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Interaction profile-based protein classification of death domain

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Abstract

**Background:** The increasing number of protein sequences and 3D structure obtained from genomic initiatives is leading many of us to focus on proteomics, and to dedicate our experimental and computational efforts on the creation and analysis of information derived from 3D structure. In particular, the high-throughput generation of protein-protein interaction data from a few organisms makes such an approach very important towards understanding the molecular recognition that make-up the entire protein-protein interaction network. Since the generation of sequences, and experimental protein-protein interactions increases faster than the 3D structure determination of protein complexes, there is tremendous interest in developing *in silico* methods that generate such structure for prediction and classification purposes. In this study we focused on classifying protein family members based on their protein-protein interaction distinctiveness. Structure-based classification of protein-protein interfaces has been described initially by Ponstingl *et al.* [1] and more recently by Valdar *et al.* [2] and Mintseris *et al.* [3], from complex structures that have been solved experimentally. However, little has been done on protein classification based on the prediction of protein-protein complexes obtained from homology modeling and docking simulation. **Results:** We have developed an *in silico* classification system entitled HODOCO (Homology modeling, Docking and Classification Oracle), in which protein Residue Potential Interaction Profiles (RPIPs) are used to summarize protein-protein interaction characteristics. This system applied to a dataset of 64 proteins of the death domain superfamily was used to classify each member into its proper subfamily. Two classification methods were attempted, heuristic and support vector machine learning. Both methods were tested with a 5-fold cross-validation. The heuristic approach yielded
a 61% average accuracy, while the machine learning approach yielded an 89% average accuracy. **Conclusion:** We have confirmed the reliability and potential value of classifying proteins *via* their predicted interactions. Our results are in the same range of accuracy as other studies that classify protein-protein interactions from 3D complex structure obtained experimentally. While our classification scheme does not take directly into account sequence information our results are in agreement with functional and sequence based classification of death domain family members.
Background

The genomic revolution has provided vast protein data resources now waiting to be transformed into usable knowledge that can be applied to solve pressing biological problems. Classification remains a favorite method for performing such transformations because of its intuitiveness and robustness against errors. Several schemes have now been proposed for automatic classification of proteins [4, 5]. They range from simple amino acid sequence comparisons, through more localized motif-based methods [6, 7 and 8] further improved by position specific scoring matrices [9] and finally to hidden Markov model profile-based methods [10]. Alternatively, structure-based classification provides a more direct means of inferring function, albeit on the much smaller structural databases [11, 12]. Recently, groups have taken an integrated approach that blends the advantages of the methods discussed above [13, 14].

The high-throughput generation of protein-protein interaction data from a few organisms has been carried out [15, 16, 17 and 18]. This wealth of experimental data requires new computational mining approaches to help us understand molecular recognition in protein-protein interaction networks. Since the generation of sequences and experimental protein-protein interactions increases faster than the 3D structure determination of protein complexes, there is tremendous interest in developing in silico methods that could predict macromolecular structures and assembly for prediction and classification purpose.

For example, computational approaches based on sequence, expression and literature abstract data have been developed to predict protein-protein interactions [19]. These methods are based on the assumption that non-homologous pairs of genes that
show correlated behavior across data from different sources should interact with each other. In addition, structure-based classification of protein-protein interfaces has been described initially by Ponstingl et al. [1] and more recently by Valdar et al. [2] and Mintseris et al. [3], from complex structures that have been solved experimentally.

The last decade has seen enormous progress in the reliability and accuracy of 3D structure-based in silico techniques including 3D structure prediction based on sequence homology and macromolecular docking. Competitions in both domains have spurred the ingenuity necessary for tackling these challenging problems [20, 21]. In this study we combine these two approaches to perform protein classification.

To efficiently dock two molecules that participate in a protein-protein or protein-ligand interaction, a certain number of steps have to be determined [22]. The process first involves an efficient search and matching algorithm that covers the conformational space, and then one or more selective scoring functions that can eliminate efficiently between native and non-native solutions. Docking algorithms are defined and classified by the extent of flexibility that they attempt to address (1) Rigid body docking, where the two molecules are rigid solid bodies, (2) Semi-flexible docking where one molecule, the receptor, is considered a rigid body while the ligand, generally smaller, is considered flexible, and finally (3) flexible docking where both molecules are considered flexible. Flexible docking is now becoming more popular because it takes into account conformational changes that generally occur when proteins interacts with each other. However, rigid body docking simulation has already been widely employed and used successfully in the docking of protein-protein complexes [22, 23]. In this method flexibility can be incorporated through a “soft belt” into which atoms from the second molecule can penetrate, reducing drastically the complexity [22] and increasing the speed
of the simulation. Rigid body docking is based on the observation that 3D protein complexes reveal a close geometric match at the interface of a receptor and a ligand. Since many false positives with better scores than the true solution are very often obtained, additional rescoring functions have been introduced to eliminate these wrong solutions [24].

