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# Ligand-based virtual screening using binary kernel discrimination

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This paper discusses the use of a machine-learning technique called binary kernel discrimination (BKD) for virtual screening in drug- and pesticide-discovery programmes. BKD is compared with several other ligand-based tools for virtual screening in databases of 2D structures represented by fragment bit-strings, and is shown to provide an effective, and reasonably efficient, way of prioritising compounds for biological screening.

**Keywords:** Binary kernel discrimination; Circular substructure; Fingerprints; Similarity searching; Virtual screening

## 1 Introduction

Biological testing in pharmaceutical and agrochemical research is resource intensive (in terms of both time and money), and the cost-effectiveness of such testing is hence increased, when compared to selecting molecules at random, if it is possible to eliminate from consideration those molecules that have low *a priori* probabilities of exhibiting the biological activity of interest. *Virtual screening* is the name given to the use of computational methods to rank a dataset in decreasing order of such probabilities, so that attention can be focussed on the molecules at the top of the ranking [1,2]. There are two main approaches to virtual screening: Ligand-based approaches use information pertaining to molecules that are already known to be active or inactive; and structure-based approaches additionally exploit information about the three-dimensional structure of the biological target of the drug or agrochemical. Here, we consider a ligand-based virtual screening technique called *binary kernel discrimination* (BKD).

Following a general description of the BKD approach, we compare BKD with several other machine learning methods that can be used for ligand-based virtual screening when heterogeneous sets of both active and inactive molecules are available for use as training data.

These comparisons involve the NCI AIDS database [3] and pesticide data from the Syngenta corporate file, and demonstrate the general effectiveness of BKD. We next consider the use of similarity searching as a virtual screening mechanism. Similarity searching is normally used to identify those database molecules that are most similar to a single active *reference structure*, using some quantitative definition of inter-molecular structural similarity [4,5]. Here, we consider similarity-based virtual screening techniques that can be used when not one but several different bioactive reference structures are available, i.e. when the training-set contains only active molecules. A comparison of several search methods that exploit multiple reference structures, this time using the *MDL drug data report* (MDDR) database [6], demonstrates the effectiveness of an approximate form of BKD. The NCI, Syngenta and MDDR experiments involve the molecules being represented by fingerprints that encode the presence or absence of 2D fragment substructures and that are generated using the Unity software for chemical information management produced by Tripos Inc. [7]. We then discuss BKD's performance when an alternative type of 2D fingerprint, based on circular substructures, is used, and compare the results obtained with those reported in a recent study of similarity-based virtual screening using a different type of circular-substructure descriptor

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[8]. Finally, we describe current work that considers the effect of using different types of kernel function (see below) on the performance of BKD.

## 2 Binary kernel discrimination

Kernel estimation methods seek to evaluate the density of points in a space surrounding a given point. In the virtual-screening context, we are interested in the distribution of known active and inactive molecules in a training-set that are in the immediate vicinity of a test-set molecule of unknown activity [9]. The BKD method makes use of this information to estimate the likelihood of activity for molecules, and these likelihoods can then be used for ranking. All machine-learning approaches involve some training stage; the output from this stage in BKD is a value for the smoothing parameter  $\lambda$  (see below) that measures the extent of the contribution from the training-set molecules as a function of their distance in chemical space from the test-set molecule.

Most kernel methods assume the use of real-valued descriptors, but Aitchison and Aitken first demonstrated the use of a kernel function specifically designed for binary descriptors in a medical application [10]. The resulting BKD method was first applied to problems in chemoinformatics by Harper *et al.* [9], who compared its performance in simulated virtual screening experiments with conventional similarity searching and with a neural network. For two molecules  $i$  and  $j$ , characterised by binary fingerprints of length  $M$  that differ in  $d_{ij}$  positions, Harper *et al.* used the kernel function  $K_\lambda(i, j)$

$$K_\lambda(i, j) = \lambda^{M-d_{ij}}(1 - \lambda)^{d_{ij}}$$

where  $\lambda$  is a smoothing parameter whose optimum value ( $0.5 \leq \lambda \leq 1.0$ ) is to be determined. Training-set molecules (i.e. those whose activity or inactivity is already known) are then ranked using the scoring function

