Maximum Common Property: A New Approach for Molecular Similarity

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Abstract

The maximum common property similarity (MCPhd) method is presented using descriptors as a new approach to determine the similarity between two chemical compounds or molecular graphs. This method uses the concept of maximum common property arising from the concept of maximum common substructure and is based on the electrotopographic state index for atoms. A new algorithm to quantify the similarity values of chemical structures based on the presented maximum common property concept is also developed in this paper. To verify the validity of this approach, the similarity of a sample of compounds with antimalarial activity is calculated and compared with the results obtained by the small molecule subgraph detector (SMSD) method. The results obtained by the MCPhd method differ significantly from those obtained by the SMSD method, improving the quantification of the similarity. A major advantage of the proposed method is that it helps to understand the analogy or proximity between physicochemical properties of the molecular fragments or subgraphs compared with the biological response or biological activity. In this new approach, more than one property can be potentially used. The method can be considered a hybrid procedure because it combines descriptor and the fragment approaches.

Keywords: Maximum Common Property; Electrotopographic State Index; Molecular similarity; Tanimoto function; Maximum Common Structure

Introduction

Molecular similarity is one of the most explored and employed concepts in cheminformatics (chemical informatics or chemoinformatics) [1]. Moreover, it is currently one of the central subjects in medicinal chemistry research [1, 2]. Molecular similarity can be evaluated using different approaches, which can be classified into two principal categories: those based on descriptors and those based on substructures [3]. To estimate similarity among molecules, it is necessary to identify those structural or chemical/physical properties that are useful to correlate and then predict the relationships among them.

Similarity calculations based on molecular descriptors use fingerprint representations [3, 4]. These representations can be codified both by topological or topographic descriptors. Topological descriptors are the most popular because the 2D representation of molecules is computationally less difficult to work with than the 3D representation [1].

This work proposes a different approach in contrast with what is rigorously known as molecular similarity or chemical similarity [1]. The descriptor and the method of reduction of the graph used contain both structural and chemical-physical information. Thus, the approach allows evaluations and comparisons to be made by accounting for not only the structure but also other properties associated with the electrostatic nature of the molecule or fragment. The methods of structural similarity in 2D are more popular and simple. However, when working with only the topology of the molecules, most of the information associated with the spatial distribution is lost, except in the molecules that are essentially flat. As opposed to 2D methods, 3D methods consider that the properties of molecules tend to be strongly associated with the spatial distribution of their atoms [5, 6]. On the other hand, the configuration of minimum energy in a molecular structure with more than one degree of rotational freedom does not condition that this configuration is responsible for the biological response.
This issue causes a dilemma for researchers: losing all three-dimensional information for the sake of simplicity in the calculations or complicating the calculations and possibly delaying the results. The possibility of obtaining large data sets is an unquestionable reality. In that case, the eventual distortion of the 3D results due to not adjusting to the required conformation must be compensated by the increase in the number of compounds. However, such voluminous processing is not currently an impediment in terms of computational cost [7, 8].

Another concept that has been used for more than two decades is the scaffold and, more recently, scaffold hopping. These concepts allow the reduction of the molecule by eliminating R-substituents from the nucleus supposedly responsible for the activity in series of compounds in the first case, and in the other case, they allow the scaffold to be determined and enable comparisons to be made between structurally different compounds [9]. In other words, this approach bears a certain similarity to the proposed method since both seek to identify structurally different compounds that may show similar biological activity.

For these reasons, the proposed similarity method is based on the molecular description with a 3D descriptor that has structural information and on the polarity of the molecular graph or its fragments defined by a chemical graph reduction method.

Furthermore, molecular similarity based on substructure allows obtaining the molecular fragment or common subgraph among pairs of compounds [10, 11]. Several similarity methods have been developed based on a group of algorithms aimed at obtaining the largest common subgraph among a pair of compounds, the maximum common subgraph (MCS) [12, 13]. To quantify the molecular similarity, this method uses the Tanimoto coefficient \( T_{CMCS} \) [14, 15].

In this work, we introduce a new concept called maximum common property (MCPhd), inspired by MCS, to quantify the similarity based on substructure, using the electrotopographic state index for atoms \( S_{state3D} \) [17], which was developed from its parent electrotopology defined by Kier and Hall [16] from the connectivity matrix of the hydrogen-depleted chemical graph as an atomic descriptor.

The rest of the paper is organized in sections as follows: Materials and Methods describes the dataset and molecular codification, the general procedure and the proposed MCPhd algorithm; Results and Discussion describes the experimental results; and finally, Conclusions presents a summary of this work.

