

# Browsing Large Scale Cheminformatics Data with Dimension Reduction

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## Abstract

*Visualization of large-scale high dimensional data is highly valuable for data analysis facilitating scientific discovery in many fields. We present PubChemBrowse, a customized visualization tool for cheminformatics research. It provides a novel 3D data point browser that displays complex properties of massive data on commodity clients. As in GIS browsers for Earth and Environment data, chemical compounds with similar properties are nearby in the browser. PubChemBrowse is built around in-house high performance parallel MDS (Multi-Dimensional Scaling) and GTM (Generative Topographic Mapping) services and supports fast interaction with an external property database. These properties can be overlaid on 3D mapped compound space or queried for individual points. We prototype the integration with Chem2Bio2RDF system using SPARQL endpoint to access over 20 publicly accessible bioinformatics databases. We describe our design and implementation of the integrated PubChemBrowse application and outline its use in drug discovery. The same core technologies are generally applicable to develop high performance scientific data browsing systems for other applications.*

## 1. Introduction

The scale of public scientific data generated by new techniques makes data utilization really challenging. Volumes of PubChem<sup>1</sup> data in cheminformatics, for example, need to be visualized towards the end of capture, curation, and analysis pipeline to support interactive studies. However, because browsing tens of millions of intrinsically high-dimensional PubChem data in one platform is not easy due to the high dimensionality and large volumes of data, we have developed a 3D data point visualization tool for drug discovery, named PubChemBrowse, to display chemical structures with various properties such as relationships among chemical, gene, and disease established through querying Chem2Bio2RDF system [1], which we shall discuss shortly.

We are applying parallel Multidimensional Scaling (MDS) and Generative Topographic Mapping (GTM) algorithms to reduce 166 structural descriptors (known as MACCS keys) of around 60 million chemical compounds in PubChem into three, aiming at providing three-dimensional graphical plot representations of the structural diversity of nearly all of the known public compounds [2, 3]. By labeling these plots with the properties of compounds, ranging from simple properties like molecular

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<sup>1</sup> NIH Public Chemical Database, <http://pubchem.ncbi.nlm.nih.gov/>

weights to complex properties such as chemical-disease relationships established through querying Chem2Bio2RDF system, we can investigate the overall properties of regions of chemical space inhabited by PubChem compounds, as well as embed new compounds into this framework.

In section 2, we give a brief overview of related work and our high performance parallel dimension reduction technologies including MDS and GTM and relational biochemical repository framework – Chem2Bio2RDF and its SPARQL query interface. Section 3 presents details of our design and implementation of an integrated system PubChemBrowse that plots a 3D scattered distribution of PubChem compounds, as well as embed new compounds retrieved in real time with various labeling capabilities. Several case studies are given in section 4 with a data set of up to 930,000 data points followed by a summary of our work and future improvements.

## **2. Data visualization and remote data access**

Large scale data visualization is an active research area in many fields of science: The Sloan Digital Sky Survey (SDSS) project [4] for astronomy, UCSC [5] and Ensemble [6] for genomics, and Google Earth for geology. Although designed with different technologies, they share one common concept: users need only a lightweight client but can access huge data that are curated and processed through intensive computation. We have used a multi-stage pipeline model to explore natural data parallelism for large-scale scientific problems. This has been demonstrated in our biological gene sequencing application from data acquisition, analysis, reduction and aggregation for visualization and implemented with both classic HPC and Cloud technologies [7]. We are taking a similar approach for the cheminformatics research, assisting drug discovery from large dataset.

In the field of modern drug discovery aiming at mining cause-and-effect relationships among chemicals, genes and diseases from a large number of data sources, using various kinds of public databases, such as PubChem for chemical compounds and structures, Comparative Toxicogenomics Database (CTD) for chemical gene information, DrugBank database for drug target association to name a few, is an interesting researching topic and high-dimensional data visualization is gaining more importance in understanding complex datasets. It is worth mentioning that D. M. Maniyar et al. [8] have introduced the use of data visualization for drug discovery and their visualization tool. We present the same type of tool for drug discovery but we integrate our system with more cutting-edge technologies: high-performance visualization algorithms based on HPC and Cloud environments, large data access by using semantic web technologies, and a lightweight 3D visualization client.

More specifically, such process typically involves three main tasks: i) visualizing multi-dimensional PubChem data in 3D space, ii) finding relationships by using various public databases, such as genes-compounds and/or diseases-compounds, and assigning labels to chemical compounds, and iii) displaying labels by using different colors or symbols in the previous 3D visualization and exploring the results. The last two steps can be repeated until we have meaningful discovery.

Authors have been researching high performance visualization algorithm development, such as parallel MDS and GTM and their interpolation extensions [2, 3], to visualize large PubChem dataset in 3D space by using our in-house 3D data point visualization tool and now we extend its functionality to access external data sources in a dynamic way.

Issues in dealing with various types of databases in drug discovery are that databases are often too big to store in local storage and it is hard to maintain consistency with the original datasets if there are frequent updates. Other issues can arise due to non-uniform data structures. Typically those databases are not compatible to each other so that elaborated work needs to be done to use them together. To overcome such problems, Chem2Bio2RDF system has been developed as an integrated repository of chemogenomic and systems chemical biology data [1]. Chem2Bio2RDF aggregates over 20 publicly accessible databases in Resource Description Framework (RDF) format and allows users to access them by using SPARQL which is a standard query language for RDF data and is a part of semantic web technology. PubChemBrowse can interact with Chem2Bio2RDF system remotely by using SPARQL query to access various databases and large number of properties in an online and uniform way.

