



# The BioKET Biodiversity Data Warehouse: Data and Knowledge Integration and Extraction

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**Abstract.** Biodiversity datasets are generally stored in different formats. This makes it difficult for biologists to combine and integrate them to retrieve useful information for the purpose of, for example, efficiently classify specimens. In this paper, we present BioKET, a data warehouse which is a consolidation of heterogeneous data sources stored in different formats. For the time being, the scopus of BioKET is botanical. We had, among others things, to list all the existing botanical ontologies and relate terms in BioKET with terms in these ontologies. We demonstrate the usefulness of such a resource by applying FIST, a combined biclustering and conceptual association rule extraction method on a dataset extracted from BioKET to analyze the risk status of plants endemic to Laos. Besides, BioKET may be interfaced with other resources, like GeoCAT, to provide a powerful analysis tool for biodiversity data.

**Keywords:** Biodiversity, Information Technology, Ontologies, Knowledge Integration, Data Mining.

## 1 Introduction

Biological diversity, or biodiversity, refers to the natural variety and diversity of living organisms [26]. Biodiversity is assessed by considering the diversity of ecosystems, species, populations and genes in their geographical locations and their evolution over time. Biodiversity is of paramount importance for a healthy environment and society, as it ensures the availability of natural resources and the sustainability of ecosystems [6, 10, 13, 16, 22, 25]. The effects of biodiversity loss on the environment, caused by habitat loss and fragmentation, pollution, climate change, invasive alien species, human population, and over-exploitation can affect all life forms and lead to serious consequences [9]. Understanding biodiversity is an essential prerequisite for sustainable development.

For many years, biodiversity datasets have been stored in different formats, ranging from highly structured (databases) to plain text files, containing plant descriptions (vocabularies and terms). Numerous data and knowledge repositories containing biodiversity and environmental information are available on the Internet as on-line and off-line resources nowadays. Data repositories store

large amounts of information depicting facts on concrete objects related to a specific domain of application, e.g., results of environmental studies or inventories of species in a geographic location. This makes it difficult for botanists or zoologists to combine and integrate them to retrieve useful information for the purpose of identifying and describing new species.

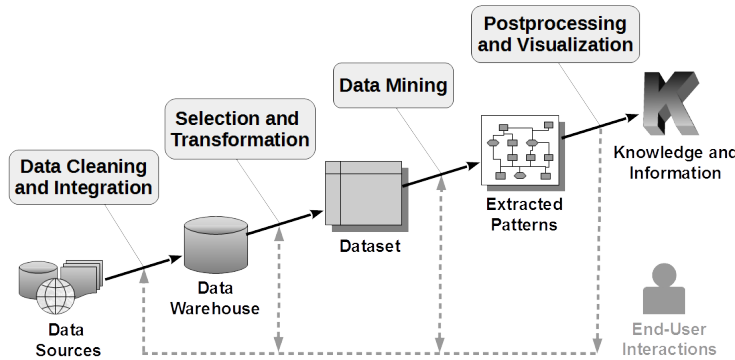
The ever increasing availability of data relevant to biodiversity makes the idea of applying data mining techniques to the study of biodiversity tempting [12].

Data mining, also known as knowledge discovery from data (KDD), is a set of concepts, methods and tools for the rapid and efficient discovery of previously unknown information, represented as knowledge patterns and models, hidden inside massive information repositories [11].

One important obstacle to the application of data mining techniques to the study of biodiversity is that the data that might be used to this aim are somewhat scattered and heterogeneous [24]. Different datasets cover different aspects of the problem or focus on some geographical areas only. None of them is complete and there is no standard format.

To overcome these limitation, we have designed and implemented BioKET, a data warehouse whose purpose is to consolidate a maximum of data sources on biodiversity in a logically organized, coherent, and comprehensive resource that can be used by the scientific community as a basis for data-intensive studies.