$$L_A(j) = \frac{\sum_{i \in \text{active}} K_\lambda(i, j)}{\sum_{i \in \text{inactive}} K_\lambda(i, j)},$$

with the optimum value of  $\lambda$  being found from repeated analysis of the training-set. The optimum is obtained by computing scores for each training-set molecule using the other training-set molecules for a number of different values of  $\lambda$  in the range 0.50 to 0.99. For each value of  $\lambda$  the sum of the ranks of the active molecules is computed. If this is plotted against  $\lambda$  a clear minimum should be observed indicating the optimum  $\lambda$ , i.e. the value that minimises the summed ranks of the actives in the training-set. It is assumed that the optimal value in the training-set is also optimal for the test-set: This is clearly a strong assumption, but the results we have obtained suggest that it does not result in poor predictive performance (and it is difficult to use a machine-learning technique like BKD without such an assumption). Once the optimum value has

been obtained, test-set molecules (i.e. those whose activity or inactivity is unknown) are compared with the actives and inactives in the training-set, and then ranked in decreasing order of the calculated  $L_A(j)$  values.

The parameter  $\lambda$  lies at the heart of the BKD procedure, in that it determines how much bigger the value of the kernel function is for similar molecules than it is for dissimilar molecules. Specifically, if  $\lambda$  takes its lower bound value of 0.5 then the kernel functions for all of the training-set molecules contribute equally to the overall score,  $L_A(j)$ , for each individual test-set molecule,  $j$ . However, as  $\lambda$  increases towards its upper bound value of unity, the overall score is increasingly dominated by the largest kernel functions. Thus, if some particular test-set molecule is highly similar to many training-set molecules there will be many significant contributions to  $L_A(j)$ . Conversely, if some particular test-set molecule resembles only a very few training-set molecules then that molecule will receive correspondingly few significant contributions to  $L_A(j)$ . The parameter  $\lambda$  hence tunes BKD to the particular characteristics of the training-set actives and inactives that are closest to each test-set molecule. It is this tuning facility that enables BKD to be a highly discriminating tool for classification or ranking.

## 3 Virtual screening with active and inactive training data

### 3.1 Virtual screening methods

**3.1.1 Similarity methods.** The simplest way of predicting the activities of a set of molecules is by computing their similarities to a training-set of known actives and inactives, i.e. a  $k$ -nearest neighbour classifier. Two approaches are reported here; experiments with other, less effective similarity approaches are discussed by Wilton *et al.* [11]. Given a test-set molecule,  $j$ , to be predicted,  $S_{\max}$  is defined to be the similarity to the most similar training-set active, i.e.

$$S_{\max}(j) = \max\{S(i, j)\} \quad i \in \text{actives};$$

while  $S_{A-I}$  is defined to be the mean similarity to all of the  $A$  training-set actives minus the mean similarity to all of the  $I$  training-set inactives, i.e.

$$S_{A-I} = \frac{1}{A} \sum_{i \in \text{actives}} S(i, j) - \frac{1}{I} \sum_{i \in \text{inactives}} S(i, j).$$

The similarities in these approaches are calculated using the Tanimoto Coefficient [4]. If  $a$ ,  $b$  and  $c$  are the numbers of bits set in one fingerprint, set in the other fingerprint, and set in both fingerprints, respectively, then the Tanimoto coefficient is defined to be

$$\frac{c}{a + b - c}$$

In these methods, as with all the others considered here, the test-set molecules are ranked in descending order of

the calculated scores, i.e. similarity values in the present context, with the expectation that the top-ranked molecules have the greatest probability of activity.

**3.1.2 Substructural analysis methods.** Substructural analysis was first described by Cramer *et al.* [12], and many different weighting schemes have been described for this purpose [13]. For each fragment or bit,  $j$ , in the binary fingerprints that characterise the training-set molecules, a weight is calculated that is a function of the numbers of active and inactive training-set molecules that have the  $j$ -th bit set. A score is then computed for a test-set molecule by summing (or otherwise combining) the weights of those bits that are set in its fingerprint. Here, we report the use of the  $R1$  and  $R2$  weights that performed well in the comparative study of Ormerod *et al.* [13]. Let  $A_j(I_j)$  be the number of active (inactive) molecules with bit  $j$  set, and let  $T_j$  be the total number of molecules with bit  $j$  set. Similarly, let  $A(I)$  be the total number of active (inactive) molecules, and let  $N_T$  be the total number of molecules in the training-set. Then the two weights are given by

$$R1 = \log\left(\frac{A_j/A}{T_j/N_T}\right), \quad R2 = \log\left(\frac{A_j/A}{I_j/I}\right).$$

**3.1.3 Binary kernel discrimination.** Finally, we used the BKD approach that was first reported by Harper *et al.* [9] and that has been described previously in this paper.

