

Ligand-Based Virtual Screening Using a Genetic Algorithm with Data Fusion

John D. Holliday¹, Nor Sani^{1,2}, Peter Willett¹

¹Information School, University of Sheffield, 211 Portobello,
Sheffield S1 4DP, United Kingdom

²Faculty of Information Science and Technology, National University of Malaysia,
43600 UKM Bangi, Malaysia

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Abstract

Substructural analysis provides a simple and effective way of ranking the 2D fingerprints representing the molecules in a database upon the basis of weights that denote a substructural fragment's contribution to the overall activity or inactivity of a molecule. A substructural analysis method has been described recently that is based on the use of a genetic algorithm (GA), with the resulting sets of weights proving to be more effective for ligand-based virtual screening than existing approaches. However, the inherently non-deterministic nature of a GA means that different runs are likely to result in different sets of weights and hence in variations in the effectiveness of screening. This paper describes the use of data fusion to combine the rankings generated in multiple GA runs, and demonstrates that the resulting fused rankings are markedly superior to GA runs on average, and in some cases can even exceed the performance of the very best individual GA run.

Introduction

Virtual screening methods play an important role in the discovery of novel bioactive molecules, such as pharmaceuticals or agrochemicals, and involve the use of computational techniques to score the molecules in a database in descending order of probability of exhibiting the desired activity. High-ranked molecules can then undergo high-throughput or *in vitro* biological testing to ascertain whether they are, in fact, active. Here, we consider ligand-based virtual screening (LBVS) methods, where the only information available for the discovery process are sets of

molecules that have been tested previously and that are hence known to be either active or inactive. Specifically, we focus on substructural analysis, which was pioneered by Cramer *et al.* in the early 1970s [1, 2] and subsequently developed by Hodes *et al.* [3, 4]: it was probably the first LBVS method to be used on a large scale, forming an important component of a National Institutes of Health programme to develop novel anticancer agents [5, 6].

A molecule in a chemical database is often described by a fingerprint, a bit-string in which bits are set to on (or off) depending upon whether a substructural feature is present (or absent) in that molecule. In substructural analysis, a weight is associated with each bit that is computed on the basis of how frequently the corresponding substructural feature occurs, or does not occur, in sets of molecules that have been tested previously for activity. The weight hence describes the probability that a molecule with that feature will prove to be active, and a score is computed for a molecule by combining the weights for all of its constituent features. The resulting scores are used to rank a database in descending order, with those at the top of the ranking then being considered for biological screening since they are assumed to have the greatest *a priori* probabilities of activity. Substructural analysis hence embodies the assumption that a given substructure can influence the activity of a molecule, regardless of the environment in which the feature occurs: this is clearly an extremely crude assumption but the approach has been found to be of value for screening in drug-discovery programmes [7, 8].

A variety of weighting schemes have been described for use in substructural analysis, as reviewed by Ormerod *et al.* [9]. In their comparison, the most generally useful was found to be one that was originally developed by Robertson and Spark Jones [10] for use in text searching (where the aim is to rank a database of documents in order of decreasing probability of relevance to a query, as against decreasing probability of activity in a biological screen in the present context) and that is analogous to LBVS systems based on naïve Bayesian classifier weights such as that used in the Pipeline Pilot software [11-13]. Rather than the detailed probabilistic models that underlie these approaches to substructural analysis, Holliday *et al.* have recently described an approach to weighting that is based on the use of a genetic algorithm (or GA) [14]. The approach proved to be highly effective in operation but it was noted that the inherently non-deterministic nature of GAs meant that different runs could result in different database rankings. This brief communication seeks to address this limitation of the previous work. The GA is described in the next section, together with the use of data fusion (*vide infra*) to combine multiple runs of the algorithm. We then report detailed experiments using three

large datasets for which bioactivity data are available; and compare the effectiveness of the basic GA with the results obtained using a range of different data fusion rules.