The main contribution of this paper is to provide a detailed account of how the BioKET data warehouse has been designed and populated, by consolidating and integrating multiple and heterogeneous sources of data. The reader should not underestimate the methodological challenges and the practical problems that had to be overcome in order to achieve that result. As all data mining practitioners agree, pre-processing, which includes data cleaning, integration, and transformation is the most time-consuming and critical phase of the data mining process [14, 15] illustrated in Figure 1.



**Fig. 1:** Main phases of a data mining process

We demonstrate the use of such resource by applying FIST, a combined biclustering and conceptual association rule extraction method already described

in the literature [17], on a dataset extracted from it to analyze the risk status of plants endemic to Laos.

## 2 The BioKET Data Warehouse

The BioKET data warehouse is the consolidation of four main data sources:

- BIOTIK [2] (Western Ghats of India and National University of Laos), which contains 652 species records;
- the herbarium from the BRAHMS repository [3] (National University of Laos, Faculty of Forestry), with 7548 species records;
- the herbarium from the NAPIS repository [18] (Lao Ministry of Public Health, Institute of Traditional Medicine), with 747 species records;
- the IUCN Red List Data [27], with 71570 species records.

These data sources are stored in different formats: BIOTIK and IUCN Red List are in HTML, while the two others use, respectively, the dBase and Paradox file formats. Integrating such diverse data sources required performing the following tasks:

1. Listing all botanical and plant ontologies available on the Internet.
2. Selecting relevant information (phenotypic/plant traits/features/characteristics).
3. Relating terms in our database with terms in these ontologies.
4. Searching for thesauri/glossaries/taxonomies of terms for plants available on the Internet.
5. Relating terms in our database with terms in these thesauri.
6. Relating terms in Plant Ontology (PO) (which seems to be the most complete ontology in Botany) with terms/definitions (e.g., Latin terms) in these thesauri.

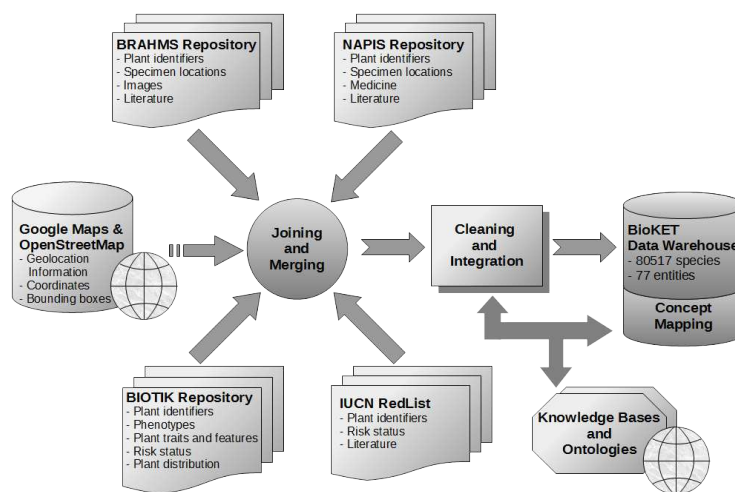
The first step was to extract data from sources and store them in a standard file format (such as an Excel spreadsheet), by using database management tools. Then, data cleaning was performed by using advanced Excel functions. The next step was to generate and link Google Maps Geocoding Service with the BIOTIK, BRAHMS, and NAPIS data by using VBA script (GoogleGeoLocation Function). The last step was to import the data thus obtained into the BioKET database, under MySQL.

A key factor for the integration and the enrichment of the data was the use of ontologies. Formal ontologies are a key for the semantic interoperability and integration of data and knowledge from different sources. An ontology may be regarded as “a kind of controlled vocabulary of well-defined terms with specified relationships between those terms, capable of interpretation by both humans and computers” [28]. From a practical point of view, an ontology defines a set of concepts and relations relevant to a domain of interest, along with axioms stating their properties. An ontology thus includes a taxonomy of concepts, a formally defined vocabulary (called a terminology), and other artifacts that help structure a knowledge base. A knowledge base that uses the terms defined in an

ontology becomes usable by and interoperable with any other system that has access to that ontology and is equipped by a logic reasoner for it [19].

