Validation of topochemical models for the prediction of permeability through the blood-brain barrier

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Recently published topochemical models for permeability through the blood-brain barrier were validated and cross-validated in the present study. Five models based on three topochemical indices, Wiener's topochemical index – a distance-based topochemical descriptor, molecular connectivity topochemical index – an adjacency-based topochemical descriptor and eccentric connectivity topochemical index - an adjacency-cum-distance based topochemical descriptor, for permeability of structurally and chemically diverse molecules through blood-brain barrier were used in the present investigation. A data set comprising 62 structurally and chemically diverse compounds was selected. This data set was divided into two sets of 31 compounds each - one to serve as the validation set and other as the cross-validation set. The values of all the three-topochemical indices in the original as well as in the normalized form for each of the 31 compounds of the validation set were computed using an in-house computer program. Resultant data was analyzed and each compound was assigned a permeability characteristic using topochemical models, which was then compared with the reported permeability through the blood-brain barrier. Accuracy of prediction of these models was calculated. The same procedure was similarly followed for the crossvalidation set. Studies revealed accuracy of prediction of the order of 70-80% during validation. Surprisingly, very high predictability of the order of 77-91% was observed during cross-validation. High predictability observed during validation as well as cross-validation authenticates topochemical models for prediction of permeability through the blood-brain barrier.

Keywords: topochemical indices, Wiener's topochemical index, molecular connectivity topochemical index, eccentric connectivity topochemical index, permeability, blood-brain barrier

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An important aspect of drug design is the consideration of the potential for penetration of the blood-brain barrier by a new candidate drug molecule (1). There has been a surge in computational efforts to compute absorption, distribution, metabolism, excretion, and toxicity properties, including blood-brain barrier (BBB) partitioning, of structurally diverse compounds, including drugs (2–5). A good example of the great utility of a predictive computational model in drug discovery is the model for predicting BBB penetration (6). Prediction of passage across the BBB is of importance for centrally acting drugs or peripherally acting drugs, which should be devoid of CNS side effects (7). The BBB is a selective barrier formed by the endothelial cells that line cerebral microvessels. It acts as a physical barrier because complex tight junctions between adjacent endothelial cells force most molecular traffic to take a transcellular route across the BBB, rather than moving paracellularly through the junctions, like in most endothelia (8). Modeling blood-brain portioning is a challenging problem both because of the paucity of data and the task of establishing a useful relation between the molecular structure and measured blood-brain partitioning (1). Experimental determination of brain-blood partitioning is time-consuming, difficult and expensive (9). A broadly applicable method for predicting the BBB permeation of candidates at an early stage of discovery would have a great impact on drug research and development (10).

Physicochemical properties and biological activities of organic compounds change in a very systematic way with changes in chemical structure (11). Topological indices have been successfully employed in developing a suitable correlation between chemical structure and biological activity by translating chemical structures into numerical descriptors (12). Topostructural and topochemical indices fall into the category normally grouped together as topological indices. Topostructural indices are topological indices that encode information about the adjacency and distance of atoms in molecular structures, irrespective of the chemical nature of the atoms involved in bonding or factors such as hybridization states and the number of core/valence electrons in individual atoms. Topochemical indices are parameters that quantify information about the topology (connectivity of atoms), as well as specific chemical properties of the atoms making a molecule (13).

The objective of the present study is to validate the recently published topochemical models (14) for the prediction of permeability through the blood-brain barrier using external validation and cross-validation sets. Validation and cross-validation of the topochemical models based on Wiener's topochemical index, molecular connectivity topochemical index and eccentric connectivity topochemical index in the original and their normalized forms for permeability through the blood-brain barrier have been investigated.

EXPERIMENTAL

Calculations of topochemical indices

Wiener's topochemical index (W_c) (15) is a modified form of the oldest and most widely used distance based topological index – Wiener's index (16) and this modified

index takes into consideration the presence as well as relative position of heteroatoms in a molecular structure. It is defined as the sum of chemical distances between all the pairs of vertices in a hydrogen suppressed molecular graph, *i.e.*:

$$W_{c} = \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} P_{i_{c}j_{c}}$$
(1)

where $P_{i,j,}$ is the chemical length of the path containing the smallest number of edges between vertex i and j in graph G, n is the maximum possible number of i and j.