Combining GA runs using data fusion

A GA provides a non-deterministic way of tackling computational problems characterised by solution spaces that are too large for exploration using conventional, deterministic algorithms. The approach has been very extensively employed, not least in applications in chemoinformatics and drug discovery [15]. In the present context, the space is that of all the possible weights that could be assigned to each of the bits comprising the 2D fingerprint used to describe each of the molecules in a database. The chromosome in the GA is a vector in which the i -th element contains the weight associated with the substructural feature denoted by the i -th bit in the fingerprint. In the experiments reported here, the fingerprint encoded the 166 MDL structural key definitions in the widely used Pipeline Pilot software, so that the chromosome contained 166 real values, each corresponding to one of the bits in the fingerprint. The score for a particular molecule was then the sum of the weights for those features that it contained (as denoted by the bits that were set to on), and a database was ranked in order of decreasing sums of scores to identify those molecules with the greatest probability of activity. A training-set that contains molecules of known activity and inactivity is used to derive the final sets of weights, with the fitness of an individual chromosome, c , being assessed by identifying the number of active molecules at the top of the ranked training-set when it is ranked using the set of weights encoded in c . Specifically, the fitness function was the enrichment factor based on the top-1% of the ranked training-set (*vide infra*): by using conventional crossover and mutation operators, the fitness function hence evolves chromosomes (i.e., sets of weights) that are able to maximise the enrichment factor, and thus the effectiveness of LBVS when those weights are subsequently applied to test-set molecules for which activity data are unavailable.

The design and implementation of the GA are described in detail by Holliday *et al.*, who found that it provided consistently better enrichments in simulated LBVS experiments than did the Robertson-Spark Jones and Pipeline Pilot weights mentioned previously [14]. That said, the non-deterministic nature of the GA means that different runs may converge on different solutions: this can be problematic if these solutions are markedly different in character since there will be little or no basis for deciding which of the available solutions should be accepted as the final output from a series of runs. In the present context, this means that

different runs will result in different sets of weights and, accordingly, different rankings and different levels of enrichment when these weights are applied to test-set molecules. Holliday *et al.* noted that this was an inherent limitation of the GA approach and reported experiments demonstrating that such variations did indeed occur in practice. For example, in one series of ten runs using the cyclooxygenase inhibitors in the MDDR database, the mean correlation coefficient between the sets of weights, when averaged over all 45 distinct pairs of runs, was only 0.79 [14]. That said, it is important to note that the principal output from a substructural analysis is not the set of weights themselves (although these could provide valuable information for, e.g., suggesting a possible topological pharmacophore); instead, it is the top-ranked test-set molecules that are identified using the weights, since it is these compounds that are being highlighted for further investigation.

In the absence of any obvious means of identifying the best from among a set of rankings, an alternative is to consider ways of combining them and we here draw upon the concept of data fusion, which is a method for combining the information gained from different sensors to achieve an effective or improved decision when compared to that achievable when only a single sensor is available [16]. Data fusion has been used in many different problem domains [17], including chemoinformatics, where it has been shown to enhance the effectiveness of similarity searching [18] and used to combine multiple runs of ligand-protein docking programs and multiple clusterings of a given dataset [19, 20]. When data fusion is used for similarity searching, the sensors that are combined are the rankings of the molecules in a database that result when different similarity measures are used to conduct a similarity search, e.g., a search might be conducted using different fingerprints or different similarity coefficients. The individual rankings are then combined using a fusion rule (*vide infra*) to obtain a single, combined ranking of the molecules in the database that represents the final output of the search. The obvious advantage of data fusion is that there is no need to choose from amongst different rankings of the same set of compounds; but there are two further advantages that have become apparent as the approach has become widely adopted for similarity searching [18]. First, use of fusion offers a much more consistent level of performance (as evaluated using a measure such as the enrichment factor) than when individual rankings are used; second, the fused level of performance is normally superior to the average individual similarity search, and may occasionally be superior to even the best individual search. This is because, as noted by Sheridan and Kearsley, there is no single similarity measure that can be expected to give the best possible results in all possible circumstances [21].

Similarity searching generates a ranking of the molecules in a database that is entirely comparable to that resulting from a substructural analysis study, and it is hence possible to consider applying the fusion rules used in similarity searching to the rankings resulting from multiple runs of the GA described above. Hence, rather than generating, e.g., ten different sets of GA-based weights and the ten consequent rankings of the molecules comprising a test-set, and then having to choose one of these as a basis for deciding which molecules should go forward for biological screening, the ten rankings are here fused to give just a single ranking to provide the input to the screening process. Our implementation of this combined GA/fusion procedure is described in the next section.

