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Molecular Interaction Fields based descriptors to interpret and compare chromatographic indexes

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Abstract

Molecular Interaction Fields (MIFs) based descriptors can be conveniently used to characterize and compare chromatographic scales. In this study, Quantitative Structure–Retention Relationships (QSRR) for eight different chromatographic systems were obtained with VolSurf+ descriptors and Partial Least Squares (PLS). A new and purpose-designed analysis tool highlights the different balance of intermolecular interactions governing solute retention, and estimates the similarity between chromatographic systems.
1. Introduction

Chromatographic indexes determined by HPLC have recently gained considerable relevance in Absorption, Distribution, Metabolism and Excretion (ADME) studies, because they are faster and easier to obtain than log P (P is the partition coefficient in octanol/water system), and can provide significant information about the compounds’ lipophilicity, with or without conversion to log P [1-4]. The large variety of stationary phases and mobile-phase additives that are now available provides the flexibility enabling chromatography to be used to adjust the properties of the partitioning phases [5], so as to directly model biological partition rather than mimicking it. However, to model biological partition with chromatographic data, it must be verified that the balance of intermolecular forces governing the two systems is the same.

In more general terms, the deconvolution of the intermolecular interactions governing chromatographic retention is a topic of great interest in many analytical fields. Modeling tools are routinely used for this purpose [2, 6]. The linear solvation energy relationship (LSER) is currently the most widely-used approach to understanding the types and relative strengths of the chemical interactions controlling retention and selectivity. One of the most generally accepted symbolic representations of the LSER model is given in Eq. 1 [7]

\[ \log k = c + eE + sS + aA + bB + vV \]  

Eq. 1

The model consists of product terms representing solute properties (descriptors), indicated by capital letters, and the complementary system properties, indicated by lower-case letters. Each product term describes the relative contribution of specific intermolecular interactions to the correlated solute property, in this case, \( \log k \) (k is the retention factor). The contribution from electron lone pair interactions is expressed by \( eE \), interactions of a dipole type by \( sS \), hydrogen-bond interactions by \( aA \) and \( bB \), and differences in cavity formation and dispersion interactions for transfer of the solute from one phase to the other, by \( vV \). The solute descriptors (often called solvatochromic descriptors) are
formally defined as excess molar refraction, $E$; dipolarity/polarizability, $S$; effective hydrogen-bond acidity, $A$; effective hydrogen-bond basicity, $B$; and McGowan’s characteristic volume, $V$.

Solvatochromic descriptors are empirical in character, and thus only available for a limited number of compounds. In general, solvatochromic descriptors are lacking for multiple functional and for ionised compounds [2, 8-10].

For this reason it appeared valuable to replace solvatochromic descriptors with more convenient descriptors of a fully computational nature, based solely on the 3D structural formula of compounds, i.e. VolSurf+ descriptors. These are based on 3D molecular fields [11, 12] and so could lead to a rational explanation of retention differences in terms of intermolecular interactions. Briefly, a 3D molecular field may be viewed as a 3D matrix, with attractive and repulsive energy values between a chemical probe and a target molecule; VolSurf+ is a tool that extracts molecular descriptors from the 3D matrix.

Figure 1 shows the visual representation of some VolSurf+ descriptors calculated for amitrole (example of target molecule present in the dataset of the investigated compounds).

Insert Figure 1.

Figure 1A shows the envelope corresponding to three VolSurf+ descriptors (W3, W5 and W7) obtained with the water probe. They represent the volume of the molecular envelope which is accessible to, and interacts attractively with water molecules at three levels of interaction energy (-1, -3 and -5kcal/mol).

Since at different energy values the contribution of the intermolecular (and intramolecular-) interactions to the single descriptor varies, VolSurf+ descriptors are better interpreted using blocks of descriptors obtained with a single probe.