## 3.2 Results and discussion

We carried out simulated virtual screening experiments on two datasets, one public and one corporate. The initial experiments used the NCI AIDS file [3], which contains molecules that have been checked for anti-HIV activity, with 1129 confirmed actives or confirmed moderately actives and 34,862 confirmed inactives. Training-sets were randomly generated, each containing 200 actives and 200 inactives, with the remaining 35,591 molecules forming the test-set: Three such training-sets were generated for the experiments. The Syngenta dataset contained 1,32,784 molecules that had been tested in various *in vivo* whole organism screens; of these 7127 were active in at least one screen, with the remaining 1,25,657 having a response in the screens less than a pre-defined threshold value. As before, three different training-sets were randomly generated, each containing 713 actives (i.e. 10% of the total actives) and 713 inactives with the remaining 1,31,358 molecules in each case forming the test-sets.

The test-set and training-set molecules were represented by 988-bit Tripos Unity fingerprints [7] and scores were calculated for each of the test-set molecules using one of the techniques described in the previous section. The test-set was then ranked, and the effectiveness of the various methods determined by noting the numbers

Table 1. Percentages of the active molecules retrieved in the top 1% and in the top 5% of the rankings for the NCI and Syngenta datasets.

Ranking Method	NCI dataset		Syngenta dataset	
	Top 1%	Top 5%	Top 1%	Top 5%
$S_{\max}$	8.2	38.9	5.9	19.1
$S_{A-I}$	10.4	21.4	6.6	19.1
$R1$	11.5	21.4	6.8	18.7
$R2$	1.5	16.5	6.9	20.6
BKD	13.7	42.7	9.8	27.1

of actives in the top 1% and the top 5% of the ranking. The results obtained with the three different training-sets were all very similar, in that though there were variations in the precise values obtained with the different test-sets there was very little difference in the relative performance of the various methods; we hence consider the results for only one of the training-sets, as detailed in table 1. Inspection of this table leads to a simple, unequivocal conclusion: That the BKD method gives rankings that are far superior to those of the other ranking methods considered here, and that this is hence the method of choice in terms of the effectiveness of virtual screening. The difference between BKD and the other approaches in the table (and the many other approaches discussed by Wilton *et al.* [11]) is particularly marked with the large Syngenta dataset.

## 4 Virtual screening with active training data

The increasing use of high-throughput screening means that large amounts of active and inactive training data are likely to become available shortly after the commencement of an agrochemical or pharmaceutical lead-discovery programme. At the start of a programme, however, the medicinal chemist may have knowledge of just a few active molecules, such as natural products, or patented, competitor molecules. In such cases, an alternative type of virtual screening can be used, based on similarity searching [4,5]. This involves matching a single bioactive reference structure against each of the database structures to identify those that are most similar (and hence most likely to exhibit the same activity as the reference structure). Here, we evaluate three distinct approaches that can be used when multiple reference structures are available, i.e. when the training-set contains only actives.

### 4.1 Virtual screening methods

**4.1.1 Modal fingerprint method.** The single fingerprint approach was first described by Shemetulskis *et al.* in their work on Stigmata [14]. The method generates a *modal fingerprint* from an input set of molecules that seeks to capture the common chemical features present in the members of this training-set. A bit  $j$  is set to “on” in the modal fingerprint if that bit is found in more than

a user-defined threshold percentage of the training-set molecules. The modal fingerprint is then used as a query and compared to the fingerprints of the molecules in the test-set. We have adopted this idea here, using the Tanimoto Coefficient to rank the molecules of the database.