**Materials and Methods**

**Sample used**

We employed a set of 4-aminobicyclo[2.2.2]octan-2-yl 4-aminobutanoates (Table 1) reported by Weis et al. [18] and evaluated compounds against the multiresistant K-1 strain of Plasmodium falciparum.

**Codification of structures**

The electrotopographic state index for atoms [17] was used to codify chemical structures. This index is defined by Eq. (1).

\[
S_{state3D} = I_i + \Delta I_{ij}
\]

where \( S_{state3D} \) is the calculated value of the atom \( i \) in the corresponding molecule and \( I_i \) is the intrinsic value of the atom \( i \) calculated with Eq. (2).

\[
I_i = \left( \frac{2N_v}{\delta} \right) + 1 / \delta
\]

where \( N \) is the principal quantum number of atom \( I \), \( \delta \) is the number of valence electrons in the molecular skeleton \((Z'\text{-}h)\) and \( \delta \) is the number of \( \sigma \) electrons in the skeleton \((\sigma - h)\). For each atom of the molecular skeleton, \( \delta \) is the number of valence electrons, \( \sigma \) is the number of electrons in \( \sigma \) orbitals and \( h \) is the number of hydrogen atoms bonded.

\( \Delta I_{ij} \) represents the disturbance of the atoms of the environment, which is calculated by Eq. (3).

\[
\Delta I_{ij} = \sum (I_i + I_j) / r_{ij}^2
\]

where the sum is over the difference of the intrinsic values of atom \( i \) with respect to each one of the other atoms in the molecule and \( r_{ij}^2 \) is the Euclidean distance between the analyzed atoms.

**Graph reduction**

The reduction of the chemical graph is carried out by the method described by Carrasco et al. [19], where the descriptor centers (CDs), rings of different orders (Rn), clusters of order 3 and 4 (C3 and C4, respectively), heteroatoms such as halogens, amino, etc. (X), and terminal groups such as methyl (M3), methylene (M2) and methyne (M) are defined. Examples of these parameters are shown in Fig. 1. This graph reduction procedure, named CALEDE, is inspired by the procedure developed by Avindon et al. [20], where each CD is assigned the total value of \( S_{state3D} \), quantified as the sum of the value of \( S_{state3D} \) of each atom that conforms to it.
Definition of the Maximum Common Property
The maximum common property (MCPhd) between two fully connected and complete (not hydrogen-depleted) \( G_1 \) and \( G_2 \) chemical graphs is defined as the maximum similarity in the chemical-physical properties represented by the index \( S_{state3D} \), which exists between subgraphs \( g_1 \) and \( g_2 \) of the molecular graphs \( G_1 \) and \( G_2 \), respectively. Both \( g_1 \) and \( g_2 \) represent the link of at least two CDs that are at a Euclidean distance \( d(E(CD_1, CD_2)) \) from their corresponding centers of mass from pairs of CDs.

To quantify the value of similarity between two compounds using the concept of the maximum common property (MCPhd), the calculation of the similarity of two compounds is assumed using the Tanimoto function or coefficient on the basis of the maximum common substructure called \( T_{MCPhd} \) [14, 15]. The \( T_{MCPhd} \) for two molecules \( A \) and \( B \) is defined as:

\[
T_{MCPhd} = \frac{|MCPhd(A, B)|_b}{|A|_b + |B|_b + |MCPhd(A, B)|_b}
\]

(4)

where \( |A|_b \) is the number of links of \( A \), \( |B|_b \) is the number of links of \( B \) and \( |MCPhd(A, B)|_b \) is the number of links of the MCS of \( A \) and \( B \). If the concept MCPhd is replaced in Eq. (4), it yields:

\[
T_{MCPhd} = \frac{|MCPhd(A, B)|_b}{|A|_b + |B|_b + |MCPhd(A, B)|_b}
\]

(5)

where \( |A|_b \) is the number of heavy atoms of \( A \), \( |B|_b \) the number of heavy atoms of \( B \) and \( |MCPhd(A, B)|_b \) the smallest number of heavy atoms among the fragments with the highest MCP between \( A \) and \( B \).

