

COSMO sim : Bioisosteric similarity based on COSMO-RS σ -profiles

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ABSTRACT. A novel approach for the quantification of drug similarity is proposed which makes use of the surface polarities, i.e. conductor surface polarization charge densities σ , as defined in the quantum chemically based Conductor-like Screening Model for Realistic Solvation (COSMO-RS). The histogram of these surface polarities, the so-called σ -profiles have been proven to be the key for the calculation of all kinds of partition and adsorption coefficients, and therefore of relevant ADME parameters as solubility, pK_a , $\log BB$, and many others. They also carry a large part of the information required for the estimation of desolvation and binding processes responsible for receptor binding and enzyme inhibition of drug molecules. Thus, a large degree of similarity with respect to the σ -profiles appears to be a necessary condition for drugs of similar physiological action. Driven by this insight, we propose a σ -profile based drug similarity measure SMS for the detection of new bioisosteric drug candidates. In several examples we demonstrate its statistical and pharmaceutical plausibility, its practicability for real drug research projects, and its unique independence from the chemical structure, which enables scaffold hopping in a natural way.

INTRODUCTION. Bioisosteric transformation is one of the most frequently used approaches to the design and optimization of compounds with biopharmacological importance. Both, scaffold hopping and group interchange converge seamlessly in this approach. Current computational approaches to bioisosteric transformation are so far *a posteriori* classifiers that deduct rules, which are then applied in a project-dependent manner. We describe the development and application of an *a priori* method for the prediction of bioisosters. We assume that bioisosters must have similar physico-chemical properties that rule their interactions with different environments like solvents, membranes, and ultimately protein receptors. These interactions define their biological effects and biopharmacological properties. We built our method, therefore, upon COSMO-RS, a general and fast methodology for the *a priori* prediction of thermophysical data. Cheap unimolecular quantum chemical calculations combined with exact statistical thermodynamics provide the information necessary for the evaluation of molecular interactions. Thus, we can represent molecules detached from their chemical structures as electronic surfaces. For the rapid comparison of COSMO surfaces, they are projected into 1D σ -profiles. Suitable similarity metrics are then developed based on σ -profiles. Empirical data of many thousand bioisosteric transformations is used to validate the method and to further optimize the free parameters of the similarity metrics. Finally, the method is applied to bioisosteric transformations of functional groups and for virtual HTS.

A number of biological targets of pharmaceutical interest are currently beyond the scopes of experimental 3D structure determination. Examples for such targets are membrane-standing proteins like ion channels and G-protein coupled receptors. The design of potent modulators for such targets is though not a hopeless task. Ligand-based design of compounds makes use of the structural information of well-characterized compounds that is used at different levels of abstraction for subsequent intermolecular similarity calculations¹⁻³. Ideally, selection of target ligands should follow the concepts of ligand efficiency⁴⁻⁶. Two scenarios of ligand-based design strategies can be envisaged. Firstly, ligands with improved potency and/or improved biophysical properties ultimately related to their ADME-Tox profiles are searched within the class of ligands described by the scaffold. Secondly, alternative structural classes are explored for backup or patent busting purposes, a concept termed scaffold hopping. Both approaches are intimately related to the concept of bioisosterism, i.e., the introduction of structural

changes into a given active compound leading to a derivative that maintains broadly the bioactivity at the given receptor in question following the well-known active analogue principle.

The search for novel and better bioisosters has been the challenge for medicinal chemistry over the past decades. On the one hand a lot of experience has been gained via trial-and-error. A survey of successful bioisosteric transformations is accessible in form of the BioSter database⁷. This is a well-suited dataset for method development in the bioisosterism field^{8,9}. *A posteriori* analyses of successful bioisosteric transformations led to the common classical and non-classical categorizations of bioisosters^{10,11}. On the other hand, some *a priori* concepts tried to rationalize bioisosterism based on physical concepts like Grimm's hydride displacement law^{12,13}. Newer non-classical definitions focus on certain aspects of the compound structures like molecular shape, topology, pharmacophoric patterns, and electronic isosters¹⁴. However, none of the currently available bioisosteric approaches is built on a solid physicochemical basis.

Ultimately, a suitable method must provide an intermolecular similarity measure that ranks proven bioisosteric pairs of compounds high while random pairs rank low. It can then be used for picking tentative bioisosters for a target ligand from compound databases. Furthermore, intermolecular similarity is used as objective function for the design and optimization of target-focused compound libraries.

MATERIALS AND METHODS

COSMO-RS. While almost all computational chemistry methods used in drug design are based either on the force-field concept or on decomposition concepts as group-contributions or fingerprints, a rather orthogonal and fundamental approach has been developed over the past 15 years which is based on quantum chemistry combined with dielectric continuum solvation and statistical thermodynamics. Being developed in an industrial computational chemistry lab, this Conductor-like Screening Model for Realistic Solvation (COSMO-RS)¹⁵⁻¹⁷ was originally considered for the quantification of environmental and technical partition behavior, before it was recognized a very generally applicable and most predictive model for fluid phase thermodynamics in the chemical engineering community. In the light of its applicability in wide ranges of chemistry its applicability to biophysical properties became apparent, and it has been proven in several papers on solubility, physiological partition properties as logBB or intestinal absorption, and pK_a during the past years¹⁸⁻²¹, and its extension towards the problem of ligand receptor binding appears attractive and is presently being exploited. While detailed descriptions of the COSMO-RS theory have been given elsewhere, the basic concept of the methods is outlined in the following.

- 1) In a first step all chemical compounds of a liquid (or pseudo-liquid) ensemble are considered as embedded and swimming in an infinite, perfect conductor. This state of the compounds can be very well treated by quantum chemical calculations combined with dielectric continuum solvation models, the most efficient and natural choice for the latter being the Conductor-like Screening Model COSMO²². For the quantum chemistry, density functional theory has been proven to provide a good compromise of computational efficiency and reliability, because the much faster semi-empirical methods suffer from severe deficiencies in the solvation electrostatics, especially if hypervalent elements as sulfur or phosphorous come into play. By such combination of methods the self-consistent state of a molecule within such virtual

conductor, i.e. its energy, its geometry and electron distribution, and the surface polarization charges of the conductor on the molecular surface, can be calculated at almost the same costs as in the gas-phase²³.

- 2) This state of molecules in a conductor has proven to be a very useful reference for the understanding of molecular behavior in the liquid. Much better suited than the traditional reference state of an isolated molecule in vacuum. The conductor surface polarization charge density σ is a very good local measure of molecular surface polarity, carrying more information than the often considered electrostatic potential (ESP). Examples of COSMO surfaces color coded by σ are given for water, caffeine, and theophylline in figure 1. The regions of strongly negative molecular polarity are displayed in red. It is important to recognize that the negative molecular regions carry positive polarization charge density σ , because σ is just compensating the molecular electrostatic field and has, thus, opposite sign. The strongly positive molecular regions carrying negative σ are colored blue, while the neutral parts of the molecules with σ close to zero appear green.

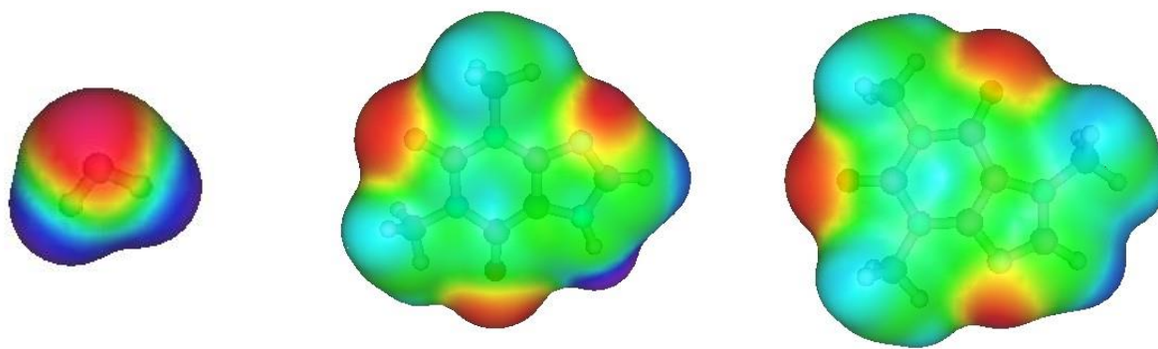


Figure 1. COSMO surfaces color coded by the polarization charge density σ for water, caffeine, and theophylline

- 3) A big conceptual advantage of the conductor reference state is the fact that the molecules together with their polarization charges now are electrostatically perfectly non-interacting, because no electric field can escape through the solute - conductor interface. As long as we leave at least a thin film of conductor between the molecules, we can build any geometrical configuration of the molecules without changing the energy of the system. By allowing for small, volume conserving and energetically irrelevant deformations of the surfaces we can finally build densely packed, liquid like systems of the molecules as schematically shown in figure 2, but still assuming an infinitely thin film of conductor separating and screening the molecules from each other. If we suppose that the small deformations required for the close packing do not significantly influence the energy and the polarization charge densities, our ensemble now still has the same energy as in the dilute state in the conductor, but each surface segment μ with polarization σ_μ of a molecule has a nearest neighbor segment ν with σ_ν , i.e. we only have direct face to face segment pairs.