In this study, rigid body docking is applied to the classification of protein-protein interactions in the death domain superfamily. We chose rigid body docking because of its higher speed. The scheme uses in silico protein-protein interaction predictions, applied to 3D protein structures built using homology modeling, as its exclusive means of performing classifications. We implemented the approach in a system called HODOCO (HOmology modeling, DOcking, and Classifying Oracle), and used Residue Pair Interaction Profiles (RPIPs) as a means to summarize protein interaction characteristics. The system was applied successfully to the problem of classifying members of the human death domain superfamily. We show that despite the limited reliability of current docking algorithm, interaction profile-based classification of this family can be obtained with 90% accuracy.

**Results and Discussion**

The goal of this study was to perform protein classification from in silico predicted protein-protein interaction. We chose to concentrate on the death domain superfamily as our model family. The superfamily consists of four families: Caspase-associated recruitment domain (CARD), death effector domain (DED), death domain (DD), and pyrin/AIM/ASC/DD-like domain (PAAD). Each domain in the superfamily has a characteristic 6-helix bundle fold called the “death fold”, and performs protein-
protein interactions between members of the same subfamily class. Interaction between superfamily members has been shown to be a functional mechanism of signal transduction in apoptosis (programmed cell death), and therefore is key to many vital life processes such as the proper maintenance of homeostasis and the immune system, as well as diseases such as neurodegenerative disorders and cancer. The characteristics of the superfamily, such as the low intrafamily sequence identity (30% on average), the high interfamily structural similarity, and the fact that family membership is primarily defined by the ability of members of the same family to interact exclusively with each other [35, 36] make it ideal for classification based on interaction profiles. The combined literature survey from the database of Protein FAMily (http://pfam.wustl.edu), and an iterative BLAST search resulted in 80 sequences, distributed moderately evenly across families. Nine had solved 3D structures (Table 1). The rest had to be modeled in silico, and those that could not be modeled to a reasonable degree of confidence were removed from the study. What remained were 64 models (i.e. \( m = 64 \)): 21 CARD, 10 DED, 21 DD, and 12 PAAD (Table 2).

Great care was taken during the model building process to ensure that the highest quality models were accepted. This degree of caution was required to avoid the risk of propagating inaccuracies throughout the system. GRAMM was chosen as the docking engine for its ability to obtain raw putative complexes that have not been further filtered, thereby allowing us to estimate a signal-to-noise ratio from the raw data and in turn allowing us to devise a procedure for reducing the search space. Docking algorithm was used to build a database that could be mined for specific complexes with properties unique to a given family. We chose to perform docking only between members of the same family as the intrafamily complexes provided a broad enough sampling to yield
high accuracy rates and limit our computational cost. Keeping in mind GRAMM's asymmetric algorithm, $21^2 + 10^2 + 21^2 + 12^2 = 1126$ docking simulations were conducted each offering 1000 putative complexes, for a total of over 1.1 million putative complexes.

In transforming the putative complexes into interaction profiles, we had to answer the question: If a pair of models forms 2000 putative complexes, how can we identify those that are most valuable for identifying family membership? Ideally, the algorithm should agree with the two known 3D protein complexes (APAF1/CASP9 and Pelle/Tube) that have been solved in the superfamily. Furthermore, we considered that a protein’s interaction signature should not only consist of its binding domains and specific binding partners, but should be characterized by a gradient of different binding specificity. Our first intuition was to mine for frequently occurring putative complexes. To visualize what we mean by this, it is possible to plot all putative complexes in a 3D environment (see Figure 4(a)). The figure shows that points represent docking hits form clusters, and that these clusters represent frequently occurring putative complexes. However, contrary to our intuition, when these clusters were compared to the two known 3D protein complexes of death domain there were no spatial proximity between the position of the model with the position of the cluster representing the different docking solutions (Figure 4(b)). These data shows that the rescoring of the protein complexes solutions obtained by docking needed to be performed to eliminate false solutions from the real one. Effectively, GRAMM's method considers geometrical fit and hydrophobicity, but does not have optimal molecular mechanics force fields and desolvation parameters that can eliminate false positives. As a result we applied a rescoring algorithm [37] to the “raw” docking results to improve the ranking. The rescoring algorithm has the luxury of considering a much broader spectrum of factors due to the reduced search space. As a
result, the list of complexes was re-ranked and a cutoff imposed with increased confidence, resulting in 11,260 complexes.