VolSurf descriptors can therefore be classified in six blocks (see Supporting Information): a) descriptors that characterize the size and shape of the solute (7 descriptors in the text, briefly called size, color-code green), b) 19 descriptors that express the solute’s interaction with water molecules (in the text indicated as water, color-code light blue), c) 5 descriptors that describe the solute’s ability to form hydrogen bond
interactions with the donor group of the probe (that mimics the chromatographic system, color-code blue, see below), d) 5 descriptors expressing the solute’s ability to form hydrogen bond interactions with the acceptor group of the probe (that mimics the chromatographic system, color-code red, see below) e) 28 descriptors describing the solute’s propensity of the solute to participate in hydrophobic interactions (in the text called DRY for short, color-code yellow), f) 18 descriptors mainly describing the imbalance between hydrophilic and hydrophobic regions (in the text called others, color-code grey). Please note that for the sake of clarity, to identify hydrogen bonding (HB) interactions (Hydrogen Bond Acceptor capability (HBA)/Hydrogen Bond Donor capability (HBD) and color code) we refer to the probe’s properties and not to the solute. This is shown in Figure 1C.

Some years ago, VolSurf descriptors and PLS were used to calculate and interpret log P in alkane/water system [13]. In 2011 the same strategy was applied to model log k30 (k30 is the isocratic retention factor) data for a series of Pt(II) derivatives [14]. It is here shown that MIFs-based descriptors can conveniently be used to qualitatively characterize any chromatographic index on the basis of the intermolecular interactions governing the system. For this purpose, QSRR models based on PLS algorithms were firstly made. The study’s end-point was to develop graphical and numerical tools at different levels of complexity that enable researchers to extract as much chemical information as possible from a series of chromatographic systems, and numerically rank their similarity/dissimilarity.

2. Methods

Frog 2.1 (http://bioserv.rpbs.univ-paris-diderot.fr/cgi-bin/Frog2) was used to build the 3D structures that were then submitted to AM1 minimization as implemented in Spartan ‘08 (Wavefunction Inc, Irvine, CA). For any compound, the lowest energy conformation as obtained from Spartan minimization was selected as the final structure, saved in mol2 format and submitted to VolSurf+ (version 1.0.4, Molecular Discovery Ltd. Pinner, Middlesex, UK, 2009, http://www.moldiscovery.com) using default
settings and four probes (OH2, DRY N1 and O probes that mimic respectively water, hydrophobic, hydrogen bond acceptor and hydrogen bond donor interaction of the compounds with the environment). PCA and PLS tools implemented in VolSurf+ were used.

3. Results and discussion

3.1. The chromatographic systems

The dataset used was that developed by Tuelp et al. [9], containing 76 structurally-diverse and complex organic structures, chiefly pesticides together with some drugs. The full list of chemical structures is in the Supporting Information of the original paper [9].

The capacity factors of the 76 compounds were determined in eight HPLC systems [9]. The specifications of the HPLC systems are in Table 1; further details are in the original paper [9]. Each system is identified by an abbreviation (Table 1), used throughout the text. Six different columns were investigated and different composition of the mobile phases were adopted. Log k values were available for four reversed phase (RP), three normal phase (NP) and one hydrophilic interaction (Hydrophilic Interaction Liquid Chromatography, HILIC) chromatographic systems (Table S1 in the Supporting Information shows the descriptive statistics of log k). Taken together retention characteristics were highly diverse as confirmed by the correlation matrix (Table S2 in the Supporting Information) that evidences a large variability in cross correlations (Pearson coefficients).

3.2. Building PLS models

To relate log k values with the chemical structures of compounds mathematically, a PLS study was performed as will be described.

Firstly, the 3D structures of the 76 molecules in the dataset were built (see Experimental Part) and, for each structure, 82 descriptors (the full list of descriptors is available in Table S3 of the Supplementary Information) were calculated by VolSurf+.
Log k values for the eight chromatographic systems [9] were then imported into VolSurf+ as dependent variables (Y) and a relation between Y and VolSurf+ descriptors (X) was sought using the standard PLS tool implemented in the software. The statistics of the best models are shown in Table 1.

To validate these models internal validation was firstly used. Whereas some researchers in the QSAR field support internal validation, others consider that internal validation is not a sufficient test to check the robustness of models, and external validation is necessary [15, 16]. In this case, however, since the sample size is small, and thus holding a portion of it back for testing would be wasteful, it was preferred to use cross-validation, with multiple rounds using different partitions (Table 1 shows $Q^2$ for the LOO procedure, but results were similar with different partitions). Most of the PLS models in Table 1 (apart from the CN-NP system, which is a limit situation) showed $R^2 > 0.6$ and $Q^2 > 0.5$, and satisfied the Tropsha et al. validation rules [17]. Finally, to further validate PLS models we randomized the order of Y values which produced unacceptable $R^2$ and $Q^2$ values (data not shown).

