Frequent Sub-Structure-Based Approaches for Classifying Chemical Compounds*

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Abstract

In this paper we study the problem of classifying chemical compound datasets. We present a sub-structure-based classification algorithm that decouples the sub-structure discovery process from the classification model construction and uses frequent subgraph discovery algorithms to find all topological and geometric sub-structures present in the dataset. The advantage of our approach is that during classification model construction, all relevant sub-structures are available allowing the classifier to intelligently select the most discriminating ones. The computational scalability is ensured by the use of highly efficient frequent subgraph discovery algorithms coupled with aggressive feature selection. Our experimental evaluation on eight different classification problems shows that our approach is computationally scalable and on the average, outperforms existing schemes by 10% to 35%.

1 Introduction

Discovering new drugs is an expensive and challenging process. Any new drug should not only produce the desired response to the disease but should do so with minimal side effects and be superior to the existing drugs in the market. One of the key steps in the drug design process is to identify the chemical compounds (widely referred to as "hit" compounds) that display the desired and reproducible behavior against the disease [29] in a biological experiment. The standard technique to discover such compounds is to evaluate them with a biological experiment, known as an assay. The 1990s saw the widespread adoption of high-throughput screening (HTS), which uses highly automated techniques to conduct the biological assays and can be used to screen a large number of compounds. Though in principle, HTS techniques can be used to test each compound against every biological assay, it is never practically feasible for the following reasons. First,

the number of chemical compounds that have been synthesized or can be synthesized using combinatorial chemistry techniques is extremely large. Evaluating this large set of compounds using HTS can be prohibitively expensive. Second, not all biological assays can be converted to high throughput format. Third, in most cases it is hard to find all the desirable properties in a single compound and chemists are interested in not just identifying the hits but studying what part of the chemical compound leads to desirable behavior, so that new compounds can be rationally synthesized.

The goal of this paper is to develop computational techniques based on classification that can be used to identify the hit compounds. These computational techniques can be used to replace or supplement the biological assay techniques. One of the key challenges in developing classification techniques for chemical compounds stems from the fact that their properties are strongly related to their chemical structure. However, traditional machine learning techniques are suited to handle datasets represented by multidimensional vectors or sequences, and cannot handle the structural nature of the chemical structures.

In recent years two classes of techniques have been developed for solving the chemical compound classification problem. The first class builds a classification model using a set of physico-chemical properties derived from the compounds structure, called quantitative structure-activity relationships (QSAR) [16, 17, 1], whereas the second class operates directly on the structure of the chemical compound and tries to automatically identify a small number of chemical sub-structures that can be used to discriminate between the different classes [3, 46, 19, 27, 8]. A number of comparative studies [43, 22] have shown that techniques based on the automatic discovery of chemical sub-structures are superior to those based on QSAR properties and require limited user intervention and domain knowledge. However, despite their success, a key limitation of these techniques is that they rely on heuristic search methods to discover these sub-structures. Even

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though such approaches reduce the inherently high computational complexity associated with these schemes, they may lead to sub-optimal classifiers in cases in which the heuristic search failed to uncover sub-structures that are critical for the classification task.

In this paper we present a sub-structure-based classifier that overcomes the limitations associated with existing algorithms. One of the key ideas of our approach is to decouple the sub-structure discovery process from the classification model construction step and use frequent subgraph discovery algorithms to find all chemical sub-structures that occur a sufficiently large number of times. Once the complete set of these sub-structures has been identified, our algorithm then proceeds to build a classification model based on them. The advantage of such an approach is that during classification model construction, all relevant sub-structures are available allowing the classifier to intelligently select the most discriminating ones. To ensure that such an approach is computationally scalable, we use recently developed [25, 27] highly efficient frequent subgraph discovery algorithms coupled with aggressive feature selection to reduce both the amount of time required to build as well as to apply the classification model. In addition, we present a sub-structure discovery algorithm that finds a set of sub-structures whose geometry is conserved, further improving the classification performance of our algorithm.

We experimentally evaluated the performance of our algorithms on eight different problems derived from three publically available datasets and compared their performance against that of traditional QSAR-based classifiers and existing sub-structure classifiers based on SUB-DUE [4] and SubdueCL [15]. Our results show that our approach, on the average, outperforms QSAR-based schemes by 35% and SUBDUE-based schemes by 10%.

The rest of the paper is organized as follows. Section 2 provides some background information related to chemical compounds, their activity, and their representation. Section 3 provides a survey on the related research in this area. Section 4 provides the details of our chemical compound classification approach. Section 5 experimentally evaluates its performance and compares it against other approaches. Finally, Section 6 provides outlines directions of future research and provides some concluding remarks.

2 Background

A chemical compound consists of different atoms being held together via bonds adopting a well-defined geometric configuration. Figure 1(a) represents the chemical compound Flucytosine from the DTP AIDS repository [10] it consists of a central aromatic ring and other elements like N, O and F. The representation shown in the figure is a typical graphical representation that most chemists work with.

Figure 1: Chemical and Graphical representation of Flucytosine

There are many different ways to represent such chemical compounds. The simplest representation is the molecular formula that lists the various atoms making up the compound; the molecular formula for Flucytosine is C4H4FN3O. However this representation is woefully inadequate to capture the structure of the chemical compound. It was recognized early on that it was possible for two chemical compounds to have identical molecular formula but completely different chemical properties [14]. A more sophisticated representation can be achieved using the SMILES [6] representation, it not only represents the atoms but also represents the bonds between different atoms. The SMILES representation for Flucytosine is Nc1nc (0) ncc1F. Though the SMILES representation is a compact it is not guaranteed to be unique, furthermore the representation is quite restrictive to work with [24].

The activity of a compound largely depends on its chemical structure and the arrangement of different atoms in 3D space. As a result, effective classification algorithms must be able to directly take into account the structural nature of these datasets. In this paper we represent each compound as undirected graphs. The vertices of these graphs correspond to the various atoms, and the edges correspond to the bonds between the atoms. Each of the vertices and edges has a label associated with it. The labels on the vertices correspond to the type of atoms and the labels on the edges correspond to the type of bonds. As an example, Figure 1(b) shows the representation of Flucytosine in terms of this graph model. We will refer to this representation as the topological graph representation of a chemical compound. Note that such representations are quite commonly used by many chemical modeling software and are referred as the connection table for the chemical compound [30].

In addition, since chemical compounds have a physical three-dimensional structure, each vertex of the graph has a 3D-coordinate indicating the position of the corresponding atom in 3D space. However, there are two key issues that need to be considered when working with the compound's 3D structure. First, the number of experimentally determined molecular geometries is limited (about 270,000 *X*-ray structures in the Cambridge Crystallographic Database compared to 15 million known compounds). As a result, the 3D geometry of a compound

needs to be computationally determined, which may introduce certain amount of error. To address this problem, we use the Corina [13] software package to compute the 3D coordinates for all the chemical compounds in our datasets. Corina is a rule- and data-based system that has been experimentally shown to predict the 3D structure of compounds with high-accuracy. Second, each compound can have multiple low-energy conformations (i.e., multiple 3D structures) that need to be taken into account in order to achieve the highest possible classification performance. However, due to time constraints, in this study we do not take into account these multiple conformations but instead use the single low-energy conformation that is returned by Corina's default settings. However, as discussed in Section 4.1.2, the presented approach for extracting geometric sub-structures can be easily extended to cases in which multiple conformations are considered as well. Nevertheless, despite this simplification, as our experiments in Section 5 will show, incorporating 3D structure information leads to measurable improvements in the overall classification performance. We will refer to this representation as the *geometric graph* representation of a chemical compound.