## 2.1. Data visualization algorithms

Among many dimension reduction algorithms, we focus on MDS [9, 10] and GTM [11] due to their popularity and theoretical strength. Although both GTM and SOM share the same objective which is to find an optimal mapping or embedding in low-dimensional space out of the data in high-dimensional space, GTM, however, finds a mapping based on probability density model, which SOM lacks of. On the other hand, MDS tries to construct a mapping in target dimension with respect to the pairwise proximity information, mostly dissimilarity or distance [2]. The details of those algorithms are beyond the scope of this paper. Users are encouraged to refer to the original papers [9-11] and our previous work [2, 3]. In the following, we briefly describe the algorithms which we used for visualization in PubChemBrowse.

**MDS** : This is a general term of the techniques to configure low dimensional mappings of the given high-dimensional data with respect to the pairwise proximity information, while the pairwise Euclidean distance within the target dimension between two points is approximated to the corresponding original proximity value. In other words, MDS is a non-linear optimization problem with respect to the mapping in the target dimension and the original proximity information. The STRESS [12] value is a well-known objective function of MDS, and Eq (1) represents STRESS function ( $\sigma$ ):

$$\sigma(X) = \sum_{i < j \leq N} w_{ij} (d_{ij}(X) - \delta_{ij})^2 \quad (1)$$

where  $w_{ij}$  is weight for distance between point  $i$  ( $x_i$ ) and  $j$  ( $x_j$ ),  $d_{ij}(X)$  represents mapping distance between  $x_i$  and  $x_j$ ,  $\delta_{ij}$  represents the given original pairwise dissimilarity value between  $x_i$  and  $x_j$ , and  $X$  denotes current low dimensional mapping set.

Among many MDS solutions, we use SMACOF algorithm which is based on Expectation Maximization (EM)-like iterative majorization method. For details of the SMACOF algorithm, please refer to [13].

**GTM** : It is an unsupervised learning algorithm for modeling the probability density of high-dimensional data and finding an optimal non-linear mapping of latent points from the target space to the data space. GTM defines an explicit probability density model based on Gaussian distribution and seeks an optimal set of parameters to maximize the log-likelihood by using Expectation-Maximization (EM) method [11]. For its explicit use of probability model, GTM is also known as a

principled alternative to Self-Organizing Map (SOM) which does not have any density model. In a nutshell, GTM aims at maximizing the following objective function:

$$\mathcal{L}(\mathbf{W}, \beta) = \operatorname{argmax}_{\mathbf{W}, \beta} \sum_{n=1}^N \ln \left\{ \frac{1}{K} \sum_{k=1}^K \mathcal{N}(x_n | y_k, \beta^{-1}) \right\}, \quad (2)$$

where  $y_k$  is a prototype point in the data space mapped by a non-linear function  $f$  with respect to a latent point  $z_k$  in the target space and a parameter set  $\mathbf{W}$  such that  $y_k = f(z_k, \mathbf{W})$ ,  $x_n$  represents a data point in the data space, and  $\mathcal{N}(x|y, b)$  denotes an isotropic Gaussian probability of  $x$  centered on  $y$  with variance  $b$ .

**MDS and GTM Interpolation :** Both are extensions to the original MDS and GTM algorithms, designed to process much larger data sets with sampling approaches. In short, we compute the mapping based on  $n$  samples out of total  $N$  data points followed by interpolating the embedding of the remaining  $(N-n)$  out-of-sample data, which allows us to save computational time. Furthermore, we can build this approach as a pleasingly parallel application [3]. With minor trade-off of approximation, the interpolation approach for MDS and GTM can compute mappings of millions of data points with modest amount of computations and memory requirement. In [3], up to 2 million PubChem data points has been visualized by using parallelized MDS and GTM interpolation algorithms.

## 2.2. Remote Data Access

We can take an advantage of Chem2Bio2RDF system for accessing multiple data sources in an online manner by sending SPARQL query and fetching results in RDF format. In this way, users can visualize data with more information-rich contexts that combine with other data sources.