The normalized Wiener topochemical index (nW_c) values were calculated as the ratio of Wiener's topochemical index value to the total number of vertices in a hydrogen suppressed molecular graph.

Molecular connectivity topochemical index (χ^A) (17, 18) is defined as the summation of the modified bond values of adjacent vertices for all the edges in the hydrogen suppressed molecular graph according to the following equation:

$$\chi^{A} = \sum_{i=1}^{n} (V_{i}^{c} V_{j}^{c})^{-1/2}$$
 (2)

where n is the number of vertices, V_i^c and V_j^c are modified degrees of adjacent vertices i and j forming the edge $\{i, j\}$ in a graph G. This is a modified form of one of the most widely used adjacency based topological indices – molecular connectivity index (19) and it takes into consideration the presence as well as relative position of heteroatom(s) in a molecular structure.

The normalized molecular connectivity topochemical index ($n\chi^A$) was calculated as the ratio of the molecular connectivity topochemical index value to that of the total number of vertices in a hydrogen suppressed molecular graph.

The eccentric connectivity topochemical index (ξ_c^c) (20) is a modified form of an adjacency-cum-distance based topological index – eccentric connectivity index (21) and this modified index takes into consideration the presence as well as relative position of the heteroatom(s) in a molecular structure. The eccentric connectivity topochemical index is defined as the summation of the product of chemical eccentricity and the chemical degree of each vertex in the hydrogen suppressed molecular graph having n vertices, that is:

$$\xi_c^c = \sum_{i=1}^n (E_{ic} V_{ic})$$
 (3)

where V_{ic} is the chemical degree of vertex i, E_{ic} is the chemical eccentricity of vertex i and n is the number of vertices in graph G.

The normalized eccentric connectivity topochemical index $(n\xi_c^c)$ was calculated as the ratio of the eccentric connectivity topochemical index value to that of the total number of vertices in a hydrogen suppressed molecular graph.

The authors made an attempt at a simpler approach to predict the permeability through BBB of diverse series of compounds using topochemical models. These reported

Table I. Topochemical models derived from a training set of 28 chemically and structurally diverse compounds (14)

Index	Range in the model	Index value	Overall accuracy of prediction (%)
Wiener's topochemical index (W _c)	Permeable Transitional Impermeable	≤ 910.056 > 910.056 - < 3004.191 ≥ 3004.191	93.8
Normalized Wiener's topochemical index (nW _c)	Permeable Transitional Impermeable	≤ 49.04 > 49.04 - < 92.27 ≥ 92.27	94.7
Molecular connectivity topochemical index (χ^A)	Permeable Transitional Impermeable	≤ 12.086 > 12.086 - < 13.744 ≥ 13.744	83.3
Eccentric connectivity topochemical index (ξ_c^c)	Permeable Transitional Impermeable	≤ 404.227 > $404.227 - < 1032.901$ ≥ 1032.901	94.1
Normalized eccentric connectivity topochemical index $(n\xi_c)$	Permeable Transitional Impermeable	≤ 19.249 > 19.249 - < 31.949 ≥ 31.949	88.9

topochemical models (Table I) were developed using a training set of 28 structurally and chemically different compounds with established CNS permeation tendency, having the predictability from 83 to 95% (14). The aforementioned topochemical models were validated by an external test set of 31 compounds and cross-validated using another external test set of 31 chemically and structurally diverse compounds.

A reliable and predictive model should be statistically significant and robust, provide accurate prediction for an external dataset not used during model development and have its application boundaries defined. Iyer *et al.* (22) have reported a BBB study on a training set of 56 structurally and chemically diverse molecules and 7 molecules for the test set. The authors have omitted one molecule (methane) from this data set for lack of sufficient computed properties and used the remaining 55 compounds of training set and 7 compounds of the test set. These 62 compounds were divided into two sets. Compounds having an odd serial number were designated as the test or validation set and those having an even number were separated as the cross-validation set. The 31 compounds for validation set are listed in Table II and 31 compounds for the cross-validation set are listed in Table III.