The meaning of the various classes in the input dataset is application dependent. In some applications, the classes will capture the extent to which a particular compound is toxic, whereas in other applications they may capture the extent to which a compound can inhibit (or enhance) a particular factor and/or active site. In most applications each of the compounds is assigned to only one of two classes, that are commonly referred to as the *positive* and *negative* class. The positive class corresponds to compounds that exhibit the property in question, whereas the compounds of the negative class do not. Throughout this paper we will be restricting ourselves to only two classes, though all the techniques described here can be easily extended to multi-class as well as multi-label classification problems.

Another important aspect of modeling chemical compounds is the naming of single and double bonds inside aromatic rings. Typically in an aromatic ring of a chemical compound though the number of single and double bonds is fixed, the exact position of double and single bonds is not fixed, this is because of the phenomenon of resonance [14]. It is worth noting that the exact position of double and single bond in an aromatic ring does not affect the chemical properties of a chemical compound. To capture this uncertainty in the position of single and double bond we represent all the bonds making up the aromatic ring with a new bond type called the aromatic bond. Another aspect of the chemical compounds is that the number of hydrogen bonds connected to a particular carbon atom can usually be inferred from the bonds connecting that carbon atom [14], therefore in our representation we do not represent the hydrogen atoms that are connected to the carbon atoms, such hydrogen atoms are referred as non-polar hydrogen atoms. Note that the above transformations are widely used by many chemistry modeling tools and are usually referred to as *structure normalization* [30].

3 Related Research

In the early 1960s, the pioneering work of Hansch et al. [16, 17] demonstrated that the biological activity of a chemical compound is a function of its physico-chemical properties. These physico-chemical properties are usually derived from the compound's structure and are called quantitative structure-activity relationships (QSAR). Examples of such physico-chemical properties include the molecular weight, total energy, dipole moment, solvent accessible area, etc. Over the years a number of different QSAR properties have been developed (GAUSSIAN contains over 50) and they are used extensively to model and analyze chemical compounds within the pharmaceutical industry. The amount of time required to compute these QSAR properties varies from property to property. Some of them can be computed very fast (e.g., molecular weight), while others require time-consuming numerical simulations (e.g., dipole-moment, total energy) that can only be performed for small datasets.

In QSAR-based classification methods, each chemical compound is transformed into a vector of numerical values, corresponding to the various QSAR properties. After this transformation any traditional classifier capable of handling numerical features can be used for the classification task. Early research on QSAR-based classification methods focused primarily on regression-based techniques [16, 12]; however, more sophisticated classifiers have also been used. For example, Andrea and Kalayeh [2] show that neural networks can achieve better accuracies over regression-based techniques, whereas An and Wang [1] report that decision tree classifiers applied on QSAR features outperform those based on neural networks and logistic regression.

The key challenge in using QSAR-based approaches stems from the fact that the classification performance relies, to a large extent, on the a priori identification of the relevant QSAR properties that capture the structure-activity relationships for the particular classification problem. Identifying this relevant set of QSAR properties requires considerable domain expertise and extensive experimentation. To overcome this problem, a different set of techniques have been developed that operate directly on the structure of the chemical compound and try to automatically identify a small number of chemical substructures that can be used to discriminate between the different classes.

One of the earlier approaches that follows this paradigm are based on inductive logic programming (ILP) [36]. In this approach the chemical compound is expressed using first order logic. Each atom is represented as a predicate consisting of atomID and the element, and a bond is represented as a predicate consisting of two atomIDs. Using this representation, an ILP system discovers rules (i.e., conjunction of predicates) that are good for discriminating the different classes. Since these rules consist of predicates describing atoms and bonds, they essentially correspond to sub-structures that are present in the chemical compounds. The pioneering work in this field was done by King et al. in the early 1990s [23, 22]. They applied an ILP system, Golem [37], to study the behavior of 44 trimethoprin analogues and their observed inhibition of Escherichia coli dihydrofolate reductase. They reported an improvement in classification accuracy over the traditional QSAR-based models. Srinivasan et al. [43] present a detailed comparison of the features generated by ILP with the traditional QSAR features used for classifying chemical compounds. They show that for some applications features discovered by ILP approaches lead to a significant lift in the performance. Besides improved classification performance, an additional advantage of these structure-based approaches is that the discovered rules (i.e., sub-structures) can be easily understood by experts and could be used to check the correctness of the model and to provide insights in the chemical behavior of the compounds.

Though ILP-based approaches are quite powerful, the high computational complexity of the underlying ruleinduction system limits the size of the dataset for which they can be applied. Furthermore, they tend to produce rules consisting of relatively small sub-structures (usually three to four atoms [5, 7]), limiting the size of structural constraints that are being discovered and hence affecting the classification performance. Another drawback of these approaches is that in order to reduce their computational complexity they employ various heuristics to prune the explored search-space [35], potentially missing sub-structures that are important for the classification task. One exception is the WARMR system [5, 7] that is specifically developed for chemical compounds and discovers all possible sub-structures above a certain frequency threshold. However, WARMR's computational complexity is very high and can only be used to discover sub-substructures that occur with relatively high fre-

One of the fundamental reasons limiting the scalability of ILP-based approaches is the first order logic-based representation that they use. This representation is much more powerful than what is needed to model chemical compounds and discover sub-structures. For this reason a number of researchers have explored the much simpler graph-based representation of the chemical compound's topology and transformed the problem of finding chemical substructures to that of finding subgraphs in this graph-based representation [3, 46, 19]. Probably the most well-known approach is the SUBDUE system [18, 4]. SUBDUE finds patterns which can effectively compress the original input data based on the minimum description length (MDL)

principle, by substituting those patterns with a single vertex. To narrow the search-space and improve its computational efficiency, SUBDUE uses a heuristic beam search approach, which quite often results in failing to find subgraphs that are frequent. The SUBDUE system was also later extended to classify graphs and was referred as SubdueCL [15]. In SubdueCL instead of using minimum description length as a heuristic a measure similar to confidence of a subgraph is used as a heuristic.

Finally, another heuristics based scheme targeted for chemical compounds is MOLFEA [24]. In this scheme each chemical compound is represented as a SMILES string, and is thought of as sequence of SMILES objects. This representation simplifies the problem to discovering frequently occurring sub-sequences.

4 Classification Based on Frequent Subgraphs

The previous research on classifying chemical compounds (discussed in Section 3) has shown that techniques based on the automatic discovery of chemical sub-structures are superior to those based on QSAR properties and require limited user intervention and domain knowledge. However, despite their success, a key limitation of both the ILP- and the subgraph-based techniques, is that they rely on heuristic search methods to discover the sub-structures to be used for classification. As discussed in Section 3, even though such approaches reduce the inherently high computational complexity associated with these schemes, they may lead to sub-optimal classifiers in cases in which the heuristic search failed to uncover sub-structures that are critical for the classification task.