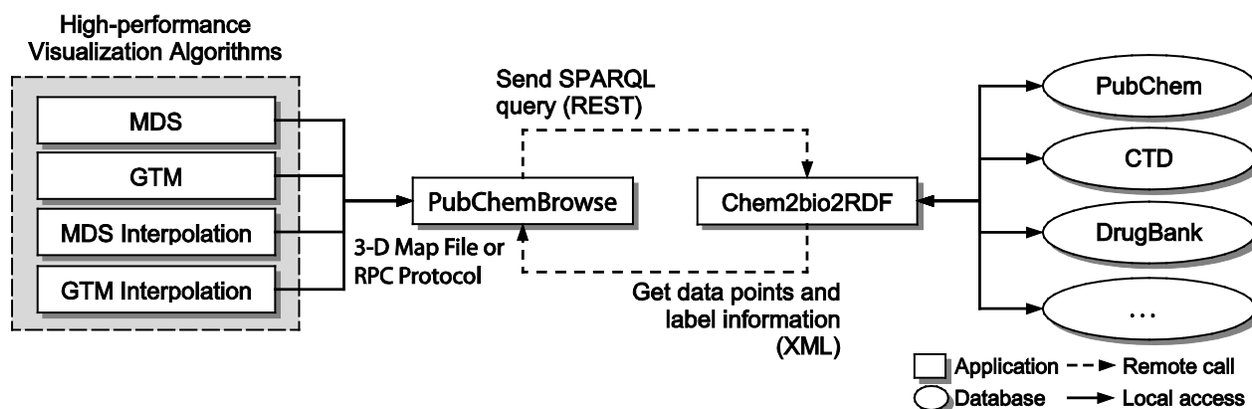
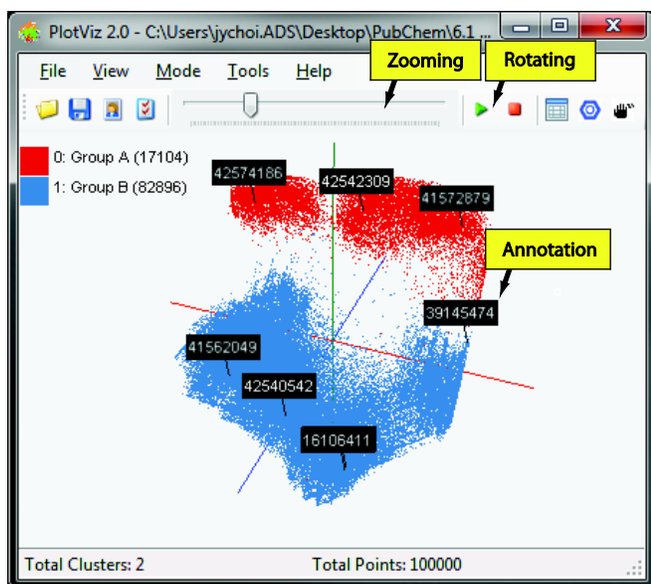
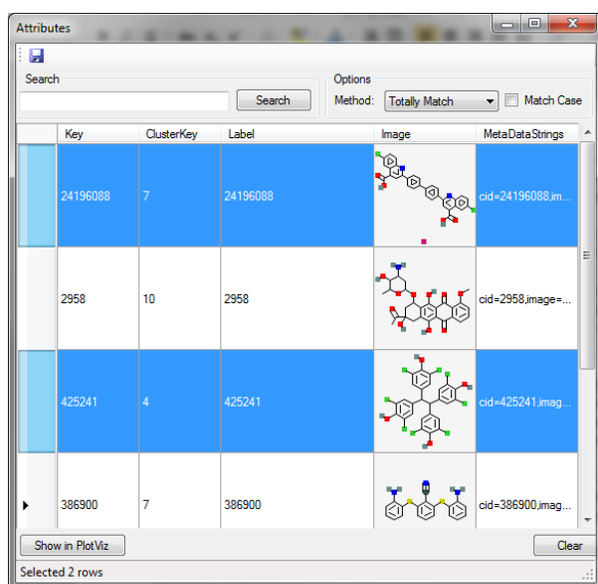


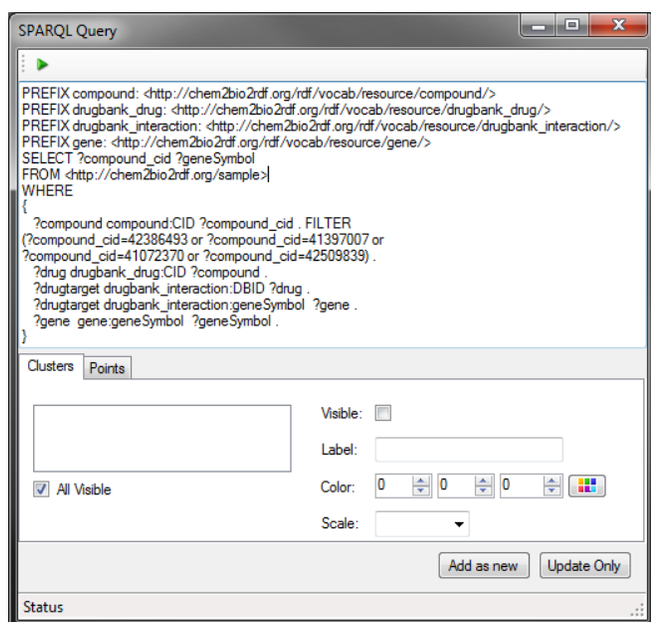
Figure 1. System architecture for PubChemBrowse



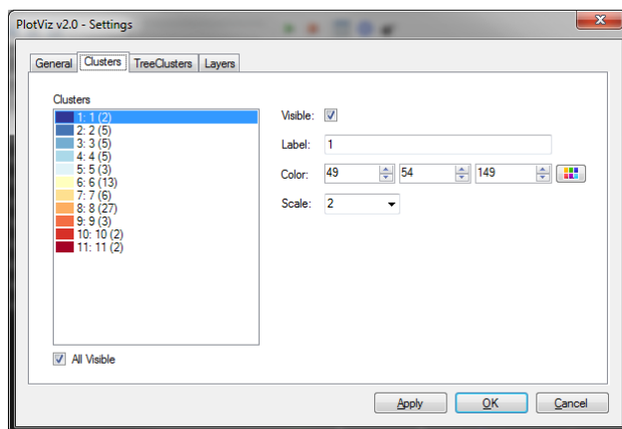
(a) Main Window



(b) A sub-window for meta-data browsing



(c) A sub-window for SPARQL



(d) A sub-window for cluster management

Figure 2. Screenshots of PubChemBrowse.

### 3. The PubChemBrowse system

We have developed PubChemBrowse that can visualize 3D data output from high-performance visualization algorithms, as well as interact with external data sources via Chem2Bio2RDF system in order to provide rich and on-line information by utilizing semantic web interfaces.