The values of Wiener's topochemical index/normalized Wiener's topochemical index were computed for each compound using an in-house computer program. Subsequently, characteristic permeability was assigned to each compound using the reported models (14), which was then compared with the reported permeability (22). Permeability was reported quantitatively as log BB value. The compounds possessing log BB values of ≤ -0.3 were considered to be permeable and compounds possessing log BB values of > -0.3 were considered to be impermeable for the purpose of the present study. Vari-

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		q		1	I	I	+	1	1	
	ugh ier	Reported	$n\xi_c^c$	+1	I	I	+1	+1	+	
	y thro n barr	Re	ν, o	+	+1	I	+1	+1	+	
	Permeability through blood-brain barrier	- F	х	+	+1	+1	+	+	+	
	Perm bloc	Predicted	nW_c	+	I	I	+1	+1	+	
		Pr	W _c	+	+1	1	+1	+1	+	
	3	nç.		24.394	32.119	40.462	19.806	30.164	14.568	
	3	ŭ		390.311	899.333	1173.39	415.918	633.454	218.52	
		χ^{A}		6.958	13.092	12.923	9.741	8.865	6.749	
	,	$n_{\rm W}$		43.682	97.853	119.938	49.955	70.448	25.747	
	147	o M		698.904	2739.88	3478.19	1049.06	1479.41	386.201	
		Compound		HN S NH NH CH ₃		H ₃ C-N _H S NH H COH ₃	E - Z	H ₃ C-N H ₃ C-N H ₃ C-N NH CH ₃	H ₃ C/N CH ₃	Ξ
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Table II. continued

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	٦		1	+	1	1	1	+
ough tier	Reported	$n\xi_c^c$	1	1	+	+1	1	+1
y thro	Ř	νς o	+1	+1	+	+1	1	+I
Permeability through blood-brain barrier	٦	х	+	+	+	+	+1	+
Perm bloc	Predicted	nW_c	+1	+1	+	+1	1	+I
	Pı	M _c	+1	+1	+	+I	ſ	+1
о ц Z	ر ر		37.168	32.624	17.561	24.104	37.094	23.822
n	J.		743.368	848.217	280.972	530.286	1075.72	595.538
∢ ;	×χ		8.197	11.795	6.953	9.694	12.915	11.459
W.	O A A 1/		62.505	88.306	31.518	57.672	112.258	67.364
W	o •		1250.09	2295.96	504.292	1268.79	3255.47	1684.1
	Compound		Br O ₂ N NH NH NH	NT HAN S	H ₂ N NH ₂	H ₂ N NH CH ₃	N ₂ O HN	23 GH ₃ NH H ₃ C NH
2	NO.		13	15	17	19	21	23

Table II. continued

NH NH NH 1809.29 75.387 11.517 695.952 2 NH NH NH NH 1309.29 75.387 11.517 695.952 2 NH NH NH 1309 2.375 1.643 11.109 2 The State of the state o		,				3	3		Pern blo	Permeability through blood-brain barrier	ity thr in bar	ough rier	
$\begin{array}{ c c c c c c c c c c c c c c c c c c c$		Compound	o M	$n_{\rm C}$	χ^{A}	, v	$n\xi_c$		redicte	p	Y	eporte	٦
2564.03 87.078 12.432 795.744 30.606 ± ± ± ± ± ± ± 1809.29 75.387 11.517 695.952 28.998 ± ± ± ± ± ± ± 18.666 3.733 2.19 20.665 4.133 + + + + + + + + + + + + + + + + + +								W	$nW_{\rm c}$		m,	$n\xi_c^c$	
1809.29	$\langle \overline{} \rangle$	\	2264.03	87.078	12.432	795.744	30.606	+I	+1	+I	+1	+I	+
18.666 3.733 2.19 20.665 4.133 + + + + + + + + + + + + + + + + + +			1809.29	75.387	11.517	695.952	28.998	+I	+1	+	+I	+I	+
18.666 3.733 2.19 20.665 4.133 +		. \	2593.08	96.04	12.553	960.121	35.56	+I	1	+I	+1	I	+
31 5.167 2.808 29 4.833 +		Butanone	18.666	3.733	2.19	20.665	4.133	+	+	+	+	+	+
9.5 2.375 1.643 11.109 2.777 +		3-methylpentane	31	5.167	2.808	29	4.833	+	+	+	+	+	+
32 5.333 2.77 31 5.167 +		2-propanol	9.5	2.375	1.643	11.109	2.777	+	+	+	+	+	+
37.268 6.211 1.963 52.919 8.82 + <td></td> <td>2-methylpentane</td> <td>32</td> <td>5.333</td> <td>2.77</td> <td>31</td> <td>5.167</td> <td>+</td> <td>+</td> <td>+</td> <td>+</td> <td>+</td> <td>+</td>		2-methylpentane	32	5.333	2.77	31	5.167	+	+	+	+	+	+
21.998 4.4 2.235 28.218 5.644 +		,1,1-trifluoro-2-chloroethane	37.268	6.211	1.963	52.919	8.82	+	+	+	+	+	+
4.333 1.444 1.309 7.443 2.481 + + + + + + + + + + + + + + + + + + +		Diethylether	21.998	4.4	2.235	28.218	5.644	+	+	+	+	+	+
98.809 12.351 2.377 177.248 22.156 + + + \pm		Ethanol	4.333	1.444	1.309	7.443	2.481	+	+	+	+	+	+
		Halothane	608.86	12.351	2.377	177.248	22.156	+	+	+	+	+I	+