The Proposed MCPhd Algorithm
Figure 2 shows the algorithm used for the calculation of similarity. The algorithm uses the following parameters: (\( G_1 \) and \( G_2 \)) two compounds or molecules, (i) the similarity threshold, (f) the similarity coefficient and (i) the index used to quantify the similarity. First, we obtain the subgraphs \( (f_1 \) and \( f_2 \)) that have a maximum common property value quantified by the index based on the parameters and similarity coefficient. These subgraphs are obtained by performing the following steps:

1. The index (i) entered as a parameter is calculated for each atom in each \( G_1 \) and \( G_2 \) graph using the Chemical Development Kit (CDK) library [22]. Lines 1 and 2 of the algorithm are shown in Fig. 2.

2. The graphs \( (G_1 \) and \( G_2 \)) on CDs are reduced, and the total index value of each one is obtained. Lines 3 and 4 of the algorithm are shown in Fig. 2.

3. The similarity matrix between the CDs obtained from the graphs \( (G_1 \) and \( G_2 \)) is constructed using the similarity coefficient introduced as a parameter, along with the distance matrix between the CDs of each graph \( (G_1 \) and \( G_2 \)) using the Euclidean distance. Line 5 of the algorithm is shown in Fig. 2.

4. The CDs from each graph \( (G_1 \) and \( G_2 \)) that meet the condition that the similarity value must be higher than the similarity threshold (\( u \)), entered as a parameter, are selected. Line 5 of the algorithm is shown in Fig. 2.

5. Finally, the CDs of each graph \( G_1 \) and \( G_2 \) are selected at a distance of 0.15, using the Canberra distance coefficient [23] and the distance matrices of the graphs. For each pair of CDs selected, a list is created where the pairs of CDs that are at a distance less than or equal to 0.15 are stored. Finally, the largest list is selected, and if there are several lists of the same size, one is selected, and all its CDs are repeated in other lists. Line 5 of the algorithm is shown in Fig. 2.

Then, with the subgraphs \( (f_1 \) and \( f_2 \)) and graphs \( (G_1 \) and \( G_2 \)) obtained, the values of the variables needed to quantify the similarity are obtained using the similarity coefficient (\( u \)) for discrete data entered as a parameter. Variable \( c \) is assigned the smallest number of heavy atoms belonging to the subgraphs \( (f_1 \) and \( f_2 \)), while variables \( a \) and \( b \) are assigned the number of heavy atoms belonging to each graph \( (G_1 \) and \( G_2 \)), respectively. Finally, these values are substituted in the similarity function to obtain the quantification of the similarity of the graphs \( (G_1 \) and \( G_2 \)). Lines 6 to 16 of the algorithm are shown in Fig. 2.

The use of the algorithm is exemplified below using the molecules 6k and 6c present in the dataset as shown in Fig. 3 and 4, respectively. We use 5 parameters \( (G_1, G_2, u, f, i) \) for its operation, where \( G_1 \) and \( G_2 \) are the molecular graphs 6k and 6c respectively, (i) is the index \( (S_{state3D}) \), (u) is the similarity threshold, and (f) is the similarity function. For this example, we will use 0.95 and the modified Tanimoto coefficient \( (T_{MCPhd}) \) as the threshold and similarity function, respectively. Then, after assigning the parameters, the following steps are performed:

1. The \( S_{state3D} \) index is calculated for each atom present in molecules 6k and 6c; these results are shown in Tables 2 and 3.
The 6k and 6c molecular graphs on CDs are reduced, and each is given the value of the total \( S_{state3D} \) index. As shown in step A of Fig. 5, molecule 6k is reduced on the CDs (R81, R52, R63, R64, R65, C36, X7), while molecule 6c is reduced on (R61, R82, R63, R64, R65, C36, M7).

The similarity matrix between the CDs of each molecule 6k and 6c is constructed using the Tanimoto coefficient (TC) for continuous data, together with the distance matrices between the CDs of each molecule (6k and 6c), as shown in step B of Fig. 5.

CDs are selected from each molecule (6k and 6c) that meet the condition that the similarity value is above the similarity threshold of 0.95. The CDs selected from molecules 6k and 6c are (R81, R52, R63, R64, R65 and C36) and (R61, R82, R63, R64, R65, C36 and M37), respectively, as shown in step C-a in Fig. 5. Furthermore, using the distance matrices of the graphs obtained in the previous step, for each pair of CDs, a list is constructed with the pairs of CDs that are at a Canberra distance less than or equal to 0.15, as shown in step C-b in Fig. 5.

From the lists of CD pairs obtained in the previous step, the following CDs are selected, namely, (R81, R52, R64 and C36) and (R61, R82, R63 and C34), corresponding to the lists (1, 3, 4 and 5) according to the larger size list with the same CDs in common.