Figure 1 illustrates our PubChemBrowse architecture. It consists of mainly 3 components: i) high-performance visualization algorithms, such as parallel MDS and GTM, which generate 3D maps for

large and high-dimensional data by utilizing parallel clustering infrastructure [2, 3], ii) an user-friendly 3D data browser which supports 3D interactions including rotation, panning and zooming and displays annotations and meta-data, and iii) SPARQL query interface which allows users to access the remote data repository Chem2Bio2RDF in order to augment new data points or update existing information in an on-line manner. More details of each component are as follows:

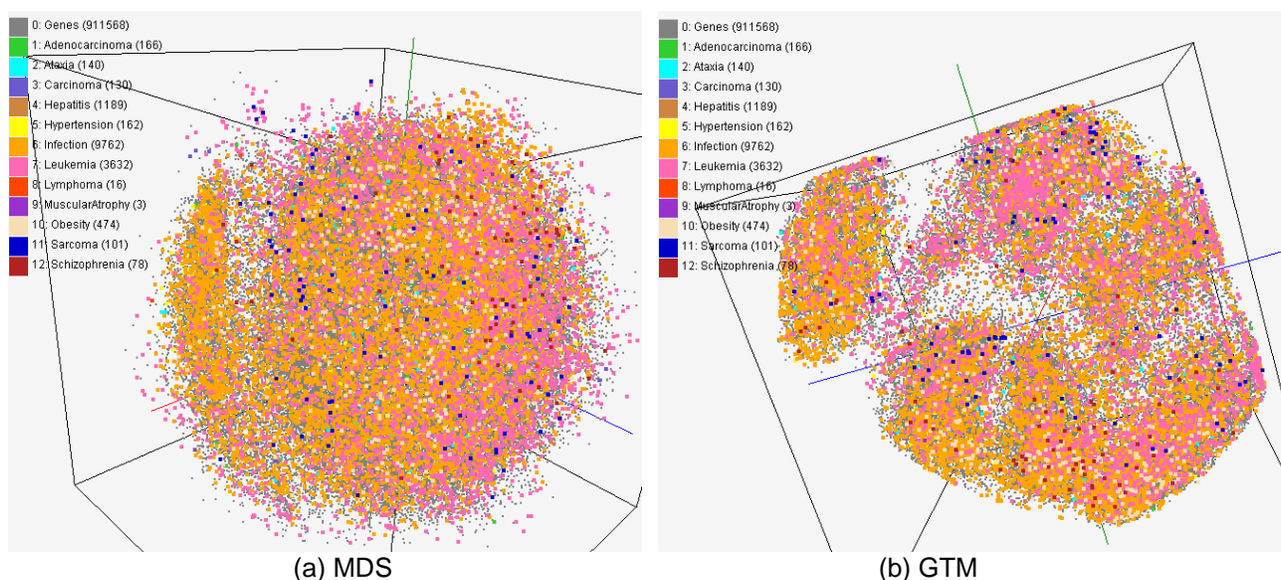
**High-performance visualization algorithms :** Currently PubChemBrowse can visualize the outputs from our in-house parallel implementations of 4 main dimension reduction algorithms: MDS, GTM, MDS interpolation, and GTM interpolation. All of them can run in high-throughput clustering infrastructure by maximally utilizing multi-core environments. Mostly those algorithms run in a batch mode but interpolation algorithms can run in an online mode too.

**3D data browser :** As shown in Figure 2 (a) and (b), one can visualize 3D data points and browse them in various ways: rotating, zooming, and viewing meta-data including chemical structures available from PubChem database.

**SPARQL query :** As shown in Figure 2 (c), our system has an interface for users to compose SPARQL query and send to Chem2Bio2RDF system remotely by using REST which is one of standard web service protocols. Then, the Chem2Bio2RDF system will process the query and return back results in XML format. Our system will parse the XML information and update or overlay the results in user's working plot.

#### 4. Applications

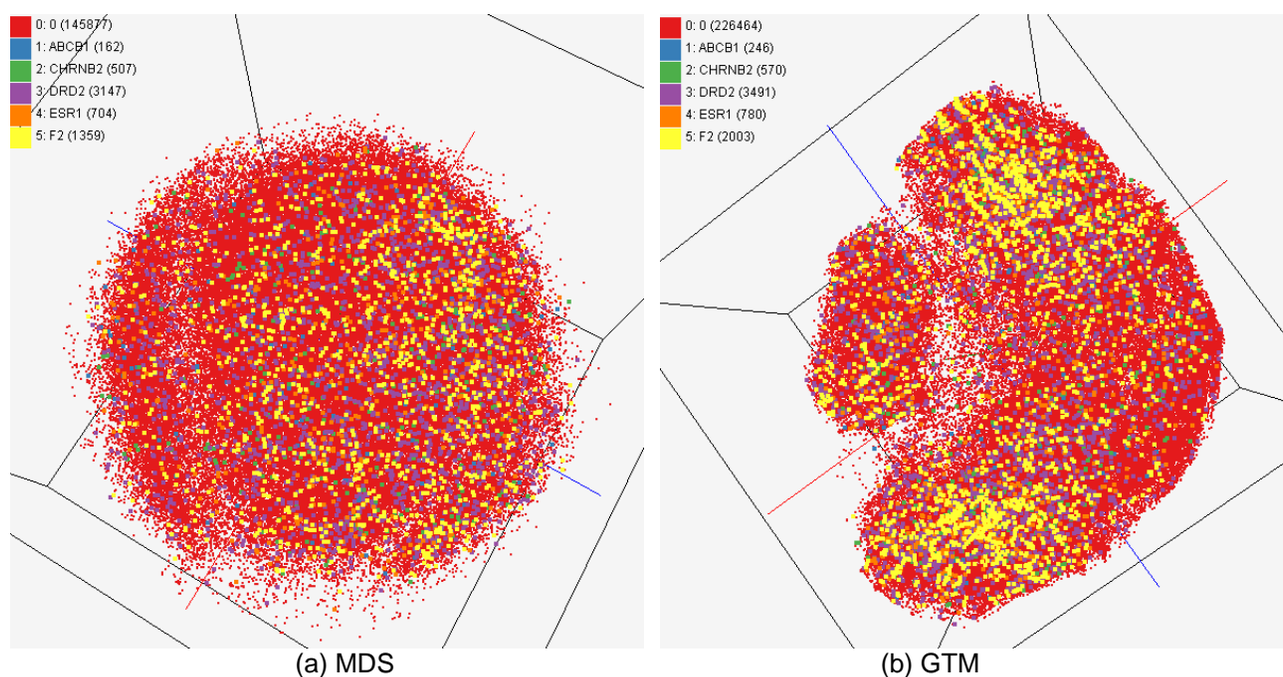
We have been using PubChemBrowse in various cheminformatics data mining research projects aiming at exploring and discovering compound-gene-disease relationships from a large number of datasets. In the following we will show examples of usages of our system in those projects.