Table II. continued

		777	744		3	3		Pern bloc	Permeability through blood-brain barrier	ty thruin bar	ough rier	
No.	Compound	o M	$n_{\rm C}$	χ^{A}	'n	nç,	Ь	Predicted	رم و	×	Reported	٦
							W	nW_{c}	χΑ	'n	пξс	
47	Hexane	35	5.833	2.914	38	6.333	+	+	+	+	+	+
49	Methylcyclopentane	26	4.333	2.894	29	4.833	+	+	+	+	+	+
51	Propanol	10.5	2.625	1.825	15.665	3.916	+	+	+	+	+	+
53	Teflurane	93.997	11.75	2.436	168.081	21.01	+	+	+	+	+1	+
55	Trichloroehtane	29.748	5.95	1.454	57.913	11.583	+	+	+	+	+	+
57	NH ²	204.264	17.022	5.692	150.44	12.537	+	+	+	+	+	I
29	O N N N N N N N N N N N N N N N N N N N	538.192	29.9	8.571	235.683	13.094	+	+	+	+	+	+
61	LA L	1628.326	62.628	11.417	603.393	23.207	+1	+1	+	+1	+1	1

(+) - Permeable compound, (-) - impermeable compound, (±) - compounds in the transitional range where permeability could not be specifically assigned.

Table III. Relationships between topochemical indices and permeability through blood-brain barrier for a cross-validation set comprising of another 31 compounds

	_		+	1	+	+	I	I
ugh rier	Keported	$n\xi_c^c$	+	1	+	+	+1	1
Permeability through blood-brain barrier	X	ņ	+	1	+	+	+1	+1
neabilit od-bra	g	χ^{A}	+	1	+	+	+	+
Perm	Predicted	nW_c	+	1	+	+	+1	+1
	H	M_{c}	+	I	+	+	+1	+1
มีก ว	3		12.145	43.007	15.363	16.745	31.31	34.655
ພິ	,		121.447	1376.22	215.087	351.653	626.207	658.446
γA	₹		4.017	14.569	5.991	10.039	8.228	7.907
$nW_{\rm c}$	ų		14.602	138.908	24.266	43.336	62.799	59.505
W	v		146.019	4445.05	339.73	910.056	1315.98	1130.6
Compound	H		H ₂ N N ₄ H		IZ Z	N - CH	NH ₂ NH NH CH ₃	Br CN
O			61	4.	9	œ	10 H ₂	12

Table III. continued

	_		I	+	1	I	+	I
ugh ier	Reported	$n\xi_c$	+1	+	+I	+1	+1	+1
Permeability through blood-brain barrier	Re	ຶ່ວ	+1	+	+	+1	+1	+1
neabilit od-bra	g	χ^{A}	+	+	+	+	+	+
Perm	Predicted	nW_c	+1	+	+	+1	+1	+1
		W	+1	+	+	+1	+1	+1
o alu	, L		27.375	17.253	20.867	31.544	24.525	24.77
m)	ù o		520.118	258.794	396.473	693.957	588.595	520.173
A.	X		8.323	962:9	8.235	9.443	10.908	9.842
W	O 11		55.61	28.436	43.609	74.163	64.835	58.072
×	ن •		1056.58	426.536	828.565	1631.59	1556.05	1219.51
Companied			NI N ^{ci} O	N ₂ HN N	H ₂ N N N N N N N N N N N N N N N N N N N	O ₂ N H ₃ C-N H ₃ C-N _N H	H ₃ C-N _H	N CHANGE OF THE
Z	ON		14	16	18	20	22	24

