Research Article

Predicting In Silico Which Mixtures of the Natural Products of Plants Might Most Effectively Kill Human Leukemia Cells?

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The aim of the analysis of just 13 natural products of plants was to predict the most likely effective artificial mixtures of 2-3 most effective natural products on leukemia cells from over 364 possible mixtures. The natural product selected included resveratrol, honokiol, chrysin, limonene, cholecalciferol, cerulenin, aloe emodin, and salicin and had over 600 potential protein targets. Target profiling used the Ontomine set of tools for literature searches of potential binding proteins, binding constant predictions, binding site predictions, and pathway network pattern analysis. The analyses indicated that 6 of the 13 natural products predicted binding proteins which were important targets for established cancer treatments. Improvements in effectiveness were predicted for artificial combinations of 2 or 3 natural products. That effect might be attributed to drug synergism rather than increased numbers of binding proteins bound (dose effects). Among natural products, the combinations of aloe emodin with mevinolin and honokiol were predicted to be the most effective combination for AML-related predicted binding proteins. Therefore, plant extracts may in future provide more effective medicines than the single purified natural products of modern medicine, in some cases.

1. Introduction

Plant-derived secondary metabolites have been used to treat acute infections, health disorders, and chronic illness for tens of thousands of years. Only during the last 100 years have natural products been largely replaced by synthetic drugs [1]. However, important anticancer agents have to be extracted from plants, due to their complex structures that often contain several chiral centers. Further, some patients show resistance to known treatments [2]. Therefore, new treatments with different modes of action are constantly sought. Plants are an abundant source of new natural products. Estimates of 200,000 natural products in plant species have been revised upward as mass spectrometry techniques have developed [3]. New databases, omics methods, and good practice standards are promising to deliver many new medicines based on plant natural products [4].

Several studies have demonstrated that mixtures in extracts from herbal medicines had anticancer potential in vitro or in vivo [5–7]. Among many other studies, aqueous extracts from willow (salix sp.; Salicaeae) leaves prevented proliferation of three cancer cell types acute myeloid leukemia (AML), acute lymphoblastic leukemia (ALL), and Ehrlich ascites carcinoma cells [7]. Therefore, the complex mixtures in crude extracts may be more effective than single purified natural products.

Leukemia was among the most common cancers throughout human history [8]. However, the greater prevalence of leukemia in the modern world may be due to the reduction of infectious diseases that lead to the increased life span for most human populations. Unfortunately, by
2. Materials and Methods

2.1. Databases and Software. The reference databases and software of Ontomine were used for predictive analysis. Ontomine was chosen because it provided an innovative cheminformatics prediction tool based on the presence or absence of chemical group(s) of a set of related natural products. Ontomine searches were performed against large and manually curated databases. They included (i) Literature searches based on experimentally determined properties from around 100,000 diverse small molecules, collected from databases, encyclopedias, and other literature followed by expert hand-curation; (ii) BioAssay Knowledgebase that was compiled from over 500 bioassay data found at NCBI-PubChem; (iii) Target Protein Knowledgebase that was compiled by curation among the ~1500 proteins from DrugBank at NCBI-PubChem (details given in Figure 1). (iv) Pathway Analysis: KEGG pathways were used as references (ftp://ftp.genome.jp/pub/kegg/); (v) Docking Algorithms were used to identify molecular binding sites and predict ligand binding constants. Ontomine databases and tools are among those used widely in this field [4].

2.2. Natural Products Selected for Prediction of the Basis of Antileukemia Activity. Thirteen commercially available, purified, natural products of plants were selected to be tested for their antileukemic (AML) properties (Table 1). Natural products had been shown to cause some cell death when incubated with a primary AML cell lines for 24 hrs at low concentrations (Supplemental Table 1 see Supplementary materials available online at http://dx.doi.org/10.1155/2013/801501). Mortality rate and dose dependence were known [7], but bioassays had not clearly identified the best therapeutics, and the in vivo analysis of all mixtures would have been costly. Therefore, in silico prediction was used to identify the best candidates for later testing in vivo. A parallel analysis was made with established drugs for AML treatments so that the predictions could be compared.

2.3. Ontomine-Based Analyses of Functional Groups. Ontomine was used to transform the structural information for chemically, biologically, or pharmacologically related molecules to a hierarchical schema of functional groups. It was used to discover patterns in the related schema and predict biological activity, toxicity, and side-effects using rules inferred from analyzing the patterns. The basic algorithms underlying Ontomine-based predictions were as follows.