As a means of further filtering the docking results, we applied the notion of sister list intersection. Recall that sister lists are those that are returned from docking model \( \mu_i \) against model \( \mu_j \) and vice versa, for \( i, j \in [1...m] \). The motivation behind sister list intersection is that, since we are only interested in domain-domain interactions, theoretically there should be no difference between the two lists. With this and the fact that the lists are predominantly false positives in mind, we argue that sister list intersection is a logical means of filtering. When implementing the algorithm, we discovered that the routine for comparing complexes was best left with a level of flexibility. In other words, complexes need not be identical to be considered on the list, they simply must be sufficiently similar (recall Figure 2). The reason for this is that if identity were to be upheld, the resulting lists would not contain enough elements to allow for classification. By combining sister list intersection then rescoring the protein complex solution obtained from the raw data we were able to reduce the 1.1 million putative complexes down to 913 informative interfaces (i.e. \( n=913 \)). When we applied this mining procedure to the solved complex APAF1/CASP9, of the 2000 putative complexes returned from GRAMM output only two informative interfaces were found, one of which was the true interface. By filtering out the majority of putative complexes we were given the advantage of not having to perform exhaustive docking for each of the new family members we wanted to classify. Instead, we limited the translation/rotation space to the informative interfaces with the understanding that they form a representative population of the important interactions.

When arriving at the definition of the RPIP, the goal was to find a profile that was
fast and easy to calculate, and that suitably described a protein’s interaction behavior so that it could be successfully classified. We should note that the RPIP was not designed to have a direct physicochemical interpretation. As mentioned in the methods section, every RPIP element is associated with an informative interface, which in turn is associated with a model. Since every model corresponds to a member of the superfamily, we can assign a family to each of the RPIP elements. In this way, the RPIP can be divided into four sections, one for each family. In the family-based classifier we hypothesized that RPIP elements corresponding to the target’s true family would, on average, be greater than the elements corresponding to the incorrect families. Figure 5 illustrates the hypothesized RPIP against a typical observed RPIP. As the figure implies, the family-based classifier performed poorly with an average 5-fold cross-validation accuracy of 61%. Alternative summary statistics for representing the families including mean, maximum and standard deviation, did not significantly improve the accuracy. Moreover, removing RPIP elements whose scores were consistently high or low across families similarly did not result in higher accuracy (data not shown). The most obvious implication of these results is that an interface that is particularly stereo-chemically favorable to one family member is not necessarily favorable to another family member.

Unlike the family based classifier, the SVM-based classifier made no assumptions about how true family membership is related to RPIP element family membership; instead it relied on machine learning to automatically find correlations in the elements. The SVM-based classifier performed significantly better with an average cross-validation accuracy rate of 89%. Figure 6 shows an example SVM output from HODOCO. The increased performance of the SVM can be attributed to its ability to detect weaker patterns in the RPIPs, than the heuristic approach taken by the family based classifier. In
particular, it can consider multiple simultaneous interaction propensities to arrive at its final decision. It is interesting to note that some sequences were consistently misclassified by the SVM. Figure 6 shows that APAF1’s true family (CARD) is predicted to be the least probable amongst the four families. In this case, misclassification is due to the fact that it effectively has very unique binding and sequence properties compared to other members of the superfamily. It forms a supra-molecular complex called the apoptosome with CASPASE-9 (CASP9) and NAC [38] which involves a protein-protein interface not present in other family members.

**Conclusions**

The goal of this work was to show that *in silico* interaction-based protein classification can be obtained reliably for the death domain superfamily. We developed a classification pipeline that allows us to obtain protein classification.

While others have used *in silico* interaction profiles to characterize the docking ability of small molecules binding to experimentally determined 3D structures [39], or to discover novel protein interactions using known 3D complexes [40], our method is unique in that it applies to 3D molecular models of proteins complexes, and it not only considers multiple binding partners, but also multiple interfaces for each partner.