Finally, the similarity value of the two molecules 6k and 6c is quantified using the modified Tanimoto coefficient \( T_{MCPhd} \), where the value of \( |MCPhd(A, B)|_b \) is the lowest number of heavy bonds present between fragments \( f_1 \) and \( f_2 \), while the values of \( |A|_b \) and \( |B|_b \) are obtained from the number of heavy atoms present in molecules 6k and 6c, respectively. With these values, it is possible to quantify the similarity between molecules 6k and 6c. In step E of Fig. 5, it can be seen that the number of heavy atoms of fragments \( f_1 \) and \( f_2 \) is 23 and 24, respectively, so the value of \( |MCPhd(A, B)|_b \) is 23, while the number of heavy atoms of molecules 6k and 6c is 36 and 37, respectively; that is, \( |A|_b = 36 \) and \( |B|_b = 37 \). Therefore, the calculated value of similarity between molecules 6k and 6c is 0.46.

Small Molecule Subgraph MCS approach
The Small Molecule Subgraph Detector (SMD) algorithm differs from previous MCS algorithms in that it uses a combination of several algorithms to find the common maximum subset and filters the results in a way that is chemically relevant because it incorporates chemical knowledge (coincidence of atom type with information sensitive and insensitive to the bond) while searching for molecular similarity. In addition, the algorithm calculates the maximum subgraph common between two molecules (A and B) by combining the power of the VFLibMCS, MCSP and CDKMCS algorithms. These algorithms are used on a case-by-case basis, depending on the molecules under consideration for the common maximum subgraph search [24]. This algorithm is implemented in the SMD tool available free of charge on the official site of the European Institute of Bioinformatics.

General Experimental Procedure
The experiments were carried out as shown in Fig. 6 based on a test of 36 compounds with a 2D structure, which have been tested experimentally in the study conducted by Weis et al. in 2014 [18]. The 3D structure of each compound was obtained through the Corina online service [25]. The 2D structures were used to calculate the molecular similarity (all against all) with the SMD algorithm, while the 3D structures were processed to calculate the \( S_{state3D} \) index for each atom and to reduce their graphs on CDs in order to apply the MCPhd algorithm to calculate the molecular similarity (all against all). Finally, the results obtained by both algorithms were compared through a screening process, which was evaluated by the percentage of success of finding structures with the same activity. The algorithms used were implemented using the JAVA language and were executed on an Intel(R) Core(TM) i7-7500U PC with 16 GB of RAM.

Results and Discussion
The molecular similarity methods used in this work, SMD and MCPhd, use different approaches to quantify the similarity between two molecular graphs or molecules. Whereas SMD uses graph isomorphism as a criterion, the similarity calculated with MCPhd is based on the criterion of analogy or proximity between the physicochemical properties of the molecular fragments or subgraphs that are compared, expressing these properties as an \( S_{state3D} \) value.

This approach places MCPhd closer to the concepts of bioisosterism than SMD. Bioisosterism means that two different molecules can provide similar biological responses if the structural aspects are phenomenologically accompanied by a physicochemical property associated with the biological response. This concept was coined by Friedman [26], extended by Burger [27] and recently used by Lassalas et al. [28] and Tahirova [29].

Using these two different approaches, different similarity values were obtained. For example, Table 4 shows
the results of the comparison with the remaining 35 molecules of the sample, with compounds 8c and 7j used as target elements since they had the minimum and maximum IC₅₀ values, respectively.

To determine whether both methods yielded significantly equal results, the nonparametric statistical test is used for two independent Mann-Whitney [30] samples with a significance level of 5%. The results of the Mann-Whitney U statistic were 243.00 and 98.00, with a value of p (bilateral asymptotic significance) of 0.00 and 0.00 for the most active compound (8c) and the least active compound (7j), respectively. It is then shown that the results obtained by both methods for both compounds were significantly different.

In addition, the results obtained by both methods had a low correlation; see Fig. 7 for the most active compound (8c) and Fig. 8 for the least active compound (7j).

To analyze the results from another perspective, a similarity function was developed for the dependent variable IC₅₀ using the Tanimoto coefficient (TcIC₅₀, column in Table 4), and subsequently, the molecular similarities calculated by both methods were correlated with this new variable. As shown in Fig. 9, the similarity results obtained by the MCPhd method for the most active compound (8c) had a slope closer to that obtained with the TcIC₅₀ variable; in addition, the results correlated better with a correlation coefficient value of Pearson r_xy = 0.84 against r_xy = 0.53, as shown in Figs. 10 and 11, respectively.