For rule or pattern detection the cluster formation was based on a similarity threshold (ST) that was calculated for each molecule using the formula:

\[
\text{No. of common functional groups in molecule 1 and 2} \times \text{Maximum functional group count in either molecule}
\]

Once ST was calculated, clusters formed if the ST was less than or equal to 0.7. Two clusters could be merged if each contained the same molecules or a subset thereof. Results were generated with confidence levels of high, medium, and low.

The Ontomine algorithm took into account presence and absence of functional groups among sets of related molecules (with similar bioactivity or toxicity) to derive rules for making predictions. This approach was different from traditional maximum common subgraphs (MCS) approaches in which only a part of molecule is taken into consideration for decision making.

2.4. Drug Target Analysis. The database used contained bioassay data for ~500 predicted binding proteins (Supplemental Figure 1). Natural product target identification was done by mining two specific knowledgebases: the “Drug Target KB” and “BioAssay KB”. BioAssay KB was generated from 493 bioassay records from NCBI and Drug Target KB (1,346 records) was generated from PubChem, DrugBank, and TTD [4]. Predicted binding proteins for 13 natural products were predicted at three confidence levels of high, medium, and low. The predicted bioassays were validated using scientific journals, and some of the bioassay targets were found as active against AML in the literature [13–24].

2.5. Reverse Docking to Predict Binding Affinities for Proteins. Docking was a computational method used to estimate binding strength among biomolecules (protein-ligand and protein-protein interactions). Traditionally docking was used as computational tool for screening databases of natural products to mine a set of a few candidate drug-like compounds. Reverse docking was a comparatively new application of docking in which a database of proteins (drug targets) was docked against a set of natural products (potentially anticancerous compound), to predict binding affinities. AutoDock4.0 software was used to carry out reverse docking over a database of ~1,100 predicted binding proteins available in the Potential Drug Target Database (PDTD) [10]. The PDTD contained more than 1,100 protein entries with known
3D structures. The PDTD covered diverse information of more than 830 known or potential drug targets, including protein and active site structures, related diseases, biological functions, and their associated regulating (signaling) pathways. Taking every ligand as a probe, reverse docking was carried out with the entire database (PDTD). The respective ligands were also prepared using AutoDock4.0. A grid of each protein’s binding pocket (site) was constructed using the Autogrid module of AutoDock4.0. Every ligand was separately docked into the binding site of each protein. The interaction energies between the ligand and the proteins were calculated in the form of docking scores. AutoDock4.0 also yielded the inhibition constant ($K_i$) for every docking calculation. The predicted binding proteins were screened using a cut off value of $<10 \mu M$ for the inhibition constant ($K_i$).

2.6. Pathway Analyses. Pathway analyses were performed on predicted binding proteins, to provide insights into potential mechanisms of natural product activity. Three statistical confidence levels were considered for pathway analyses. Statistical analysis used Fishers Exact Test to identify over-represented pathways.

2.7. Gene Expression Network Analysis. Protein-protein interaction network analyses can provide important information about disease mechanisms and help to select candidates for genes and their encoded proteins underlying
Table 1: Bioassay targets are confirmed as AML active proteins by literature searches and associated binding proteins are predicted by Ontomine.

| S. no | Ontomine predicted bioassays                  | Confidence level | Compound(s)                              | Reference no |
|-------|-----------------------------------------------|------------------|------------------------------------------|--------------|
| 1     | Guanine nucleotide binding protein, alpha activating polypeptide-O | Medium           | Honokiol                                 | [13]         |
|       |                                               | Low              | Cholecalciferol                           |              |
|       |                                               |                  | Limonene                                 |              |
|       |                                               |                  | Retinyl palmitate                         |              |
| 2     | Multidrug resistance protein 1               | Low              | Mevinolin                                 | [14]         |
|       |                                               |                  | Limonene                                 |              |
|       |                                               |                  | Resveratrol                               |              |
|       |                                               |                  | Chrysin                                  |              |
| 3     | Myeloid or lymphoid or mixed-lineage leukaemia protein | High            | Aloe emodin                               | [15]         |
|       |                                               |                  | Mevinolin                                 |              |
|       |                                               | Low              | Barbaloin                                 |              |
|       |                                               |                  | Limonene                                 |              |
|       |                                               |                  | Retinyl palmitate                         |              |
|       |                                               |                  | Cholecalciferol                           |              |
|       |                                               |                  | Chrysin                                  |              |
|       |                                               |                  | L-Ascorbic acid 6-palmitate              |              |
| 4     | Runt-related transcription factor 1 isoform AML1c | Medium           | Honokiol                                 | [16]         |
|       |                                               |                  | Barbaloin                                 |              |
|       |                                               |                  | Mevinolin                                 |              |
|       |                                               | Low              | Limonene                                 |              |
|       |                                               |                  | L-Ascorbic acid 6-palmitate              |              |

disease. Networks can include transcript and protein co-expression data. Here AML was compared to normal white blood cell transcript abundance data from published studies [12, 25] available within the NCBI-GEO database (Gene Expression Omnibus) (Accession: GSE17054, GSE9476). The algorithm described [25] for finding significant subnetwork/modules in gene interaction network was used to find transcripts which are altered in abundance in AML samples when compared to normal samples. This algorithm focused on finding small networks, which would be easier to be interpreted and validated. It computes P values for subnetworks, which helped identify significant subnetworks. This analysis produced lists of significant sub-networks (Supplemental Table 2).