Experimental details and results

Our approach has been evaluated in simulated LBVS experiments using three common databases that contain both structural and bioactivity information: the MDL Drug Data Report database (MDDR); the World Of Molecular Bioactivity database (WOMBAT); and the European Bioinformatics Institute's ChEMBL database. The MDDR and WOMBAT datasets used here are described in detail by Gardiner *et al.* [22]: the MDDR dataset contains eleven activity classes and 102,514 molecules, and the WOMBAT dataset contains 14 activity classes and 138,127 molecules. A total of 15 activity classes were obtained from the much larger ChEMBL database by choosing activities that matched one of the classes in the MDDR and WOMBAT datasets; and then each of the 1,352,681 molecules in version 18 of the database was recorded as being active in a specific class if there was a measured pIC₅₀ for that activity of ≥ 5.0 and if there was an associated confidence score of 9; all other molecules were then assumed to be inactive for that class. The molecules in the three datasets were characterised using MDL fingerprints, which encode 166 important substructural fragments and which were generated using Pipeline Pilot software. The training-set for a particular activity class was generated by randomly selecting 10% of the actives and 10% of the inactives, with the remaining 90% of the database providing the test-set for which virtual screening was carried out. The GA was run on the training-set, weights determined for each of the 166 fragments comprising a fingerprint, and then these weights used to rank the molecules in the test-set. For example, there are 982 molecules noted as being renin inhibitors in the ChEMBL dataset, so the training-set contained 98 of these active molecules and 135,170 other, inactive molecules, with the resulting weights being used to score the remaining 1,217,413 test-set molecules.

In their original paper, Holliday *et al.* reported parameterization experiments that were undertaken to optimize the effectiveness of the GA, and suggested the following settings: roulette-wheel selection; one-point crossover with a crossover rate of 0.95; real-value mutation with a mutation rate of 0.01; a population of 200 chromosomes; and 500 iterations of the GA [14]. These parameters were used to obtain all of the rankings discussed here, with the effectiveness of a ranking of the test-set molecules being measured using the enrichment factor as follows. Assume that a database contains A active molecules, so that selecting 1% of the molecules at random would yield $0.01A$ actives, and that the top-1% of the ranking resulting from some virtual screening method (such as the GA considered here) yields a actives. Then the enrichment factor is $a/0.01A$, with a value greater than unity meaning that the virtual screening method has resulted in an increase in the number of actives when compared with the number obtained by random selection.

The GA was run ten times on each activity class in each dataset, and the ten resulting rankings in each case were then combined into a single ranking using one of the fusion rules listed in Table 1, where $S_i(m_j)$ denotes the score for the j -th molecule in the i -th of the n GA rankings. To illustrate the operation of a fusion rule, the Sum rule simply aggregates all of the n scores, and then ranks the test-set molecules in decreasing order of these sums of scores; in like vein, the Med, Max and Min rules rank the test-set molecules on the basis of the median, maximum or minimum scores. These four rules can also be applied if the scores, $S_i(m_j)$, are converted to the corresponding ranks, $R_i(m_j)$. Thus, if we take the Sum formulation there are two fusion rules: SumS and SumR for fusing the scores and the ranks, respectively; and similarly so for Med, Max and Min. However, the final rule in Table 1, the RKP (from *reciprocal rank*) rule, which was first developed to rank the outputs of text search engines [23], is applicable only to the ranks. Assume that a threshold – here the top 1% – is applied to each of the ten rankings and that a molecule m_j appears in p of these: then the molecule is scored by adding the reciprocal of its rank positions in those p rankings. There was thus a total of nine different fusion rules that could be used to combine the sets of ten GA rankings for each activity class. It is worth noting in passing that the Sum (when applied to ranks) and RKP rules involve mathematical operations on ordinal data: this is mathematically inappropriate but has been found to be effective in operation in both chemical and textual applications [18, 23] and these rules have hence been included in the present study.

Table 1. Fusion rules

Fusion Rule	Formula
Sum	$\sum_{i=1}^{10} S_i(m_j)$
Max	$\max\{S_1(m_j), \dots, S_i(m_j), \dots, S_{10}(m_j)\}$
Med	$\text{med}\{S_1(m_j), \dots, S_i(m_j), \dots, S_{10}(m_j)\}$
Min	$\min\{S_1(m_j), \dots, S_i(m_j), \dots, S_{10}(m_j)_1\}$
RKP	$\sum_{i=1}^p \frac{1}{R_i(m_j)}$

The results that were obtained for the 15 ChEMBL activity classes are shown in Table 2. For each of the 15 activity classes, we list first the mean enrichment factor when averaged over the ten GA runs that were carried out for this activity class, with each of the nine subsequent columns listing the enrichment factor obtained when the fusion rule listed at the top of the column was used to fuse the ten GA rankings. The comparable sets of results for the much smaller MDDR and WOMBAT datasets are shown in Tables 3 and 4, respectively.