Analogous behavior occurred with the similarity results obtained with the less active compound (7j). The slope for MCPhd was nearest to TcIC₅₀ than that for SMSD (Fig. 12), and r_xy for MCPhd vs TcIC₅₀ was greater than SMSD vs TcIC₅₀, as seen in Fig. 13 and 14.

To generalize these results, the similarity obtained with both methods of the rest of the 17 compounds selected as active by Baptista [31] was correlated against TcIC₅₀. The results showed (Table 5) that compared to the SMSD method, the MCPhd method improves the correlation coefficient in 65% of cases.

To perform a more exhaustive study comparing the molecular similarity results obtained by both methods, the following steps were performed: (1) The similarity is calculated with both methods for all compounds (one against all); (2) the results up to or equal to the similarity thresholds (0.90, 0.80 and 0.70) are selected for each method; and (3) in each method, the threshold with the highest percentage of success in finding structures with the same activity is selected as the best threshold, and its results are compared.

As a result, a threshold of 0.90 was selected for the SMSD method because 65% of structures with the same activity (active-active and inactive-inactive) were found out of 116 pairs; a threshold of 0.70 was selected for the MCPhd method because the percentage of finding structures with the same activity is 67% of 92 pairs found. Tables 6 and 7 show the results for the SMSD and MCPhd methods, respectively, which validate the selection.

If we analyze the results obtained with the best similarity threshold in each method, it can be inferred that the percentage of finding structures with the same activity (active-active and inactive-inactive) obtained using the MCPhd method (67%) was better than the results with SMSD (65%) by 2%. Analyzing only the active-active pairs, the increase was 11% (34% for the SMSD method against 45% for MCPhd). These results suggested, once again, that the MCPhd method improved the similarity results obtained by the SMSD method.

As a last criterion, the 41 (MCPhd) and 39 (SMSD) pairs of compounds classified in the active-active category shown in Tables 6 and 7 were compared for both methods. To do so, relationship graphs (Figs. 15 and 16) were drawn for the 17 compounds present in the 41 and 39 active-active pairs for each method.

Figures 15 and 16 show different behaviors in the relationships between the 17 compounds when the SMSD method was used compared to using the MCPhd method. Thus, for example, whereas SMSD classified compounds by separating them by the different families of bicyclo-butanoates (6, 7 and 8 in Table 1), the MCPhd method put almost all of them in the same cluster. This result implies that the MCPhd method enabled establishing similarity relationships between compounds even from different families. The reason is that in addition to the structural information content provided by the electrotopographical state index for atoms, it includes electrostatic information content.

**Conclusions**

This work proposed a new approach that uses the 3D structure of molecules with physical-chemical information to estimate the molecular similarity between chemical compounds. The method has been favorably compared with the standard SMSD method and shows
better performance in obtaining structures with the same activity using similarity cutoff values during the screening process. Furthermore, the proposal shows the ability to find similar compounds among different families. This strongly suggest the possibility of applying the MCPhd method for isosteric studies.

Finally, the proposal presented in this paper provides a promising method for extending this method to be used in the construction of QSAR models for molecular activity prediction.

Declarations:
Acknowledgements
Not applicable.

Author’s contributions
Authors contributed equally to this work. All authors read and approved the final manuscript.

Funding
This work was supported in part by Project TIN2015-66108-P of the Spanish Ministry of Science and Innovation, by Project 1264182-F of the Andalusian Regional Government and by Project PP2019-Submod-1.2 of the Cordoba University.

Availability of data and materials
The data described in this article are available in Mendeley Data, doi:10.17632/corina.demo. Accessed 18 Feb 7

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Figures

Figure 1 Examples of Descriptor Centers (CD) employed in the fragmentation of the chemical graphs.

Algorithm: CDH(a, G, u, f, i)

1. input: Two graph G1 and G2, the similarity threshold (u), the coefficient of similarity (f) and index (i)
2. exit: Two similar fragments (f1, i) and quantification of similarity
3. - g1 = calculateTopographicIndices(G1)
4. - g2 = calculateTopographicIndices(G2)
5. - reducedGraph G1 = g1, getReducedGraph()
6. - reducedGraph G2 = g2, getReducedGraph()
7. - frag(f1, i, index) = getFragmentsAndCDH(g1, graphG1, u, f, i)
8. - if f1 != null and f1 != null then
9. - atom(f1) = getHeaetAtomsMCP(f1)
10. - atom(f2) = getHeaetAtomsMCP(f2)
11. - c = min(atom(f1), atom(f2))
12. - if c < (a+b-c) then
13. - frag(f1, f2, index) = index
14. - return frag(f1, f2, index)
15. - else
16. - return frag(f1, f2, o)

Figure 2 algorithm of the algorithm to calculate similarity.
Figure 5 Example of applying the MCPhd algorithm to the 6k and 6c molecules belonging to the dataset.