2.8. Final Protein Target Selection. The objective of the protein target selection was to analyze data generated through chemoinformatics and structural informatics method, and select relevant and important predicted binding proteins, which would be used for selecting potential drug/drug combination. Detailed literature searches were conducted for predicted binding proteins generated by the earlier analyses (Target Discovery/Identification). The disease pathways annotations from KEGG were used to select cancer related predicted binding proteins along with additional literature searches. The literature searches were used to support annotations from KEGG and also to provide additional information about target protein whose role in carcinogenesis has been established recently [13–24].

Predicted binding proteins were selected based on following criteria:

1. Target protein should be predicted as related with compounds of interest (natural products estimated from Reverse Docking or Ontomine).
2. Predicted binding proteins should be related with AML/cancer (KEGG annotation and/or literature).

2.9. Validation of Existing Drugs for AML. Seventeen existing drugs for AML were included in the analysis to (i) understand the mode of action of drugs; (ii) predict activity profiles; and (iii) use them as benchmarks for the natural products analysis. The following drugs were used as benchmark sets in analysis; natural products 6-mercaptopurine; 6-thioguanine; L-malate and vincristine; and synthesized drugs, amonafide, belinostat, clofarabine, cytarabine, daunorubicin, etoposide, fludarabine, gencitabine, idarubicin, mitoxantrone, paclitaxel, prednisone, and tipifarnib. Reverse docking for 13 natural products along with 17 benchmark drugs was carried out with a target database (PDTD) of ~1060 protein
Table 2: Protein targets for natural products previously reported in the literature.

| S. no | Ontomine predicted compound(s) | Bioassay targets | References |
|-------|-------------------------------|------------------|------------|
| 1     | Mevinolin                     | Multidrug resistance protein 1 | [17]       |
| 2     | Limonene                      | Multidrug resistance protein 1 | [18]       |
| 3     | Resveratrol                    | Multidrug resistance protein 1 | [19]       |
| 4     | Chrysin                       | Multidrug resistance protein 1 | [20]       |
| 5     | Cholecalciferol                | Myeloid or lymphoid or mixed-lineage leukaemia protein | [21]       |

Figure 2: Aloe emodin was predicted to be bound at the centre surrounded by the amino acid residues from the active site of 2GDZ. It can be seen that aloe emodin forms 4 hydrogen bonded interactions with the residues of the active site. These hydrogen bonded interactions are partly responsible for stabilizing the ligand-protein complex. In addition the hydrophobic interactions present between the ligand and the protein are displayed as wireframe spheres.

2.10. Drug Combination Selection. Searching potential interactions among 13 natural products required a factorial design to consider all possible combinations (78 pairs or 286 sets of 3 natural products). The search algorithm was designed to find potential nodes in the tree structure where root nodes represented singlet drugs and processes. Finding interactions involved moving up the tree to explore combinations. This approach helped to avoid false positives by limiting the search space to significant nodes and processes. Combiscores (hereafter Cscores) were used in drug/drug-combination ranking and selection. The Cscore was derived by analyzing target profile by Ontomine for drug combinations where

\[
Cscore = (0.2 \times (Ac - (Am + Ah)) + (0.4 \times Am) \\
+ (0.3 \times Ah)) - 0.1 \times comm + sp,
\]

where \(Ac\) was the number of predicted binding proteins related to cancer; \(Am\) was the number of predicted binding proteins related to AML; \(Ah\) was the number of predicted binding proteins detected as hubs in interaction networks from gene-expression network analysis; \(comm\) was the number of common target protein(s) shared by constituent drugs in combination; \(sp\) was the specificity score defined as (the number of cancer-related predicted binding proteins minus the number of protein not related to cancer)/(Total no. of predicted binding proteins for drug/combination).

Dscore was derived by analyzing target profile obtained through docking analysis for drug combinations where

\[
Dscore = ((0.2 \times (Ac - (Am + Ah)) + (0.4 \times Am)) \\
+ (0.3 \times Ah)) - 0.1 \times comm) N^{-1};
\]

\(N\) was the number of proteins used for forward docking (\(N = 59\)).