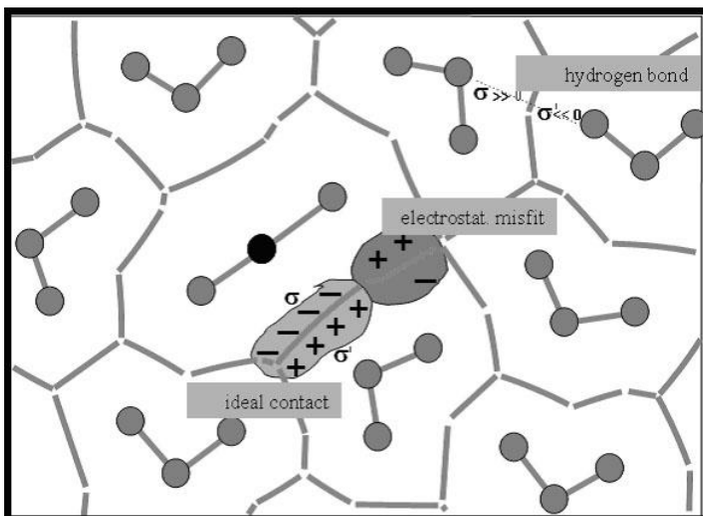


Figure 2. Schematic visualization of COSMO-RS contacts and interactions on the molecular cavity

- 4) Since in nature there is no conductor between the molecules, in the next step the thin film of conductor between the surface segments has to be removed. As shown in the COSMO-RS papers in more detail, we can do this segment pair by segment pair. In this way we can consider the energy difference going along with the removal of the conductor between a segment pair (μ, n) as a local surface interaction of the neighboring molecules, and we can quantify its electrostatic and hydrogen bond contributions per unit surface area as

$$e_{misfit}(\sigma, \sigma') \cong \frac{\alpha'}{2}(\sigma + \sigma')^2 \quad (1)$$

and

$$e_{hb}(\sigma, \sigma') \cong c_{hb}(T) \min[0, \sigma\sigma' + \sigma_{hb}^2] \quad (2)$$

where σ and σ' are the polarization charge densities of the interacting surfaces. While the electrostatic misfit part can be quite accurately derived from theoretical arguments, the hydrogen bond energy term must be considered as an empirical but physically plausible expression. Given the very few parameters in these formulae, the interaction energy of a fixed configuration of our ensemble of molecules relative to its conductor reference state can, thus, be calculated as an integral of all local pair-wise surface interactions.

- 4.) For a liquid system we have to calculate the thermodynamic averages of the relevant configurations of our ensemble. Normally this requires the generation of large numbers of such ensembles, as done in MC or MD simulations. But due to the local pair-wise surface interactions description, the thermodynamics can be reduced to an ensemble of independently interacting surface segments. For this purpose we just need the surface polarization charge density distribution $p^X(\sigma)$, the so-called σ -profile, of each compound X which tells us how much surface of which polarity σ is available on the surface of compound X. Since these σ -profiles are of central importance for our new similarity approach, a collection of σ -profiles is shown in figure 3. These σ -profiles turned out to be very useful fingerprints of molecules. Next we denote the normalized σ -profile of a solvent S by $p_S(\sigma)$. For a pure solvent $p_S(\sigma)$ is just the normalized σ -profile of the solute molecule, and for a mixture it is trivially generated from the

mole fraction weighted σ -profiles of the components. Based on $p_S(\sigma)$ the statistical thermodynamics of the interacting surface pairs can be solved efficiently and exactly from the integral equation

$$\mu_S(\sigma) = -kT \ln \left\{ \int p_S(\sigma') \exp \left[-\frac{a_{\text{eff}} e_{\text{int}}(\sigma, \sigma') - \mu_S(\sigma')}{kT} \right] d\sigma' \right\} \quad (3)$$

where $\mu_S(\sigma)$ is the chemical potential of an additional surface segment in the ensemble S, and a_{eff} is the size of an effective contact segment. The function $\mu_S(\sigma)$, called σ -potential further on, has to be derived recursively from eq. 3 due to its appearance on both sides of the equation. The σ -potential describes how much a solvent S likes additional surface of polarity σ . As explained in more detail elsewhere these σ -potentials express a wide range of important aspects of the interaction capabilities of the solvent, including electrostatics, hydrogen bonding, and hydrophobicity. Finally the chemical potential of a compound X in solvent S is calculated from the σ -profile of the solute and the σ -potential of the solvent as

$$\mu_S^X = \int \mu_S(\sigma) p^X(\sigma) d\sigma + \mu_{S,\text{comb}}^X \quad (4)$$

where $\mu_{S,\text{comb}}^X$ is a usually small correction for size effects of solute and solvent, thus, a combinatorial contribution, depending on volumes and surface areas of solute and solvent. Hence the important part of the chemical potential of a solute X in a solvent S is expressed as a surface integral of the solvent σ -potential over the surface of the solute X, since $p^X(\sigma)d\sigma$ just is the amount of surface area of polarity σ on the surface of X.

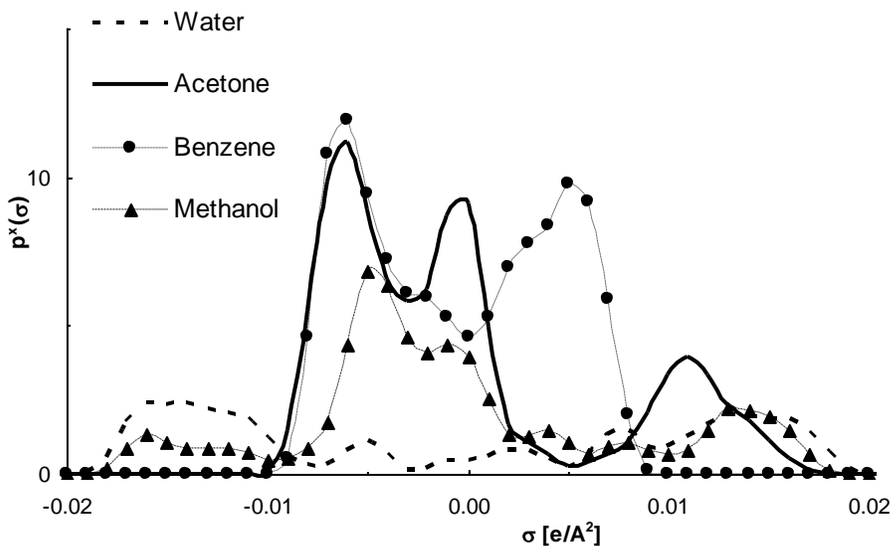


Figure 3. Solvent σ -Profiles. These profiles show the amount of molecular surface in a given interval of polarization charge density σ .

In this way COSMO-RS gives access to the chemical potentials of almost arbitrary compounds in almost arbitrary liquid phases using the σ -profile of the solute as the only information. While chemically

well defined phases as water, octanol, alkane, etc. and the corresponding partition coefficients are directly accessible by the COSMO-RS theory, partitioning between physiological phase as blood and brain and other ADME properties can be derived from the σ -profiles with a slightly more empirical extension, the σ -moment approach. Finally it has been shown that even drug solubility can be calculated from the σ -profiles, using the chemical potential μ_X^X of the drug in its virtual liquid state as the most important input.

Since solubility and ADME properties can be well described from the σ -profile alone, we can expect that two compounds with similar σ -profiles should have similar ADME characteristics. But we must be aware that the binding of a drug to a receptor involves constraints regarding to the 3D-relations of the various polar, hydrogen bonding, and hydrophobic interaction sites. This information is not included in the σ -profile any more. Nevertheless, it is at least a necessary while not sufficient requirement for strongly binding ligands of a receptor that they have roughly the same amount of surface area available for the various interaction modes. Thus, it is at least plausible that a drug candidate with a similar σ -profile as a known strongly binding ligand has a good chance to be strongly binding as well. Summarizing the above considerations, we consider it as a plausible assumption that similarity of σ -profiles should be a powerful measure for the assessment of drug similarity.

APPLICATION OF COSMO-RS TO PROTEIN-LIGAND INTERACTIONS. COSMO-RS provides access to chemical potentials (Gibbs free energies) of almost arbitrary compounds in almost arbitrary phases. This grants direct access to important chemo-physical and bio-physical properties like phase partition (e.g. n-octanol water partition as expressed in $\log P$) and thermodynamic solubility in different solvents (e.g. intrinsic aqueous molar solubility $\log S$). COSMO-RS provides furthermore a sound basis for the computation of chemical potentials in mixed phases based on statistical thermodynamics (COSMO $_{therm}$). This allows the treatment of more complex compartment partitions including extracellular matrix / cellular membranes, intestinal lumen / blood, blood / brain.

Non-covalent reversible protein-ligand interactions can well be described by the thermodynamic cycle given in Figure 4 for ligand A binding to receptor P forming the ligand-receptor complex AP²⁴. The complex formation is determined by the binding free energy $\Delta G_{\text{binding}}$ under thermodynamic equilibrium conditions.

$$\Delta G_{\text{binding}} = \Delta G_{\text{cosmo}} - \Delta G_{A,\text{aq}} - \Delta G_{P,\text{aq}} + \Delta G_{AP,\text{aq}} \quad (5)$$

Binding free energy $\Delta G_{\text{binding}}$ and complex dissociation constant K_D can be interconverted following

$$\Delta G_{\text{binding}} = -RT \ln K_D \quad (6)$$

where R is the gas constant and T is the temperature.

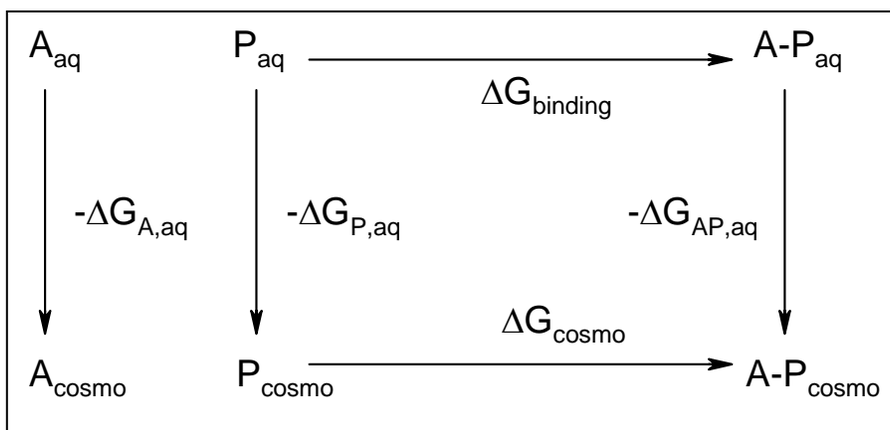


Figure 4. Thermodynamic cycle for the equilibrium binding free energy $\Delta G_{\text{binding}}$ of ligand A binding to receptor P forming the non-covalent complex AP.

The rigorous statistical thermodynamic treatment of ligand, receptor, and ligand-receptor complex as homogeneous pseudo-liquid phases represents the only approximation. Obviously, at the atomic level, protein-ligand interactions can be hampered by steric hindrance due to the anisotropic character of the protein although electronic complementarity in the COSMO concept is given and would consistently lead to energetically favorable interaction.