**4.1.2 Data fusion method.** *Data fusion* is the name given to a range of techniques that combine inputs from different sensors, with the expectation that using multiple information sources enables more effective decisions to be made than if just a single sensor is employed. The approach has been used in many different fields; when applied to chemoinformatics applications (where it is sometimes referred to as *consensus scoring*) the fusion is effected by combining the results of several database searches using different descriptors or scoring functions [15]. In conventional applications of data fusion, a single active reference structure is searched against a database in several different ways; in the present context, we have several different reference structures that are all searched against a database in exactly the same way (specifically using 2D fingerprints with the Tanimoto coefficient). We have fused the similarity scores,  $S(i,j)$  ( $1 \leq i \leq n$ , the number of active reference structures) for some molecule  $j$  using the  $S_{\max}$  fusion rule discussed in the first section of the paper, i.e. the test-set molecules were ranked on the basis of

$$\max\{S(i,j)\} \quad i \in \text{actives}$$

**4.1.3 Substructural analysis method.** A weighting scheme for substructural analysis normally requires access to training-set data for both actives and inactives. In the present context, however, we do not have access to all of the necessary information as the training-set consists of just active molecules. However, if we restrict our attention to those weighting schemes that do not make explicit use of information about the inactives and also make the assumption that the overall characteristics of the training-set are mirrored by those of the entire database that is to be searched, then we can use the  $R1$  weight. The equation for

the weight is as given previously; here, however,  $T_j$  is the total number of molecules in the database with bit  $j$  set and  $N_T$  is the total number of molecules in the database (rather than the total number of molecules in the training-set with bit  $j$  set and the total number of molecules in the training-set, respectively, as is the case in conventional substructural analysis).

**4.1.4 Binary kernel discrimination.** An analogous approximation can be used to enable BKD to be used when only actives are available for training purposes. Specifically, we make the assumption that the overall characteristics of the inactives are approximated with a high degree of accuracy by the overall characteristics of the entire database that is to be searched. If this assumption is accepted then a training-set can be generated by taking the set of reference structures and adding to it molecules randomly selected from the database, with the expectation that most, if not all, of these added molecules are inactive. Since actives are inherently very rare, both this expectation and that underlying the  $R1$  approximation are not unreasonable.

## 4.2 Results and discussion

We have evaluated the various approaches above by means of simulated virtual screening searches on the MDDR database [6]. After removal of duplicates and molecules that could not be processed using local software, 1,02,535 molecules were available for searching and these were represented by Unity fingerprints [7]. These molecules were searched using the 11 sets of active molecules from MDDR that are listed in table 2; full details of these experiments are reported by Hert *et al.* [16]. A rough guide to the diversity of each of the chosen sets of bioactives is provided by matching each compound with every other in its activity class, calculating similarities using Unity fingerprints and the Tanimoto coefficient, and computing the mean of these intra-set similarities. The resulting similarity scores are listed in the second column of table 2, where it will be seen that the renin inhibitors are the most homogeneous and the cyclooxygenase inhibitors are the most heterogeneous.

Table 2. Mean percentage of active molecules retrieved by multiple-reference methods over the top 5% of the ranked test-set for the MDDR datasets.

Activity class	Self-similarity	Modal	Data fusion	Substructural analysis	BKD
5HT3 antagonist	0.35	30.3	49.0	29.3	52.3
5HT1A agonist	0.34	21.9	37.2	30.1	38.2
5HT Reuptake inhibitor	0.34	39.6	49.7	33.1	45.8
D2 antagonist	0.34	27.1	37.4	27.5	38.7
Renin inhibitor	0.57	88.8	88.6	52.9	93.3
Angiotensin II AT1 antagonist	0.40	73.6	80.4	43.4	84.5
Thrombin inhibitor	0.41	49.4	58.6	35.6	63.1
Substance P antagonist	0.39	36.8	47.1	36.5	58.4
HIV protease inhibitor	0.44	53.5	61.6	34.1	68.5
Cyclooxygenase inhibitor	0.26	11.0	26.5	19.2	33.2
Protein kinase C inhibitor	0.32	35.6	48.0	35.6	49.4
Average over all classes	0.38	42.5	53.1	34.3	56.8



For each of the 11 activity classes, 10 active molecules were selected for use as the training-set. The selections were done at random, subject to the constraint that no pairwise similarity in a group exceeded 0.80 (using Unity fingerprints and the Tanimoto coefficient). Each searching method was repeated 10 times using different training sets, and in each search, a note was made of the percentage of the active molecules (i.e. those in the same class as those in the training-set) that occurred in the top 5% of the ranking resulting from that search. The results presented in table 2 are the mean recall values, averaged over each set of 10 searches (very similar results to those listed here were obtained if the top 1%, rather than the top 5%, of the rankings were evaluated).