Combiscore was the statistic derived from combining Cscore and Dscore:

\[
Combiscore = (0.6 \times Cscore) + (0.4 \times Dscore).
\]

Computing the scores weights was associated with parameters, to alter their relative importance. For example predicted binding proteins related with AML were given more weight than proteins associated with cancers in general. For single natural products the process began by computing the Combiscore for each of the 13 natural products. The mean Combiscore was calculated (1.36) and natural products with Combiscore greater or equal to 1.36 were selected as potentially useful.

For pairs of natural products all possible binary combinations were considered, with constraint of considering only those combinations which started from a selected singlet. A Combiscore was computed for each pair and ranked. The mean Combiscore was calculated. Finally, potential combinations that had a Combiscore greater or equal to the mean Combiscore (4.25) were reported.

For sets of three natural products combinations the same process was followed with the Combiscore threshold set...
Table 3: List of drugs predicted to form useful mixtures by significant Cscores, Dscores, and Combiscores.

(a) Singlet analysis

| Compound            | Cscore   | Dscore  | Combiscore |
|---------------------|----------|---------|------------|
| Aloe emodin         | 9.40909  | 0.02711 | 5.65630    |
| Chrysin             | 8.2      | 0.04406 | 4.93762    |
| Honokiol            | 6.13333  | 0.05423 | 3.70169    |
| Mevinolin           | 5.35     | 0.02033 | 3.21813    |
| Resveratrol         | 4.55925  | 0.01356 | 2.74097    |
| L-ascorbic acid 6-palmitate | 2.376470588 | 0 | 1.42588 |
| Cholecalciferol     | 2.25     | 0.02542 | 1.36016    |
| Limonene            | 2.23076  | 0.01864 | 1.34591    |
| Cerulenin           | 2        | 0.03898 | 1.21559    |
| Retinyl palmitate   | 1.4      | 0.00508 | 0.84203    |
| Aloe sin            | -1       | 0.02542 | -0.58983   |
| Salicin             | -10      | 0.05254 | -5.97898   |
| Barbaloin           | -10      | 0.03728 | -5.98508   |

(b) List of binary combinations selected by analysis

| Compound                        | Cscore   | Dscore  | Combiscore |
|---------------------------------|----------|---------|------------|
| Aloe emodin + honokiol          | 10.33333 | 2.12543 | 7.05016    |
| Aloe emodin + Aloe sin          | 8.83333  | 2.58813 | 6.33525    |
| Aloe emodin + cholecalciferol   | 10.25384 | 0.37627 | 6.30281    |
| Aloe emodin + salicin           | 8.52075  | 2.80677 | 6.23516    |
| Aloe emodin + mevinolin         | 9.52307  | 1.15423 | 6.17554    |
| Aloe emodin + l-ascorbic acid 6-palmitate | 8.968421053 | 1.63728 | 6.03597    |
| Aloe emodin + cerulenin         | 9.56153  | 0.61016 | 5.98099    |
| Aloe emodin + barbaloin         | 8.52075  | 1.93220 | 5.88534    |
| Aloe emodin + retinylpalmitate  | 8.94545  | 0.86440 | 5.71305    |
| Aloe emodin + limonene          | 2.25     | 0.02542 | 1.36016    |
| Aloe emodin + resveratrol       | 7.91034  | 2.31359 | 5.67163    |
| Chrysin + l-ascorbic acid 6-palmitate | 8.711764706 | 0.50339 | 5.42841    |
| Chrysin + cerulenin             | 8.78461  | 0.28475 | 5.38466    |
| Chrysin + salicin               | 7.83846  | 1.67288 | 5.37222    |
| Aloe emodin + chrysin           | 7.87704  | 1.41864 | 5.29368    |
| Aloe emodin + barbaloin         | 7.83846  | 0.79830 | 5.02239    |
| Mevinolin + chrysin             | 7.77704  | 0.56949 | 4.89402    |
| Cholecalciferol + chrysin       | 6.88148  | 1.69830 | 4.80821    |
| Honokiol + resveratrol          | 7.69629  | 0.35423 | 4.75947    |
| Honokiol + limonene             | 7.775    | 0.16610 | 4.73144    |
| Mevinolin + resveratrol         | 6.824489796 | 1.46440 | 4.68045    |
| Honokiol + chrysin              | 7.609836066 | 0.16610 | 4.63234    |
| Honokiol + cholecalciferol      | 7.433333333 | 0.16610 | 4.52644    |
| Mevinolin + honokiol            | 6.385712486 | 1.27627 | 4.34193    |
| Honokiol + cerulenin            | 7.030769231 | 0.16610 | 4.28490    |
| Mevinolin + limonene            | 6.885714286 | 0.30508 | 4.25346    |
| Mevinolin + cholecalciferol      | 6.572727273 | 0.30508 | 4.06567    |
| Honokiol + salicin              | 6.033333333 | 0.847457627 | 3.958983051 |
| Mevinolin + salicin             | 5.15     | 1.957627119 | 3.873050847 |