Figure 6 General Experimental procedure.
Figure 7 Correlation between MCPhd and SMSD similarity among compound 8c and the others.

Figure 8 Correlation between MCPhd and SMSD similarity among compound 7j and the others.
Figure 9 Comparison of molecular similarity among compound 8c and the others.

Figure 10 Correlation between MCPhd and TcIC<sub>50</sub> similarity among compound 8c and the others.

Figure 11 Correlation between SMSD and TcIC<sub>50</sub> similarity among compound 8c and the others.
Figure 12 Comparison of molecular similarity among compound 7j and the others.

Figure 13 Correlation between MCPhd and $TcIC_{50}$ similarity among compound 7j and the others.

Figure 14 Correlation between SMSD and $TcIC_{50}$ similarity among compound 7j and the others.

Figure 15 Relationship graph of active compounds obtained by the MCPhd method with a molecular similarity greater than 0.70.

Figure 16 Relationship graph of active compounds obtained by the SMSD method with a molecular similarity greater than 0.90.
Tables
### Table 1: Compounds set*

| a  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.35 |
|----|--------------------------|--------------------------|----------------------|
|    | H₃C⁻N⁻CH₃               | N⁻N⁻                    | 6a  IC<sub>50</sub> = 0.35 |
| b  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.46 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7a  IC<sub>50</sub> = 0.46 |
| c  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.24 |
| d  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.71 |
| e  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.106 |
| f  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.06 |
| g  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.55 |
| h  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.19 |
| i  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.70 |
| j  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.70 |
| k  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.37 |
| l  | NR<sup>1</sup>R<sup>2</sup> | NR<sup>3</sup>R<sup>4</sup> | IC<sub>50</sub> = 0.37 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6c  IC<sub>50</sub> = 0.54 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7b  IC<sub>50</sub> = 0.26 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8b  IC<sub>50</sub> = 0.12 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6e  IC<sub>50</sub> = 0.71 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7c  IC<sub>50</sub> = 0.06 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8c  IC<sub>50</sub> = 0.19 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6d  IC<sub>50</sub> = 0.106 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7d  IC<sub>50</sub> = 0.55 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8d  IC<sub>50</sub> = 0.19 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6f  IC<sub>50</sub> = 0.47 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7f  IC<sub>50</sub> = 0.25 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8f  IC<sub>50</sub> = 0.09 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6g  IC<sub>50</sub> = 0.52 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7g  IC<sub>50</sub> = 0.40 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8g  IC<sub>50</sub> = 0.35 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6h  IC<sub>50</sub> = 0.37 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7h  IC<sub>50</sub> = 0.29 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8h  IC<sub>50</sub> = 0.20 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6i  IC<sub>50</sub> = 0.18 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7i  IC<sub>50</sub> = 0.26 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8i  IC<sub>50</sub> = 0.18 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6j  IC<sub>50</sub> = 2.16 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7j  IC<sub>50</sub> = 2.25 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8j  IC<sub>50</sub> = 1.43 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6k  IC<sub>50</sub> = 0.86 |
|    | N⁻N⁻                    | N⁻N⁻                    | 7k  IC<sub>50</sub> = 1.91 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8k  IC<sub>50</sub> = 0.51 |
|    | N⁻N⁻                    | N⁻N⁻                    | 6l  IC<sub>50</sub> = nd |
|    | N⁻N⁻                    | N⁻N⁻                    | 7l  IC<sub>50</sub> = 0.17 |
|    | N⁻N⁻                    | N⁻N⁻                    | 8l  IC<sub>50</sub> = 0.09 |

*Taken from reference Weiss*
Table 2 Result of the Sstate3D calculation for each atom of the 6k molecule.