To overcome this problem, we developed a classification algorithm for chemical compounds that uses the graph-based representation and limits the number of substructures that are pruned a priori. The key idea of our approach is to decouple the sub-structure discovery process from the classification model construction step, and use frequent subgraph discovery algorithms to find all chemical sub-structures that occur a sufficiently large number of times. Once the complete set of such sub-structures has been identified, our algorithm then proceeds to build a classification model based on them. To a large extent, this approach is similar in spirit to the recently developed frequent-itemset-based classification algorithms [32, 31, 9] that have been shown to outperform traditional classifiers that rely on heuristic search methods to discover the classification rules.

The overall outline of our classification methodology is shown in Figure 2. It consists of three distinct steps: (i) feature generation, (ii) feature selection, and (iii) classification model construction. During the feature generation step, the chemical compounds are mined to discover the frequently occurring sub-structures that correspond to either topological or geometric subgraphs. These sub-

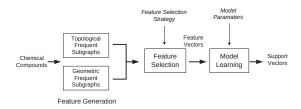


Figure 2: Frequent Subgraph Based Classification Framework

structures are then used as the features by which the compounds are represented in the subsequent steps. During the second step, a small set of features is selected such that the selected features can correctly discriminate between the different classes found in the dataset. Finally, in the last step each chemical compound is represented using these set of features and a classification model is learned. The rest of this section describes these three steps in detail.

4.1 Feature Generation

Our classification algorithm finds sub-structures in a chemical compound database using two different methods. The first method uses the topological graph representation of each compound whereas the second method is based on the corresponding geometric graph representation (discussed in Section 2). In both of these methods, our algorithm uses the topological or geometric connected subgraphs that occur in at least $\sigma\%$ of the compounds to define the sub-structures.

There are two important restrictions on the type of the sub-structures that are discovered by our approach. The first has to do with the fact that we are only interested in sub-structures that are connected and is motivated by the fact that connectivity is a natural property of such patterns. The second has to do with the fact that we are only interested in frequent sub-structures (as determined by the value of σ) as this ensures that we do not discover spurious sub-structures that will in general not be statistically significant. Furthermore, this minimum support constraint also helps in making the problem of frequent subgraph discovery computationally tractable.

4.1.1 Frequent Topological Subgraphs

Developing frequent subgraph discovery algorithms is particularly challenging and computationally intensive as graph and/or subgraph isomorphisms play a key role throughout the computations. Despite that, in recent years, four different algorithms have been developed capable of finding all frequently occurring subgraphs with reasonable computational efficiency. These are the AGM algorithm developed by Inokuchi et al [19], the FSG algorithm developed by members of our group [25], the chemical substructure discovery algorithm developed by Borgelt and

Berthold [3], and the gSpan algorithm developed by Yan and Han [46]. The enabling factors to the computational efficiency of these schemes have been (i) the development of efficient candidate subgraph generation schemes that reduce the number of times the same candidate subgraph is being generated, (ii) the use of efficient canonical labeling schemes to represent the various subgraphs; and (iii) the use of various techniques developed by the data-mining community to reduce the number of times subgraph isomorphism computations need to be performed.

In our classification algorithm we find the frequently occurring subgraphs using the FSG algorithm. FSG takes as input a database D of graphs and a minimum support σ , and finds all connected subgraphs that occur in at least σ % of the transactions. FSG, initially presented in [25], with subsequent improvements presented in [27], uses a breadth-first approach to discover the lattice of frequent subgraphs. It starts by enumerating small frequent graphs consisting of one and two edges and then proceeds to find larger subgraphs by joining previously discovered smaller frequent subgraphs. The size of these subgraphs is grown by adding one-edge-at-a-time. The lattice of frequent patterns is used to prune the set of candidate patterns and it only explicitly computes the frequency of the patterns which survived this downward closure pruning. Despite the inherent complexity of the problem, FSG employs a number of sophisticated techniques to achieve high computational performance. It uses a canonical labeling algorithm that fully makes use of edge and vertex labels for fast processing, and various vertex invariants to reduce the complexity of determining the canonical label of a graph. These canonical labels are then used to establish the identity and total order of the frequent and candidate subgraphs, a critical step of redundant candidate elimination and downward closure testing. It uses a sophisticated scheme for candidate generation [27] that minimizes the number of times each candidate subgraph gets generated and also dramatically reduces the generation of subgraphs that fail the downward closure test. Finally, for determining the actual frequency of each subgraph, FSG reduces the number of subgraph isomorphism operations by using TID-lists [11, 41, 48, 47] to keep track of the set of transactions that supported the frequent patterns discovered at the previous level of the lattice. For every candidate, FSG takes the intersection of TID-lists of its parents, and performs the subgraph isomorphism only on the transactions contained in the resulting TID-list. As the experiments presented in Section 5 show, FSG is able to scale to large datasets and low support values.

4.1.2 Frequent Geometric Subgraphs

Topological sub-structures capture the connectivity of atoms in the chemical compound but they ignore the 3D shape (3D arrangement of atoms) of the sub-structures. For certain classification problems the 3D shape of the

sub-structure might be essential for determining the chemical activity of a compound. For instance, the geometric configuration of atoms in a sub-structure is crucial for its ability to bind to a particular target [29]. For this reason we developed an algorithm that findss all frequent sub-structures whose topology as well as geometry is conserved.

There are two important aspects specific to the geometric subgraphs that need to be considered. First, since the coordinates of the vertices depend on a particular reference coordinate axes, we would like the discovered geometric subgraphs to be independent of these coordinate axes, i.e., we are interested in geometric subgraphs whose occurrences are translation, and rotation invariant. This dramatically increases the overall complexity of the geometric subgraph discovery process, because we may need to consider all possible geometric configurations of a single pattern. Second, while determining if a geometric subgraph is contained in a bigger geometric graph we would like to allow some tolerance when we establish a match between coordinates, ensuring that slight deviations in coordinates between two identical topological subgraphs do not lead to the creation of two geometric subgraphs. The amount of tolerance (r) should be a user specified parameter. The task of discovering such r-tolerant frequent geometric subgraphs dramatically changes the nature of the problem. In traditional pattern discovery problems such as finding frequent itemsets, sequential patterns, and/or frequent topological graphs there is a clear definition of what a pattern is, given its set of supporting transactions. On the other hand, in the case of r-tolerant geometric subgraphs, there are many different geometric representations of the same pattern (all of which will be r-tolerant isomorphic to each other). The problem becomes not only that of finding a pattern and its support, but also finding the right representative for this pattern. The selection of the right representative can have a serious impact on correctly computing the support of the pattern. For example, given a set of subgraphs that are r-tolerant isomorphic to each other, the one that corresponds to an outlier will tend to have a lower support than the one corresponding to the center. These two aspects of geometric subgraphs makes the task of discovering the full fledged geometric subgraphs extremely hard [26].