(c) List of triple combinations selected by analysis

| Compound                       | Cscore   | Dscore  | Combiscore |
|--------------------------------|----------|---------|------------|
| Aloe emodin + mevinolin + honokiol | 12.00864198 | 1.866101695 | 7.951625863 |
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(c) Continued.

| Compound                                      | Cscore  | Dscore  | Combiscore  |
|-----------------------------------------------|---------|---------|-------------|
| Aloe emodin + honokiol + cholecalciferol       | 11.74146341 | 2.125423729 | 7.89504754 |
| Aloe emodin + honokiol + cerulenin            | 11.49268293 | 2.047457627 | 7.714592807 |
| Aloe emodin + cholecalciferol + chrysin       | 11.78888889 | 1.418644068 | 7.64079096 |
| Aloe emodin + honokiol + limonene            | 11.15064935 | 2.125423729 | 7.540559102 |
| Aloe emodin + honokiol + chrysin              | 11.15064935 | 1.777966102 | 7.401576051 |
| Aloe emodin + Aloe sin + cholecalciferol      | 10.45384615 | 2.588135593 | 7.30756193 |
| ...                                           | ...     | ...     | ...         |
| Mevinolin + chrysin + barbaloin               | 9.87704918 | 0.994915254 | 6.32419561 |
| Aloe emodin + resveratrol + cerulenin         | 9.046376812 | 2.23559322  | 6.322063735 |
| Mevinolin + chrysin + l-ascorbic acid 6-palmitate | 10.06507937 | 0.7        | 6.319047619 |
| Mevinolin + chrysin + cerulenin               | 10.11343284 | 0.569491525 | 6.29586312 |
| Aloe emodin + mevinolin + l-ascorbic acid 6-palmitate | 9.523076923 | 1.377966102 | 6.265032595 |
| Aloe emodin + mevinolin + retinylpalmitate    | 9.733333333 | 0.991525424 | 6.236610169 |
| Mevinolin + honokiol + limonene              | 9.433333333 | 1.276271186 | 6.170508475 |
| Honokiol + limonene + resveratrol            | 10.02258065 | 0.354237288 | 6.155243302 |
| Mevinolin + honokiol + resveratrol           | 9.46705882 | 1.140677966 | 6.135094716 |
| Aloe emodin + Aloe sin + barbaloin           | 8.833333333 | 2.069491525 | 6.12779661 |
| ...                                           | ...     | ...     | ...         |

greater or equal to the mean Combiscore (6.15). Higher Combiscores indicated better drug combinations.

3. Results

3.1. Bioassays. Ontomine analysis identified probable targets for each of the 13 natural products among proteins that were potential bioassay targets (Table 2). In total 618 proteins were predicted to be bioassay targets at various confidence levels (157 at high confidence, 91 at medium confidence and 370 at low confidence). Table 1 showed the four proteins previously reported to be bioassays targets of the natural products. Multiple (4–8) natural products were predicted to interact with bioassays targets, guanine nucleotide binding protein, alpha activating polypeptide O, multidrug resistance protein 1, myeloid or lymphoid or mixed-lineage leukaemia protein, and runt-related transcription factor isoform 1 (AML1c). Clearly Ontomine identified a larger set of bioassays than what has been previously reported. Those bioassay targets could be contributing to the effects or side-effects of the natural products.