Table 2. Enrichment factor of actives when using nine different fusion rules on the ChEMBL activity classes

Activity Class	GA	RKP	MaxR	MedR	SumR	MinR	MedS	SumS	MaxS	Mins
5HT1A agonists	39.7	40.2	40.5	40.5	40.7	40.2	39.5	40.5	40.6	40.5
5HT3 antagonists	55.1	59.4	56.3	57.8	58.3	53.7	53.1	57.3	56.3	55.7
5HT reuptake inhibitors	34.5	35.1	35.4	33.4	34.5	35.1	33.0	34.7	35.8	33.9
Acetylcholinesterase inhibitors	36.8	37.1	40.9	33.8	35.8	41.5	41.2	38.2	35.8	31.1
AT1 antagonists	81.4	84.2	84.2	85.3	85.3	81.1	82.1	84.2	85.3	84.2
Cyclooxygenase inhibitors	38.3	40.0	39.2	40.0	39.2	37.6	39.2	37.6	39.2	35.2
D2 antagonists	31.8	32.5	33.4	31.5	33.0	34.0	31.7	33.1	30.7	30.6
Factor Xa inhibitors	47.4	50.0	49.6	47.0	48.5	47.9	47.6	49.1	50.0	47.3
HIV protease inhibitors	64.9	68.4	67.5	65.9	67.3	66.8	63.8	67.7	66.6	65.0
Matrixmetalloprotease inhibitors	68.8	69.7	69.4	68.0	69.4	69.1	68.0	69.1	69.9	67.1
Phosphodiesterase inhibitors	40.2	41.5	44.5	39.3	41.5	42.4	39.3	42.8	40.6	45.9
Protein kinase C inhibitors	58.6	60.0	58.4	58.4	58.4	59.5	57.4	58.4	59.0	59.0
Renin inhibitors	56.6	58.7	59.4	56.5	58.3	58.4	56.1	58.9	56.9	56.3
Substance P antagonists	70.1	73.0	73.2	70.5	71.8	72.4	71.4	71.7	72.8	70.5
Thermolysin inhibitors	46.8	48.3	48.7	48.3	48.4	48.0	47.5	48.8	48.3	45.4

Table 3. Enrichment factor of actives when using nine different fusion rules on the MDDR activity classes

Activity Class	GA	RKP	MaxR	MedR	SumR	MinR	MedS	SumS	MaxS	MinS
5HT1A agonists	19.2	20.2	19.8	19.1	19.9	19.4	19.6	19.9	19.6	19.6
5HT3 antagonists	41.2	43.6	43.43	41.8	42.8	42.5	42.7	43.3	42.7	41.7
5HT reuptake inhibitors	17.9	18.9	18.6	18.3	19.2	19.2	19.5	18.9	18.9	18.0
AT1 antagonists	48.0	49.0	49.0	49.5	48.9	48.8	48.5	48.9	49.4	47.9
Cyclooxygenase inhibitors	29.3	30.2	30.6	30.8	30.8	29.6	29.0	30.2	30.8	30.9
D2 antagonists	17.4	18.3	18.0	14.3	17.1	19.4	19.7	17.4	19.4	14.0
HIV protease inhibitors	47.0	49.0	49.0	47.7	48.7	47.7	45.3	48.9	49.5	47.9
Protein kinase C inhibitors	29.0	31.1	30.9	28.4	29.4	30.4	29.7	30.6	31.4	26.7
Renin inhibitors	70.4	72.2	72.5	71.8	72.0	71.3	71.9	71.8	72.3	69.3
Substance P antagonists	29.2	30.9	30.4	28.9	30.2	30.0	28.9	30.5	30.5	28.9
Thermolysin inhibitors	48.3	49.5	49.8	49.0	49.9	49.1	47.6	49.9	49.0	48.8

Table 4. Enrichment factor of actives when using nine different fusion rules on the WOMBAT activity classes