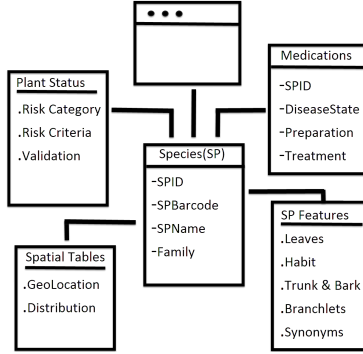
It was thus important to construct a map among all the concepts in all the data sources and all the considered ontologies. It is worth noting that (i) some concepts are not equally represented in all the sources, (ii) some are represented in some sources and not in others and (iii) other concepts are not represented at all. The mapping process works as follows: the textual descriptors of plants are segmented into small chunks, which are then matched with the labels of concepts in the target ontology. For instance, from the descriptor “evergreen tree up to 8 m”, we can infer that “evergreen” is related to “shedability”, “up to 8m” is related to “height” and “tree” is related to “plant type”. In the process, new concepts may be generated (e.g., from the textual descriptor “branches ascending or horizontal”, where “branch”, “branch ascending”, and “branch horizontal” match concepts in the ontology, a new concept “branch ascending or horizontal”, subsumed by “branch” and subsuming the latter two is generated). The plant record can thus be automatically enriched with a large number of “implicit” fields, inferred from the ontology. We have designed a relational data base of concepts that make it possible to relate concepts represented in different ways.

The result of this integration process — the BioKET data warehouse — is schematized in Figure 2. BioKET contains 77 relationship entities and a total of 80,517 records.



**Fig. 2:** An Overview of the BioKET Data Integration Process

As pointed out by many researchers (see, e.g., [1]), to conserve organisms, whether plant or animal, one important step to take is to identify rare and endangered organisms in a given geographical area or country. The integration of geographical information from Geographical Information Systems (GIS) with



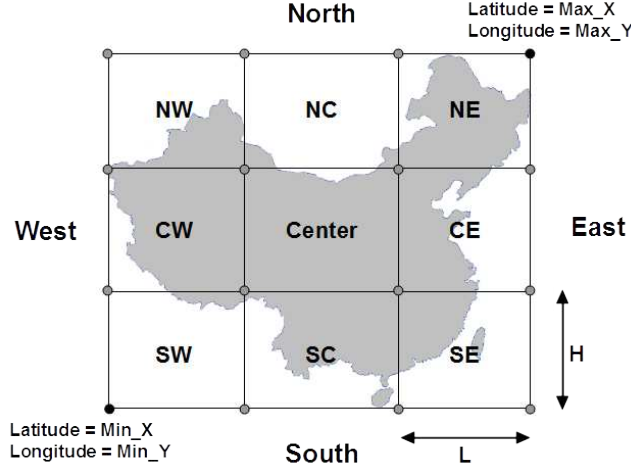
**Fig. 3:** Overview of the BioKET Datawarehouse Entity-Relationship Model

species data, and its use in data mining studies is the object of the biodiversity informatics project of the W. P. Fraser Herbarium (SASK) [20]. The participants in such project developed an integrated bio-geography GIS model, using Google Maps API, based on data mining concepts to map and explore flora data. This allows data to be explored on a map and analyzed in several ways to reveal patterns showing relationships and trends that are not discernible in other representations of information.