3.2. Reverse Docking. Reverse docking was predicted with structures of natural products docked against protein databases. Using the structures of the natural product to interrogate the database of protein structures identified 17 proteins predicted to bind natural products with a \( K_i \) less than 1 \( \mu \)M. All 17 were predicted to be related to AML by network analysis. They were ALOX12 that encoded arachidonate 12-lipoxygenase; AR for the androgen receptor; BCL2L1 and BCL2L2 for the B-cell lymphoma 2-like proteins; CDC42 encoding the cell division cycle 42 (GTP binding protein, 25 kDa); DUSP3 for the dual specificity phosphatase 3; GSK3B encoded the glycogen synthase kinase 3 beta; IGF1R for the insulin-like growth factor 1 receptor; KLF5 for the ruppel-like factor 5 (intestinal); MAPK1 for the mitogen-activated protein kinase 1; MMP14 for the matrix metallopeptidase 14 (membrane inserted) protein; NFkB1 that encoded the nuclear factor of kappa light polypeptide gene enhancer in B cells; RHOA, the ras homolog gene family, member A; RUNX1, the runt-related transcription factor 1; SMAD3, for mothers against decapentaplegic homolog family member 3; and STAT1 and STAT3 the signal transducer and activator of transcription 91kDa protein and acute-phase response factor, respectively. In Figure 1 three of these proteins (STAT3, MAPK1, IGF1R) were found within a small node of just fifteen interacting proteins. Further, two proteins were involved in B-cell regulation (BCL-2, KFB1). In total reverse docking for 13 natural products and 17 drugs to ~1,060 protein structures yielded 92 targets with inhibition constants \( (K_i) < 10 \mu \)M. The reverse docking results were then analysed in conjunction with the Ontomine results. These include targets which either belong to a subset of reverse docking and study or else they are identified as important targets by Ontomine. A total of 95 targets were finally identified (Supplemental Table 3).

3.3. Docking. Among the 95 binding proteins identified by Reverse Docking, the protein structures which could be used for docking were available for 59 targets. These 59 structures were finally used for docking against the 14 ligand structures under consideration along with the benchmark ligands. The docking parameters were kept the same as those used for reverse docking.

Among the ligand-protein binding interactions of significance was aloe emodin predicted to bind to the protein 15-hydroxyprostaglandin dehydrogenase type 1 (2GDZ; Figure 2). Aloe emodin was predicted to bind at the center of
the structure surrounded by the amino acid residues from the active site of 2GDZ. It can be seen that aloe emodin forms 4 hydrogen bond interactions with the residues of the active site. These hydrogen bond interactions were predicted to stabilize the ligand-protein complex. The human 15-hydroxyprostaglandin dehydrogenase type 1 has been reported to be elevated in abundance and activity in AML cell lines [7].

3.4. Combinations of Natural Products and Drugs for AML Treatments. The Cscore, Dscore and Combiscores were all significant for aloe emodin, chrysin, honokiol, mevinolin, resveratrol, l-ascorbic acid 6-palmitate, and cholecalciferol (Table 3(a)). Considering pairs of natural products with predicted synergistic interactions, 25 combinations were found (Table 3(b)). Twelve pairs included aloe emodin suggesting that this natural product would work well in many mixtures. There were 7 pairs that included honokiol and 6 for chrysin. Even natural products not predicted to be effective alone, like limonene, could be effective in paired mixtures. Considering sets of three natural products with predicted synergistic interactions 15 combinations were found (Table 3(b)). Eleven drug combinations included aloe emodin, again suggesting that this natural product would work well in many mixtures. There were 5 mixtures that included honokiol but

| Singles | Couples | Triples |
|---------|---------|---------|
| AE      | HK      | 7.1     |
| AE      | AL      | 6.3     |
| AE      | MV      | 6.2     |
| AE      | LA      | 6.0     |
| AE      | CR      | 6.0     |
| AE      | BB      | 5.9     |
| AE      | RP      | 5.7     |
| AE      | LM      | 5.7     |
| AE      | CH      | 4.9     |
| HK      | 4.3     |
| MV      | 3.7     |
| RS      | 3.2     |
| LA      | 2.7     |
| CF      | 1.4     |
| LM      | 1.3     |
| CR      | 1.2     |
| RP      | 0.8     |
| AL      | 0.6     |
| SA      | 0.0     |
| BB      | -6      |
| AE      | HK      | 7.9     |
| AE      | CF      | 7.7     |
| AE      | CR      | 7.6     |
| AE      | CH      | 7.5     |
| AE      | LM      | 7.4     |
| AE      | AL      | 7.3     |
| AE      | CF      | 7.3     |
| AE      | SA      | 7.3     |
| AE      | HK      | 7.3     |

| AE      | MV      | HK      |
|---------|---------|---------|
| AM      | HK      | 6.1     |
| AE      | AL      | 6.1     |
| AE      | SA      | 6.1     |
| AE      | MV      | 6.1     |
| AE      | LA      | 6.1     |
| AE      | CR      | 6.1     |
| AE      | BB      | 6.1     |
| AE      | RP      | 6.1     |
| AE      | LM      | 6.1     |
| AE      | RS      | 6.1     |

**Figure 3:** Drug combination selections. Values inside beige squares were the Combiscores. Each black box was the mean used to derive the cutoff.
only 3 for chrysin. Again natural product not predicted to be effective alone like limonene could be effective in multiple mixtures (Figure 3).