Activity Class	GA	RKP	MaxR	MedR	SumR	MinR	MedS	SumS	MaxS	MinS
5HT1A agonists	55.7	58.7	57.6	58.5	57.4	57.0	53.10	57.4	56.5	55.7
5HT3 antagonists	43.5	44.4	44.4	43.4	44.4	43.9	43.43	44.4	46.5	43.4
Acetylcholinesterase inhibitors	51.1	51.4	52.1	50.3	51.9	51.0	53.20	51.2	49.9	54.3
AT1 antagonists	80.7	83.0	83.0	83.7	83.4	83.13	82.52	83.0	82.4	83.6
Cyclooxygenase inhibitors	67.4	67.6	67.6	67.4	67.4	67.43	66.9	67.6	67.6	67.8
D2 antagonists	40.8	41.9	41.0	42.7	41.8	40.90	42.0	41.4	41.4	41.3
Factor Xa inhibitors	45.7	47.8	45.0	41.2	43.0	47.89	43.3	44.6	40.4	45.7
HIV protease inhibitors	53.6	57.3	57.1	51.3	54.7	55.47	55.5	55.7	56.4	54.2
Matrixmetalloprotease inhibitors	63.4	64.6	65.0	64.6	64.6	63.68	63.8	64.6	65.0	62.6
Phosphodiesterase inhibitors	49.3	51.9	52.1	49.8	51.7	51.68	49.8	51.9	52.8	48.3
Protein kinase C inhibitors	74.0	72.7	73.4	71.9	71.9	71.88	72.7	71.9	75.0	71.1
Renin inhibitors	78.8	80.6	80.6	80.6	80.8	80.33	77.5	81.0	81.0	81.7
Substance P antagonists	49.2	51.6	52.6	49.8	52.8	50.80	50.8	52.2	51.0	51.4
Thermolysin inhibitors	56.1	58.8	58.6	55.2	57.8	58.58	58.8	57.8	58.1	55.2

Discussion and conclusions

Two factors are immediately obvious from inspection of the results in Table 2. First, the mean values for the ten GA runs are normally slightly less than the fused scores. With nine fusion rules and 15 activity classes the mean GA run value can be compared with a fusion value 135 times: of these comparisons, the enrichment factor for the fused run exceeded the mean-GA factor 99 times. Second, while the differences between the various fusion rules are quite small (as illustrated for the MDDR activity classes in Figure 1) there are variations in performance between the rules, and it is hence reasonable to consider which is the most generally effective.

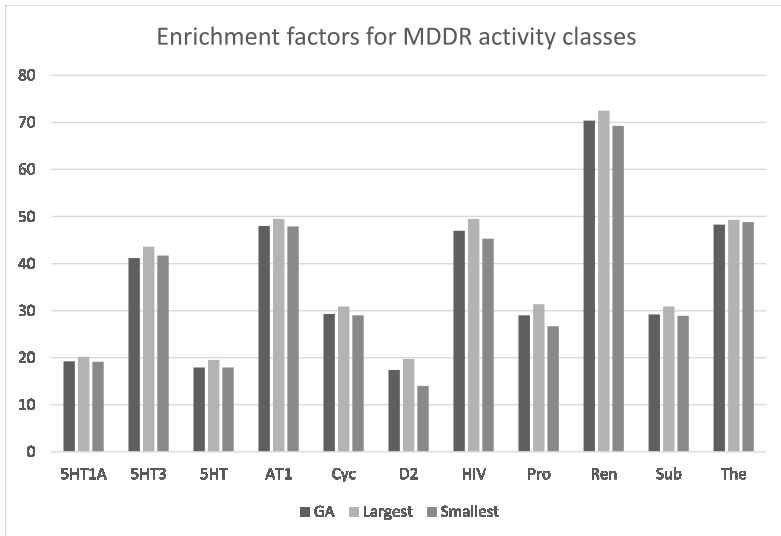


Figure 1. Variations in enrichment factor for the eleven activity classes in the MDDR dataset. For each class (as denoted by the initial characters of the class name listed in the left-hand column of Table 3) the three columns denote first the mean enrichment factor for the GA, and then the largest and the smallest enrichment factors for that class over the complete set of fusion rules.

The variations were analyzed using the Kendall Coefficient of Concordance, W , which provides a means of quantifying the degree of association between k different rankings of n different objects [24]. In the present context, $k=15$ and $n=10$ since each of the 15 activity classes enables us to rank the 9 fusion rules and the GA in decreasing order of enrichment factor. The value of W lies between zero and unity, and its significance can be checked using

the χ^2 test of statistical significance, since $\chi^2 = k(n-1)W$ with $n-1$ degrees of freedom. In the case of the ChEMBL results shown in Table 2, the computed value for W is 0.38, with an associated value for χ^2 of 51.30 that is significant ($p \leq 0.01$). If a significant value is obtained then an overall ranking of the n objects is given by their mean ranks when averaged over the k rankings as described by Siegel and Castellan [24]. Allocating scores of 9 (for the highest enrichment factor for a particular activity class) down to 0 (for the lowest such factor) then the mean scores for the ten approaches are listed in the column headed ChEMBL in Table 5, this corresponding to the following ordering of fusion rules

RKP > MaxR > SumS, MaxS > SumR > MinR > MedR > MinS > Mean GA > MedS, i.e., the RKP rule gave the best overall level of performance across the 15 activity class rankings.