The BioKET data warehouse integrates geographical information and 8,947 species out of the 80,517 total species have descriptions of specimen location and risk status that may differ depending on the area considered. This information is described at different levels of precision, from continent to specific places such as cities or villages. For example, *Cratoxylum formosum* grows up in Myanmar, South China, Thailand, Indochina, and Laos (Khammouan) [2]. This species is also reported in the Lower Risk/Least Concern category by IUCN Red List data [27]. The integration of geolocation information allows to explore species properties in different areas using the GeoCAT (Geospatial Conservation Assessment Tool) platform [8]. GeoCAT is based on Google Maps to explore geographical information if coordinates, i.e., latitudes and longitudes, are provided. We already linked Google Maps with the terms of geographical information of each source (BIOTIK, BRAHMS, and NAPIS) and extracted their coordinates into BioKET database system. Google Maps does not support coordinates of directions (South, North, East, West, etc.) like “South China”, but Google Bounding Box (BBox) coordinates are provided. We propose to improve this issue by calculating the coordinates for each direction (Figure 4) from Google BBox coordinates.

In the geolocation domain, the BBox of an area on Earth is defined by two points corresponding to the minimal and maximal longitudes and latitudes of the area [4]. Figure 4 shows the 13 partitions of an area: the 9 elementary partitions and the North, South, East and West partitions that result of merging the 3 corresponding elementary partitions, e.g., NW, NC and NE for North. This

multilevel partitioning allows to represent location related properties of species, such as risk status or abundance for instance, at different area covering levels.



**Fig. 4:** Thirteen partitions for an area on Earth

Formulas to calculate the BBox of each partition are given in Table 1. These computations use the  $L$  and  $H$  values computed from the minimal ( $Min\_X$ ,  $Min\_Y$ ) and maximal ( $Max\_X$ ,  $Max\_Y$ ) longitude and latitude coordinates of the BBox of the partitioned area as follows:

$$L = \frac{(Max\_X - Min\_X)}{3}, \quad H = \frac{(Max\_Y - Min\_Y)}{3}.$$

This computation of partitions can be applied to all objects defined by a geolocation bounding box, from continents to cities. For example, using the BBox of China, that is  $\{73.4994137, 18.1535216, 134.7728100, 53.5609740\}$ , the BBox of Southern China will be computed as  $\{73.4994137, 18.1535216, 93.9238791, 53.5609740\}$ .

### 3 BioKET Experimental Analysis

For experimental purpose, we constructed a dataset containing information on the 652 species extracted from the Biotik repository. This information is represented as 1834 binary attributes describing morphological and environmental properties (characteristics of part of the plant, size, habitat, exudation, etc.) and risk status of species. Extracting knowledge patterns can then provide support to relate increases and decreases in risk status to environmental factors impacting specific species (climate change, pollution, etc.). They can also help taxonomists to analyze the different types of species in an ecosystem, e.g., associating species

**Table 1:** Partition Bounding Box Computations

Area	Min_Long	Min_Lat	Min_Long	Max_Lat
South	$Min\_Y$	$Min\_X$	$Min\_Y + H$	$Max\_X$
North	$Min\_Y + 2H$	$Min\_X$	$Max\_Y$	$Max\_X$
West	$Min\_Y$	$Min\_X$	$Max\_Y$	$Max\_X + L$
East	$Min\_Y$	$Min\_X + 2L$	$Max\_Y$	$Max\_X$
SW	$Min\_Y$	$Min\_X$	$Min\_Y + H$	$Min\_X + L$
SC	$Min\_Y$	$Min\_X + L$	$Min\_Y + H$	$Min\_X + 2L$
SE	$Min\_Y$	$Min\_X + 2L$	$Min\_Y + H$	$Max\_X$
CW	$Min\_Y + H$	$Min\_X$	$Min\_Y + 2H$	$Min\_X + L$
Center	$Min\_Y + H$	$Min\_X + L$	$Min\_Y + 2H$	$Min\_X + 2L$
CE	$Min\_Y + H$	$Min\_X + 2L$	$Min\_Y + 2H$	$Max\_X$
NW	$Min\_Y + 2H$	$Min\_X$	$Max\_Y$	$Min\_X + L$
NC	$Min\_Y + 2H$	$Min\_X + L$	$Max\_Y$	$Min\_X + 2L$
NC	$Min\_Y + 2H$	$Min\_X + 2L$	$Max\_Y$	$Max\_X$

with specific features and risk categories, and their viability, or growth rate, in some particular areas.

