Volume 13, No. 2, 2022, p. 3467-3478 https://publishoa.com ISSN: 1309-3452

Fourteen Degree-based Topological Indices on twelve Penicillin Class of Drugs with QSPR Analysis

C.Tamilarasi¹, F. Simon Raj²

¹Hindustan Institute of TechnologyandScience, Chennai, Tamilnadu, India;

Email: tamilbabu4864@gmail.com

²Hindustan Institute of TechnologyandScience, Chennai, Tamilnadu, India;

E mail: simon240906@gmail.com

Received: 2022 March 15; Revised: 2022 April 20; Accepted: 2022 May 10.

Abstract- This research article aims to establish the values of fourteen notable topological indices that are defined on some chemical structures belonging to the Penicillin class of drugs, which are a type of antibiotic derived from Penicillin fungi. This is performed to predict the Physicochemical properties of these compounds through Quantitative Structure-Property Relationships (QSPR). Also, it is observed that these characteristics have a good correlation with the Physico-chemical characteristics of the selected chemical structures of the penicillin class of compounds. The chemical properties of the selected compounds that this paper discusses include Polarizability, Log Octanol-Water Partition Coefficient(Log K_{ow}), Molecular Volume, Molar Refractivity, Log P, Soil Absorption Coefficient (Koc), Log Koc.

MSC2010codes 05C10, 05C30, 05C92, 05C90,05C09

Keywords- Topological index, degree-based topological index, Quantitative Structure-Property Relationships, Penicillin, Physico-chemical properties.

1IntroductionChemical graph theory is one of the key branches of graph theory and playsa significant role in applied mathematics, which deals with chemical structures.Atoms representvertices and chemical bonds represent edges of the

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chemical structures in chemical graph theory. The numeralvalue associated to the vertices as well as edges in the molecular graph is termed as topological index/molecular descriptor. One of the main qualities of the topological indices is to predict the Physicochemical Properties of the respected chemical compounds through QSPRstudy^{2,9,10,12,13}. In this paper, some of the properties of the Penicillin class of drugs are taken for QSPR analysis with prominent fourteen degreebased topological indices which are given good correlation with them.

Penicillinwas the first true antibiotic to be discovered, which helped to fight bacterial infections such as whooping cough, ear infection, Salmonella, and Tuberculosis. The Scottish bacteriologist Alexander Fleming first observed that colonies of the bacterium Staphylococcus aureus failed to grow in certain areas of its culture that had been accidentally contaminated by the mold Penicillium notatum which was green in color. This is the well-known Penicillium notatum, which was finally discovered in 1928.

Wiener index, the first topological index(distance-based)wasintroduced by Harold Wiener in 1947.It is defined as the summation of the shortest distance between every single vertex of graph G. The progress of degree-based topological indices^{3,4}was started in 1975 by Milan Randic.Degree-based as well as distancebased topological indices are widely used in predicting the physicochemical properties^{7,11} of a class of compounds. Based on that, this paper is established. Here, fourteen notable degree-based topological indices are taken with twelve penicillin classes of drugs in the prediction task of properties through QSPR analysis.

2 **Preliminaries**In this part. some supportivenotions are mentioned here, like, molecular graph G, degree of a vertex, topological index, degree-based topological indices, and basic formulas of fourteen degree-based topological indices. Let the molecular graph be the two-dimensional model of the chemical compound and is denoted as graph G(V,E) holds two nonempty sets V(set of all atoms as vertices)& *E*(set of all chemical bonds as edges). The degree of vertex v is the number of vertices adjacent to the vertex v.

The basic definitions of the fourteen topological indices are as follows:

Definition 2.1One of the important degree-based topological indices whichhave a highly predictive ability of Physico-chemicalproperties of chemical

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compounds is the Augmented Zagreb Index (Az(G)). It was introduced by Furtula*etal* and is denoted as follows:

$$Az(G) = \sum_{u,v \in E} \left(\frac{dudv}{du + dv - 2} \right)^{3}$$

Definition 2.2 Estrada *et al.* introduced the degree-based topological index named Atom Bond connectivity index(ABC(G)) which is a good predictor for the stability of alkanes and strain energy and is denoted as follows:

$$ABC(G) = \sum_{u,v \in E} \sqrt{\frac{du + dv - 2}{dudv}}$$

Definition 2.3TheGeometricArithmetic Index $(GA(G))^1$ is the well-studied degree-based topological index and is denoted as follows:

$$GA(G) = \sum_{u,v \in E} \frac{2\sqrt{dudv}}{du+dv}$$

Definition 2.4 Harmonic $Index(H(G))^{15}$

$$H(G) = \sum_{u,v \in E} \frac{2}{du + dv}$$

Definition 2.5 Forgotten $\text{Index}(F(G))^{14}$ was defined to be used in the analysis of drug molecular structures which was introduced by B. Furtula*et al.* and is denoted as follows:

 $F(G) = \sum_{u,v \in E} du^2 + dv^2$

Definition 2.6 The Zagreb indices $M_1(G)$, $M_2(G)$, and $M_3(G)$ are the numerical

measures to pronounce the structureproperty relationships. $M_1(G)$ is the oldest degree-based topological index that was initiated in 1972 and the related versions were developed later.

$$M_{1}(G) = \sum_{u,v \in E} (du + dv)$$
$$M_{2}(G) = \sum_{u,v \in E} (du \, dv)$$
$$M_{3}(G) = \sum_{u,v \in E} |du - dv|$$

Definition 2.7TheRandic Index⁵ is one of the most powerful degree-based topological indices to study the structureproperty relationships of graphs and is denoted as follows:

$$R(G) = \sum_{u,v \in E} \frac{1}{\sqrt{du \, dv}}$$

Definition 2.8 Sum Connectivity Index¹⁶is denoted as follows:

$$S(G) = \sum_{u,v \in E} \frac{1}{\sqrt{du + dv}}$$

Definition 2.9 In 2013, Shirdel*et al.* proposed novel indices such as the First Hyper-Zagreb $index(H_1(G))$, Second Hyper-Zagreb $index(H_2(G))$ and are denoted as follows:

$$H_1(G) = \sum_{u,v \in E} (du + dv)^2$$
$$H_2(G) = \sum_{u,v \in E} (dudv)^2$$

Definition 2.10 FORAN index(FR(G)) holds the properties of the Forgotten index as well as the Randic index, and it is

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suitable for predicting the relationship between molecular structures of drugs and their physicochemical properties. It is denoted as follows:

$$FR(G) = \sum_{u,v \in E} du \sqrt{\frac{du}{dv}} + dv \sqrt{\frac{dv}{du}}$$

Definition 2.11Symmetric Division $DegIndex(SDD(G))^{6}$ was defined asD.Vukicevicin 2010 by the order of its vertex degrees with the maximum degree and the minimum degree and is denoted as follows:

$$SDD(G) = \sum_{u,v \in E} \frac{\max(du,dv)}{\min(du,dv)} + \frac{\min(du,dv)}{\max(du,dv)}$$

Based on the above fourteen degree-based topological indices, we computed twelve penicillin classes of drugs. Table 1 and its continuation show the values of fourteen topological indices with the chemical structures of the penicillin class of compounds. Table 2 shows the results of the properties.

Figure 1 Molecular structures of Penicillin class of drugs

Ampicillin

Oxacillin



Amoxicillin



Nafcillin







Carbenicillin



Penicillin



Flucloxacillin



Dicloxacillin



Ticarcillin



Pheneticillin

Cloxacillin

Methicillin

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Penicillin	Az(G)	ABC(G)	GA(G)	H(G)	F(G)	$M_1(G$	$M_2(G)$
class))
of drugs							
Amoxicillin	209.9397	19.6809	25.5873	10.885	390	138	168
				7			
Ampicillin	206.5647	18.8644	24.7615	10.585	370	132	161
				7			
Carbenicillin	221.3303	20.3475	26.6277	11.419	398	142	173
				1			
Dicloxacillin	276.1237	24.5956	32.3278	13.657	504	176	221
				1			
Nafcillin	274.1024	22.8761	30.7981	13.052	453	163	203
				4			
Oxacillin	239.522	22.3192	29.7213	12.552	440	158	195
				5			
Penicillin	195.6454	18.2454	18.0254	10.057	375	129	158
				1			
Ticarcillin	230.7237	21.1999	27.2766	11.523	448	158	190
				7			
Cloxacillin	267.3487	23.9189	31.3298	13.123	498	172	215
				7			
Methicillin	230.5803	21.5430	26.7407	11.552	398	142	175
				5			
Flucloxacillin	266.8772	23.8715	31.4533	13.219	480	170	211
				0			
Phenethicillin	267.1741	19.4049	25.7213	11.052	378	136	164
				3			
					1		

Table 1.It shows the calculated values of twelvePenicillin class of drugs with fourteen degree-based topological indices.