4. Discussion

Analysis of 13 natural products inferred that some interactions might be useful and novel. Target profiling by Ontomine, Docking, and Gene Expression Network analysis indicated that the natural products were predicted to bind to proteins which were important targets for cancer treatments. Pathway analyses indicated statistical overrepresentation of cancer-related pathways among drug targets for aloe emodin, cerulenin, chrysin, honokiol, mevinolin, and resveratrol.

Mixtures of natural products were predicted to be more effective than single products, as reported experimentally [7, 8, 26]. Improvements in predicted effectiveness for mixtures of natural products could be attributed to drug synergism due to increase in relevant targets or improved specificity of drug constituents. This increase in predicted effectiveness was not likely to be derived from random effects like pooling of result of individual drugs, since the analysis accounted for important factors for drug combinations like target relevance to cancer/AML, specificity, and common targets among drug constituents, while calculating scores are used for ranking drug combination.

The parallel analysis on benchmarked drugs which exist in the market for AML treatment was significant since the efficacy of these drugs was supported by many publications [2, 8, 10, 27]. Combination analysis on drugs was also successful in discovering well-known combinations like aminofide plus cytarine and daunorubicin plus prednisone. That aloe emodin plus mevinolin plus honokiol was identified as the best combination that has interesting clinical implications. Analysis of this mixture with respect to drug specificity, targeting AML-related proteins and targeting cancer-related hubs, will be a priority for future laboratory and clinical research. Several other combinations have potential to treat drug-resistant cancers in the future. In future analyses in silico more complex drug combination search algorithms can be applied to incorporate dose, absorption, and excretion as suggested [27].

Conflict of Interests

The authors of the paper do not have any conflict of interests nor any direct financial relation with the National CFIDS Foundation, 103 Aletha Road, Needham, MA 02492, USA.

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References

[1] M. Wink, A. W. Alfermann, R. Franke et al., “Sustainable bioproduction of phytochemicals by plant in vitro cultures: anticancer agents,” Plant Genetic Resources, vol. 3, no. 2, pp. 90–100, 2005.
[2] S. Mishra, B. Zhang, J. M. Cunnick, N. Heisterkamp, and J. Groffen, “Resistance to imatinib of Bcr/Abl P190 lymphoblastic leukemia cells,” Cancer Research, vol. 66, no. 10, pp. 5387–5393, 2006.
[3] R. Mungur, A. D. M. Glass, D. B. Goodenow, and D. A. Lightfoot, “Metabolite fingerprinting in transgenic Nicotiana tabacum altered by the Escherichia coli glutamate dehydrogenase gene,” Journal of Biomedicine and Biotechnology, vol. 2005, no. 2, pp. 198–214, 2005.
[4] D. J. Barlow, A. Buriani, T. Ehrman, E. Bosisio, I. Eberini, and P. J. Hylands, “In-silico studies in Chinese herbal medicines’ research: evaluation of in-silico methodologies and phytochemical data sources, and a review of research to date,” Journal of Ethnopharmacology, vol. 140, pp. 526–534, 2012.
[5] H. Sheridan, L. Krenn, R. Jiang et al., “The potential of metabolic fingerprinting as a tool for the modernisation of TCM preparations,” Journal of Ethnopharmacology, vol. 140, pp. 482–491, 2012.
[6] X. Chen, Z. J. Zhan, X. W. Zhang, J. Ding, and J. M. Yue, “Sesquiterpene lactones with potent cytotoxic activities from Vernonia chinensis,” Planta Medica, vol. 71, no. 10, pp. 949–954, 2005.
[7] H. A. El-Shemy, A. M. Aboul-Enein, K. M. Aboul-Enein, and K. Fujita, “Willow leaves’ extracts contain anti-tumor agents effective against three cell types,” PLoS ONE, vol. 2, no. 1, article e178, 2007.
[8] F. Cervantes, F. Passamonti, and G. Barosi, “Life expectancy and prognostic factors in the classic BCR/ABL-negative myeloproliferative disorders,” Leukemia, vol. 22, no. 5, pp. 905–914, 2008.
[9] H. A. El-Shemy, M. A. M. Aboul-Soud, A. A. Nassr-Allah, K. M. Aboul-Enein, A. Kabash, and A. Yagi, “Antitumor properties and modulation of antioxidant enzymes’ activity by Aloe vera leaf active principles isolated via supercritical carbon dioxide extraction,” Current Medicinal Chemistry, vol. 17, no. 2, pp. 129–138, 2010.
[10] M. Campillos, M. Kuhn, A. C. Gavin, L. J. Jensen, and P. Bork, “Drug target identification using side-effect similarity,” Science, vol. 321, no. 5886, pp. 263–266, 2008.
[11] X. Gao, Y. X. Xu, G. Divine, N. Janakiraman, R. A. Chapman, and S. C. Gautam, “Disparate in vitro and in vivo antileukemic effects of resveratrol, a natural polyphenolic compound found in grapes,” Journal of Nutrition, vol. 132, no. 7, pp. 2076–2081, 2002.
[12] A. D. Kinghorn, E. J. Carcache de Blanco, H.-B Chai et al., “Discovery of anticancer agents of diverse natural origins,” Pure and Applied Chemistry, vol. 81, no. 1051, 1063 pages, 2009.
[13] J. H. Won, J. S. Park, H. H. Ju, S. Kim, H. Suh-Kim, and S. H. Ghil, “The alpha subunit of Go interacts with promyelocytic leukemia zinc finger protein and modulates its functions,” Cellular Signalling, vol. 20, no. 5, pp. 884–891, 2008.
[14] N. Fazlina, A. Maha, A. L. Zarina et al., “Assessment of P-gp and MRP1 activities using MultiDrugQuant Assay Kit: a preliminary study of correlation between protein expressions and its functional activities in newly diagnosed acute leukaemia patients,” The Malaysian Journal of Pathology, vol. 30, no. 2, pp. 87–93, 2008.
[15] M. S. Brassesco, A. P. Montaldi, D. E. Gras et al., “Cytogenetic and molecular analysis of MLL rearrangements in acute lymphoblastic leukaemia survivors,” Mutagenesis, vol. 24, no. 2, pp. 153–160, 2009.
[16] M. Buchert, C. Darido, E. Lagerqvist et al., “The symplekin/ZONAB complex inhibits intestinal cell differentiation by the repression of AML1/Runx1,” Gastroenterology, vol. 137, no. 1, pp. 156–164, 2009.