### 3.1 Conceptual Bicluster Extraction

This dataset was analyzed using the FIST approach which is based on the frequent closed itemsets framework. FIST extracts minimal covers of conceptual association rules and biclusters jointly.

Conceptual biclusters of the form  $\{I_1 \ V_1\}$  associate to a maximal set of instances  $I_1$ , a maximal set of variable values  $V_1$  that are common to all instances. In other words, a bicluster is a sub-matrix associating a subset of rows and a subset of columns such that all these rows have a similar value for each of these columns. Conceptual biclusters are partially ordered according to the inclusion relation and form a lattice. This hierarchical organization allows to explore groups of instances (species) and properties (characteristics) at different levels of abstraction: the highest biclusters in the lattice regroup a large number of properties shared by small groups of instances; the lowest biclusters regroup small set of properties that are common to large group of instances.

Conceptual association rules are rules with the form  $\{V_1 \longrightarrow V_2, I_1, support, confidence, lift\}$  where  $V_1$  and  $V_2$  are sets of variable values (properties) and  $I_1$  is the set of instances (species) supporting the rule. Statistical measures computed for each rule are:

- $support = P(V_1 \cup V_2)$  (or  $count(V_1 \cup V_2) = |I_1|$  if given as an absolute number) evaluates the scope, or weight, of the rule in the dataset. It corresponds to the proportion of instances containing  $V_1$  and  $V_2$  among all instances.
- $confidence = \frac{P(V_1 \cup V_2)}{P(V_1)}$  evaluates the precision of the rule. It corresponds to the proportion of instances containing  $V_2$  among those containing  $V_1$ . Rules with  $confidence = 1$ , that have no counter-example in the dataset, are called *exact* rules. Rules with  $confidence < 1$  are called *approximate* rules.

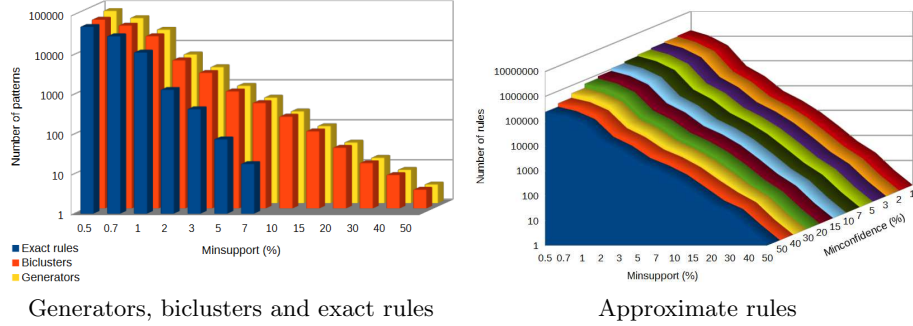


- $lift = \frac{P(V_1 \cup V_2)}{P(V_1)P(V_2)}$  corresponds to the correlation between occurrences of  $V_1$  and  $V_2$ :
  - $lift > 1$  means there is positive correlation between  $V_1$  and  $V_2$ ,
  - $lift = 1$  means  $V_1$  and  $V_2$  are independent,
  - $lift < 1$  means there is a negative correlation between  $V_1$  and  $V_2$ .

Extraction parameters are the *minsupport* threshold, which corresponds to the minimal number of supporting instances required for a rule to be considered valid, and the *minconfidence* threshold, which corresponds to the minimal *confidence* required in order to consider a rule valid.

Experiments were conducted on a Dell PowerEdge R710 server with 2 Intel Xeon X5675 processors at 3.06 GHz, each possessing 6 cores, 12 MB cache memory, 24 GB of DDR3 RAM at 1333 MHz and 2 Hot Plug SAS hard disks of 600 GB at 15000 rounds/min with RAID 0 running under the 64 bits CentOS Linux operating system.