Table III. continued

Table III. continued

	q		+	+	+	+	+	1	+
ugh ier	Reported	$n\xi_c^c$	+	+	+	+	+	+	+
Permeability through blood-brain barrier	Re	m,	+	+	+	+	+	+	+
ermeability through blood-brain barrier	_	$\chi^{\rm A}$	+	+	+	+	+	+	+
Perme bloo	Predicted	nW_c	+	+	+	+	+	+	+
	Pre	W _c	+	+	+	+	+	+	+
3.	ης°		4.8	2.777	6.249	11.62	17.58	14.012	16.258
3.	, v		24	11.109	45	127.821	334.016	266.232	341.41
•	χ^{A}		2.414	1.643	3.394	5.094	8.999	8.962	10.133
1447	$n_{\rm C}$		4	2.375	9	16.319	41.028	31.949	42.374
1 4 7	o M		20	9.5	42	179.511	779.532	607.024	889.849
	Compound		Pentane	Propanone	Toluene	LO CH	O Z D C F	O Z Z Z Z	J. J.
;	So.		20	52	54	26	28	09	62

(+) - Permeable compound, (-) - impermeable compound, (±) - compounds in the transitional range where permeability could not be specifically assigned.

ous researchers including Iyer *et al.* (22) and Abraham *et al.* (23) had reported that compounds with log BB values of > 0.3 are readily permeated into the brain whereas compounds with values <-1 are poorly permeated into the brain. The cut-off value consi- dered for the present study was the average value of the ranges reported by earlier researchers (22, 23). Accuracy of prediction of permeable and impermeable ranges as well as the overall degree of prediction of the validated model were also calculated. A similar procedure was followed for the molecular connectivity topochemical index and the eccentric connectivity topochemical index/normalized eccentric connectivity topochemical index.

The aforementioned procedure was similarly followed during cross-validation of the second set of 31 compounds. The results are summarized in Tables II and III.

RESULTS AND DISCUSSION

Among the pharmacokinetics issues in the design of new drugs, prediction of the BBB permeability is a crucial factor (24). The relationship of topochemical models based on Wiener's topochemical index, molecular connectivity topochemical index and eccentric connectivity topochemical index with permeability through BBB was evaluated for prediction of permeability through the blood-brain barrier.

All the 62 compounds reported by Iyer *et al.* (22) were employed for validation study of topochemical models. These compounds were divided equally into two different groups to constitute validation and cross-validation sets. Compounds having odd serial numbers were designated as validation set while those having even numbers were categorized as cross-validation set.

The accuracy of prediction for the test set and for the cross-validation set are shown in Table IV. The methodology used in the present study relates to validation of topochemical models. These models have high potential for providing permeable compounds through exploitation of permeable ranges in the models derived from topochemical indices. These models are unique and differ widely from conventional QSAR models. Both systems of modeling have their advantages and limitations. In the present case, the modeling system adopted has the distinct advantage of identification of narrow permeable ranges, which may be erroneously skipped during routine regression analysis in conventional QSAR modeling. Since the ultimate goal is to provide permeable compounds, these permeability ranges can play a vital role in providing permeable compounds (14).

Retrofit analyses of the data (Tables I to IV) for validation and cross-validation sets reveal that the compounds were classified either as permeable or impermeable using the aforementioned models. A transitional range between permeable and impermeable ranges is ideal because it simply reveals the gradual change in permeability from the permeable range to an impermeable range. The overall accuracy of prediction during validation was found to vary from 70 to 80% (Table IV). However, the overall accuracy of prediction during cross-validation was found to be from 77 to 91% (Table IV). Four out of five models revealed overall accuracy of prediction > 87% during cross-validation.