The relatively low $R^2$ values might be due to the complexity and molecular flexibility of the dataset, whereas the small difference between $R^2$ and $Q^2$ supports the stability of the models. Some outliers were identified for 6 out of 8 systems. An in-depth analysis of the reasons for the presence of outliers (most of them probably related to the flexibility of compounds) is outside the scope of this study, but it is worth noting that the inclusion/exclusion of outliers slightly altered PLS statistics, but did not alter the chemical interpretation of the PLS models (see below).

### 3.3. Mechanistic interpretation of PLS models

The main drawback of PLS models is the difficulty of interpreting them from the chemical standpoint. Variable Importance in Projection (VIP) plots are often used to overcome this limit [13]. The VIP represents the value of each predictor in fitting the PLS model; the VIP values in the plot summarise the overall contribution of each X-variable to the PLS model, summed over all components and weighted...
according to the Y variation accounted for by each component (Eq. 2). The sum of squared VIP values is equal to the number of descriptors [15].

\[
VIP_j = \left( \frac{ \sum_{f=1}^{F} w_{jf}^2 \cdot SSY_f \cdot J}{SSY_{total} \cdot F} \right)^{1/2}
\]

\textit{Eq 2}

Where \( w_{jf} \) is the weight value for variable \( j \) component \( f \), \( SSY_f \) is the sum of squares of explained variance for the \( f \)th component and \( J \) the number of variables. \( SSY_{total} \) is the total sum of squares explained of the dependent variable, and \( F \) is the total number of components. The weights in a PLS model reflect the covariance between the independent and dependent variables and the inclusion of the weights is what allows VIP to reflect not only how well the dependent variable is described but also how important that information is for the model of the independent variables. Note that \( \sum_{j=1}^{F} VIP_j^2 = J \).

As explained in the Introduction, the best interpretation of VolSurf+ descriptors is achieved when they are grouped into six blocks. Therefore, to obtain a fully optimized interpretation of the PLS models (Table 1) BlockRelevance (BR) was defined as the ratio of the sum of the squared VIP values of a given block of descriptors and the number of those descriptors (Eq. 3).

\[
BR_i = \left( \sum_{j=1}^{N_i} VIP_j^2 \right) / N_i
\]

\textit{Eq 3}

where \( i \) is the number of blocks (6 here, Table 2), \( N \) is the number of descriptors for any block, VIP is the value of each predictor fitting the PLS model.

BR shows the relevance of a certain block of descriptors to a given chromatographic system; the higher the value of BR, the more important is that block. The set of BR values can therefore be used as a first set of numbers able to characterize a chromatographic system.
Table 2 lists the BR values for the 8 systems investigated. For example, C18 is mainly characterized by the size block of descriptors, \((BR = 2.35)\) which is at least twice as important as any other block. A similar trend is found for Ph. The CN-RP system is almost equally characterized by HBD (Fig. 1C), size and water descriptors; the situation is similar for OH-RP. HILIC is mainly characterized by size and HBD. The contribution of water and size descriptors is most significant for OH-NP, whereas HBD properties are the mainly determinant of the NH2-NP system.

From BR analysis, it is known, for example, that a given block of solutes descriptors characterizes a system (e.g. size for C18, Table 2) but not if the corresponding intermolecular interactions promote the retention either in the stationary or in the mobile phase. The sign of the PLS coefficient is informative: a positive coefficient means that an increase of the descriptor considered causes an increase in log \(k\) (and thus the retention in stationary phase increases); the reverse is true for negative coefficients. Depending on the sign of the PLS coefficient, BR was broken down into BR (+) relating to retention in the stationary phase, and BR (-), relating to retention in the mobile phase (Eq. 4).

\[ BR_i = BR_i (+) + BR_i (-) \quad \text{Eq. 4} \]

BR (+) and BR (-) for the eight investigated systems are illustrated in Fig. 2; this is a significant tool to compare chromatographic systems.