DATA SETS. A virtual screening database of >3.5 Mio unique compounds was compiled from various in-house, medchem, and vendor databases. *COSMOfrag*²⁵ was used to calculate approximate σ -profiles for all compounds. This database is dynamically maintained and updated with new screening compounds. The search for bioisosters of groups and entire molecules was performed with this database.

The BioSter database⁷ contains data on successful bioisosteric transformations for various receptors and chemical classes. This dataset was previously used for the evaluation of methods to distinguish between bioisosteric pairs and random pairs of molecules^{8,9}. These molecular pairs were extracted from the database and approximate σ -profiles were calculated using *COSMOfrag*. Obvious prodrugs, ambiguous pairs, and molecules, for which no σ -profiles could be calculated, like ions, were removed from the set. A list of random pairs was prepared from the same set of molecules by scrambling the right column of the pair table. The final sets contained 6041 bioisosteric pairs and 5823 random pairs. All similarity and energy calculations were run on these two sets.

BIOPHYSICAL CONCEPT OF BIOISOSTERIC TRANSFORMATIONS. Medicinal chemistry uses bioisosteric transformation as a tool for the replacement of certain functional groups with alternatives that display higher potency, better specificity, improved safety and other pharmacological properties or for the design of novel bioactive compounds via scaffold hopping for backup and patent busting purposes like therapeutic copies of commercialized drugs²⁶. Common concepts deduce rules from datasets that describe previously successful bioisosteric transformations and these rules are then applied to prioritize the synthesis and characterization of analogues²⁷. Our approach attempts to provide a biophysical fundament for bioisosteric transformations and will allow as such the *a priori* prediction of bioisosters.

In ligand-based drug design approach the receptor is assumed to be constant while the ligand is the variable. The bioisosteric transformation of ligand A to ligand B, both binding to receptor P, can be

formulated as a thermodynamic cycle (see Figure 5). For ligand-based drug design purposes, A can be regarded as the target-ligand while B is the result of the bioisosteric transformation applied. This approach allows the calculation of the energetic cost of the bioisosteric transformation $\Delta\Delta G_{AP,BP, \text{binding}}$.

$$\Delta\Delta G_{AP,BP, \text{binding}} = \Delta\Delta G_{AP,BP, \text{cosmo}} - \Delta\Delta G_{A,B, \text{aq}} - \Delta\Delta G_{P, \text{aq}} + \Delta\Delta G_{AP,BP, \text{aq}} \quad (7)$$

All terms of this thermodynamic cycle are directly accessible via COSMO-RS and COSMOtherm under the principle assumption of interacting pseudo-liquid molecules. The protein desolvation term $\Delta\Delta G_{P, \text{aq}}$ cancels out. For compounds A and B with identical σ -profiles, the mixed phase desolvation term $\Delta\Delta G_{AP,BP, \text{aq}}$ becomes zero, as well as the protein-ligand interaction terms $\Delta\Delta G_{AP,BP, \text{cosmo}}$ do.

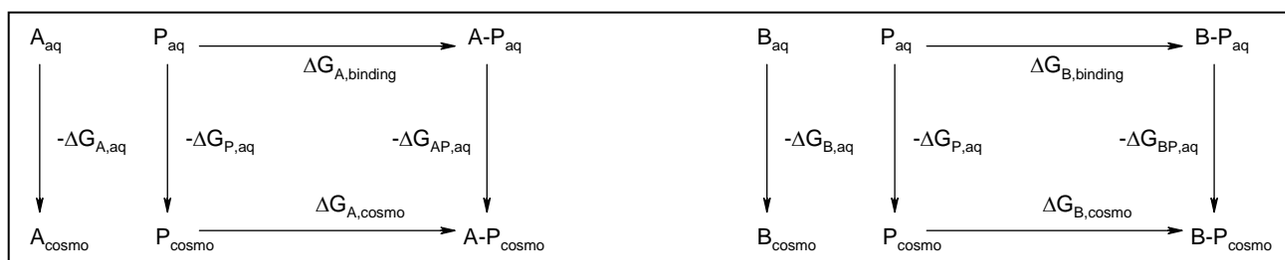


Figure 5. Thermodynamic cycle for the bioisosteric transformation of target ligand A to B both targeting receptor P.

The success of a bioisosteric transformation can, thus, be quantified prior synthesis by computing $\Delta\Delta G_{AP,BP, \text{binding}}$. For practical purposes, successful bioisosteric transformations should not exceed 1-3 kcal/mol (about two log orders of magnitude in the binding constant K_D). The final goal is to gain access to the estimated energetic cost of a bioisosteric transformation to be used to rank different transformations in virtual screening.

COSMOtherm defines the energetics of molecular interactions on the basis of pairwise σ interactions, thus, the unimolecular σ -profiles of the ligands in question define ultimately the energetic contributions. Consistently, the more similar the σ -profiles of A and B become, the more the ligand-dependent contributing terms cancel out. The next chapter focuses on different intermolecular similarity methods based on σ -profiles.

SIMILARITY COEFFICIENTS. COSMOtherm represents the σ -profiles by 61 real values for the relevant σ -range from $\sigma = -3 \text{ e/nm}^2$ to $\sigma = 3 \text{ e/nm}^2$. We initially started from this 61 bin representation which represents a one-dimensional structure-free holographic electronic profile. We term σ -profile based similarity methods COSMOsim.

A large set of coefficients is available^{1,28} to calculate intermolecular distances based on binary, integral, or floating point vectors containing chemical information. One of the widely used coefficients is the Tanimoto coefficient, which has been found useful for distance measures of binary and integral vectors encoding molecular descriptors. In order to compare σ -profiles we started with examining the

suitability of the Tanimoto coefficient. Usually, the binary variant of the Tanimoto coefficient is used. However, extension to non-negative floating point values is straightforward:

$$T_c = \frac{\sum_{i=1}^l 1 - \frac{|N_{A,i} - N_{B,i}|}{N_{A,i} + N_{B,i}}}{l} \quad (8)$$

The Tanimoto coefficient T_c is the intermolecular similarity, where l is the number of bins, $N_{A,i}$ and $N_{B,i}$ are the surface areas corresponding to bin i in molecule A and B, respectively.

Equation 8 reveals that the usability of the Tanimoto coefficient suffers when many bins are non-zero and few bins differ largely. In such cases very high similarities are computed that neglect substantial differences. In order to illustrate these consequences, a congeneric set of n-alcohols (n-propanol through n-hexadecanol) was prepared, the corresponding σ -profiles were computed for all compounds in the set (Figure 6), and T_c was calculated using n-propanol as target ligand. Despite the very different molecular sizes of n-propanol and n-hexadecanol high, physically implausible, T_c values result (Figure 7).

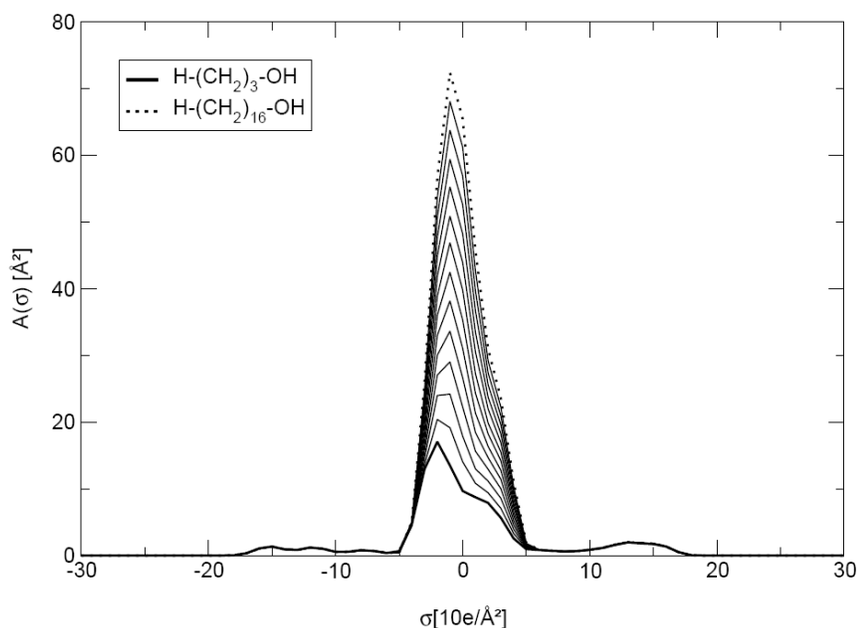


Figure 6. σ -profiles of a congeneric series of n-alcohols.

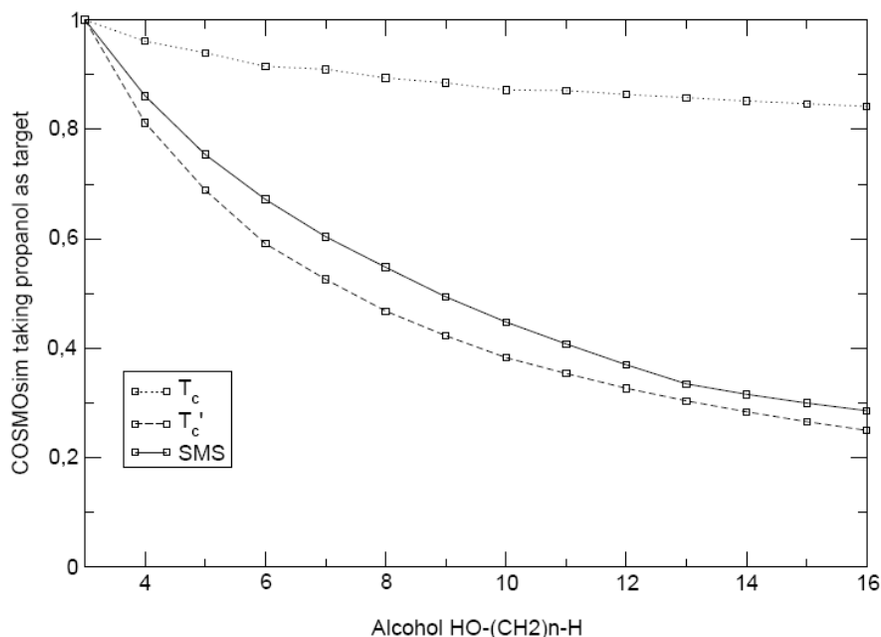


Figure 7. σ -profile based similarities of a congeneric series of n-alcohols compared to n-propanol using different COSMOsim coefficients.