| Molecule 6k | Atom Number | Sstate3D | Atom Number | Sstate3D |
|-------------|-------------|----------|-------------|----------|
| C           | 1           | 0.27653  | N           | 19       | -1.4592  |
| C           | 2           | 4.24474  | C           | 20       | 3.38006  |
| C           | 3           | -2.11427 | C           | 21       | 3.15142  |
| C           | 4           | 4.25231  | C           | 22       | -0.12941 |
| C           | 5           | 0.49396  | C           | 23       | 3.09432  |
| C           | 6           | 0.52371  | C           | 24       | 3.32881  |
| C           | 7           | 0.28273  | O           | 25       | 4.22234  |
| C           | 8           | 4.2063   | N           | 26       | -1.30847 |
| C           | 9           | -0.3793  | C           | 27       | 3.2697   |
| C           | 10          | 3.51969  | C           | 28       | 2.9831   |
| C           | 11          | 2.98731  | C           | 29       | 3.01061  |
| C           | 12          | 2.89649  | C           | 30       | 3.54754  |
| C           | 13          | 2.99502  | C           | 31       | -0.38433 |
| C           | 14          | 3.5348   | C           | 32       | 3.54001  |
| C           | 15          | -0.08644 | C           | 33       | 2.99825  |
| C           | 16          | -0.10545 | C           | 34       | 2.90069  |
| O           | 17          | 5.5644   | C           | 35       | 2.99544  |
| C           | 18          | 3.75612  | C           | 36       | 3.53921  |

Table 3 Result of the Sstate3D calculation for each atom of the 6c molecule.

| Molecule 6c | Atom Number | Sstate3D | Atom Number | Sstate3D |
|-------------|-------------|----------|-------------|----------|
| C           | 1           | 0.3202   | C           | 20       | 3.314    |
| C           | 2           | 4.17197  | C           | 21       | 2.96595  |
| C           | 3           | -2.19195 | N           | 22       | -2.22688 |
| C           | 4           | 4.25211  | C           | 23       | 2.94306  |
| C           | 5           | 0.37013  | C           | 24       | 3.25012  |
| C           | 6           | 0.50352  | C           | 25       | 8.9881   |
| C           | 7           | 0.21735  | N           | 26       | -1.46255 |
| C           | 8           | 4.23211  | C           | 27       | 3.52712  |
| C           | 9           | -0.24053 | C           | 28       | 2.94719  |
| C           | 10          | 3.50889  | C           | 29       | 2.85191  |
| C           | 11          | 3.00512  | C           | 30       | 2.95634  |
| C           | 12          | 2.94959  | C           | 31       | 3.54763  |
| C           | 13          | 3.09722  | C           | 32       | -0.13708 |
| C           | 14          | 3.69435  | C           | 33       | 3.3945   |
| O           | 15          | -0.48635 | C           | 34       | 2.9161   |
| O           | 16          | -0.31876 | C           | 35       | 2.85145  |
| O           | 17          | 5.6053   | C           | 36       | 2.96557  |
| N           | 18          | 3.73854  | C           | 37       | 3.52722  |

| Atom Number | Sstate3D | Atom Number | Sstate3D |
|-------------|----------|-------------|----------|
| N           | 19       | -1.66555    |          |

Table 4 Molecular similarity values of the more active and inactive compounds with the rest of the sample.

| Molecule 6c | Molecule 7j | Target | IC50 | Molecule 8l | Molecule 9c | Molecule 10j |
|-------------|-------------|--------|------|-------------|-------------|-------------|
| SMSD        | MCPhd      | TcIC50 |      | SMSD        | MCPhd      | TcIC50      |
| 1.00        | 1.00       | 8c     | 0.05 | 0.61        | 0.42       | 0.02        |
| 0.97        | 0.93       | 87c    | 0.87 | 0.32        | 0.59       | 0.03        |
| 0.90        | 0.73       | 8f    | 0.73 | 0.99        | 0.51       | 0.04        |
| 0.65        | 0.55       | 3c    | 0.106| 0.60        | 0.33       | 0.05        |
| 0.37        | 0.31       | 65c   | 0.14 | 0.12        | 0.26       | 0.08        |
| 0.70        | 0.67       | 86c   | 0.18 | 0.61        | 0.25       | 0.09        |
| 0.93        | 0.75       | 87c   | 0.18 | 0.61        | 0.49       | 0.09        |
| 0.87        | 0.40       | 35c   | 0.19 | 0.86        | 0.60       | 0.09        |
| 0.92        | 0.67       | 33c   | 0.19 | 0.65        | 0.34       | 0.09        |
| 0.90        | 0.52       | 85c   | 0.20 | 0.64        | 0.33       | 0.09        |
| 0.92        | 0.44       | 38c   | 0.24 | 0.80        | 0.46       | 0.12        |