To overcome this problem we developed a simpler, albeit less discriminatory, representation for geometric subgraphs. We use a property of a geometric graph called the *average inter-atomic distance* that is defined as the average Euclidean distance between all pairs of atoms in the molecule. Note that the average inter-atomic distance is computed between all pairs of atoms irrespective of whether a bond connects the atoms or not. The average inter-atomic distance can be thought of as a geometric signature of a topological subgraph. The geometric subgraph consists of two components, a topological subgraph and an

interval of average inter-atomic distance associated with it. A geometric graph contains this geometric subgraph if it contains the topological subgraph and the average interatomic distance of the embedding (of the topological subgraph) is within the interval associated with the geometric subgraph. Note that this geometric representation is also translation and rotation invariant, and the width of the interval determines the tolerance displayed by the geometric subgraph. We are interested in discovering such geometric subgraphs that occur above $\sigma\%$ of the transactions and the interval of average inter-atomic distance is bound by r.

Since a geometric subgraph contains a topological subgraph, for the geometric subgraph to be frequent the corresponding topological subgraph has to be frequent, as well. This allows us to take advantage of the existing approach to discover topological subgraphs. We modify the frequency counting stage of the FSG algorithm as follows. If a subgraph g is contained in a transaction t then all possible embeddings of g in t are found and the average interatomic distance for each of these embeddings is computed. This list of average inter-atomic distances is added to the g.aiadList such that at the end of the frequent subgraph discovery each topological subgraph has a list of average inter-atomic distances associated with it. Each one of the average inter-atomic distances corresponds to one of the embeddings i.e., a geometric configuration of the topological subgraph. This algorithm can be easily extended to cases in which there are multiple 3D conformations associated with each chemical compound (as discussed in Section 2), by simply treating each distinct conformation as a different chemical compound.

The task of discovering geometric subgraphs now reduces to identifying those geometric configurations that are frequent enough, *i.e.*, to identify intervals of average inter-atomic distance such that each interval contains the minimum number geometric configurations (σ) and the width of the interval is less than the tolerance threshold (r). This task can be thought of as 1D clustering on the vector of average inter-atomic distances such that each cluster contains items above the minimum support and the spread of each cluster is bounded by the tolerance r. Note that not all items will belong to a valid cluster as some of them will be infrequent. In our experiments we set the value of r to be equal to half of the minimum distance between any two pairs of atoms in the compounds.

To find such clusters we perform agglomerative clustering on the vector of average inter-atomic distance values. The distance between any two average inter-atomic distance values is defined as the difference in their numeric values. To ensure that we get the largest possible clusters we use the maximum-link criterion function for deciding which two clusters should be merged [21]. The process of agglomeration is continued until the interval containing all the items in the cluster is below the tolerance threshold (r). When we reach a stage where further agglomeration

would increase the spread of the cluster beyond the tolerance threshold, we check the number of items contained in the cluster. If the number of items is above the support threshold, then the interval associated with this cluster is considered as a geometric feature. Since we are clustering one-dimensional datasets, the clustering complexity is low. Some examples of the distribution of the average inter-atomic distance values and the associated clusters are shown in Figure 3. Note that the average inter-atomic distance values of the third example are uniformly spread and lead to no geometric subgraph.

Note that this algorithm for computing geometric subgraphs is approximate in nature for two reasons. First, the average inter-atomic distance may map two different geometric subgraphs to the same average inter-atomic distance value. Second, the clustering algorithm may not find the complete set of geometric subgraphs that satisfy the r tolerance. Nevertheless, as our experiments in Section 5 show the geometric subgraphs discovered by this approach improve the classification accuracy of the algorithm.

4.1.3 Additional Considerations

Even though FSG provides the general functionality required to find all frequently occurring sub-structures in chemical datasets, there are a number of issues that need to be addressed before it can be applied as a black-box tool for feature discovery in the context of classification. One issue deals with selecting the right value for the σ , the support constraint used for discovering frequent substructures. The value of σ controls the number of subgraphs discovered by FSG. Choosing a good value of σ is especially important for the dataset containing classes of significantly different sizes. In such cases, in order to ensure that FSG is able to find features that are meaningful for all the classes, it must use a support that depends on the size of the smaller class.

For this reason we first partition the complete dataset, using the class label of the examples, into specific class specific datasets. We then run FSG on each of these *class datasets*. This partitioning of the dataset ensures that sufficient subgraphs are discovered for those class labels which occur rarely in the dataset. Next, we combine subgraphs discovered from each of the *class dataset*. After this step each subgraph has a vector that contains the frequency with which it occurs in each class.

4.2 Feature Selection

The frequent subgraph discovery algorithms described in Section 4.1 discovers all the sub-structures (topological or geometric) that occur above a certain support constraint (σ) in the dataset. Though the discovery algorithm is computationally efficient, the algorithm can generate a large number of features. A large number of features is detrimental for two reasons. First, it could increase the time required to build the model. But more importantly, a

large number of features can increase the time required to classify a chemical compound, as we need to first identify which of the discovered features it contains before we can apply the classification model. Determining whether a compound contains a particular feature or not can be computationally expensive as it may require a subgraph isomorphism operation. This problem is especially critical in the drug discovery process where the classification model is learned on a small set of chemical compounds and it is then applied on large chemical compound libraries containing millions of compounds.

One way of solving this problem is to follow a heuristic subgraph discovery approach (similar in spirit to previously developed methods [4, 15]) in which during the subgraph discovery phase itself, the discriminatory ability of a particular subgraph is determined, and the discovery process is terminated as soon as a subgraph is generated that is less discriminatory than any of its subgraphs. By following this approach, then the total number of features will be substantially reduced, achieving the desired objective. However, the limitation with such an approach is that it may fail to discover and use highly discriminatory subgraphs. This is because the discriminatory ability of a subgraph does not (in general) consistently increase as a function of its size, and subgraphs that appear to be poor discriminators may become very discriminatory by growing their size. For this reason, in order to develop an effective feature selection method, we use a scheme that first finds all frequent subgraphs and then selects among them a small set of discriminatory features. The advantage of this approach is that during feature selection all frequent subgraphs are considered irrespective of when they were generated and whether or not they contain less or more discriminatory subgraphs.

The feature selection scheme is based on the sequential covering paradigm used to learn rule sets [33]. To apply this algorithm we assume that each discovered substructure corresponds to a rule, with the class label of the sub-structure as the target attribute, such rules are referred as class-rules in [32]. The sequential covering algorithm takes as input a set of examples and the features discovered from these examples, and iteratively applies the feature selection step. In this step the algorithm selects the feature that has the highest estimated accuracy. After selecting this feature all the examples containing this feature are eliminated and the feature is marked as selected. In the next iteration of the algorithm the same step is applied, but on a smaller set of examples. The algorithm continues in an iterative fashion until either all the features are selected or all the examples are eliminated.