A straightforward solution to this problem appeared to be the utilization of the relative molecular sizes. The molecular size is here defined as the molecular surface area (COSMO surface) that equals the sum over all bins of the σ -profile. We call the novel similarity measure Tanimoto prime coefficient, T_c' :

$$T_c' = T_c \frac{\min(S_A, S_B)}{\max(S_A, S_B)} \quad (9)$$

Where T_c is the Tanimoto coefficient, S_A and S_B are the COSMO surface areas of compounds A and B, respectively.

The bin-based Tanimoto similarity of σ -profiles suffers from a few theoretical weaknesses. Firstly, its values quite strongly depend on the fineness of the discretization of the σ -range, while a theoretically robust similarity definition should be almost independent of the discretization level. Secondly, since only the ratio of deviation and sum of the two σ -values of a bin enters the definition of bin similarity, bins with small values of the similarity may have the same influence on the Tanimoto similarity coefficient as bins with large values, while under physical aspects bins with small values should be less important. Finally, the bin-based similarity definition totally disregards the physical neighborhood relation of σ -bins. If one compound has a high intensity in bin i while the other has a high intensity in bin $i+1$, the Tanimoto coefficient considers this as dissimilarity, while such slightly shifted peaks would still cause rather similar physicochemical behavior.

Based on these considerations we defined another similarity measure that we call σ -match similarity (SMS). The basic idea behind it is the matching of most similar surface area pairs, starting with the most polar segments present in either of the two σ -profiles and proceeding to the least polar ones. In the first

step the most polar piece of surface available in the two σ -profiles is matched with the same area of the most polar surface segment of the same polarity sign of the other compound. Now the minimum area a_{seg} of the two matched surface segments is subtracted from both σ -profiles and a contribution $dSMS$ as given in eq. 10 is added to the raw similarity measure SMS_0 :

$$dSMS(a_{seg}, \sigma, \sigma') = a_{seg} \left(1 + \frac{c}{4} (\sigma + \sigma')^2 \right) \exp \left\{ \frac{(\sigma - \sigma')^2 \left(1 + \frac{b}{4} (\sigma + \sigma')^2 \right)}{a^2} \right\} \quad (10)$$

where σ and σ' are the two σ -values matched in this step. In order to better understand this expression it is useful to consider it first for the two parameters b and c set to zero. In this case the formula apparently gives a contribution identical to the matched area, if the two matched σ -values are identical. Otherwise the contribution to the raw similarity index is reduced by a Gaussian function. Hence the similarity strongly decreases with increasing mismatch of the two σ -values. The parameter a is a measure for the σ -mismatch tolerance. Repeating the described procedure until no surface area is left in one of the σ -profiles, we finally get a raw similarity coefficient SMS_0 . At the end of the procedure we will have a residual surface area a_{res} left in the bigger of the two compounds. The maximum value of SMS_0 can be the value of the smaller of the two components. This can only be achieved if the two σ -profiles are identical. By the two parameters b and c we can introduce two different ways to increase the sensitivity of the σ -similarity measure in the polar σ regions, which might be useful considering the fact that especially the hydrogen bond interactions contribute very strongly in the polar regions and make these more important for drug similarity than the less polar regions. A positive value of b decreases the σ -tolerance in the polar regions, while a positive value of c increases the weight of surface area of polar segments compared with less polar segments.

If we calculate the maximum achievable values of the raw similarity of the two compounds, i.e. their raw self-similarities, and denote them by SMS_1 and SMS_2 , respectively, then we can define our final expression for the σ -match similarity coefficient SMS by

$$SMS = \frac{SMS_0 + da_{res}}{\sqrt{SMS_1 + SMS_2}} \quad (11)$$

where d is a parameter responsible for the treatment of the residual surface area a_{res} : For $d = 0$ a_{res} is considered as maximal dissimilar, for $d = 1$ it is treated as maximal similar.

The presented definition of the σ -match similarity fulfills all requirements of a similarity metrics, i.e. it is unity if applied to identical compounds, asymptotically takes a value of 0 for very dissimilar compounds, and it is commutative with respect to the compounds. Furthermore, it is rather independent of the bin discretization of the σ -range, reasonable weights its contributions according to the surface areas, and introduces a mismatch tolerance with respect to σ . Reasonable values for the four parameters a , b , c , and d of the SMS definition will be presented in the next chapter.

SMS PARAMETER OPTIMISATION. A genetic algorithm was used in order to optimize the four free variables a , b , c , and d in the SMS similarity calculation using empirical data of the Bioster database. The target function for the GA-optimization was the maximization of the separation of random

versus bioisosteric pairs. Therefore, the corresponding similarity values were split into 51 bins covering the SMS similarity range from 0 to 1. The overlapping gray zone g was defined as follows and the GA was used to maximize $1-g$.

$$g = \sum_{i=0}^{50} \min\left(\frac{n_{bioisoster,i}}{m_{bioisoster}}, \frac{n_{random,i}}{m_{random}}\right) \quad (12)$$

200 generations of GA optimization with 20 unique individual parameter sets per generation were performed starting with random values for a ($1.0 \leq a \leq 5.0$), b ($0 \leq b \leq 0.01$), and c ($0 \leq c \leq 0.01$). For each individual parameter set, the intermolecular similarity values were recalculated for the bioisoster and random data sets, and the individual g values were calculated thereupon. The individual solutions were then sorted by g , and a next generation of individual parameter sets was created by applying crossover and mutation to the binary genomic representations of the better-than-average solutions.

In a first approach d was held fixed to 0 while in a second independent run d was also optimized within ($0 \leq d \leq 1$). In order to check for the possible effects of the four parameters, the GA was re-run but this time to maximize g , thus, leading to the worst model possible. The resulting parameters along with the parameter-free metrics T_c and T_c' are shown in Table 1. The optimized parameter sets lead to practically indistinguishable separation values. To limit the number of free parameters, the three parameters model with d set to zero was used throughout the article for the COSMOsim SMS calculations.

Table 1. Overview on SMS parameters and statistics on separation of bioisosteric and random molecular pairs.

model	a^1	b^2	c^3	d^4	g^5	bioisosters avg SMS \pm stdev	random avg SMS \pm stdev
worst	1.096	0.009430	0.001310	0.993	0.745	0.570 \pm 0.155	0.471 \pm 0.142
best 3 parameters	2.533	0.000350	0.009960	fixed 0	0.424	0.697 \pm 0.191	0.382 \pm 0.207
best 4 parameters	2.561	0.000124	0.009990	0.130	0.424	0.713 \pm 0.180	0.416 \pm 0.197
T_c	-	-	-	-	0.406	0.726 \pm 0.126	0.528 \pm 0.112
T_c'	-	-	-	-	0.395	0.636 \pm 0.165	0.369 \pm 0.145

¹ σ -tolerance, ² σ -tolerance of polar surface, ³ weighting of polar surface, ⁴ size tolerance, ⁵ gray zone overlap

ENERGETIC COST OF BIOISOSTERIC TRANSFORMATION. A bioisosteric transformation of A to B can hardly be performed in practice without affecting the electronic nature of the compound. Such changes will have a more or less drastic effect on the various energetic contributions to the relative binding free energies (eq. 5). For practical purposes, a bioisosteric transformation can be regarded as

successful if B does not lose more than around one to two log orders of magnitude in binding affinity. A factor of 10 to 100 in K_D corresponds to a change in binding free energy by 1.3 to 2.7 kcal/mol at room temperature.

It is conceptually straightforward to suppose that the change in binding free energies upon bioisosteric transformation becomes *small* as the intermolecular similarity approaches unity. The underlying distance metric, however, can have substantial influence on the meaning of *small*. In order to give a qualitative estimation on the energetic cost of bioisosteric transformation $\Delta\Delta G_{A,B, \text{binding}}$ in different similarity metrics concepts one would ideally compute all terms of the thermodynamic cycle given in Figure 5. This is currently not possible with reasonable efforts.

In order to get still a rough estimation for the energetic cost, the desolvation term $\Delta\Delta G_{A,B, \text{aq}}$ can be rapidly computed with reasonable accuracy (~ 0.5 kcal/mol) as reported previously¹⁹. The difference in free energy of ligand desolvation provides an estimate for the energetic cost of the bioisosteric transformation for one of the three terms of Equation 8. ΔG_{aq} was calculated for every molecule in the bioisoster and random data sets using COSMOtherm²⁹. Subsequently, $\Delta\Delta G_{A,B, \text{aq}}$ was calculated for all pairs.

For a qualitative assessment of the energetic cost the intermolecular similarity values were plotted against the $\Delta\Delta G_{A,B, \text{aq}}$. Funnel-shaped curves are obtained for each similarity coefficient (see Figures 8 - 10), which shows that the solvation free energy differences do become *small* as the intermolecular similarity approaches unity. As expected, the funnel-shape depends on the metrics. In order to get a statistically meaningful quantitative estimation of the energetic cost of the bioisosteric transformation, the molecular pairs were binned according to their intermolecular similarities. For each of the 21 bins, the mean averages and standard deviations of $\Delta\Delta G_{A,B, \text{aq}}$ were calculated. The results are presented in Figure 11.

Based on the quantitative energetic assessment, reasonable similarity thresholds were derived for the various distance metrics at the 1 kcal/mol, 1.5 kcal/mol, and 2 kcal/mol levels. Additionally, the number of bioisosteric and random pairs retrieved at these levels were extracted from the similarity calculations.