A large number of searches were carried out to identify the best parameter settings for the various methods discussed above (and also several other, less effective methods that are described by Hert *et al.* [16]). The resulting settings were then used in the main experiments, the results of which are detailed in the body of table 2. Inspection of this table shows that fusion of the similarity scores and BKD are the clear methods of choice, consistently out-performing modal fingerprints and sub-structural analysis.

With some minor exceptions, the performance of all of the methods tends to increase as the self-similarity of the active molecules increases. The correlation with intra-class similarity is not unexpected; what is of importance is that good screening performance is obtained even with quite diverse activity classes (such as the protein kinase C inhibitors and the D2 antagonists). The worst results are obtained with the most diverse set of actives, i.e. the cyclooxygenase inhibitors; even here, however, the data fusion and BKD runs represent 5.3-fold and 6.6-fold enrichments, respectively, over a random ranking of the dataset (with average enrichment factors for these two approaches over all classes being 10.6 and 11.3, respectively).

In the final set of experiments, we sought to quantify the benefit that can be achieved using multiple reference structures, when compared with the single reference structures that characterise conventional similarity searching. This was done by using every single active molecule in turn in each of the 11 chosen activity classes as the reference structure, and recording the minimum, mean and maximum performance, as detailed in table 3. The mean values correspond to the performance that might be expected using an average, individual reference structure and are clearly much lower than the figures reported in table 2 for the BKD and data fusion methods (an average over all classes of 30.6% as against 56.8 and 53.1%, respectively). Thus, the use of 10 actives, rather than just one, results in an increase of over two-thirds in the numbers of actives retrieved. Perhaps the most interesting figures in table 3 are those listed under "Maximum". These represent the best single similarity searches possible from the many hundreds of individual bioactive molecules (this number ranges from 349 for the 5HT

Table 3. Percentage of active molecules retrieved by single similarity searches over the top 5% of the ranked test-set for the MDDR datasets.

Activity class	Mean	Maximum	Minimum
5HT3 antagonist	21.2	41.0	1.9
5HT1A agonist	18.4	39.3	2.5
5HT Reuptake inhibitor	24.0	42.7	1.4
D2 antagonist	17.4	35.6	0.3
Renin inhibitor	80.5	93.2	3.0
Angiotensin II AT1 antagonist	48.0	81.7	3.6
Thrombin inhibitor	33.5	63.6	0.6
Substance P antagonist	26.9	57.7	0.5
HIV protease inhibitor	37.6	63.7	1.9
Cyclooxygenase inhibitor	9.4	21.1	0.3
Protein kinase C inhibitor	19.4	46.1	0.7
Average over all classes	30.6	53.2	1.5

reuptake inhibitors up to 1236 for the substance P antagonists). If we consider the average over all activity classes, it will be seen that this upperbound is only fractionally better than the data fusion result in table 2 and is actually worse than the BKD figure. Thus, on average, picking any 10 active reference structures and combining them using BKD (or the  $S_{\max}$  data fusion approach) will enable searches to be carried out that are comparable to even the best possible conventional similarity search using a single active reference structure.

## 5 Use of circular substructures

Thus far, we have considered the use of a single type of 2D fingerprint, specifically those produced by the Unity software [7]. We have recently reported a detailed evaluation of 15 different types of 2D fingerprint using the MDDR datasets discussed in the previous section [17]. This comparison demonstrated the general effectiveness of the fingerprints produced by Scitegic Inc. [18], which are based on circular substructures. A circular substructure is a fragment descriptor where each atom is represented by a string of extended connectivity codes that are calculated using a modification of the Morgan Algorithm [19]. The best results in the study were obtained with the Scitegic extended connectivity fingerprints (ECFPs). The initial code assigned to an atom is based on the number of connections, the element type, the charge, and the mass. This code, in combination with the bond information and with the codes of its immediate neighbour atoms is hashed to produce the next order code, which is mapped into an address space of size  $2^{32}$ , and the process iterated until the required level of description has been achieved. Here, the ECFP\_4 fingerprints were used, where the numeric code denotes the diameter in bonds up to which features are generated.