The numbers of patterns extracted, i.e., generators, biclusters, and rules, are shown in Figure 5. For this experiment, the *minsupport* threshold was varied between 50% (326 species) and 0.5% (3 species). The *minconfidence* threshold was varied between 50% and 1%. It should be noted that the vertical axes are on a logarithmic scale.



**Fig. 5:** Number of Patterns Generated by FIST.

### 3.2 Extracted Pattern Evaluation

In this section, we present some interesting conceptual association rules obtained from FIST. We would like to stress that these results mainly depend on the data collected within BioKET, which, as far as we know, is the only data warehouse consolidating different biodiversity information sources. These rules make it possible to estimate the risk status of a plant species according to IUCN RedList categories (*Lower Risk, Endangered, Least Concern, Vulnerable, Critically Endangered, Rare, Data Deficient, Rare & Threatened, Possibly Extinct*) with respect to their characteristics and *vice-versa*. For this experiment, the *minsupport*

threshold was set to 1%, which corresponds to 6 species in the dataset, and the *minconfidence* threshold was set to 50%.

One of the obtained rules with the highest lift (11.75) is

INFL:pedicels up to 3 mm long, BBT:Twigs terete, INFL:axillary  $\Rightarrow$  RS:Lower Risk. (1)

According to this rule, of the six species with pedicels up to 3 mm long, twigs terete, and axillary inflorescence, 66,67% belong to the *lower risk* category. The identified species are *Cratoxylum cochinchinense*, *Cratoxylum formosum*, *Engelhardtia serrata*, *Engelhardtia spicata*, *Irvingia malayana*, and *Knema globularia*. This result is corroborated, for example, by information from Singapore flore.<sup>1</sup>

The following rule states, with 83.33% confidence, that a plant species classified as *critically endangered* has simple leaves:

RS:Rare  $\Rightarrow$  LEAVES:Leaves simple. (2)

This rule is corroborated, for example, by [7], which describes *Gaultheria paucinervia*, a new species restricted to the eastern slopes of Mt. Kinabalu in Sabah State, Borneo, Malaysia, which has been confused with *Gaultheria borneensis* Stapf, but differs in its more erect habit and larger stature, longer nonappressed leaf trichomes, purple (vs. white) fruiting calyx, and lower elevation range, among other features. *Gaultheria paucinervia* has not yet been assessed for the IUCN Red List, (but is in the Catalogue of Life: *Gaultheria paucinervia* P.W. Fritsch & C.M. Bush apparently). Besides, by taking into account the features in the geographical data source, the FIST algorithm finds the rule

RS:Rare, GEO:Western Ghats  $\Rightarrow$  LEAVES:Leaves simple, (3)

which identifies species *Bentinckia condapanna*, *Drypetes malabarica*, *Glycosmis macrocarpa*, *Holigarna grahamii*, *Lasianthus jackianus*, *Pittosporum dasycaulon*, and *Vepris bilocularis*, all found in the Western Ghats.

The following rule states, with 79.59% confidence, that a plant species classified as *Vulnerable* has simple leaves:

RS:Vulnerable  $\Rightarrow$  LEAVES:Leaves simple. (4)

This result is corroborated, for example, by [23, 29]. In [23], the author discusses the applicability of the Accelerated Pioneer-Climax Series (APCS) method for restoring forests to degraded areas in Southern Vietnam using many local species such as *Hopea odorata* directly concernend by the above rule and which has been identified as *vulnerable* in the IUCN red list. Wickneswari [29], instead, proposes a document which can help the readers to understand the entire life cycle of *Hopea odorata* Roxb in Malaysia, Vietnam, Cambodia, and Thailand.

<sup>1</sup> URL: <http://florasingapura.com/Home.php>. The aim of this site is to to bridge the gap between the terse technical descriptions of plants found in various botanical text books and what is observed in the Singapore forests.