Figure 2 clearly distinguishes RP from NP systems, as shown by the different trends of the two blocks of VolSurf+ descriptors expressing size and interaction with the water probe. In RP systems, the size
block contributes significantly to increasing interaction with the stationary phase, whereas the same block is significant and of opposite sign in NP systems. Interestingly, for HILIC the contribution of the size block is very important, but of negative sign, distinguishing this system from both RP and NP, as might be expected [18]. The block of water descriptors is of negative sign in RP systems, but positive for NP and HILIC systems. This confirms that hydrophilic compounds prefer the mobile phase in RP systems, whereas they have greater affinity for the stationary phase in NP and HILIC systems.

The blocks size, HBA and (although in minor extent) HBD enable the four RP systems to be classified. The contribution of the size descriptors varies in the following order: C18-Ph>CN-RP>OH-RP. The HBD trend is similar in the four systems, but the reverse is true for HBA. Ph and OH-RP show opposite trends in the HBA block (red in Figure 2, negative for Ph and positive for OH-RP). This confirms that HBA properties are mainly expressed in the stationary phase of the OH-RP system, and in the mobile phase of Ph system.

The three NP systems investigated share similar trends for size, water and dry blocks. The most significant differences are again found for HB-related descriptors. The HBA block (red in Fig. 2) is large and positive for NH$_2$-NP, about half the size and negative for CN-NP, with OH-NP in an intermediate situation. The HBD block shows the same trend for the three systems, being particularly important for CN-RP. To sum up, in qualitative terms NH$_2$-NP is different from CN-NP and from OH-NP, which are much closer to one another (see the quantitative analysis below).

As expected the HILIC system shows a unique profile. As mentioned above, the block of size descriptors is the most important whereas the block of HBA is slightly less so. These findings show that two polar compounds with similar size can be separated by the HILIC system investigated (Table 1) on the basis of their HBD properties. This supports experimental evidence about the application of HILIC systems containing high amounts of acetonitrile in the mobile phase to the separation of amines and peptides. [18]. It should be stressed that application of this analysis method to different HILIC systems could, significantly, improve a thorough understanding of retention behavior of HILIC, thus enhancing their applicability domain.
OH-RP and OH-NP are a pair of chromatographic systems characterized by the same column but different mobile phase composition (polar for the RP system and apolar for the NP system; see Table 1). As Figure 2 shows, in the presence of the same stationary phase, different mobile phases can completely alter the balance of intermolecular forces responsible for retention.

OH-NP and NH$_2$-NP are a pair of systems that use the same mobile phase (Table 1) but different stationary phases. Figure 2 highlights the different performances of the two columns in terms of intermolecular interactions, and suggests that this analysis could be particularly important when new columns are launched on the market.

Finally, a further requirement is a numerical value with which to quickly rank the diversity of chromatographic systems. For this purpose, a simple match parameter called LipophilicityMatch (LM) (Eq. 5), was defined

\[ LM_{i,j} = \sum_{f=1}^{82} f(c_i^f ; c_j^f) \]  
\[ \text{Eq. 5} \]

where

\[ f(c_i^f ; c_j^f) = 1 \text{ for } c_i^f \cdot c_j^f \geq 0 \]

\[ = 0 \text{ for } c_i^f \cdot c_j^f < 0 \]

and c is the coefficient and I and j are a pair of chromatographic systems

LM values for the eight systems investigated are reported in Table 3.

As expected, the RP and NP systems have little similarity (not above 33%). Among the RP systems investigated, LM values range from 85% to 76%. Among the NP systems, LM values range from 66 to 77% confirming the qualitative results discussed above. The LM analysis also confirms that the HILIC system is more similar to NP (about 70%) than to RP (about 28%) systems.
3.4. How to individuate the chromatographic system that best mimics the octanol/water system

A way of comparing log $P_{oct}$ with log $k$ values could facilitate the implementation of relevant log $k$ descriptors in running research programs in the pharmaceutical industry. This comparison can easily be determined by LM as described below. For the above mentioned dataset of compounds a good PLS model ($R^2 = 0.82$, $Q^2 = 0.70$, $LV = 3$) was obtained using log $P_{oct}$ (data by Tuelp et al. [9]) as dependent variable and VolSurf+ descriptors as independent variables. Figure 3A shows the trend of BR (+) and BR (-) for the octanol/water system. The size block is large and positive. The remaining blocks are less important. LM were calculated and shown in Figure 3B. As expected, the highest values were found for RP systems. Given the LM definition (Eq. 5), a minute comment of the numerical differences should be avoided.