The Scitegic software represents a molecule by a list of integers, each describing a molecular feature and each ranging from  $-2^{31}$  to  $2^{31}$ . These integer lists were normalised in two ways, referred to as method-A and method-B. In method-A, all the features present in the database were enumerated, so that each feature was given

as its new code its rank in the sorted list of codes, with the length of the resulting fingerprints being the number of distinct features in the database. In method-B, the integers describing a molecule were hashed to a bit-string of length 1024 bits. This inevitably means that collisions occur, with the result that method-B loses some of the structural information that is retained by method-A; however, the latter representation is dependent on the precise database that is being processed.

Bender *et al.* have recently used the 11 MDDR datasets of tables 2 and 3 to test an alternative machine-learning technique based on atom environment fingerprints, information-gain-based feature selection, and a naïve Bayesian classifier [8]. The atom environments are circular fragments centred on each heavy atom in a molecule and encoding, like the Scitegic fragments, the pattern of atoms within a fixed radius of the central atom [20]. The environments are generated for each molecule in a training-set and the feature-selection procedure is next used to identify those environments that are best able to discriminate between the sets of active and inactive molecules in the training-set. The most discriminating environments are then used in a Bayesian classifier routine that ranks the test-set molecules in decreasing probability of activity, on the assumption that the presence and absence of individual environments are statistically independent of each other. Bender *et al.* compare their procedure, called MOLPRINT 2D, with the BKD MDDR results discussed in the previous section, and show that MOLPRINT 2D is noticeably superior [8]. Based on these comparisons (and on comparisons with two other published accounts of virtual screening), Bender *et al.* suggest that the superior performance of MOLPRINT 2D is mainly due to the different descriptors that are used. We believe that this is indeed the case, as discussed further below.

Table 4 details the mean percentage of active molecules retrieved in the top 5% of the ranked test-set in multiple-reference searches using MOLPRINT 2D and using BKD with the Scitegic circular substructures, these results being taken from [8] and [17], respectively. The situation is the

same as for the results in table 2: the training-set contained 10 actives and 100 (assumed) inactives, and the results quoted are mean values averaged over 10 randomly selected training-sets. The MOLPRINT 2D results are those obtained when the feature-selection procedure was used to identify the 250 most discriminating fragments [8]. It will be seen that there is very little difference between the two approaches: BKD performs better for seven of the datasets and MOLPRINT 2D for the remaining four, but there are only two cases (thrombin and protein kinase C) where the difference in mean recall is at all marked. Thus, given a comparable level of molecular description, the similarity calculations required for BKD seems to be as effective as those required for MOLPRINT 2D.

## 6 Use of different similarity coefficients

The kernel function suggested by Aitchison and Aitken [10] is based on the number of bit positions at which two binary strings differ, i.e. the Hamming distance; this was adopted by Harper *et al.* in their application of BKD to virtual screening, and has been used for all of the experiments reported thus far. In principle, however, there is no reason why one could not use an alternative measure of the distance (or similarity) between the fingerprints representing two molecules. Many different similarity coefficients have been used previously for similarity and diversity calculations based on 2D fingerprints [4,21], with these experiments demonstrating that the Tanimoto coefficient generally, but not consistently, gives the best levels of performance, with the Hamming distance normally being notably inferior. We have hence carried out a preliminary series of experiments in which the Hamming distance exponent in the kernel function  $K_{\lambda}(i,j)$  is replaced by one minus one of the similarity coefficients listed in table 5, rescaled by the fingerprint length, i.e.  $M \times (1 - \text{Coefficient})$ . Each of these similarity coefficients is based on two or more of the following variables:  $a$  is the number of bits set to one in the first molecule,  $b$  the number of bits set to one in the second fingerprint,  $c$  the number of bits set to one in both of the fingerprints (so  $c \leq \min\{a,b\}$ ), and  $d$  the number of bits that are not set to one in either of the fingerprints. Thus,  $M$ , the length of each fingerprint, is given by  $(a + b + d - c)$ .