Table 2 detailed the enrichment factors for the ChEMBL dataset, the largest of the three considered here. Very similar pictures of behavior were observed with the MDDR and WOMBAT datasets as shown in Tables 3 and 4, with statistically significant levels of agreement being obtained for the comparison of the various fusion rules in both cases: the resulting mean ranks are included in the appropriate columns of Table 5. The final column of this table gives the mean score for each of the ten types of ranking when taken across all three datasets, where it will be seen that the RKP rule gives the best overall results and that the mean GA results are inferior to those resulting from every one of the fusion rules.

Table 5. Kendall W analysis using nine fusion rules and mean GA scores for the MDDR, WOMBAT and ChEMBL databases

Fusion Rule	Database			Mean
	MDDR	WOMBAT	ChEMBL	Score
RKP	7.0	6.5	6.7	6.7
MaxR	6.4	6.3	6.6	6.4
MaxS	6.6	5.3	5.7	5.9
SumS	5.7	5.3	5.7	5.6
SumR	5.6	5.0	5.6	5.4
MinR	4.0	3.8	4.7	4.2
MedR	3.0	3.4	3.2	3.2
MedS	3.5	3.5	1.9	3.0
MinS	2.0	3.9	2.5	2.8
Mean GA	1.3	2.1	2.4	1.9

The finding that RKP is the most effective fusion rule here is in line with a previous study that found it was also the best for fusing the outputs of multiple similarity searches [25]. In fact, this fusion rule gives results that are sometimes comparable to those obtained with the very best individual GA run (rather than the mean-GA results considered thus far). Of the 15 ChEMBL activity classes, the best GA enrichment factor exceeded the RKP factor for seven classes while the converse applied for another seven classes (and there was one class - the AT1 antagonists - where the factors were the same). The results are less striking for the other two datasets: for MDDR, the best GA factor exceeded the RKP factor for seven classes while the converse applied for four classes; and for WOMBAT, the corresponding figures were nine and three (with two classes where the factors were the same)

In our previous paper [14], we demonstrated that a GA was able to produce dataset rankings that were noticeably superior in terms of enrichment to those resulting from established weighting schemes for substructural analysis. However, it was also demonstrated that the GA's non-deterministic nature meant that there was some degree of variation in the weights generated in multiple runs and, consequently, in the effectiveness of screening that could be expected if the approach was to be used in practice. In this short paper, we have suggested the use of data fusion as a way of combining the rankings resulting from multiple runs of the GA. The resulting, fused rankings are consistently better than the average effectiveness of screening (as measured by the enrichment factor) and in some cases are comparable with the maximum GA effectiveness. The simplicity and the effectiveness of this joint procedure hence suggest itself as a useful addition to existing methods for LBVS.

That said, there are two inherent limitations to the use of data fusion. The first is the need to specify the nature of the fusion rule that is used to combine the various rankings of the molecules comprising a database. The second is the identification of suitable weights and their assignment to each ranking as a means of specifying the relative importance or effectiveness of each individual measure. It is, however, common to assign each measure the same weight, as was the case in the work reported here. This uniform weighting approach can be enhanced by the use of machine learning techniques but these require extensive training data that is unlikely to be available during the early stages of a drug discovery programme when LBVS is most valuable [18]. An alternative approach that does not suffer from these limitations involves the use of partial ordering methods, as exemplified by the Hasse diagram. This provides a simple way of ordering objects where multiple criteria can be used without the need to specify a fusion rule or to assign weights to the rankings that are being combined, and has already found some application in chemoinformatics [26-28]. There is, however, the problem that they

are only suitable for use where small numbers of objects are involved, whereas applications of the sort considered here involve databases containing hundreds of thousands or millions of molecules. The development of efficient algorithms for the generation of Hasse diagrams could hence enhance still further the attractiveness of the GA approach to LBVS.

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