In future, there is much further work that can be done. Similar to [39], RPIPs could be used to cluster models rather than classify them. Here the goal would be to find alternative “families” based on interaction data, without regard to sequence homology. For example, discovering groups of models, sharing a common interaction interface or that are an outlier with a unique interface. Such a property has been highlighted by the constant misclassification of the CARD domain of APAF1 obtained in this study that has unique binding properties in the family.
It is very important to note that this study is only as powerful as the methods it builds upon. In particular, docking is still a highly active area of research with much work remaining to be done on model flexibility, solvent simulation, and force field optimization. Similarly, model building of the protein complex remains a difficult procedure requiring great care, making it difficult to accurately automate [41]. Our results have shown that despite the introduction of error in the classification pipeline due to the reliability of the underlying tools, interaction-profile based protein classification can be obtained with confidence. The fact that multiple parties have independently begun researching the potential of interaction profiles suggests that it may become a popular method for biological data mining in the future.

Methods

HODOCO is an *in silico* protein characterization and classification system consisting of three parts, a model building subsystem, a classifier building subsystem, and the classifier itself. An overview of the three major components is shown in Figure 1. When building the classifier the input to the system is a set of 3D structures, termed models (µ₁ through µₘ), that belong to the superfamily of interest. Since 3D structures for the majority of proteins are unsolved, a major aspect of the system is the model building subsystem (Figure 1(a)). Once the models are built, docking is performed to find putative complexes, which are then used to find what we call informative interfaces (ι₁ through ιₙ to be discussed later). The informative interfaces are directly related to calculating the RPIPs, which in turn are used to build the classifier (Figure 1(b)). The model building subsystem and the classifier building subsystem have been designed to be fully independent so that improvements to their respective
underlying tools can be applied to one without necessitating changes to the other. The final subsystem, the classifier, however is tightly bound to the second subsystem, and its distinction is mostly conceptual (Figure 1(c)). Each step in the system is now explained in detail.

**Superfamily Dataset Collection**

Three parallel approaches were used to obtain the set of human death domain protein sequences examined in this study. First, a literature survey was conducted to identify an incomplete set of protein sequences from well-known family members. Second, the Pfam database was consulted to extract a set of protein sequences from members of the CARD, DED, DEATH and PAAD/DAPIN/PYRIN families. Third, the sequences found in the previous two methods were pooled to conduct a distant homologue search via an iterative BLAST procedure [25].

**3D-structure Modeling and Alignment**

Atomic coordinates from solved 3D protein structures were retrieved from the Protein Data Bank [26] and used for docking studies where available. Atomic coordinates from the remaining family members were obtained by homology modeling. Each amino acid sequence (the target sequence) from the death domain superfamily was submitted into the Polish metaserver [27]. Once Pair-wise alignments between the target sequence and a template were generated, a pair-wise alignment with the best score and the template’s structure were submitted into the MODELLER program [28] to generate homology models for the target sequence. The default parameters for MODELLER were used, while the “loop-modelling” option was enabled. A total of 6 models were generated for each target, and the models were refined by molecular dynamics with simulated
annealing (a functionality in MODELLER) to improve the quality of the model. All 6 models were verified for favorable geometrical and stereochemical properties using Verify3D [29] and PROCHECK [30], and the rms deviation between the model and the template from which the model originated. From these criteria the best one was selected as the representative model for the target. Low quality models were discarded if they exhibited an RMSD greater than 1.5Å on the total main chain atoms with the structure template from which they were built. All remaining models were then used as input to the classification analysis, as if they were the original input to the system. Models were structurally superimposed on a reference model (ASC) using the Combinatorial Extension method [31] for docking studies. We refer to the models as $\mu_1$ through $\mu_m$, where $m$ is the number of models.