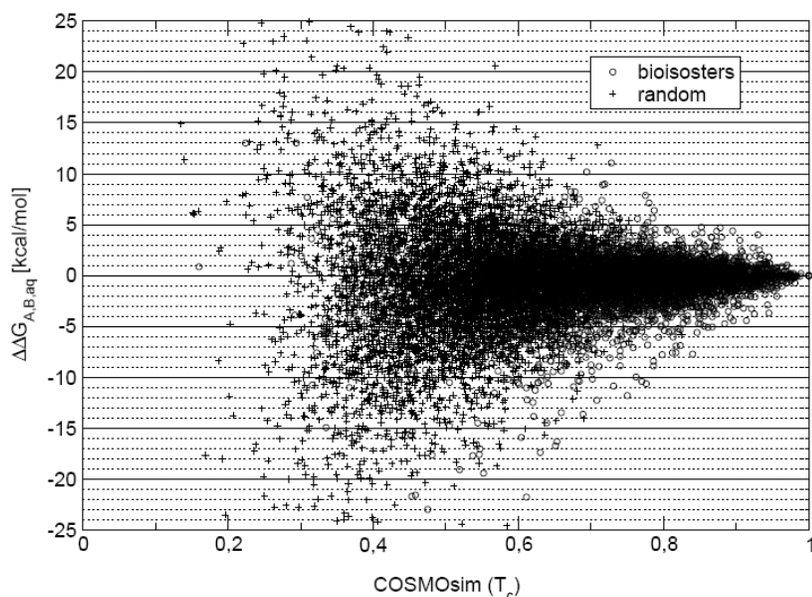


Figure 8. Free energy of solvation changes upon bioisosteric transformation of compound A into B for bioisosteric and random pairs using the COSMOSim Tanimoto coefficient.

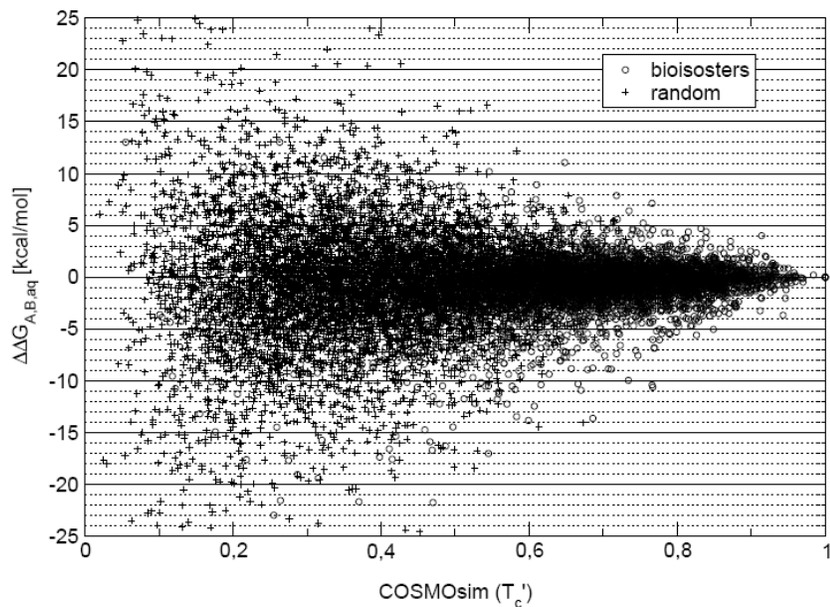


Figure 9. Free energy of solvation changes upon bioisosteric transformation of compound A into B for bioisosteric and random pairs using the COSMOsim Tanimoto prime coefficient.

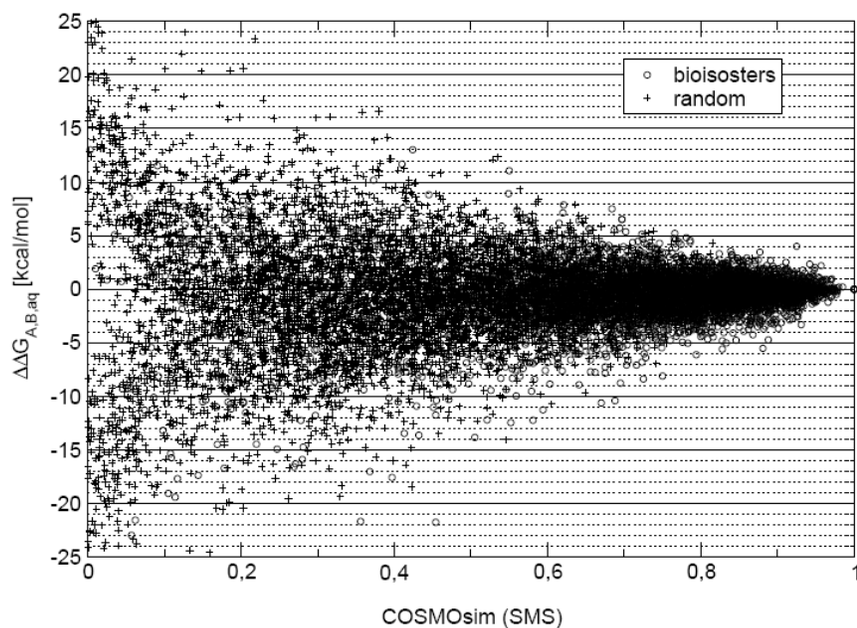


Figure 10. Free energy of solvation changes upon bioisosteric transformation of compound A into B for bioisosteric and random pairs using the COSMOsim SMS coefficient.

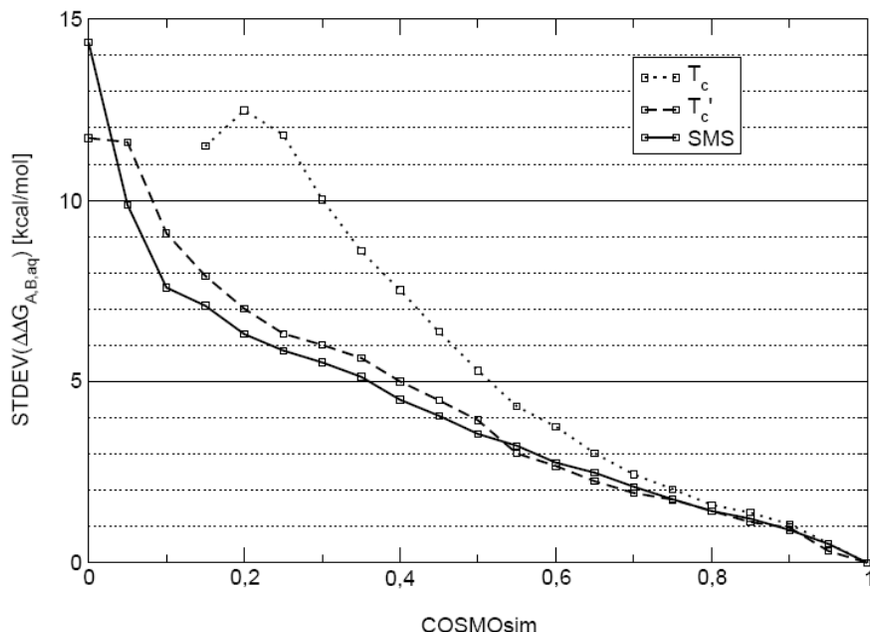


Figure 11. Standard deviation of free energy of solvation changes upon bioisosteric transformation of compound A into B for bioisosteric and random pairs using various COSMOsim coefficients.

PROPER AND IMPROPER BIOISOSTERS. Commonly, bioisosters are seen as pairs of compounds that show comparable activity on the target in question. The energetic cost of the corresponding bioisosteric transformation is low for all bioisosters. We have shown that for compounds with very similar σ -profiles, i.e., COSMOsim approaching unity, the energetic cost of the bioisosteric transformation $\Delta\Delta G_{A,B, \text{binding}}$ becomes small as a result of the different energetic contribution terms according to Eq. 7 becoming small. The reverse is, however, not necessarily given: not all fairly equipotent ligands need to have similar σ -profiles. Consistently, the COSMOsim values of such ligand pairs are low. The reason can be found in the different energetic contribution terms to $\Delta\Delta G_{A,B, \text{binding}}$ that sum up to zero although the single terms do considerably differ from zero and have opposite signs. This distinct energetic behavior gives rise to the definition of two principle classes of bioisosters:

- 1) Proper Bioisosters: All contributions to the energetic cost of the bioisosteric transformation according to Eq. 7 are small. They have overall similar physicochemical properties and their similarity is principally receptor-independent. A hypothetical example is given below:

$$\Delta\Delta G_{AP, BP, \text{binding}} = 0 \text{ kcal} = 0 + 0 + 0 + 0 \text{ kcal}$$

- 2) Improper Bioisosters: The energetic cost of the bioisosteric transformation is small but the discrete contributions are not small. They have different biophysical properties and their similarity is receptor-dependent. A hypothetical example is given below:

$$\Delta\Delta G_{AP, BP, \text{binding}} = 0 \text{ kcal} = 5 - 8 + 15 - 12 \text{ kcal}$$

The application of COSMOsim methods in vHTS will, therefore, yield proper bioisosters. They can fail at the target receptor for steric mismatch or repulsion reasons, which are, though, beyond pseudo-liquid treatment. We are, therefore, currently extending COSMOsim towards the third dimension and first results were recently published³⁰.

SEPARATION OF BIOISOSTERIC FROM RANDOM MOLECULAR PAIRS. The general applicability of COSMOsim was evaluated aiming at the numeric separation of the two sets of molecular pairs, bioisosteric and random. The different coefficients, T_c , T_c' , and SMS with optimized parameters were applied to each molecular pair. Subsequently, the resulting intermolecular similarities were evaluated in the normalized and cumulative histograms (Figures 12 – 13).

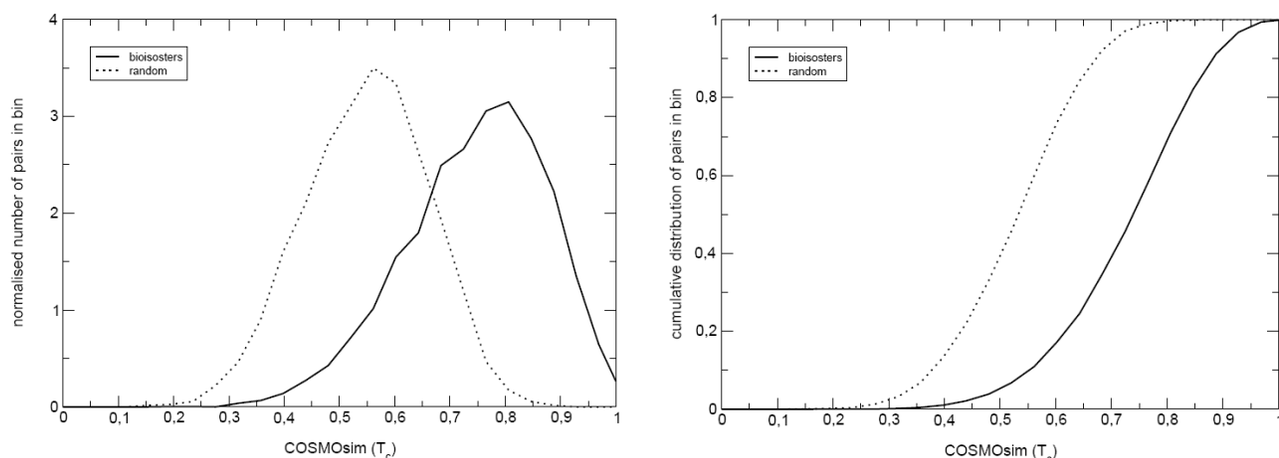


Figure 12. Separation of known bioisosters from random pairs resulted from the application of COSMOsim Tanimoto coefficient to the BioSter dataset (left: normalized distribution, right: cumulative normalized distribution).