**Figure 3. Visualization of chemical-disease relationships based on the data in Comparative Toxicogenomics Database (CTD). About 930,000 chemical compounds are visualized as points in 3D space, computed by MDS (a) and GTM (b).**

**CTD data for chemical and disease relations:** The environment plays an important role in the etiology of many human diseases and chemicals are one component of environment. However, the relation between chemicals and diseases are poorly known [14]. As a part of a research process to find cause-and-effect correlation between chemical compounds and diseases by using public databases, we visualized about 930,000 biologically interesting compounds as points in a 3D space by using our system. Figure 3 shows an example of usages of PubChemBrowse in this study for finding potential correlations. More specifically, we computed X-Y-Z positions processed by either MDS (a) or GTM (b) by using MACCS keys, so that the distance between any two points in the 3D plot reflects structural similarities of two compounds. On top of this bare map which has no additional information except positions, we overlaid disease information mined from CTD with different colors as labels by using SPARQL queries to Chem2Bio2RDF system.

The relation between diseases based on their associated chemicals could be observed from the visualization. It is not surprising that the disease chemicals in Figure 3 are scattered in the chemical spaces instead of being clustered in a particular area, as chemicals have a variety of ways to cause disease via directly or indirectly interacting with disease related genes and their structures are not necessarily similar.

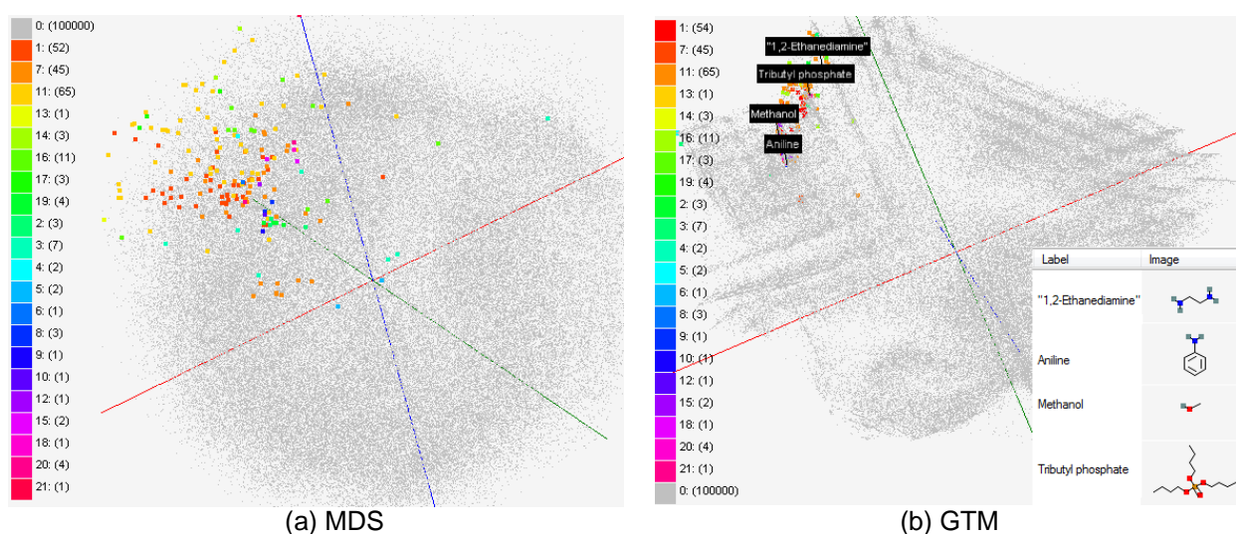


**Figure 4. Visualization of target relations by using multiple data sources in Chem2Bio2RDF. About 234,000 points are visualized in 3D space. Each point represents a chemical compound in PubChem database and its configuration is computed by using MDS (a) and GTM (b).**

**Chem2Bio2RDF :** Many public databases are appeared recently to integrate chemical gene/protein relation data (including various interactions such as activation, inhibition and expression) and the study of which is a major task in either academia or pharmaceutical company. Chem2Bio2RDF dedicates to integrating them in a semantic manner and well classifying their relations, after which a variety of scientific questions with respect to chemicals are raised. For example, it is reported that the chemicals of public databases have little overlap [15], the examination of their differences is of interest in order to optimize their utilization. Another interesting question is to investigate the

differences of chemicals between different targets. Chemicals with similar structures are prone to interact with same targets. The study of relations between targets based on their active compounds could act as a complementary tool to study biological function [16]. These differences could be observed from our system, in which all the compounds are projected into 3D space based on their structural similarity and they are labeled (or colored) either by their original data databases or by their targets that would be retrieved via SPARQL.

Figure 4 shows an example to study the target relations. We visualized 234,000 chemical compounds reported in major journal literature as compounds possibly related to one of 5 targets (ABCB1, CHRN2, DRD2, ESR1, and F2). Positions of compounds are computed by MDS (a) or GTM (b) with the same method used in the previous work and colors are assigned based on their targets retrieved remotely through Chem2Bio2RDF system.