[17] D. Yamasaki, T. Nakamura, N. Okamura et al., “Effects of acid and lactone forms of 3-hydroxy-3-methylglutaryl coenzyme A reductase inhibitors on the induction of MDR1 expression and function in LS180 cells,” European Journal of Pharmaceutical Sciences, vol. 37, no. 2, pp. 126–132, 2009.

[18] T. Nabekura, T. Yamaki, and S. Kitagawa, “Effects of chemopreventive citrus phytochemicals on human P-glycoprotein and multidrug resistance protein 1,” European Journal of Pharmacology, vol. 600, no. 1–3, pp. 45–49, 2008.

[19] M. Bobrowska-Hägerstrand, M. Lillås, L. Mrówczyńska et al., “Resveratrol oligomers are potent MRP1 transport inhibitors,” Anticancer Research, vol. 26, no. 3, pp. 2081–2084, 2006.

[20] R. Zhao, T. J. Raub, G. A. Sawada et al., “Breast cancer resistance protein interacts with various compounds in vitro, but plays a minor role in substrate efflux at the blood-brain barrier,” Drug Metabolism and Disposition, vol. 37, no. 6, pp. 1251–1258, 2009.

[21] N. Niitsu, Y. Hayashi, and Y. Honma, “Downregulation of MLL-CBP fusion gene expression is associated with differentiation of SN-1 cells with t(11;16)(q23:p13),” Oncogene, vol. 20, no. 3, pp. 375–384, 2001.

[22] Z. Gao, H. Li, H. Zhang et al., “PDTD: a web-accessible protein database for drug target identification,” BMC Bioinformatics, vol. 9, article 104, 2008.

[23] D. L. Stirewalt, S. Meshinchi, K. J. Kopecky et al., “Identification of genes with abnormal expression changes in acute myeloid leukemia,” Genes Chromosomes and Cancer, vol. 47, no. 1, pp. 8–20, 2008.

[24] R. Majeti, M. W. Becker, Q. Tian et al., “Dysregulated gene expression networks in human acute myelogenous leukemia stem cells,” Proceedings of the National Academy of Sciences of the United States of America, vol. 106, no. 9, pp. 3396–3401, 2009.

[25] S. Nacu, R. Critchley-Thorne, P. Lee, and S. Holmes, “Gene expression network analysis and applications to immunology,” Bioinformatics, vol. 23, no. 7, pp. 850–858, 2007.

[26] H. A. El-Shemy, A. M. Aboul-Enein, M. I. Aboul-Enein, S. I. Issa, and K. Fujita, “The effect of willow leaf extracts on human leukemic cells in vitro,” Journal of Biochemistry and Molecular Biology, vol. 36, no. 4, pp. 387–389, 2003.

[27] D. Calzolari, S. Bruschi, L. Coquin et al., “Search algorithms as a framework for the optimization of drug combinations,” PLoS Computational Biology, vol. 4, no. 12, Article ID e1000249, 2008.