The following rule, whose lift is 1.189 and whose support is 4.14%, states, with 55.1% confidence, that a plant species classified as *vulnerable* has both *glabrous* and *simple* leaves:

$$\text{RS:Vulnerable} \Rightarrow \text{LEAVES:glabrous, LEAVES:Leaves simple.} \quad (5)$$

Indeed, [21], proposing a deep and comprehensive botanical study of two rock outcrops in India, corroborates this rule.

Another interesting rule with a support of 3.37% and a lift of 1.07, states, with 59.46% confidence, that a plant species classified as having a *lower risk* has alternate leaves:

$$\text{RS:Lower Risk} \Rightarrow \text{LEAVES:alternate.} \quad (6)$$

This result is corroborated, for example, by results obtained by Craenel [5]. Species concerned include *Aglaia elliptica*, *Aphanamixis polystachya*, and *Prunus arborea*. As seen for Rule 3, the integration of geolocation information with multiple heterogeneous biological data can show common properties related to species with a specific risk status and/or in a specific area. For instance, the following rule with a lift of 4.26 states that 88.9% of species having a *lower risk* in the Indochina geographic region (i.e., 8 species) have leaves with entire margin:

$$\text{RS:Lower Risk, GEO:Indochina} \Rightarrow \text{LEAVES:Margin entire.} \quad (7)$$

Another example of such rules is the following, showing that 88.2% of *endangered* species in Western Ghats have alternate leaves:

$$\text{RS:Endangered, GEO:Western Ghats} \Rightarrow \text{LEAVES:Alternate.} \quad (8)$$

This rule, whose lift is 1.96, concerns 15 species. Such patterns can help comparisons between different geographical areas, at different levels of abstraction. For instance, considering the Malaysia geographic region, a part of Indochina, only 61.5% of species having a *lower risk* have leaves with entire margin as stated by the following rule, whose lift is 2.95 and which concerns 8 species:

$$\text{RS:Lower Risk, GEO:Malaysia} \Rightarrow \text{LEAVES:Margin entire.} \quad (9)$$

If we consider the Agasthyamalai area, lying at the extreme southern end of the Western Ghats mountain range along the western side of Southern India, we can see from the following rule that only 50% of *endangered* species in this area have alternate leaves, whereas the percentage is of 88.2% in the whole Western Ghats:

$$\text{RS:Endangered, GEO:Agasthyamalai} \Rightarrow \text{LEAVES:Alternate.} \quad (10)$$

This rule, which has a lift of 4.27, concerns 10 species.

All the above rules have been constructed from the consolidation of data from the four data sources presented above. Although some of the species are not yet included in the IUCN red list, combining information from the three other data sources allowed us to infer their risk status using the rules constructed by

FIST. This is the case, e.g., for the species related to Rule 3, with the sole exception of *Bentinckia condapanna*, whose risk category is explicitly in IUCN. Indeed, *Glycosmis macrocarpa*'s taxon has not yet been assessed for the IUCN Red List, but is listed in the Catalogue of Life as *Glycosmis macrocarpa* Wight. The same holds for *Drypetes malabarica* (in the Catalogue of Life as *Drypetes malabarica* (Bedd.) Airy Shaw), *Lasianthus jackianus* (in the Catalogue of Life as *Lasianthus jackianus* Wight), *Pittosporum dasycaulon* (in the Catalogue of Life as *Pittosporum dasycaulon* Miq), and *Vepris bilocularis* (in the Catalogue of Life as *Vepris bilocularis* (Wight & Arn.) Engl.).

## 4 Conclusion

We presented BioKET, a data warehouse obtained by consolidation of a number of heterogeneous data sources on biodiversity. As far as we know, this is the first data warehouse containing that amount of heterogeneous data which can be used for conducting data-intensive studies about biodiversity. For the moment, the scopus of BioKET is botanical, but we plan to integrate other types of data.

We have demonstrated the usefulness of BioKET by applying FIST, an existing conceptual biclustering method, on a dataset extracted from BioKET to analyze the risk status of plants endemic to Laos. The evaluation of the extracted patterns against the botanical literature shows that meaningful knowledge can be inferred from BioKET.

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