**Figure 5. MDS (a) and GTM (b) visualization of 215 solvents (colored and/or labeled) with 100k PubChem dataset (colored in grey) to navigate chemical space. Users can see images of compound structures in PubChemBrowse. 4 structures of solvents are shown as an example.**

**Solvent screening :** Inspired by the work of C. Lipinski and A. Hopkins [17], we have studied to draw a large chemical space consisting of by sub regions of similar compounds. In an example shown in Figure 5, we have visualized 215 solvents used in a pharmaceutical pre-screening process [18] along with 100 thousand randomly selected chemical compounds by using MACCS keys. As a result, our tool can display a solvent space (a sub region for solvents) separated from other chemicals. This shows that our tool can help users to navigate the large chemical space with visualization.

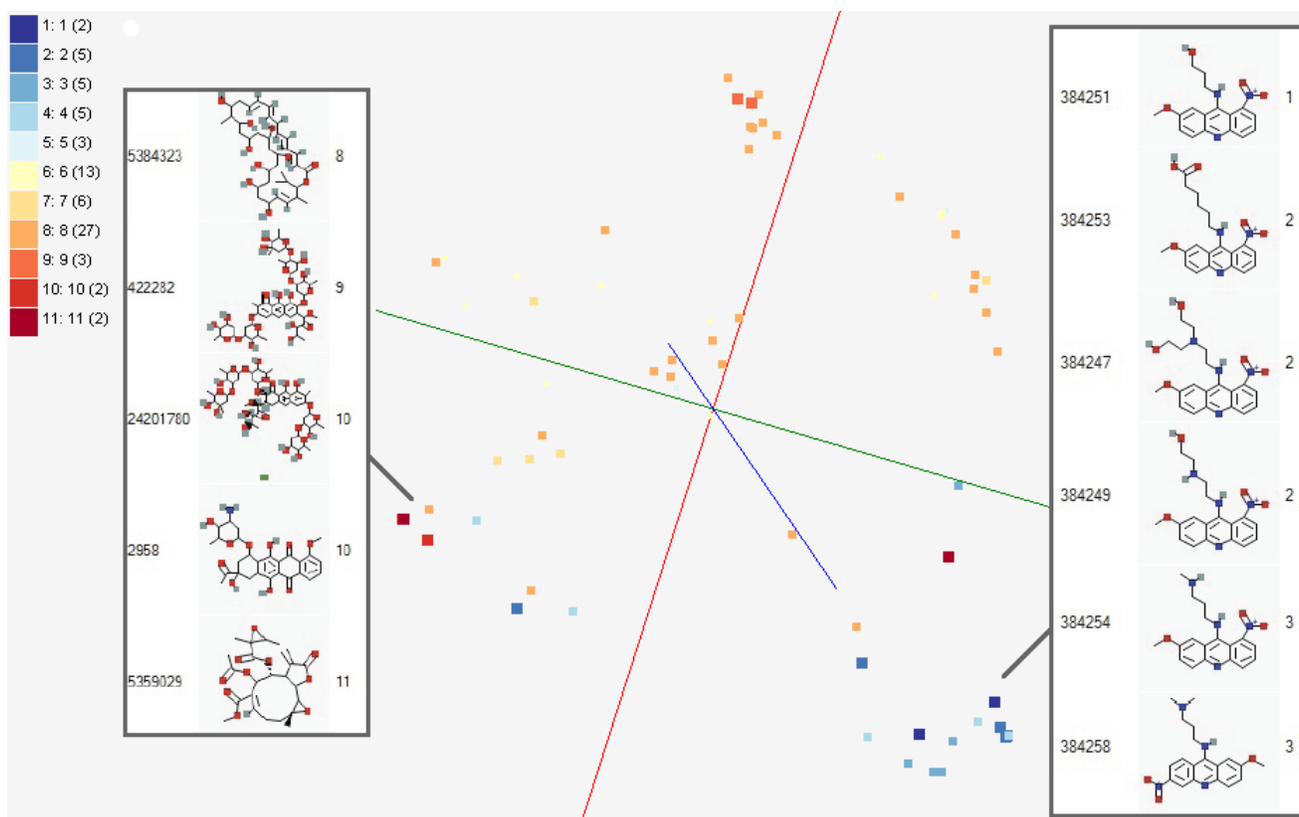


Figure 6. GTM visualization of bioassay activities.