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Table 1 continuation

Penicillin	$M_3(G)$	R(G)	S(G)	$HM_1(G)$	$HM_2(G)$	FR(G)	SDD(G)
class							
of drugs							
Amoxicillin	28	11.6308	11.9878	774	1326	159.0796	68.2500
Ampicillin	24	11.2370	11.6827	692	1177	150.0692	64.5822
	26	12.1475	12.5909	744	1267	162.4651	69.9166
Carbenicillin							
	34	14.5521	15.1634	946	1731	200.0976	84.5000
Dicloxacillin							
Nafcillin	25	13.6832	14.3792	859	1503	181.0681	76.0833
Oxacillin	26	13.2201	13.8936	830	1427	177.3061	74.9167
Penicillin	29	10.7570	11.1611	691	1212	150.9228	64.1666
Ticarcillin	32	12.4311	12.8607	828	1520	178.3217	75.6666
Cloxacillin	34	13.9867	14.6495	928	1671	197.7770	86.9998
Methicillin	26	12.2236	12.6400	748	1297	226.3317	68.5833
	30	14.0415	14.7101	902	1575	192.8531	81.5833
Flucloxacilli							
n							
	26	11.7201	12.1690	706	1168	126.4385	66.9167
Phenethicilli							
n							

Table 2. Itshows the Penicillin class of drugs with their physicochemical properties

Penicillin	Polarizabilit	Log	Molar	Molar	Log	K _{oc}	Log
class of	у	K _{ow}	Volume	Refractivit	Р		k _{oc}
drugs	10^{-24} cm^3			у			
				Cm ³			
	36.3	0.97	236.2	91.5	0.61	865.5	115.7
Amoxicilli							

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n							
	35.7	1.45	239.3	89.9	1.35	534.4	105.4
Ampicillin							
	36.7	1.19	246.2	92.6	1.01	2169	113.0
Carbenicil							
lin							
	44	3.86	290.1	111.0	3.02	9004	106.5
Dicloxacil							
lin							
Nafcillin	43.7	3.79	289.9	110.3	3.52	1448	109.6
Oxacillin	40.2	2.57	268.5	101.3	2.05	3294	105.8
Penicillin	34.2	1.85	235.2	86.3	1.67	421.4	102.5
	36	1.01	236.4	90.7	0.69	1176	117.3
Ticarcillin							
	42.1	3.22	279.3	106.2	2.53	5446	106.2
Cloxacilli							
n							
	37.4	2.20	262.6	94.4	1.27	50.31	99.3
Methicilli							
n							
	42.1	3.42	283.7	106.3	2.60	9004	104.4
Flucloxaci							
llin							
	36.7	2.29	258	92.7	2.22	292.3	104.0
Phenethici							
llin							

3 Results and discussions

Part 1 Here, the indices are correlated with the penicillin class of drugs and are

presented in table 3.High correlation coefficients are highlighted in the table and establish linear regression models for the highlighted values through QSPR analysis.

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Topological	Polarizability	Log	Molar	Molar	Log P	K _{oc}	Log
Index	10^{-24} cm^3	K _{ow}	Volume	Refractivity			K _{oc}
				Cm ³			
$A_Z(G)$	0.8566	0.8404	0.8952	0.8593	0.8155	0.6169	0.4897
ABC(G)	0.9277	0.7935	0.8844	0.9271	0.6491	0.8135	0.6709
GA(G)	0.8885	0.6753	0.8268	0.8870	0.5719	0.7057	0.6488
H(G)	<u>0.9694</u>	0.8373	0.9242	<u>0.9689</u>	0.7335	0.8078	0.7167
F(G)	0.8664	0.7263	0.7783	0.8649	0.6091	0.8376	0.7799
$M_1(G)$	0.8959	0.7408	0.8129	0.8945	0.6294	0.8210	0.7708
$M_2(G)$	0.9201	0.7798	0.8430	0.9188	0.6663	0.8342	0.7670
$M_3(G)$	0.3654	0.3021	0.2720	0.3646	0.1785	0.6449	0.5569
R(G)	<u>0.9558</u>	0.8160	0.9023	0.9552	0.7055	0.8289	0.7370
S(G)	<u>0.9623</u>	0.8299	0.9110	<u>0.9617</u>	0.7266	0.8243	0.7421
$HM_1(G)$	0.8841	0.7133	0.7795	0.8824	0.5902	0.8320	0.7847
$HM_2(G)$	0.8264	0.6630	0.7121	0.8243	0.5353	0.8024	0.7524
FR(G)	0.5563	0.4655	0.5608	0.5557	0.2305	0.4579	0.1602
SDD(G)	0.8508	0.7016	0.7663	0.8495	0.5841	0.8378	0.7868

Table 3.It shows correlation coefficients between the physicochemical properties and indices.

Part 2In this section, suitable linear regression⁸ models(based on the highlighted values of table 3) of fourteen topological indices, and physicochemical properties of twelve penicillin class of drugs are established.

Linear regression models(QSPR analysis)

Linear regression model $N = n(Topological index) + n^{\circ}$

where N = physical property, TI = Topological index, n = slope & n^{\circ} = intercept.

Using this, we obtained the following.