In this paper we use a computationally efficient implementation of sequential covering algorithm known as CBA [32], this algorithm proceeds by first sorting the features based on confidence and then applying the sequential covering algorithm on this sorted set of features. One of

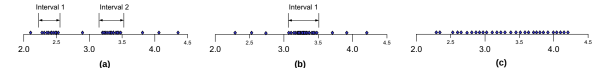


Figure 3: Some examples of the one-dimensional clustering of average inter-atomic distance values.

the advantages of this approach is that it requires minimal number of passes on the dataset, hence is very scalable. To obtain a better control over the number of selected features we use an extension of the sequential covering scheme known as *Classification based on Multiple Rules* (CMAR) [31]. In this scheme instead of removing the example after it is covered by the selected feature, the example is removed only if that example is covered by δ selected features. The number of selected rules increases as the value of δ increases, an increase in the number of features usually translates into an improvement in the accuracy as more features are used to classify a particular example. The value of δ is specified by the user and provides a means to the user to control the number of features used for classification.

4.3 Classification Model Construction

Given the frequent subgraphs discovered in the previous step, our algorithm treats each of these subgraphs as a feature and represents the chemical compound as a frequency vector. The ith entry of this vector is equal to the number of times (frequency) that feature occurs in the compound's graph. This mapping into the feature space of frequent subgraphs is performed both for the training and the test dataset. Note that the frequent subgraphs were identified by mining only the graphs of the chemical compounds in the training set. However, the mapping of the test set requires that we check each frequent subgraph against the graph of the test compound using subgraph isomorphism. Fortunately, the overall process can be substantially speeded up by taking into account the frequent subgraph lattice that is also generated by FSG. In this case, we traverse the lattice from top to bottom and only visit the child nodes of a subgraph if that subgraph is isomorphic to the chemical compound.

Once the feature vectors for each chemical compound have been built, any one of the existing classification algorithms can potentially be used for classification. However, the characteristics of the transformed dataset and the nature of the classification problem itself tends to limit the applicability of certain classes of classification algorithms. In particular, the transformed dataset will most likely be high dimensional, and second, it will be sparse, in the sense that each compound will have only a few of these features, and each feature will be present in only a few of the compounds. Moreover, in most cases the positive class will be much smaller than the negative class, making it unsuitable for classifiers that primarily focus on optimizing

the overall classification accuracy.

In our study we built the classification models using support vector machines (SVM) [44], as they are well-suited for operating in such sparse and high-dimensional datasets. Furthermore, an additional advantage of SVM is that it allows us to directly control the cost associated with the missclassification of examples from the different classes [34]. This allow us to associate a higher cost for the missclassification of positive instances; thus, biasing the classifier to learn a model that tries to increase the true-positive rate, at the expense of increasing the false positive rate.

5 Experimental Evaluation

We experimentally evaluated the performance of our classification algorithm and compared it against that achieved by earlier approaches on a variety of chemical compound datasets. The datasets, experimental methodology, and results are described in subsequent sections.

5.1 Datasets

We used three different publicly available datasets to derive a total of eight different classification problems. The first dataset was initially used as a part of the Predictive Toxicology Evaluation Challenge [42] which was organized as a part of PKDD/ECML 2001 Conference¹. It contains data published by the U.S. National Institute for Environmental Health Sciences, the data consists of bioassays of different chemical compounds on rodents to study the carcinogenicity (cancer inducing) properties of the compounds [42]. The goal being to estimate the carcinogenicity of different compounds on humans. Each compound is evaluated on four kinds of laboratory animals (male Mice, female Mice, male Rats, female Rats), and is assigned four class labels each indicating the toxicity of the compound for that animal. There are four classification problems one corresponding to each of the rodents and will be referred as P1, P2, P3, and P4.

The second dataset is obtained from the National Cancer Institute's DTP AIDS Anti-viral Screen program [10, 24] ². Each compound in the dataset is evaluated for evidence of anti-HIV activity. The screen utilizes a soluble formazan assay to measure protection of human CEM cells from HIV-1 infection [45]. Compounds able to provide at least 50% protection to the CEM cells were re-

¹http://www.informatik.uni-freiburg.de/~ml/ptc/.

²http://dtp.nci.nih.gov/docs/aids/aids_data.html.

tested. Compounds that provided at least 50% protection on retest were listed as moderately active (CM, confirmed moderately active). Compounds that reproducibly provided 100% protection were listed as confirmed active (CA). Compounds neither active nor moderately active were listed as confirmed inactive (CI). We have formulated three classification problems on this dataset, in the first problem we consider only *confirmed active* (CA) and moderately active (CM) compounds and then build a classifier to separate these two compounds; this problem is referred as H1. For the second problem we combine moderately active (CM) and confirmed active (CA) compounds to form one set of active compounds, we then build a classifier to separate these active and confirmed inactive compounds; this problem is referred as H2. In the last problem we only use confirmed active (CA) and confirmed inactive compounds and build a classifier to categorize these two compounds; this problem is referred as H3.

The third dataset was obtained from the Center of Computational Drug Discovery's anthrax project at the University of Oxford [40]. The goal of this project was to discover small molecules that would bind with the heptameric protective antigen component of the anthrax toxin, and prevent it from spreading its toxic effects. A library of small sized chemical compounds was screened to identify a set of chemical compounds that could bind with the anthrax toxin. The screening was done by computing the binding free energy for each compound using numerical simulations. The screen identified a set of 12,376 compounds that could potentially bind to the anthrax toxin and a set of 22,460 compounds that were unlikely to bind to the chemical compound. The average number of vertices in this dataset is 25 and the average number of edges is also 25. The classification problem for this dataset was given a chemical compound classify it in to one of these two classes, i.e, will the compound bind the anthrax toxin or not. This classification problem is referred as A1.

	Toxic.	Aids	Anthrax	Class Dist. (% +ve class	;)
N	417	42,687	34,836	Toxicology	
\bar{N}_A	25	46	25	P1: Male Mice	38.3%
\bar{N}_B	26	48	25	P2: Female Mice	40.9%
\bar{L}_A	40	82	25	P3: Male Rats	44.2%
\bar{L}_B	4	4	4	P4: Female Rats	34.4%
$\max N_A$	106	438	41	AIDS	
$\min N_A$	2	2	12	H1: CA/CM	28.1%
$\max N_B$	1	276	44	H2: (CA+CM)/CI	3.5%
$\min N_B$	85	1	12	H3: CA/CI	1.0%
				Anthrax	
				A1: active/inactive	35%

Table 1: The characteristics of the various datasets. N is the number of compounds in the database. \bar{N}_A and \bar{N}_B are the average number of atoms and bonds in each compound. \bar{L}_A and \bar{L}_B are the average number of atomand bond-types in each dataset. $\max N_A/\min N_A$ and $\max N_B/\min N_B$ are the maximum/minimum number of atoms and bonds over all the compounds in each dataset.

Some important characteristics of these datasets are

summarized in Table 1. The right hand side of the table displays the class distribution for different classification problems, for each problem the table displays the percentage of positive class found in the dataset for that classification problem. Note that both the DTP-AIDS and the Anthrax datasets are quite large containing 42,687 and 34,836 compounds, respectively. Moreover, in the case of DTP-AIDS, each compound is also quite large having on an average 46 atoms and 48 bonds.