Table 5 Correlation results between both methods for the active compounds in the sample.

| Molecule | IC50 | txy | b | c |
|----------|------|----|---|---|
| 8c       | 0.50 | 0.48| 0.52| 0.80|
| 7c       | 0.06 | 0.40| 0.07| 0.72|
| 8f       | 0.09 | 0.58| 0.57| 0.04|
| 8l       | 0.09 | 0.56| 0.51| 0.60|
| 6c       | 0.106| 0.53| -0.13| 0.10|
| 8b       | 0.12 | 0.69| 0.47| 0.53|
| 7l       | 0.17 | 0.62| 0.07| 0.17|
| 6l       | 0.18 | 0.60| -0.19| 0.14|
| 8l       | 0.18 | 0.55| 0.37| 0.33|
| 8d       | 0.19 | 0.74| 0.25| 0.02|
| 8e       | 0.19 | 0.73| 0.51| 0.30|
| 8h       | 0.20 | 0.76| 0.26| 0.35|
| 8a       | 0.24 | 0.65| 0.12| 0.08|
| 7f       | 0.25 | 0.52| 0.15| 0.17|
| 7b       | 0.26 | 0.41| 0.15| 0.18|
| 7r       | 0.26 | 0.51| 0.14| 0.09|
| 7r       | 0.28 | 0.46| 0.02| -0.13|

a*: SMSD vs MCPhd, b*: SMSD vs TcIC50, c*: MCPhd vs TcIC50.
Table 6 Comparison of the observed and predicted by SMSD for several similarity thresholds.

| Threshold | Real-Predicted | Pairs* | % | Predicted | Pairs* | % |
|-----------|----------------|--------|---|-----------|--------|---|
| 0.90      | Active-Active  | 39     | 14| Correct   | 75     | 65|
|           | Inactive-Inactive | 36 | 11 | Incorrect | 41     | 35|
|           | Active-Inactive | 19     | 17| Incorrect | 41     | 35|
|           | Inactive-Active | 19     | 17| Incorrect | 41     | 35|
|           | Total          | 116    | 100 | Total     | 116    | 100|
| 0.80      | Active-Active  | 52     | 28| Correct   | 106    | 57|
|           | Inactive-Inactive | 54 | 29 | Incorrect | 81     | 43|
|           | Active-Inactive | 48     | 26| Incorrect | 81     | 43|
|           | Inactive-Active | 33     | 18| Incorrect | 81     | 43|
|           | Total          | 187    | 100 | Total     | 187    | 100|
| 0.70      | Active-Active  | 63     | 20| Correct   | 121    | 57|
|           | Inactive-Inactive | 22 | 27 | Incorrect | 91     | 43|
|           | Active-Inactive | 46     | 23| Incorrect | 91     | 43|
|           | Inactive-Active | 43     | 20| Incorrect | 91     | 43|
|           | Total          | 212    | 100 | Total     | 212    | 100|

*Pairs*-Number of predicted pairs

Table 7 Comparison of the observed and predicted by MCPhd for several similarity thresholds.

| Threshold | Real-Predicted | Pairs* | % | Predicted | Pairs* | % |
|-----------|----------------|--------|---|-----------|--------|---|
| 0.90      | Active-Active  | 14     | 33| Correct   | 26     | 62|
|           | Inactive-Inactive | 12 | 29 | Incorrect | 16     | 38|
|           | Active-Inactive | 8      | 19| Incorrect | 16     | 38|
|           | Inactive-Active | 8      | 19| Incorrect | 16     | 38|
|           | Total          | 42     | 100 | Total     | 42     | 100|
| 0.80      | Active-Active  | 16     | 36| Correct   | 28     | 62|
|           | Inactive-Inactive | 12 | 27 | Incorrect | 17     | 38|
|           | Active-Inactive | 9      | 20| Incorrect | 17     | 38|
|           | Inactive-Active | 9      | 20| Incorrect | 17     | 38|
|           | Total          | 45     | 100 | Total     | 45     | 100|
| 0.70      | Active-Active  | 41     | 45| Correct   | 62     | 67|
|           | Inactive-Inactive | 23 | 35 | Incorrect | 30     | 33|
|           | Active-Inactive | 15     | 16| Incorrect | 30     | 33|
|           | Inactive-Active | 15     | 16| Incorrect | 30     | 33|
|           | Total          | 92     | 100 | Total     | 92     | 100|

*Pairs*-Number of predicted pairs