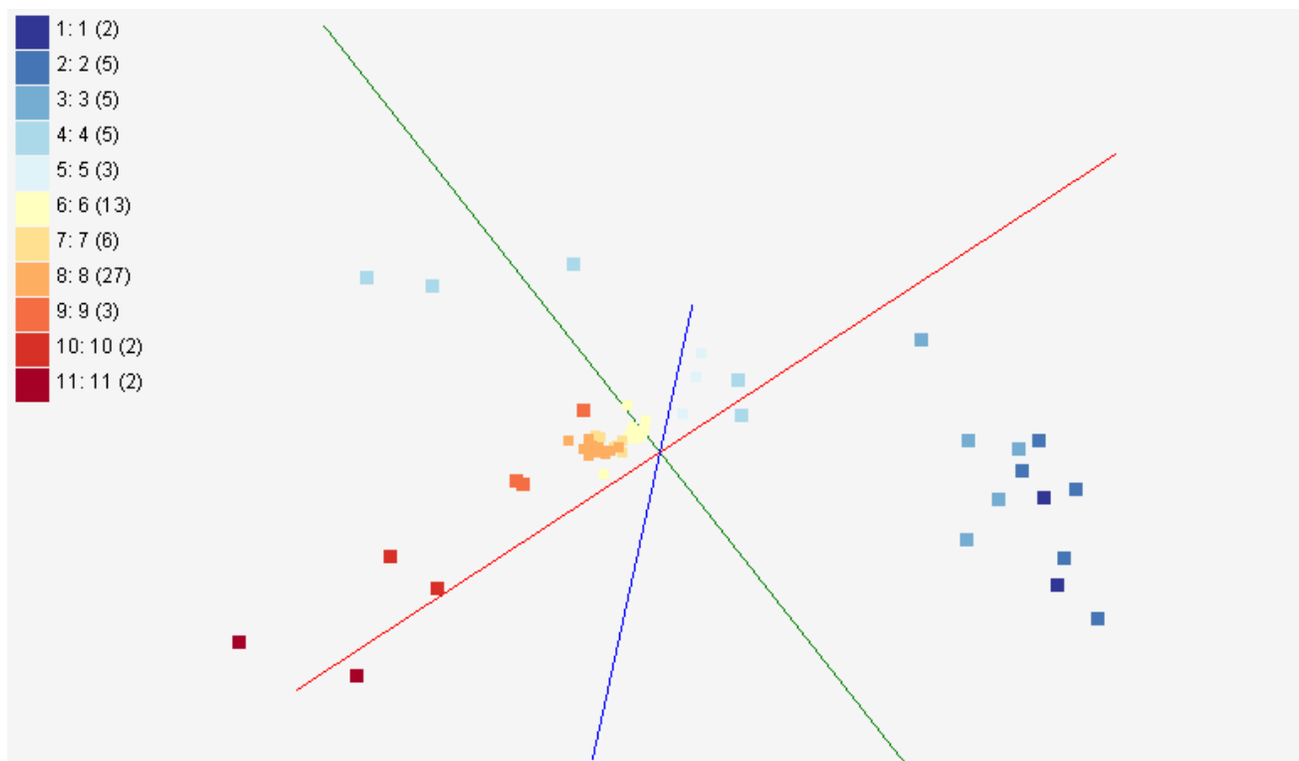


Figure 7. MDS mapping for activity cliffs, based on SALI pairwise matrix.

**Activity Cliffs :** This is a research project to study activity cliffs with SALI index [19] for applications in polypharmacology and chemogenomics. In this study, visualization of bioassay activities and SALI values in 3D chemical space is necessary. Figure 6 shows a visualization example of bioassay activities of 73 chemical compounds in 3D chemical space by using MACCS keys, processed by GTM. Figure 7 depicts the MDS result showing 3D SALI space, based on pairwise SALI indices. Note that SALI space mapping can be only generated by MDS. Unlike GTM, MDS can generate a mapping based on pairwise proximity information.

**Other Life Science Applications :** Although PubChemBrowse is mainly developed to provide special functions for cheminformatics applications, it can process and visualize various other life science datasets. As examples, we present two life science applications. One is the visualization of health data to study child obesity by using the PubChemBrowse system. The dataset consists of children's health-related features, such as Body Mass Index (BMI), blood pressures, and so on, as well as environmental factors, like greenness of neighborhoods, incomes of households, etc. We prepare a pairwise dissimilarity matrix of 10 thousand points based on 98 features as an input and generate a 3D mapping constructed by MDS as shown in Figure 8. Another example is the visualization of biological sequence data. We produce a 3D mapping of 30 thousand metagenomics sequence data based on the pairwise sequence alignment scores measured by a well-known sequence alignment algorithm called Smith-Watermann [20]. Figure 9 shows the 3D mapping result of 30 thousand sequence data for the metagenomics study, also configured by MDS algorithm.

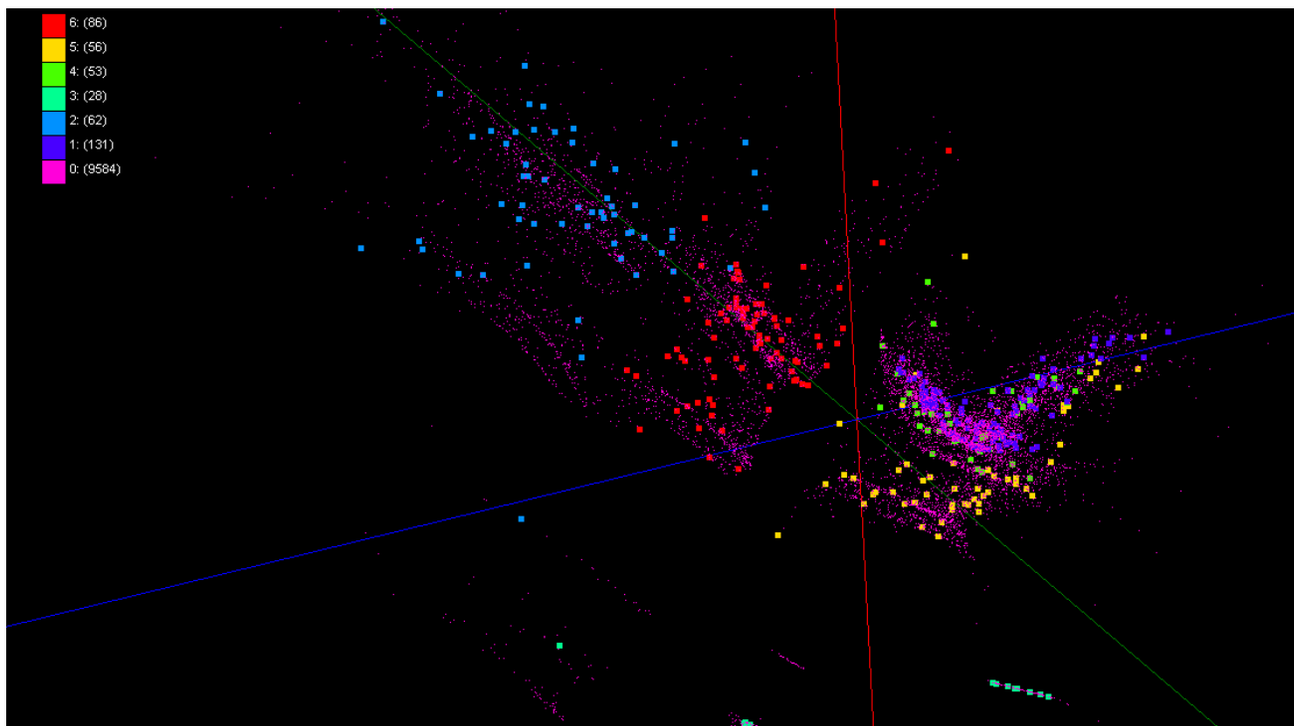


Figure 8. A 3D mapping of the 10 thousand children obesity data.