5.2 Experimental Methodology & Metrics

The classifications results were obtained by performing 5-way cross validation on the dataset, ensuring that the class distribution in each fold is identical to the original dataset. For the SVM classifier we used SVMLight library [20]. All the experiments were conducted on a 1500MHz Athlon MP processor having a 2GB of memory.

Since the size of the positive class is significantly smaller than the negative class, using *accuracy* to judge a classifier would be incorrect. To get a better understanding of the classifier performance for different cost settings we obtain the ROC curve [38] for each classifier. ROC curve plots the false positive rate (*X*-axis) versus the true positive rate (*Y*-axis) of a classifier; it displays the performance of the classifier without regard to class distribution or error cost. Two classifiers are compared in terms of the area under their respective ROC curves, a larger area under ROC curve indicating better performance. The area under the ROC curve will be referred by the parameter *A*.

5.3 Results

Varying Minimum Support The key parameter of the proposed frequent sub-structure-based classification algorithm is the choice of the minimum support (σ) used to discover the frequent sub-structures (either topological or geometric). To evaluate the sensitivity of the algorithm on this parameter we performed a set of experiments in which we varied σ from 10% to 20% in 5% increments. The results of these experiments are shown in the left sub-table of Table 2 for both topological and geometric sub-structures.

From Table 2 we observe that as we increase σ , the classification performance for most datasets tends to degrade. However, in most cases this degradation is gradual and correlates well with the decrease on the number of sub-structures that were discovered by the frequent sub-graph discovery algorithms. The only exception is the H2 problem for which the classification performance (as measured by ROC) degrades substantially as we increase the minimum support from 10% to 20%. Specifically, in the case of topological subgraphs, the performance drops from 70.1 down to 59.0, and in the case of geometric subgraphs it drops from 76.0 to 58.1.

These results suggest that lower values of support are

Dset.	$\sigma=10$	0.0%	$\sigma = I$	15.0%	$\sigma = 20.0\%$		Optimized σ			
	Торо.	Geom.	Торо.	Geom.	Торо.	Geom.	Торо.	Geom.	Per class	Time _p
	A N_f	$A N_f$	A N_f	A N_f	A N_f	A N_f	$A N_f$	$A N_f$	σ	(sec)
P1	66.0 1211	65.5 1317	66.0 513	64.1 478	64.4 254	60.2 268	65.5 24510	65.0 23612	3.0, 3.0	211
P2	65.0 967	64.0 1165	65.1 380	63.3 395	64.2 217	63.1 235	67.3 7875	69.9 12673	3.0, 3.0	72
P3	60.5 597	60.7 808	59.4 248	61.3 302	59.9 168	60.9 204	62.6 7504	64.8 10857	3.0, 3.0	66
P4	54.3 275	55.4 394	56.2 173	57.4 240	57.3 84	58.3 104	63.4 25790	63.7 31402	3.0, 3.0	231
H1	81.0 27034	82.1 29554	77.4 13531	79.2 8247	78.4 7479	79.5 7700	81.0 27034	82.1 29554	10.0, 10.0	137
H2	70.1 1797	76.0 3739	63.6 307	62.2 953	59.0 139	58.1 493	76.5 18542	79.1 29024	10.0, 5.0	1016
H3	83.9 27019	89.5 30525	83.6 13557	88.8 11240	84.6 7482	87.7 7494	83.9 27019	89.5 30525	10.0, 10.0	392
A1	78.2 576	79.0 492	78.2 484	77.6 332	77.1 312	76.1 193	81.7 3054	82.6 3186	5.0, 3.0	145

Table 2: Varying minimum support threshold (σ). "A" denotes the area under the ROC curve and " N_f " denotes the number of discovered frequent subgraphs.

in general better as they lead to better classification performance. However, as the support decreases, the number of discovered sub-structures and the amount of time required also increases. Thus, depending on the dataset, some experimentation may be required to select the proper values of support that balances these conflicting requirements (*i.e.*, low support but reasonable number of substructures).

In our study we performed such experimentation. For each dataset we kept on decreasing the value of support down to the point after which the number of features that were generated was too large to be efficiently processed by the SVM library. The resulting support values, number of features, and associated classification performance are shown in the right sub-table of Table 2 under the table header "Optimized σ ". Note that for each problem two different support values are displayed corresponding to the supports that were used to mine the positive and negative class, respectively. Also, the last column shows the amount of time required by FSG to find the frequent subgraphs and provides a good indication of the computational complexity at the feature discovery phase of our classification algorithm.

Comparing the ROC values obtained in these experiments with those obtained for $\sigma=10\%$, we can see that as before, the lower support values tend to improve the results, with measurable improvements for problems in which the number of discovered sub-structures increased substantially. In the rest of our experimental evaluation we will be using the frequent subgraphs that were generated using these values of support.

Varying Misclassification Costs Since for each classification problem instance the number of positive examples is in general much smaller than the number of negative examples, we performed a set of experiments in which the misclassification cost associated with each positive example was increased to match the number of negative examples. That is, if n^+ and n^- is the number of positive and negative examples, respectively, the misclassification cost β was set equal to (n^-/n^+-1) (so that $n^-=\beta n^+$). We refer to this value of β as the "EqCost" value. The classification performance achieved by our algorithm using either topological or geometric subgraphs for $\beta=1.0$ and $\beta=EqCost$ is shown in Table 3. Note

that the $\beta = 1.0$ results are the same with those presented in the right subtable of Table 2.

Dataset	,	Торо	Geom		
	$\beta = 1.0$	$\beta = EqCost$	$\beta = 1.0$	$\beta = EqCost$	
P1	65.5	65.3	65.0	66.7	
P2	67.3	66.8	69.9	69.2	
P3	62.6	62.6	64.8	64.6	
P4	63.4	65.2	63.7	66.1	
H1	81.0	79.2	82.1	81.1	
H2	76.5	79.4	79.1	81.9	
H3	83.9	90.8	89.5	94.0	
A1	81.7	82.1	82.6	83.0	

Table 3: The area under the ROC curve obtained by varying the misclassification cost. " $\beta=1.0$ " indicates the experiments in which each positive and negative example had a weight of one, and " $\beta=\text{EqCost}$ " indicates the experiments in which the misclassification cost of the positive examples was increased to match the number of negative examples.

From the results in this table we can see that, in general, increasing the misclassification cost so that it balances the size of positive and negative class tends to improve the classification accuracy. When $\beta = EqCost$, the classification performance improves for four and five problems for the topological and geometric subgraphs, respectively. Moreover, in the cases in which the performance decreased, that decrease was quite small, whereas the improvements achieved for some problem instances (e.g., P4, H1, and H2) was significant. In the rest of our experiments we will focus only on the results obtained by setting $\beta = EqCost$.

Feature Selection We evaluated the performance of the feature selection scheme based on sequential covering (described in Section 4.2) by performing a set of experiments in which we varied the parameter δ that controls the number of times an example must be covered by a feature, before it is removed from the set of yet to be covered examples. Table 4 displays the results of these experiments. The results under the column labeled "Original" shows the performance of the classifier without any feature selection. These results are identical to those shown in Table 3 for $\beta = EqCost$ and are included here to make comparisons easier.