1. Augmented Zagreb Index

Polarizability = 0.1021(Az(G)) + 14. 212 Log Kow = 0.0309 (Az(G)) - 5.1145Molar Volume = 0.6717 (Az(G)) + 98.9103

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> Molar Refractivity = 0.2591(Az(G)) +35.4633 Log P = 0.0263(Az(G)) - 4.4479

2. Atom Bond Index

Polarizability = 1.4956(ABC(G)) + 6.7448Molar Volume = 8.9797 (ABC(G)) + 68.2343Molar Refractivity =

3.78171(ABC(G)) + 16.8166

Koc = 1244.774 (ABC(G)) - 23836.4

3. Geometric-Arithmetic Index

Polarizability = 0.7673(GA(G)) + 17.6341 Molar Volume = 4.4964 (GA(G)) +

136.6615

Molar Refractivity = 1.9381(GA(G)) + 44.4099

4. Harmonic Index

Polarizability = 0.7677(H(G)) + 17.6341 Molar Volume = 4.4964 (H(G)) + 136.6615

Molar Refractivity = 7.1243(H(G)) + 13.0583

Koc = 2228.133 (H(G)) - 23683.4

5. Forgotten Index

Polarizability = 0.0608(F(G)) + 12.7484 Molar Refractivity = 0.1536(F(G)) + 32.0659 Koc = 55.8028 (F(G)) - 21056.3

6. First Zagreb Index

Polarizability = $0.1847(M_1(G)) +$ 10.8079 Molar Volume = $1.0553 (M_1(G)) +$ 100.7412 Molar Refractivity = $0.4666(M_1(G)) +$ 27.1598 $\text{Koc} = 160.6362 (M_1(G)) - 20500.9$ 7. Second Zagreb Index Polarizability = $0.1406(M_2(G)) +$ 12.5793 Molar Volume = $0.8114 (M_2(G)) +$ 109.4039 Molar Refractivity = $0.3553(M_2(G)) +$ 31.6220 $\text{Koc} = 120.9964 (M_2(G)) - 19716.7$ 8. Randic Index Polarizability = 2.6713(R(G)) + 5.0046 Log Kow = 0.7039 (R(G)) - 6.5758Molar Volume = 15.8801 (R(G)) +59.7903 Molar Refractivity = 6.7561(R(G)) +12.4161

Koc = 2198.565(R(G)) - 24972.1

9. Sum Connectivity Index

Polarizability = 2.6713(S(G)) + 5.0046 Log Kow = 0.6561(S(G)) - 6.3137Molar Volume = 14.694(S(G)) + 67.111

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> Molar Refractivity = 6.2326(S(G)) + 15.763Koc = 2003.8(S(G)) - 23556

10. First Hyper Zagreb Index

Polarizability = $0.0333(HM_1(G))$ + 11.9924 Molar Refractivity = $0.0841(HM_1(G))$ + 30.1714 Koc = 29.7338(HM_1(G)) - 21097.2

11. Second Hyper Zagreb Index

Polarizability = $0.0147(HM_2(G)) + 18.0735$ Molar Refractivity = $0.0371(HM_2(G))$

+45.5608

 $\text{Koc} = 13.5559(HM_2(G)) - 16253.1$

12. Symmetric Division Deg Index

Polarizability = $0.0147(HM_2(G)) + 18.0735$

Molar Refractivity = $0.0371(HM_2(G))$

```
+45.5608
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 $Koc = 13.5559(HM_2(G)) - 16253.1$

The above twelve indices are highly correlated with some specific physicochemical properties of the Penicillin class of compounds(Polarizability, Log Octanol-Water Partition Coefficient(Log K_{ow}), Molecular Volume, Molar Refractivity, Log P, Soil Absorption Coefficient (Koc), Log Koc).

Hence, the QSPR analysis is done with the specific properties. The third Zagreb index and the FORAN index have moderately correlated properties. Geometric-Arithmetic index. Harmonic index, Randic index, and Sum Connectivity index have a very high correlation with the properties, especially with Polarizability and Molecular Refractivitythat are underlined and highlighted in table 3. Figures 2 to 6 give apiece of solid information regarding this.



Figure 2 shows the correlation of H(G)With Polarizability of Penicillin class of compounds.



Figure 3 shows the correlation of H(G)WithMolar Refractivity of Penicillin class of compounds.

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Figure 4 shows the correlation of R(G)With Polarizability of Penicillin class of compounds.



Figure 5 shows the correlation of S(G)With Polarizability of Penicillin class of compounds.



Figure 6 shows the correlation of S(G)WithMolar Refractivity of Penicillin class of compounds.

Conclusion-In this paper, a QSPR analysis was established between fourteen degree-based topological indices and twelve

penicillin classes of drugs for the suitable pairs(topological index, property). In the field of pharmaceutical sciences, the properties of chemical compounds are very much required to establish new products. Instead of spending time, space for work, laboratory and the cost of research,topological indices are good alternators without these requirements. This theoretical approach obtained in this paper has promising aspects in designing new drugs.

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