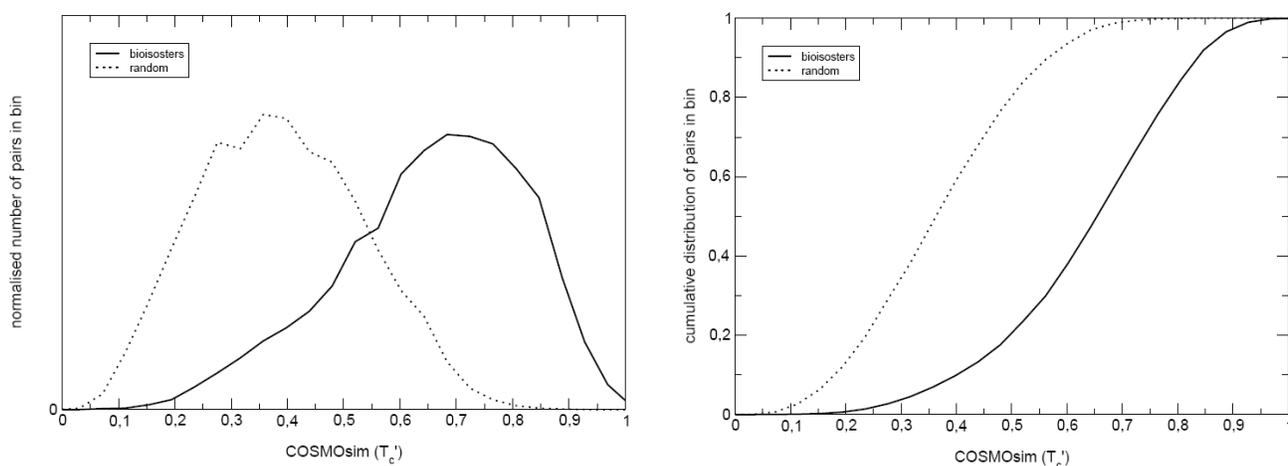


Figure 13. Separation of known bioisosters from random pairs resulted from the application of COSMOsim Tanimoto prime coefficient to the BioSter dataset (left: normalized distribution, right: cumulative normalized distribution).

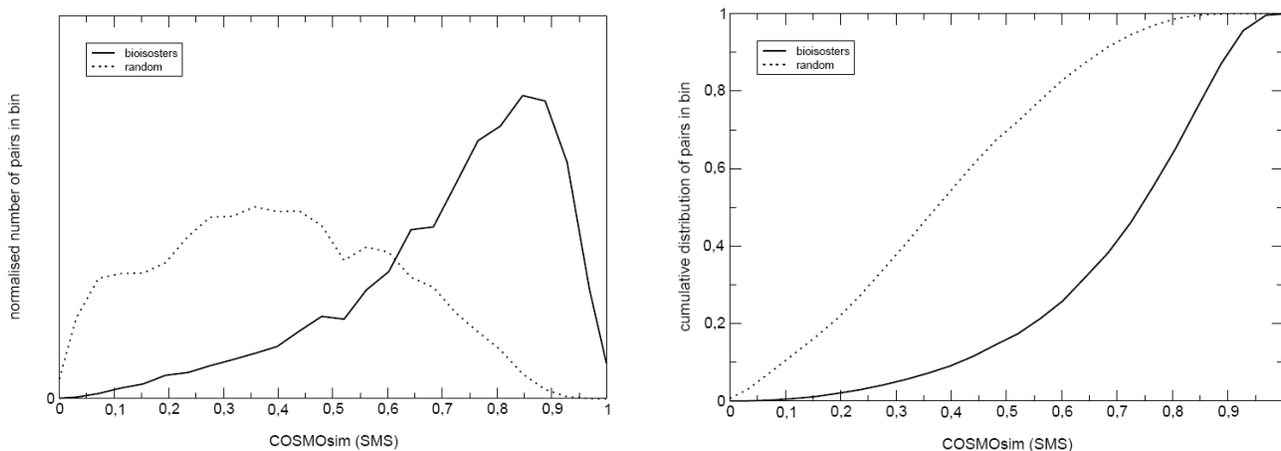


Figure 14. Separation of known bioisosters from random pairs resulted from the application of COSMOsim SMS coefficient to the BioSter dataset (left: normalized distribution, right: cumulative normalized distribution).

GROUP EXCHANGE TRANSFORMATIONS WITH COSMOsim. Group exchange applications are useful for the selection of starting materials to be employed in the next generation of compounds maintaining the scaffold (and therefore minimizing the chance of steric repulsion and maximizing the suitability of the pseudo-liquid treatment). We have selected four small molecules that are found incorporated in a plethora of natural compounds and xenobiotics and for which many bioisosteric conformations are known, i.e., cyclohexane, naphthalene, thiazole, and propionic acid. The σ -profiles of these compounds were calculated and the σ -profile database was screened for the most similar molecules according to the COSMOsim SMS coefficient. In order to make comparison to well-established 2D fingerprint similarity techniques possible, we calculated the Tanimoto coefficient based on 1024 bit Daylight keys³¹, often just (wrongly) called Tanimoto-similarity, for the top-ranking compounds. The results are given in Figures 15 to 18.

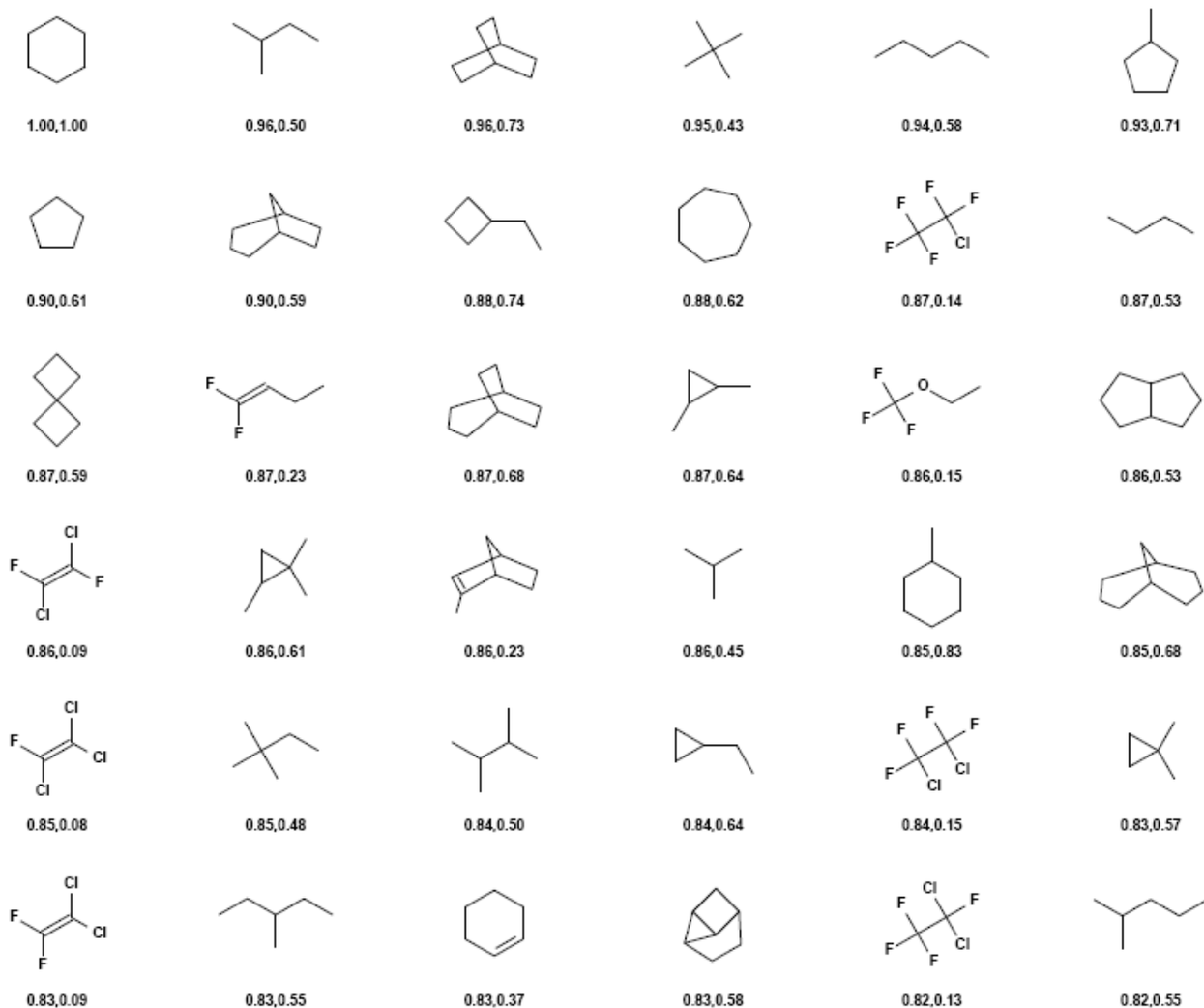


Figure 15. Bioisosters as resulted from vHTS using cyclohexane as target. COSMOsim SMS coefficients are given below the picture (left) along with the corresponding Daylight key Tanimoto coefficient (right).