Two key observations can be made by studying the results in this table. First, as expected, the feature selec-

Topologica	Topological Features									
Dataset.	Ori	ginal	$\delta = 1$		$\delta = 5$		$\delta = 10$		$\delta = 15$	
	A	N_f	A	N_f	A	N_f	A	N_f	A	N_f
P1	65.3	24510	65.4	143	66.4	85	66.5	598	66.7	811
P2	66.8	7875	69.5	160	69.6	436	68.0	718	67.5	927
P3	62.6	7504	68.0	171	65.2	455	64.2	730	64.5	948
P4	65.2	25790	66.3	156	66.0	379	64.5	580	64.1	775
H1	79.2	27034	78.4	108	79.2	345	79.1	571	79.5	796
H2	79.4	18542	77.1	370	78.0	1197	78.5	1904	78.5	2460
H3	90.8	27019	88.4	111	89.6	377	90.0	638	90.5	869
A1	82.1	3054	80.6	620	81.4	1395	81.5	1798	81.8	2065
Geometric	Feature	S								
Dataset.	Ori	ginal	$\delta = 1$		δ =	= 5	δ =	= 10		= 15
	A	N_f	A	N_f	A	N_f	A	N_f	A	N_f
P1	66.7	23612	68.3	161	68.1	381	67.4	613	68.7	267
P2	69.2	12673	72.2	169	73.9	398	73.1	646	73.0	265
P3	64.6	10857	71.1	175	70.0	456	71.0	241	66.7	951
P4	66.1	31402	68.8	164	69.7	220	67.4	609	66.2	819
H1	81.1	29554	80.8	128	81.6	396	81.9	650	82.1	885
H2	81.9	29024	80.0	525	80.4	1523	80.6	2467	81.2	3249
H3	94.0	30525	91.3	177	92.2	496	93.1	831	93.2	1119
A1	83.0	3186	81.0	631	82.0	1411	82.4	1827	82.7	2106

Table 4: Results obtained using feature selection based on sequential rule covering. " δ " specifies the number of times each example needs to be covered before it is removed, "A" denotes the area under the ROC curve and " N_f " denotes the number of features that were used for classification.

tion scheme is able to substantially reduce the number of features. In some cases the number of features that was selected decreased by almost two orders of magnitude. Also, as δ increases, the number of retained features increases; however, this increase is gradual. Second, the overall classification performance achieved by the feature selection scheme when $\delta \geq 5$ is quite comparable to that achieved with no feature selection. The actual performance depends on the problem instance and whether or not we use topological or geometric subgraphs. In particular, for the first four problems (P1, P2, P3, and P4) derived from the PTC dataset, the performance actually improves with feature selection. Such improvements are possible even in the context of SVM-based classifiers as models learned on lower dimensional spaces will tend to have better generalization ability [9]. Also note that for some datasets the number of features decreases as δ increases. This is because the features that were selected have higher average support.

Topological versus Geometric Subgraphs various results shown in Tables 2-4 also provide an indication on the relative performance of topological versus geometric subgraphs. In almost all cases, the classifier that is based on geometric subgraphs outperforms that based on topological subgraphs. For some problems, the performance advantage is marginal whereas for other problems, geometric subgraphs lead to measurable improvements in the area under the ROC curve. For example, if we consider the results shown in Table 3 for $\beta = EqCost$, we can see the geometric subgraphs lead to improvements that are at least 3% or higher for P2, P3, and H3, and the average improvement over all eight problems is 2.6%. As discussed in Section 4.1.2, these performance gains are due to the fact that conserved geometric structure is a better indicator of a chemical compounds activity than just its topology.

Property	Dim.	Property	Dim.
Solvent accessible area	\mathring{A}^2	Moment of Inertia	none
Total accessible area	\mathring{A}^2	Total energy	kcal/mol
Total accessible volume	\mathring{A}^3	Bend energy	kcal/mol
Total Van der Waal's area	\mathring{A}^2	Hbond energy	kcal/mol
Total Van der Waal's volume	\mathring{A}^3	Stretch energy	kcal/mol
Dipole moment	Debye	Nonbond energy	kcal/mol
Dipole moment comp. (X, Y, Z)	Debye	Estatic energy	kcal/mol
Heat of formation	Debye	Torsion energy	kcal/mol
Multiplicity	Kcal	Quantum total charge	eV

Table 5: QSAR Properties.

5.4 Comparison with Other Approaches

We compared the performance of our classification algorithm against the performance achieved by the QSAR-based approach and the approach that uses the SUBDUE system to discover a set of sub-structures.

Comparison with QSAR As discussed in Section 3 there is a wide variety of QSAR properties each of which captures certain aspects of a compounds chemical activity. For our study, we have chosen a set of 18 QSAR properties that are good descriptors of the chemical activity of a compound and most of them have been previously used for classification purposes [1]. A brief description of these properties are shown in Table 5. We used two programs to compute these attributes, the geometric attributes like solvent accessible area, total accessible area/vol, total Van der Waal's accessible area/vol were computed using the program SASA [28], the remaining attributes were computed using Hyperchem software.

We used two different algorithms to build classification models based on these QSAR properties. The first is the C4.5 decision tree algorithm [39] that has been shown to produce good models for chemical compound classification based on QSAR properties [1], and the second is the SVM algorithm that was used to build the classification models in our frequent sub-structure-based approach.

Since the range of values of the different QSAR properties can be significantly different, we first scaled them to be in the range of [0, 1] prior to building the SVM model. We found that this scaling resulted in some improvements in the overall classification results. Note that C4.5 is not affected by such scaling.

Datase	et -	SVM	C4.	5	Freq. Sub. Prec.		
		A	Precision	Recall	Торо	Geom	
P	1	60.2	0.4366	0.1419	0.6972	0.6348	
P	2	59.3	0.3603	0.0938	0.8913	0.8923	
P	3	55.0	0.6627	0.1275	0.7420	0.7427	
P	4	45.4	0.2045	0.0547	0.6750	0.8800	
Н	1	64.5	0.5759	0.1375	0.7347	0.7316	
H	2	47.3	0.6282	0.4071	0.7960	0.7711	
H	3	61.7	0.5677	0.2722	0.7827	0.7630	
A	1	49.4	0.5564	0.3816	0.7676	0.7798	

Table 6: Performance of the QSAR-based Classifier.

Table 6 shows the results obtained by the QSAR-based methods for the different datasets. The values shown for SVM correspond to the area under the ROC curve and can be directly compared with the corresponding values obtained by our approaches (Tables 2–4). Unfortunately, since C4.5 does not produce a ranking of the training set based on its likelihood of being in the positive class, it is quite hard to obtain the ROC curve. For this reason, the values shown for C4.5 correspond to the precision and recall of the positive class for the different datasets. Also, to make the comparisons between C4.5 and our approach easier, we also computed the precision of our classifier at the same value of recall as that achieved by C4.5. These results are shown under the columns labeled "Freq. Sub. Prec." for both topological and geometric features and were obtained from the results shown in Table 3 for $\beta = EqCost$. Note that the QSAR results for both SVM and C4.5 were obtained using the same costsensitive learning approach.