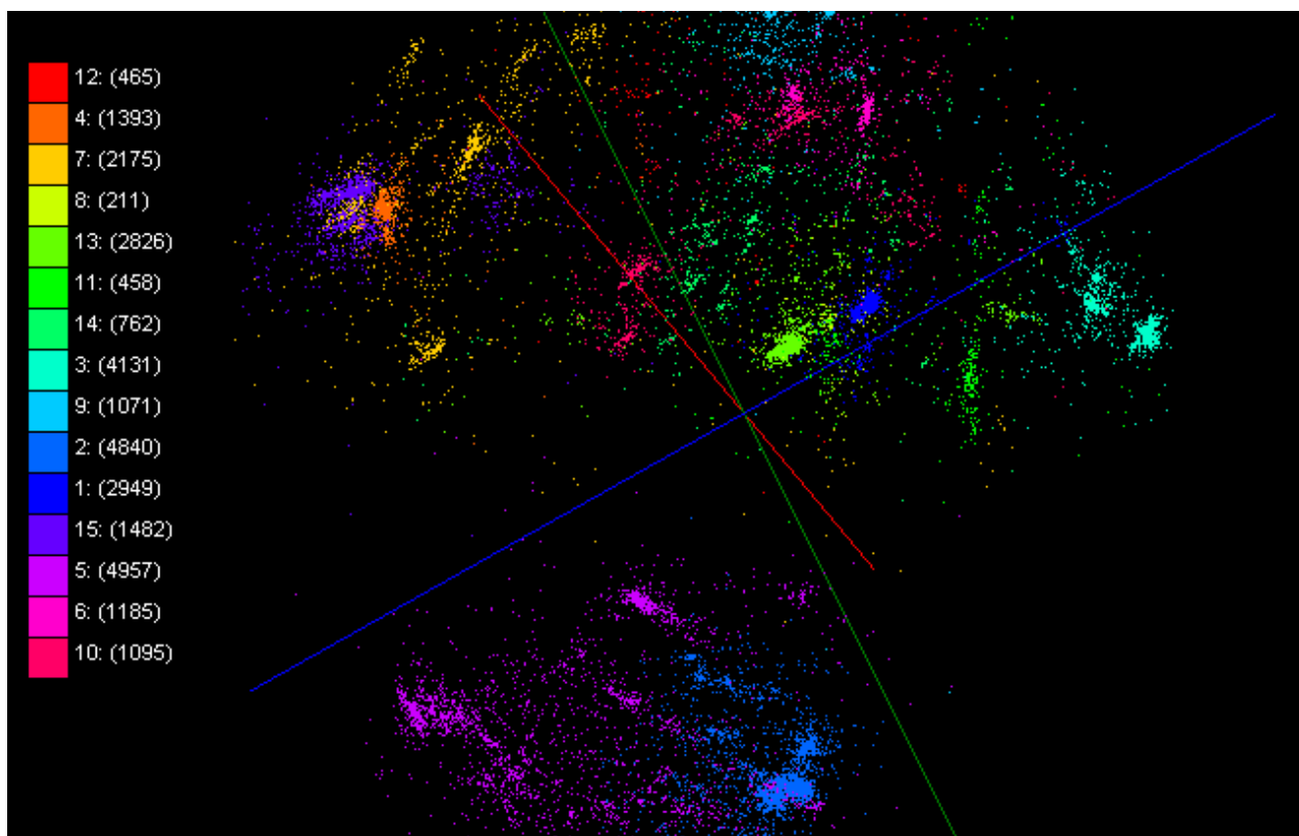


Figure 9. A 3D mapping result of the 30 thousand metagenomics sequences.

## 5. Conclusion

In this paper we discuss our design and implementation of PubChemBrowse, an integrated 3D data point visualization tool customized for browsing massive cheminformatics data. PubChemBrowse is designed to utilize our in-house high-performance visualization algorithms and be able to interact with external Chem2Bio2RDF system by using SPARQL to overlay information available from publicly accessible multiple bioinformatics databases. Further, we can embed new points and label various bioinformatics related datasets to assist on-going data mining research projects for drug-discovery.

One of the problems observed in PubChemBrowse is that plots can be cluttered as the number of points increases. This is a typical issue in visualizing large number of points because it is complicated and difficult to find structured mappings, or the amount of information needs to be visualized is large. Another problem is that there exist obvious hardware limits to prevent rendering very large number of points in a client side. As for the future work to overcome those problems, we will integrate cloud services into our system to display entire compounds as a hierarchical structure so that users can easily zoom in and zoom out on a specific region with different levels of details. We will also continue to extend our system to integrate with more external systems transparently via semantic web interfaces.

## 6. Acknowledgement

We appreciate Microsoft for their technical support. This work was made possible using the computing use grant provided by Amazon Web Services which is titled "Proof of concepts linking FutureGrid users to AWS". This work is partially funded by Microsoft "CRMC" grant and NIH Grant Number RC2HG005806-02. This document was developed with support from the National Science Foundation (NSF) under Grant No. 0910812 to Indiana University for "FutureGrid: An Experimental, High-Performance Grid Test-bed." Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the NSF.

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