WHOLE MOLECULE TRANSFORMATIONS WITH COSMOsim. A virtual screening application of COSMOsim was performed. The target ligand, chlorpromazine, is a member of the tricyclic antidepressants class. The most similar compounds of the σ -profile database, in terms of SMS, are given in Figure 19 along with the corresponding Daylight fingerprint T_c values.

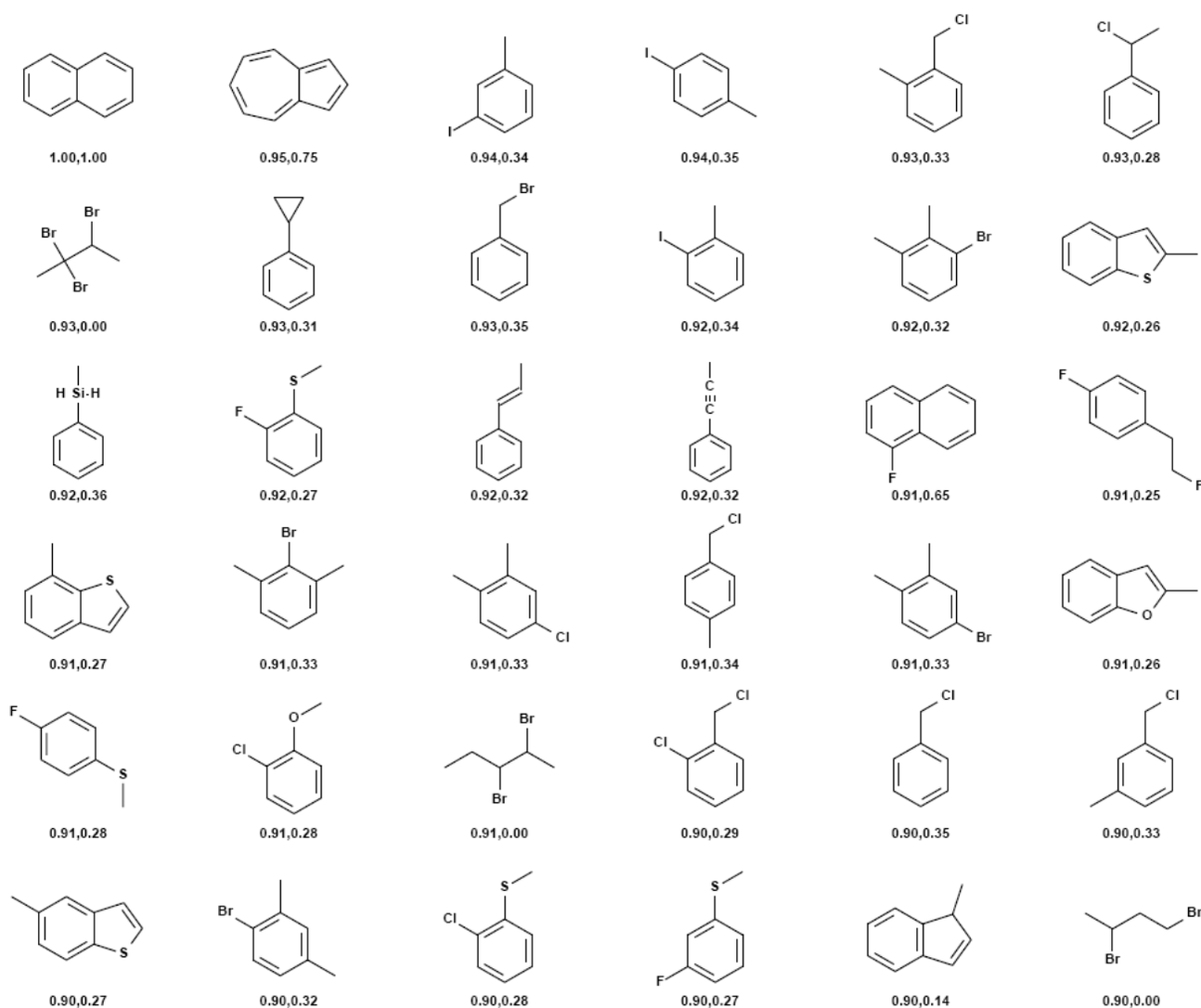


Figure 16. Bioisosters as resulted from vHTS using naphthalene as target. COSMOsim SMS coefficients are given below the picture (left) along with the corresponding Daylight key Tanimoto coefficient (right).

RESULTS AND DISCUSSION

COSMO-RS and COSMO $therm$ provide the basis for a rigorous treatment of the energetic contributions protein-ligand interactions. The only necessary assumption is that ligand, receptor, and the non-covalent ligand-receptor complex behave as pseudo-liquids or isotropic phases. This implies that there is no steric repulsion between ligand and receptor upon complex formation, which is certainly valid for many potent, highly efficient ligands⁴. Steric mismatch and repulsion and their energetic consequences are, thus, beyond the scope of COSMO sim .

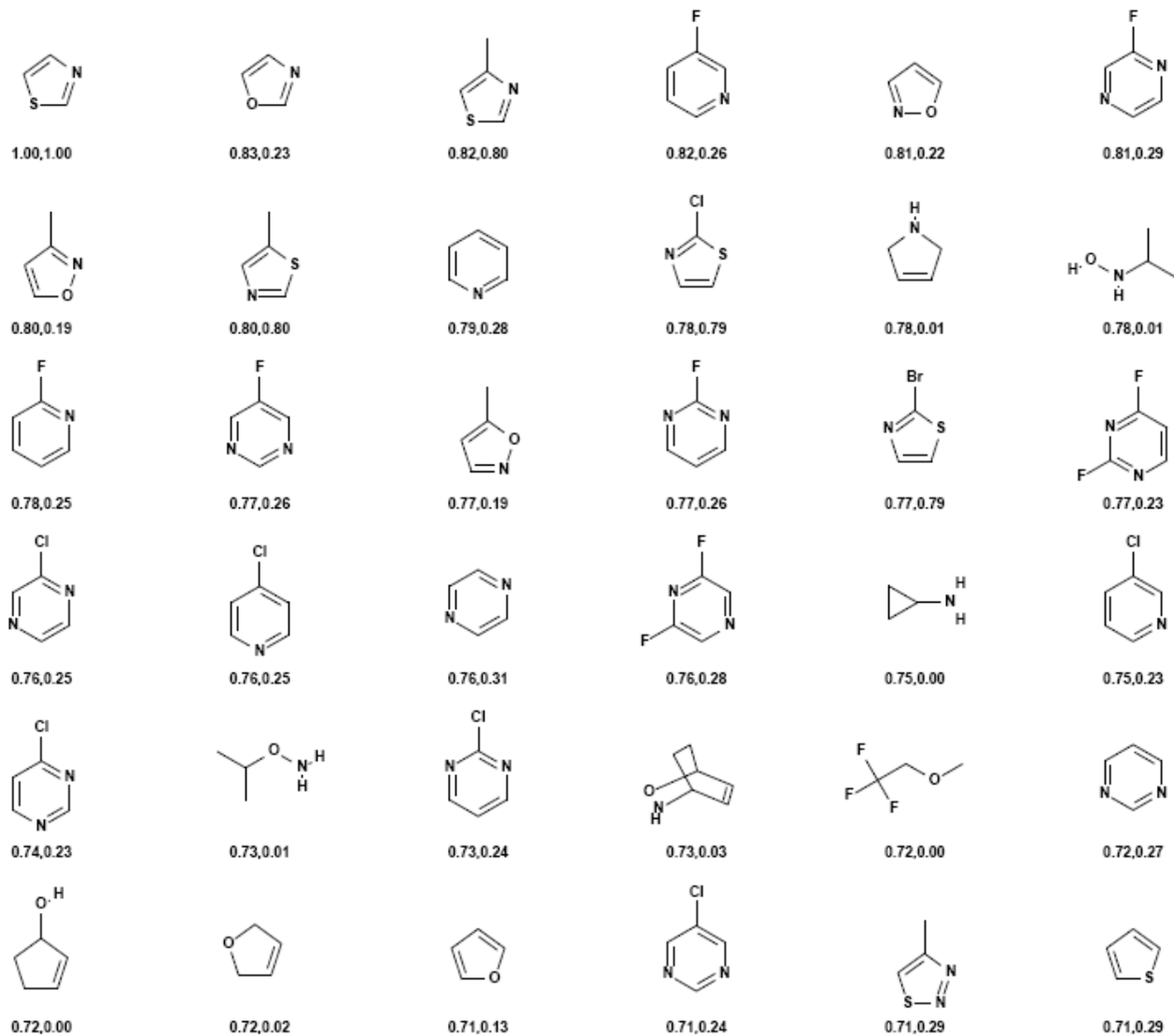


Figure 17. Bioisosters as resulted from vHTS using thiazole as target. COSMOsim SMS coefficients are given below the picture (left) along with the corresponding Daylight key Tanimoto coefficient (right).

	t_{sim}	bioisosters	random	t_{sim}	Bioisosters	random	t_{sim}	bioisosters	random
SMS	0.883	846	0	0.789	2380	120	0.715	3379	357
T_c'	0.891	191	0	0.789	1152	6	0.689	2522	72
T_c	0.906	356	0	0.823	1477	2	0.755	2740	83

For practical applications, this energetic cost has to be low in the order of a few kcal/mol to not lose too much binding affinity. For a large number of successful bioisosteric transformations and random molecular pairs we could show that bioisosters do indeed have more similar σ -profiles than random molecular pairs (see Table 2). The simple Tanimoto similarity coefficient was apparently improved by the introduction of the relative molecular sizes as outlined in the alcohol example (see Figure 6). The number of bioisosteric pairs retrieved at the same energetic cost (see Table 1) was though slightly lower for the Tanimoto prime coefficient. The neighborhood of bins in the σ -profile and the resulting biophysical impact is completely neglected by both Tanimoto based coefficients. Therefore, we developed the sigma match similarity (SMS) metrics that accounts for the biophysical relevance of neighboring bins, the distinct importance of polar versus apolar regions, and differences in molecular size.