**Model interaction prediction**

Putative complexes were predicted through computational docking using GRAMM [32]. GRAMM performs an exhaustive 6-dimensional search of all translations and rotations between a given pair of macromolecules and returns a list of high scoring complexes based on rigid-body geometric fit and hydrophobicity. It should be mentioned that, due to GRAMM's algorithm, the order of the models is important. Specifically, the list of complexes returned from docking molecule $A$ against molecule $B$ is not guaranteed to be the same as the list returned from docking molecule $B$ against molecule $A$. We refer to these related lists as *sister lists*. GRAMM's parameters chosen such that the two known death domain superfamily complexes, human caspase recruitment domains of APAF1 with pro-Caspase-9 (PDB: 3YGS) and drosophila death domains of PELLE with TUBE (PDB: 1D2Z) had the best rank possible when docking their respective individual monomers were: Matching mode=generic, grid step=1.5, repulsion attraction=5,
attraction double range=0, potential range type= atom_radius, projection=black and white, representation=all, number of matches to output=1000 and angle of rotation=10. Each model within each family was docked against every other model within the same family and the top 1000 complexes from each docking simulation were retained in a MySQL database for further filtering. It could be argued that the GRAMM parameters giving the best rank, when obtained from the docking of the individual monomers of the two known protein complexes (bound form), could not be optimal when each death domain superfamily member are docked against each others (unbound form). Effectively docking from unbound monomers has to consider conformational changes of the protein partners upon binding that do not occur when a protein monomer belongs to a known protein complex. To limit the complexity of the problem and allow comparison between our simulations we used the low resolution docking parameters of GRAMM. In such a procedure flexibility is handled through a “soft belt” into which atoms from the second molecule can penetrate reducing drastically the complexity [22], and increasing the speed of the simulation.

**Mining for informative interfaces**

The database of resulting complexes was mined for those with the maximum information gain with respect to family classification. We refer to the mined complexes as *informative interfaces*, since each complex defines an interface between the component models. We labeled the informative interfaces $\iota_1$ through $\iota_n$, $n$ the number of interfaces. The mining procedure consisted of two parts: (i) Rescoring the complexes and (ii) Taking the intersection of sister lists, described shortly. All mining was performed via a combination of shell scripts, Perl scripts and C++ programs.
**Rescoring of complexes:** The first mining technique was to rescore each complex using the software Rpdock—a member of the 3D-Dock suite [33] and reject those below a threshold. Rpdock uses evidence gathered empirically to quantify the probability of a complex's existence and returns a score (RPScore) based on the results. The algorithm uses residue pair potentials across protein interfaces [34] as the basis for the score. Ranked via this alternative scoring system, the top 10 complexes from each docking experiment were retained, while the other 990 complexes were rejected.

**Intersection of sister lists:** The second mining technique applied took advantage of the input-model order dependence of GRAMM. Every pair of sister lists was examined for complexes found in both lists. More precisely, every pair-wise combination of complexes from all sister lists was considered in turn. Recall that complexes in sister lists are composed of the same two models (call them $\mu_i$ and $\mu_j$, $i, j \in [1..m]$). If after the two instances of $\mu_i$ were structurally superimposed, the distance between the two instances of $\mu_j$ fell below a given threshold then both complexes were retained. Complexes never retained in this way were rejected (see Figure 2). Note that the two instances of $\mu_j$ need not be exactly structurally superimposed. We refer to $\mu_i$ as the *aligned model* and $\mu_j$ as the *orientation-independent model*. The informative interface is then the interface that lies between these models.

**RPIP construction**

The set of informative interfaces was used to generate vectors of RPScores; one vector per model. Each RPScore was calculated as follows.

1. Given the 3D structure of a sequence of interest (the target), and an informative interface, $\tau_i$ ($i \in [1..n]$), consisting of an aligned model and an orientation independent
model, the aligned model was replaced with the target ensuring preservation of orientation. (Recall that all models were previously structurally superimposed to a reference model, thereby normalizing rotations across models). We refer to this modified complex as the **hybrid complex**.

2. Rpdock was used to calculate the RPScore of the hybrid complex, and the result was stored in element $i$ of the target’s RPIP. Note that the RPScore of the hybrid complex could be grossly different from that of the unmodified complex.

   For a given model, the collection of RPScores resulting from performing the above calculation on each of the informative interfaces was placed into a vector and termed the model’s Residue Potential Interaction Profile, or RPIP (see Figure 3). Note that there is a one-to-one correspondence between RPIP elements and informative interfaces. To avoid bias, all informative interfaces that involved the model itself were assigned a null RPScore and ignored in all future steps.

**Building the classifier**

Two methods of classification were attempted. First, it was postulated that the RPIP elements pertaining to a model’s true family would be, on average, greater than those not. Thus, a classifier was built that compared the median RPScores across RPIP elements for each of the four families, and made a prediction based on the greatest mean. This classifier was termed the **family-based classifier**.