Insert Figure 3.

4. Conclusions

The QSRR analysis performed in this study demonstrates the goodness of the VolSurf+ descriptors, for describing the mechanism of chromatographic retention at the molecular level. The PLS models based on VolSurf+ descriptors were interpreted via custom-designed parameters (BR and LM) to fully exploit the interpretative potential of the VolSurf+ descriptors in quali- and quantitative terms. The potential applications of this kind of analysis are numerous. In drug discovery programs, the reliability of lipophilicity indexes determined on immobilized biomacromolecule stationary phases could be verified, and biomimetic chromatographic partition systems may prove to be better models than log $P$ for use in ADME studies, or otherwise. From a more general point of view, manufacturers, for example, could compare new and old products, definitively managing end-capping effects and identifying columns with equivalent properties. In proteomic studies, metabolites could be identified and/or classified by their different behavior on chromatographic systems, revealing their differing balance of intermolecular interactions.
Acknowledgement

GE and GC thank Dr. Fenner for providing additional experimental details to published results

References


Figure captions

Figure 1. Amitrole (a molecule present in the dataset): visual representation of some VolSurf+ descriptors obtained with different probes. A) water (descriptors: W3, W5 and W7), B) DRY (probe with hydrophobic properties, descriptors: D1, D4 and D8) and C) HB probes: N1 with hydrogen bonding donor properties (in the text called HBD for short, color code: blue) forms HB interactions with solute’s hydrogen bonding acceptor (HBA) properties and O with HBA properties (in the text called HBA for short, color code: red) forms HB interactions with solute’s HBD properties.

A

Water probe

W3 = 272.9 (-1 kcal/mol)  
W5 = 65.5 (-3 kcal/mol)  
W7 = 8.3 (-5 kcal/mol)

B

DRY probe

D1 = 63.0 (-0.2 kcal/mol)  
D4 = 17.8 (-0.8 kcal/mol)  
D8 = 4.5 (-1.6 kcal/mol)

C

Solute HBA  
Probe HBD (N1)

Solute HBD  
Probe HBA (O)
Figure 2. A graphical tool to compare chromatographic systems: BR (+) and BR (-) for the eight investigated systems are reported. Details about the six blocks of descriptors (size, water, DRY, HBA, HBD, others) are in the text.
Figure 3. A) BR (+) and BR (-) for the octanol/water system (data by Tuelp et al. [9], PLS model: $R^2 = 0.82$, $Q^2 = 0.70$, LV = 3, see text for details) and B) LM values to numerically compare octanol/water and the eight chromatographic systems.
Table 1. Final PLS models (n = number of observations, $R^2$ = cumulative determination coefficient, $Q^2$ = cross-validated correlation coefficient, LV = number of latent variables, RMSE = root mean square of the errors)

<table>
<thead>
<tr>
<th>System</th>
<th>Abbreviation</th>
<th>Column</th>
<th>Mobile phase</th>
<th>n</th>
<th>$R^2$</th>
<th>$Q^2$</th>
<th>LV</th>
<th>RMSE</th>
<th>Missing data</th>
<th>Excluded</th>
</tr>
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<td>C18</td>
<td>YMC-Pack-Pro C18 RS</td>
<td>MeOH/Water 75/25</td>
<td>72</td>
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<td>0.264</td>
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<td>MeOH/Water 64/36</td>
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<td>0.70</td>
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<td>NH$_2$-NP</td>
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Table 2. BlockRelevance (BR) for the eight investigated systems. In parentheses the number of descriptors for the block.

<table>
<thead>
<tr>
<th>System</th>
<th>Size (7)</th>
<th>Water (19)</th>
<th>DRY (28)</th>
<th>HBA (5)</th>
<th>HBD (5)</th>
<th>Others (18)</th>
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<td>1.37</td>
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Table 3. LipophilicityMatch (LM) values for the eight systems investigated

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<tr>
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