Investigations on the use of topochemical indices on a test set comprising structurally and chemically diverse molecules have led to successful validation of topochemical

Table IV. Evaluation of topochemical models using validation set and cross-validation set

1	Nature of	Number of compoin the range	Number of compounds in the range	Number of predicted	Number of compounds predicted correctly	Percent accuracy (%)	curacy (%)	Overall accuracy prediction (%)	Overall accuracy of prediction (%)
TIIGEY	the model	Validation set	Cross-vali- -dation set	Validation set	Cross-vali- -dation set	Validation set	Cross-vali- -dation set	Validation set	Cross-vali- -dation set
Į /M		18	22	14	20 NA	77.8	90.9	000	
٥ ١	Iransinonai Impermeable	2	0 1	1NA 2	1	100.0	100.0	0.00	C.17
	Permeable	18	22	14	20	77.8	6.06		
nW_{c}	nW _c Transitional	6	_	NA	NA	NA	NA	77.3	87.5
	Impermeable	4	7	8	1	75.0	50.0		
	Permeable	26	29	18	22	69.2	75.9		
ΧΑ	Transitional	5	1	NA	NA	NA	NA	69.2	7.6.7
	Impermeable	0	1	NA	1	NA	100.0		
	Permeable	18	22	14	20	77.8	6.06		
m,	Transitional	11	∞	NA	NA	NA	NA	80.0	91.3
	Impermeable	7	1	7	1	100.0	100.0		
	Permeable	15	20	12	19	80.0	95.0		
$n\xi_c^c$	$n\xi_c^c$ Transitional	10	∞	NA	NA	NA	NA	76.2	91.3
	Impermeable	9	3	4	2	2.99	2.99		

NA – Not applicable

models, which are highly beneficial for prediction of permeability through the blood-brain barrier. The overall accuracy of prediction of models for the validation set varied from a minimum of 70% for a model based on the molecular connectivity topochemical index to a maximum of 80% in case of models based upon Wiener's topochemical index and eccentric connectivity topochemical index. Surprisingly, these topochemical models also confirm the high prediction potential during cross-validation from a minimum of 77% for a model based on molecular connectivity topochemical index to a maximum of 91% in case of models based upon Wiener's topochemical index, eccentric connectivity topochemical index and normalized eccentric connectivity topochemical index. The results clearly reveal that the aforementioned topochemical models bear high predictability and can be utilized for permeability prediction of drugs and drug-like molecules.

CONCLUSIONS

Models derived from the topochemical indices can be used for fast screening of virtual libraries having millions of molecules and providing potent therapeutic agents with high permeability through the blood-brain barrier.

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SAŽETAK

Validacija topokemijskih modela za predviđanje permeabilnosti kroz krvno-moždanu barijeru

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U ovom su radu validirani i unakrsno validirani nedavno objavljeni topokemijski modeli za permeabilnost kroz krvno-moždanu barijeru. Predviđanje prolaska kroz krvno-moždanu barijeru strukturno i kemijski različitih molekula provedeno je na pet modela koji se temelje na tri topološka indeksa, Wienerovom topološkom indeksu, topološkom indeksu molekularne povezanosti i topološkom indeksu ekscentrične povezanosti. Ukupno 62 spoja podijeljena su u dva seta koji su sadržavali 31 spoj. Jedan set upotrebljen je za validaciju, a drugi za unakrsnu validaciju. Vrijednosti svih triju topoloških indeksa u početnom setu i u normaliziranom setu su računate pomoću kompjutorskog programa. Rezultati su analizirani i svakom spoju je pridružena teorijska vrijednost permeabilnosti, koja je zatim uspoređivana s objavljenim eksperimentalnim podacima za permeabilnost kroz krvno-moždanu barijeru. Točnost predviđanja bila je između 70 i 80%. Isti postupak je proveden za unakrsno validacijski set, a točnost je bila iznenađujuće velika (77–91%), što ukazuje da se upotrebljeni topokemijski modeli mogu upotrijebiti za predviđanje permeabilnost kroz krvno-moždanu barijeru.

Ključne riječi: topokemijski indeksi, Wienerov topološki indeks, topološki indeks molekularne povezanosti, topološki indeks ekscentrične povezanosti, permeabilnost, krvno-moždana barijera

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