The four free parameters of the SMS were optimized to achieve maximum separation of bioisosters from random pairs. A closer investigation of the parameter sets obtained from the GA optimization as given in Table 1 reveals the following trend:

- improved metrics are obtained with a balanced σ -tolerance ($a_{best3}=2.533$, $a_{best4}=2.561$) while its neglect leads to a bad separation ($a_{worst}=1.096$)
- a harsh decrease of the σ -tolerance in the polar region leads to bad separation ($b_{worst}=0.009430$) while a slight decrease yields a good separation ($b_{best3}=0.000350$, $b_{best4}=0.000124$)
- the similarity of polar regions must be weighted higher than that of apolar regions ($c_{best3}=0.009960$, $c_{best4}=0.009990$) otherwise the separation suffers ($c_{worst}=0.001310$)
- molecular size should not change for a bioisosteric transformation otherwise the separation suffers ($d_{worst}=0.993$)

These findings are in accordance with previous findings concerning the size of bioisosteric groups¹⁰ and the general importance of polar groups for binding⁶ and solvation. The most striking difference in SMS as opposed to the Tanimoto coefficients is, however, the introduction of the σ -tolerance parameter a that does indeed lead to a great improvement of the method. The application of the various coefficients to the bioisoster and random pair molecule sets show that all COSMOsim coefficients can be used for the prediction of bioisosters. However, the SMS metrics is to be preferred over the Tanimoto coefficient based measures because it is build upon a sound biophysical concept. Moreover, SMS retrieves about twice the time bioisosters at the same energetic cost for the bioisosteric transformation as expressed in the mean square change in solvation $\Delta\Delta G_{A,B,solv}$ (see Table 1). COSMOsim is extremely fast (29689/28078/9869 compounds per second on a single CPU 3GHz Pentium 4 for T_c / T_c' /SMS) and is therefore well suited for the virtual HTS of billions of compounds. We recommend performing explicit

calculation at DFT BP-SVP-COSMO-SP using a bioactive conformation of the target ligand in order to improve the quality of the target σ -profile. Moreover, explicit calculations can be run over night on the top ranking compounds since the corresponding DFT-based σ -profiles provide a basis for the calculation of biophysically relevant properties like pK_a , logP, logS, etc. with higher accuracy^{19,20,32}.

The assessment of the group transformation applications reveals clearly that COSMOsim perceives molecular similarity in a way medicinal chemists do. The results given in Figures 15 and 16 underline the apolar aliphatic and aromatic characters and sizes of cyclohexane and naphthalene bioisosters, respectively. These simple bioisosteric transformations are, though, not at all perceived by the Daylight key (substructure) based similarities. Similar results are expected for Unity or MDL keys but one has to keep in mind that these methods were mainly developed as hash keys and bioisosterism is beyond their scope although they are widely used for this purpose³³. The two polar group examples reveal further advantages of the structure-free comparison COSMOsim. Well-known bioisosters of thiazole (see Figure 17) like oxazole and pyridine appear in the top-ranking list as expected while the structure-based approach would miss them in every virtual screening. It is important to stress that the same structure-based approach would likely fail to rank larger compounds containing thiazole and pyridine as functional groups as well. The propionic acid example (see Figure 18) paints a good picture on how electronics and molecular size are reflected in COSMOsim. Obviously, quite a few hits are carboxylic acids of similar size, i.e., trivial structure derivatives and as such well perceived by the Daylight key method. COSMOsim, however, retrieves additional well-known bioisosters like carbamic and hydroxamic acids, enoles, phosphinic, boronic, and sulphenic acids, and acidic heterocycles like tetrazoles and oxadiazolones. To our knowledge, COSMOsim is the only method that retrieves the latter non-classical bioisosters *a priori*.

The bioisosteric transformation of chlorpromazine should exemplify the suitability of COSMOsim to the discovery and design of therapeutic copies. Chlorpromazine is one of the oldest tricyclic antidepressants found more than 50 years ago by chance³⁴. Apart from its antidopaminergic activity, it antagonizes furthermore histamine, 5-hydroxytryptamine, acetylcholin, and cannabinoid receptors. Since its discovery, a plethora of bioisosters has been discovered and designed that focus on one or the other CNS activity. The top ranking compounds in Figure 19 show clearly that many of the strategies employed in medicinal chemistry are perceived by COSMOsim, i.e, small variations around the same scaffold, ring substitutions, side chain length variations, and various approaches to rigidify the molecules²⁶. Again, the structure-based screening would have omitted most of the chlorpromazine analogues found by COSMOsim. This example shows impressively that bioisosteric transformations introducing only small electronic changes can yield a huge variety of distinct proper bioisosters.

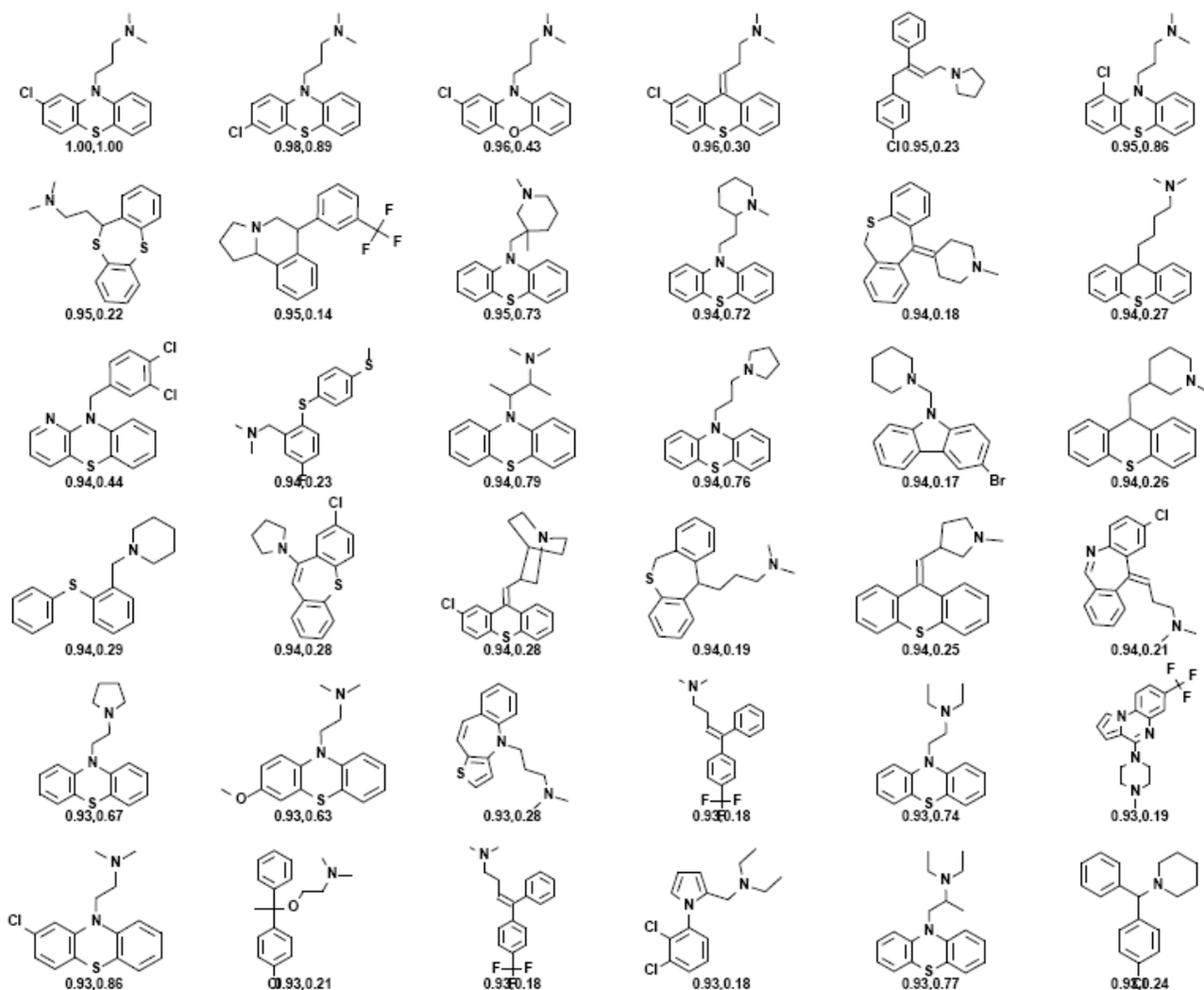


Figure 19. Bioisosters as resulted from vHTS using chlorpromazine as target. COSMOsim SMS coefficients are given below the picture (left) along with the corresponding Daylight key Tanimoto coefficient (right).

The energetic cost of a bioisosteric transformation can be measured experimentally in a biological assay after chemical synthesis. In virtual screening target-focused solutions are ranked according the corresponding fitness values, for example *COSMOsim* values. It is important to know what the energetic cost of a bioisosteric transformation can be and how this cost depends on the intermolecular *COSMOsim* similarity. Unfortunately, the Bioster database does not contain any information on binding or inhibition constants that could serve to calculate the energetic cost of the corresponding bioisosteric transformations. It can be assumed, however, that bioisosteric pairs do not differ by more than two log orders of magnitude in their inhibition which corresponds roughly to 2,7 kcal/mol. This value is in the order of magnitude of the solvation energy changes at relevant similarity levels according to Table 1. We can, therefore, conclude that the energetic changes in solvation of the protein-ligand complexes $\Delta\Delta G_{AP,BP,aq}$ and the protein-ligand binding $\Delta\Delta G_{AP,BP,cosmo}$ are of the same magnitude. Consistently, bioisosteric transformations with high *COSMOsim* will likely lead to proper bioisosters of no additional steric repulsion is introduced. Also, the energetic cost drops as the proper bioisosteric pair becomes more similar.

It should be underlined that other physicochemical molecular properties (logP, logS, etc.) follow the same thermodynamic cycle for a bioisosteric transformation shown in Figure 5 by substituting the protein receptor P with another pure or mixed isotropic or anisotropic phase of interest (water, octanol, blood, brain, etc.). Successful bioisosteric transformations will, therefore, yield compounds with overall similar properties with exception of their chemical structure.

For isotropic phases steric repulsion does not exist. Anisotropic phases like proteins can, however, display the steric mismatch and repulsion issues and consistently a proper bioisosteric transformation will eventually not work for a given receptor (while it might well work for another receptor). The biological assay remains to provide the ultimate proof. COSMOsim was successfully employed in a variety of projects for group transformations and target-focused library design where it already delivered a variety of potent inhibitor families for different targets.

ACKNOWLEDGEMENTS

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