The second method used the RPIPs to build four support vector machines one for each family. This classifier was termed the **SVM-based classifier**. The software SVM light was used in this study (http://svmlight.joachims.org). Each machine was trained to discriminate members of one family, so that all four machines would have to be used to make a final prediction. A linear kernel was used when building the machines to avoid
undue distortion of the underlying RPScores.

Testing

To test the accuracy of HODOCO we conducted five runs of 5-fold cross-validation for each classification method. Generally, \( k \)-fold cross-validation randomly divides the target dataset into \( k \) equal-sized partitions, iteratively using one partition for testing and the other partitions for training. Method accuracy is measured as the average number of correctly classified models divided by the total number of models over a series of cross-validation runs. Referring back to Figure 1(c), a target is classified by building its model, building its RPIP and finally applying one of the classification methods to the RPIP. Note that it is assumed that the unknown sequence has been previously screened to be a member of the superfamily.

Author’s contributions

MS performed the first docking experiments of the project and developed clustering methods to classify protein based on protein-protein interactions, DL contributed to the writing of the manuscript and to data analysis by implementing the support vector machine. FP performed part of the docking experiments, design the necessary experiments to complete this work, provided guidance to mine the data and contributed to the writing of the manuscript.

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**Figures**

**Figure 1 - The HODOCO system architecture.** (a) The process for determining the missing 3D structures of family members from their amino-acids sequence; (b) The classification engine creation pipeline; and (c) The process for classifying new family members. Light boxes represent computation steps. Within each computation step is a description (above the line) and a list of programs that the step uses (below the line). Dark boxes represent data sources and/or sinks. Edge labels represent the data passed from one step to the next.

**Figure 2 - Intersection of sister lists.** Complex 1 resulted from docking $\mu_i$ against $\mu_j$, $i, j \in [1..m]$. Complex 2 resulted from docking $\mu_j$ against $\mu_i$. $\mu_i$ is represented as a cross with an arrowhead; $\mu_j$ as a cross with a bulb. The different shades of gray are used to distinguish instances of the same model. In this case, if we align the two complexes by superimposing both instances of $\mu_i$, then the distance between the centers of the two instances of $\mu_j$ is less than or equal to the cutoff, $d_c$, so both complexes are retained. Note that the instances of $\mu_j$ are not superimposed; only their centers coincide. Here, $\mu_i$ is the aligned model and $\mu_j$ is the orientation-independent model. When we align the two complexes by superimposing both instances of $\mu_j$, then the distance between the centers of $\mu_i$ is greater than the cutoff. However, once a complex is retained, it is never again rejected.

**Figure 3 - Construction of an RPIP.** Each sphere represents the model ($\mu$) of a death domain superfamily member and each plane represents an informative interface ($\iota$). Note that every interface has an associated model, but that the number of models ($m$) may be different than the number of interfaces ($n$). Here, $\iota_1$, $\iota_i$ and $\iota_n$ are associated with $\mu_1$, $\mu_j$ and $\mu_m$, respectively. The target’s RPIP is the collection of RPScores calculated for each interface $\iota_i$, for all $i \in [1...n]$.

**Figure 4 – Unfiltered Predicted complexes.** (a) Clusters of predicted complexes for each of the four families. Receptor models are shown as $\alpha$-carbon traces in red. Each gray point indicates the center position of a GRAMM hit. (b) Predicted complexes vs. the known complex for APAF1/CASP9. The receptor (APAF1) of the predicted complexes is shown as $\alpha$-carbon trace in gray; the ligand center points are shown as gray points with direction vectors. The known complex is shown in blue. Note that the known (blue) and predicted (gray) APAF1’s have been aligned, and that the known CASP9 center is not within the dense regions of predicted centers.

**Figure 5 - RPIP and family based classifier.** This RPIP was generated from a CARD family member. The hypothesis of the family based classifier stated that RPIP elements corresponding to the target’s true family should on average be greater than the others. The series in gray illustrates the predicted model where the RPScores corresponding to the CARD family (indices 0 through 304) should be generally greater than the rest. Note, however, that the observed values in black do not agree with the hypothesis, instead showing a generally even trend across all families.

**Figure 6 - Classification of death domain family members using SVM.** Each RPIP ($x$-axis) has four SVM scores associated with it; one from each family. The family with the highest score is the predicted family. Here, correct classifications include 19/21
CARD members; 8/10 DED members, with one near-tie; 21/21 DD members; and 10/12 PAAD members.

**Tables**

**Table 1 - Human death domain superfamily members with known structures.** PDB codes are followed by chain identifiers; an underscore represents the only chain (http://www.rcsb.org).