Searches were carried out using the NCI AIDS database with a new training-set of 200 actives and 200 inactives, and with the remaining molecules forming the test-set. The right-hand columns of table 5 list the percentages of active molecules retrieved in the top 1% and the top 5% of the ranked test-set. There is little difference in recall between the coefficients when just the top 1% is considered. However, when the top 5% is considered, the Hamming distance is noticeably superior to the simple matching coefficient, with the three association coefficients—the Tanimoto, Rogers-Tanimoto and Russell-Rao Coefficients—giving comparable, and markedly inferior results. It would hence seem that the normalisation that

Table 4. Mean percentage of active molecules retrieved by multiple-reference methods over the top 5% of the ranked test-set using circular substructure descriptors for the MDDR datasets.

Activity class	MOLPRINT 2D	BKD	
		ECFP_4A	ECFP_4B
5HT3 antagonist	66.6	65.3	63.4
5HT1A agonist	57.1	58.7	57.5
5HT Reuptake inhibitor	46.1	50.3	49.0
D2 antagonist	53.7	55.2	54.2
Renin inhibitor	95.7	96.7	97.0
Angiotensin II AT1 antagonist	95.1	98.0	97.8
Thrombin inhibitor	66.2	74.8	74.1
Substance P antagonist	68.4	67.3	65.5
HIV protease inhibitor	76.0	80.8	80.8
Cyclooxygenase inhibitor	34.7	34.4	32.7
Protein kinase C inhibitor	54.6	49.6	50.3
Average over all classes	64.9	66.5	65.7

Table 5. Effect of similarity coefficient on the percentages of active molecules retrieved in the top 1% and top 5% of the rankings for the NCI dataset.

Coefficient name	Formula	Top 1%	Top 5%
Hamming distance	$a + b - 2c$	12.8	43.7
Tanimoto coefficient	$\frac{c}{a+b-c}$	10.4	21.4
Rogers-Tanimoto	$\frac{c+d}{2a+2b-3c+d}$	12.0	20.6
Russell-Rao	$\frac{c}{a+b+d-c}$	12.1	20.5
Matching coefficient	$\frac{c+d}{a+b+d-c}$	12.9	43.6

characterizes the three association coefficients may adversely affect the operation of the resulting kernel functions. It was also noted that the Hamming distance and simple matching coefficients gave far more well-marked optima for  $\lambda$  during the training stage than was the case with the other coefficients. Further studies of this behaviour are currently in progress.

Experiments were also carried out using sets of 200 renin and thrombin inhibitors from the MDDR datasets discussed previously. In the case of the thrombin inhibitors, we observed a similar pattern of behaviour to that in the NCI experiments, *viz* the Hamming distance being noticeably superior to the simple matching coefficient, with the Tanimoto, Rogers-Tanimoto and Russell-Rao coefficients giving comparable, and markedly inferior, results. It is hence clear that such coefficients are not appropriate for use in this kernel function, but we do not have any theoretical rationale for this behaviour at present. With the renin inhibitors, however, there was very little difference between the various coefficients. We ascribe this behaviour to the fact that these actives are structurally far more homogeneous (as discussed earlier with respect to table 2) than the other two datasets studied here; the active molecules are hence all very similar to each other and almost any machine-learning technique can be expected to produce a high level of performance.

## 7 Conclusions

In this paper we have discussed the application of a machine-learning technique, BKD, to the prediction of biological activity in virtual screening programmes. Our results suggest that BKD provides an attractive focus for future research: It is effective in operation, in both exact and approximate forms; it has been applied successfully to both pharmaceutical and agrochemical datasets; and it is efficient in operation. For example, using the 2D fingerprints for the 132K-molecule Syngenta dataset, with programs written in C and run on a Silicon Graphics R12000 processor, training took about 36 s for each value of  $\lambda$  that was tested, and the subsequent scoring of the test-set (comprising 98.9% of the entire dataset) using the optimal value of  $\lambda$  took about 10 CPU hours. The method is hence clearly applicable to very large chemical databases.

We can regard a kernel function as a new type of similarity measure and it is hence of interest to consider the three components of a chemical similarity measure [4], *viz* the structure representation, the similarity coefficient and the weighting scheme that are used. We have already started to investigate the use of different types of similarity coefficient (as discussed above). In the future, we hope additionally to investigate the use of weighted fingerprints, with the bits being assigned weights derived from substructural analysis of training-set molecules, and the use of alternative types of structure representation.

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