Comparing both the SVM-based ROC results and the precision/recall values of C4.5 we can see that our approach substantially outperforms the QSAR-based classifier. In particular, our topological subgraph based algorithm does 35% better compared to SVM-based QSAR and 72% better in terms of the C4.5 precision at the same recall values. Similar results hold for the geometric subgraph based algorithm. These results are consistent with those observed by other researchers [43, 22] that showed that sub-structure based approaches outperform those based on QSAR properties.

Comparison with SUBDUE & SubdueCL Finally, to evaluate the advantage of using the complete set of frequent sub-structures over existing schemes that are based on heuristic sub-structure discovery, we performed a series of experiments in which we used the SUBDUE system to find the sub-structures and then used them for classification. Specifically, we performed two sets of experiments. In the first set, we obtain a set of sub-structures using the

Dataset		Subd	ue.		SubdueCL.			
	A	N_f	$Time_p$	A	N_f	$Time_p$		
P1	61.9	1288	303sec	63.5	2103	301sec		
P2	64.2	1374	310sec	63.3	2745	339sec		
P3	57.4	1291	310sec	59.6	1772	301sec		
P4	58.5	1248	310sec	60.8	2678	324sec		
H1	74.2	1450	1,608sec	73.8	960	1002sec		
H2	58.5	901	232,006sec	65.2	2999	476,426sec		
Н3	71.3	905	178,343sec	77.5	2151	440,416sec		
A1	75.3	983	56,056sec	75.9	1094	31,177sec		

Table 7: Performance of the SUBDUE and SubdueCL-based approaches.

standard MDL-based heuristic sub-structure discovery approach of SUBDUE [18]. In the second set, we used the sub-structures discovered by the more recent SubdueCL algorithm [15] that guides the heuristic beam search using a scheme that measures how well a subgraph describes the positive examples in the dataset without describing the negative examples.

Even though there are a number of parameters controlling SUBDUE's heuristic search algorithm, the most critical among them are the width of the beam search, the maximum size of the discovered subgraph, and the total number of subgraphs to be discovered. In our experiments, we spent a considerable amount of time experimenting with these parameters to ensure that SUBDUE was able to find a reasonable number of sub-structures. Specifically, we changed the width of the beam search from 4 to 50 and set the other two parameters to high numeric values. Note that in the case of the SubdueCL, in order to ensure that the subgraphs were discovered that described all the positive examples, the subgraph discovery process was repeated by increasing the value of beam-width at each iteration and removing the positive examples that were covered by subgraphs.

Table 7 shows the performance achieved by SUBDUE and SubdueCL on the eight different classification problems along with the number of subgraphs that it generated and the amount of time that it required to find these subgraphs. These results were obtained by using the subgraphs discovered by either SUBDUE or SubdueCL as features in an SVM-based classification model. Essentially, our SUBDUE and SubdueCL classifiers have the same structure as our frequent subgraph-based classifiers with the only difference being that the features now correspond to the subgraphs discovered by SUBDUE and SubdueCL. Moreover, to make the comparisons as fair as possible we used $\beta = EqCost$ as the misclassification cost. We also performed another set of experiments in which we used the rule-based classifier produced by SubdueCL. The results of this scheme was inferior to those produced by the SVM-based approach and we are not reporting them

Comparing SUBDUE against SubdueCL we can see that the latter achieves better classification performance, consistent with the observations made by other researchers [15]. Comparing the SUBDUE and SubdueCL-

based results with those obtained by our approach (Tables 2–4) we can see that in almost all cases both our topological and geometric frequent subgraph-based algorithms lead to substantially better performance. This is true both in the cases in which we performed no feature selection as well as in the cases in which we used the sequential covering based feature selection scheme. In particular, comparing the SubdueCL results against the results shown in Table 4 without any feature selection we can see that on the average, our topological and geometric subgraph based algorithms do 9.3% and 12.2% better, respectively. Moreover, even after feature selection with $\delta = 15$ that result in a scheme that have comparable number of features as those used by SubdueCL, our algorithms are still better by 9.7% and 13.7%, respectively. Finally, if we compare the amount of time required by either SUBDUE or SubdueCL to that required by the FSG algorithm to find all frequent subgraphs (last column of Table 2) we can see that despite the fact that we are finding the complete set of frequent subgraphs our approach requires substantially less time.

6 Conclusions and Directions for Future Research

In this paper we presented a highly-effective algorithm for classifying chemical compounds based on frequent substructure discovery that can scale to large datasets. Our experimental evaluation showed that our algorithm leads to substantially better results than those obtained by existing QSAR- and sub-structure-based methods. Moreover, besides this improved classification performance, the sub-structure-based nature of this scheme provides to the chemists valuable information as to which sub-structures are most critical for the classification problem at hand. For example, Figure 4 shows the three most discriminating sub-structures for the PTC, DTP AIDS, and Anthrax datasets that were obtained by analyzing the decision hyperplane produced by the SVM classifier. A chemist can then use this information to understand the models and potentially use it to design better compounds.

The classification algorithms presented in this paper can be improved along three different directions. First, as already discussed in Section 2 our current geometric graph representation utilizes a single conformation of the chemical compound and we believe the overall classification performance can be improved by using all possible low-energy conformations. Such conformations can be obtained from existing 3D coordinate prediction software and as discussed in Section 4.1.2 can be easily incorporated in our existing framework. Second, our current feature selection algorithms only focus on whether or not a particular sub-structure is contained in a compound and they do not take into account how these fragments are distributed over different parts of the molecule. Better feature selection algorithms can be developed by taking this information into account to ensure that the entire

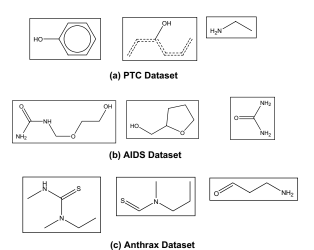


Figure 4: The three most discriminating sub-structures for the PTC, AIDS, and Anthrax datasets.

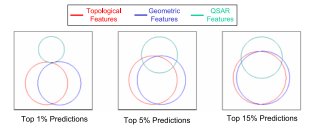


Figure 5: Venn diagrams displaying the relation between the positive examples that were correctly classified by the three approaches at different cutoff values for the Anthrax dataset. The different cutoffs were obtained by looking at only the top 1%, 5%, and 15% of the ranked predictions. Each circle in the Venn diagram corresponds to one of the three classification schemes and the size of the circle indicates the number of positive examples correctly identified. The overlap between two circles indicates the number of common correct predictions.

(or most of) molecule is covered by the selected features. Third, even though the proposed approaches significantly outperformed that based on QSAR, our analysis showed that there is a significant difference as to which compounds are correctly classified by the sub-structure- and QSARbased approaches. For example, Figure 5 shows the overlap among the different correct predictions produced by the geometric, topological, and QSAR-based methods at different cutoff values for the Anthrax dataset. From these results we can see that there is a great agreement between the substructure-based approaches but there is a large difference among the compounds that are correctly predicted by the QSAR approach, especially at the top 1% and 5%. These results suggest that better results can be potentially obtained by combining the substructure- and QSAR-based approaches.

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