**Table 2 - Gene names and RefSeq IDs of the sequences used in this study.** Data from the UCSC Genome Browser [42] April 2003 assembly. *Gene names followed by parentheses have more than one death effectors domain. The number in parentheses is the first amino acid of the domain according to Pfam release 10.0.
| Family | Gene Name | PDB Code            |
|--------|-----------|---------------------|
| CARD   | APAF1     | 1c15:A 2ygs:A 3ygs:C 1cy5:A 1cww:A |
| CARD   | CASP9     | 3ygs:P              |
| CARD   | ICEBERG   | 1dgn:A              |
| CARD   | CRADD     | 3crd:_              |
| DED    | FADD      | 1a1w:_ 1a1z:_       |
| DEATH  | FADD      | 1e3y:A 1e41:A 1fad:A |
| DEATH  | NGFR      | 1ngr:_              |
| DEATH  | TNFRSF6   | 1ddf:_              |
| DEATH  | TNFRSF1A  | 1ich:A              |
| **Total:** |           | **9**               |

**Table 1**
| CARD Name | RefSeq ID   | DED Gene Name | RefSeq ID   | DD Gene Name | RefSeq ID   | PAAD Gene Name | RefSeq ID   |
|-----------|-------------|---------------|-------------|--------------|-------------|----------------|-------------|
| APAF1    | NM_001160   | CASP8(19)     | NM_001230   | ANK1         | NM_000037   | AIM2           | NM_004833   |
| ASC      | NM_013258   | CASP8(114)    | NM_001230   | ANK2         | NM_001148   | ASC            | NM_013258   |
| BCL10    | NM_003921   | CASP8(2)      | NM_001228   | ANK3         | NM_001149   | CIAS1          | NM_004895   |
| BIRC2    | NM_001166   | CASP8(100)    | NM_001228   | CRADD        | NM_003805   | DEFCAP         | NM_014922   |
| BIRC3    | NM_001165   | CFLAR(1)      | NM_003879   | DAPK1        | NM_004938   | MEFV           | NM_000243   |
| CARD10   | NM_014550   | CFLAR(92)     | NM_003879   | FADD         | NM_003824   | NALP10         | NM_176821   |
| CARD11   | NM_032415   | DEDD          | NM_004216   | IRAK1        | NM_001569   | NALP2          | NM_017852   |
| CARD12   | NM_021209   | DEDD2         | NM_133328   | IRAK2        | NM_001570   | NALP8          | NM_176811   |
| CARD14   | NM_024110   | FADD          | NM_003824   | IRAK3        | NM_007199   | PYC1           | NM_152901   |
| CARD6    | NM_032587   | PEA15         | NM_003768   | LRDD         | NM_018494   | PYPAAF4        | NM_134444   |
| CARD9    | NM_022352   |               |             | MYD88        | NM_002468   | PYPAAF5        | NM_138329   |
| CASP1    | NM_001223   |               |             | NGFR         | NM_002507   | RNO2           | NM_033297   |
| CASP2    | NM_001224   |               |             | RIPK1        | NM_003804   |                |             |
| CASP4    | NM_001225   |               |             | THOC1        | NM_005131   |                |             |
| CASP5    | NM_004347   |               |             | TNFRSF10A    | NM_003844   |                |             |
| CASP9    | NM_001229   |               |             | TNFRSF10B    | NM_003842   |                |             |
| CRADD    | NM_003805   |               |             | TNFRSF1A     | NM_001065   |                |             |
| DEFCAP   | NM_014922   |               |             | TNFRSF21     | NM_014452   |                |             |
| ICEBERG  | NM_021571   |               |             | TNFRSF25     | NM_003790   |                |             |
| RIPK2    | NM_003821   |               |             | TNFRSF6      | NM_000043   |                |             |
| TUCAN    | NM_014959   |               |             | TRADD        | NM_003789   |                |             |

**Total:**
- CARD: 21
- DED: 10
- DD: 21
- PAAD: 12

*Table-2*
(a) Model Building

(b) Creating the Classifier

(c) Classifying
Complex 1

Complex 2

Complex Alignment w.r.t. $\mu_i$

$d \leq d_c$

Complex Alignment w.r.t. $\mu_j$

$d > d_c$

Figure 2
Figure 3
(a) Family Clusters

(b) APAF1/CASP9 Complex
Figure 5
Figure 6