k + 1 \]

- The diameter number of \( \text{PAH}_n \) is equal to \( D(\text{PAH}_n) = e(H) = 4k + 1 \).
- The radius number of \( H_k \) is equal to \( R(H_k) = 2k + 1 \).

Now, we are in a position to state and prove our main results of this paper. Therefore, the eccentric connectivity index \( \xi \) of polycyclic aromatic hydrocarbon \( \text{PAH}_n \) is equal to:

\[ \xi(\text{PAH}_n) = \sum_{v \in V(\text{PAH}_n)} d_v \times e(v) \]

\[ = \sum_{\alpha_{ij}} d_{\alpha_{ij}} e(\alpha_{ij}) + \sum_{j > i} d_{\beta_{ij}} e(\beta_{ij}) + \sum_{j = i} d_{\gamma_{ij}} e(\gamma_{ij}') \]

\[ = \sum_{\alpha_{ij}} \left( \sum_{j = 1}^{k} d_{\alpha_{ij}} e(\alpha_{ij}) + \sum_{j > i} d_{\beta_{ij}} e(\beta_{ij}) + \sum_{j = i} d_{\gamma_{ij}} e(\gamma_{ij}') \right) \]

Thus the eccentric connectivity index of polycyclic aromatic hydrocarbon \( \text{PAH}_n \) is

\[ \xi(\text{PAH}_n) = 6n^3 + 33n^2 + 9n. \]

**FIGURE 4** The general representation of ring-cut method of circumcoronene series of benzenoid \( H_k \) for all integer number \( k \geq 2 \) [31-34]
Example 1. Consider Benzene PAH₁ (Figure 1), by using the above results, the eccentric connectivity index of Benzene is equal to $\xi(\text{PAH}_1)=102$.

Example 2. Consider Coronene PAH₂ (Figure 1), by using the above results, one can see that the eccentric connectivity index of the second member of polycyclic aromatic hydrocarbon is equal to $\xi(\text{PAH}_2)=630$.

Conclusion

In graph theory, we have many invariant polynomials for a graph $G$. In this research, we focus on the structure of Polycyclic Aromatic Hydrocarbons (PAHs). We have computed the eccentric connectivity index of this hydrocarbon molecule.

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