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Names are key to the big new biology

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Those who seek answers to big, broad questions about biology, especially questions emphasizing the organism (taxonomy, evolution and ecology), will soon benefit from an emerging names-based infrastructure. It will draw on the almost universal association of organism names with biological information to index and interconnect information distributed across the Internet. The result will be a virtual data commons, expanding as further data are shared, allowing biology to become more of a 'big science'. Informatics devices will exploit this 'big new biology', revitalizing comparative biology with a broad perspective to reveal previously inaccessible trends and discontinuities, so helping us to reveal unfamiliar biological truths. Here, we review the first components of this freely available, participatory and semantic Global Names Architecture.

The value of taxonomy to a biology that is changing

'New Biology' is a vision [1] of a discipline evolving to become considerably more data-intensive as it accommodates increasing amounts of under-analyzed data from high-throughput molecular and environmental technologies, and from large-scale digitization programs such as the Biodiversity Heritage Library (BHL, http://www. biodiversitylibrary.org/). In addition, there is pressure on scientists to make their data publicly available. More of the biological community will have access to on-line resources. Biology will shift towards the data-intensive 'big sciences' [2,3]. Web services that use names to index and organize information about organisms will be a critical part of this 'big new biology'.

This change will require an organizational framework that is able to manage billions of pieces of information about our current catalogue of 2 200 000 or so living and expired species. The information will be distributed across thousands of Web-sites. Three devices have the potential to organize information on all species. The first might use information from the molecular machinery that is common to all organisms. The second, phyloinformatics [4], would call on hypotheses through which the ancestor-descendent relationships within all life are explored. While the logics are appealing, neither phylogenetic nor genetic analyses have been applied to the majority, let alone all, species. Today, they would fail as comprehensive information management devices. Fortunately, the third option, taxonomy, extends to all formally described species and so offers a lifewide axis by which all biological information might be

Taxonomy is supported by 5000-10 000 professional taxonomists worldwide, http://www.gti-kontaktstelle.de/ taxonomy_E.html [7]. This 'team' [8] is united by principles founded in the codes of nomenclature. Taxonomists discover and describe biodiversity, arrange species into classifications with sensitivity to phylogenetic insights, are aware of all of the literature that bears on the identity of the taxa, and provide services to those who rely on authoritative information. However, many taxonomists feel unable to meet the expectations of the discipline, home institutions or exasperated users [9,10], and even believe that taxonomy as a scientific discipline is in danger of extinction [5,11,12]. Others argue that the 'information age' offers new opportunities to serve those who depend on taxonomic knowledge [6,13–15], and that using taxonomy to manage on-line biological information can reinvigorate the discipline [16]. A small community of innovative taxonomists, computer scientists and data managers (collectively 'biodiversity informaticians') are pursuing this vision and are building data standards, information exchange protocols, resources and services that can bring distributed data together as a virtual pool. Taxonomists use their expertise to add taxonomic principles, practices and knowledge as 'Taxonomic Intelligence', ensuring that the products are sensitive to the character of biology [17,18].

Taxonomy has two special features that suit it for re-use in biodiversity informatics. The first is the system of scientific names. Their almost universal use allows them to be treated as metadata to index biodiversity-related information, much as names are used in the index of a book. Secondly, classification schemes transform lists of names into organizational structures (ontologies) that group data, permit generalizing statements, allow users to infer properties of taxa, to expand or focus searches, or to browse information in a biologically relevant fashion. The value of names as metadata and classifications as ontologies led to the vision of a names-based infrastructure to serve biology [19]. This approach is used in major life-wide projects such as the Encyclopedia of Life, http://www. eol.org [20,21]. Now this approach is being transformed into a 'Global Names Architecture' (GNA) that aims to make the informatics potential of names and hierarchies freely available through the Internet.

To be effective in information management, GNA must overcome an array of challenges. It must index all references to organisms. It must bring together information on

organized [5,6].



Opinion

the same taxon even when different names are used to refer to it, and it will need to 'know' when the same name refers to more than one taxon. The system must be dynamic, adapting to changes in nomenclature, phylogeny or taxonomy [18]. To scale to the task, it must automatically draw on new information as it is published in authoritative on-line sources, a process that will be made possible by the widespread adoption of agreed protocols, standards and identifiers [22].

GNA will initially serve three areas of biology with interests in names. The first is taxonomy. Taxonomists use 'names' as tokens for concepts of species (and other taxa) and compile lists of names to catalogue and discriminate all approximately 1 900 000 named extant species and 250 000 named extinct species [23,24]. Species are indefinite objects and taxonomists necessarily dispute where their boundaries lie. Their views are referred to as taxonomic concepts [22,25–27]. The architecture must be able to discriminate competing concepts, and link all of them to specimens, georeferenced data, publications and other usages that inform the concepts.

The second area deals with names from the perspective of the Codes of Nomenclature. In this context, the meaning of a name derives from 'nomenclatural acts' that begin with the creation of a new name and include subsequent actions that refine or change it. The results are compiled as nomenclators: definitive listings of code-governed names, their orthography and bibliographic citations.

The third area, biodiversity informatics, is broader than taxonomy and nomenclature. Informaticians need to keep track of any string of alphanumeric characters that was used to refer to taxa. The strings include scientific names, vernacular names (which in some contexts are the formally preferred names, e.g. the Australian Standard Fish Names http://www.fishnames.com.au/) and surrogates for names. Surrogates include provisional names and specimen, culture or strain numbers which refer to a taxon. 'SAR-11' ('SAR' refers to the Sargasso Sea) was a surrogate name given in 1990 to an important member of the marine plankton. Only a decade later did it become known as *Pelagibacter ubique* [28].

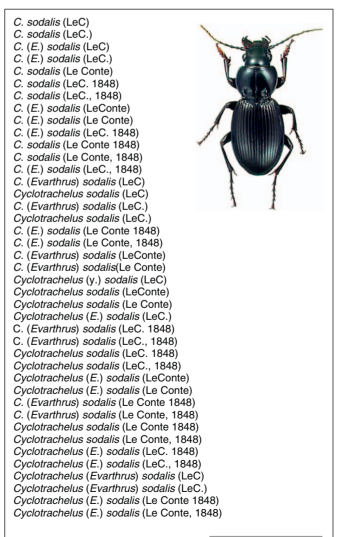
The names problems

The needs of taxonomists and nomenclators can be satisfied with relatively minor modifications of traditional practices. But, the biodiversity informaticians are encountering unfamiliar problems that confound the merger of distributed data. They require a more innovative system.

The largest problem is that of 'many-names-for-onespecies', where data on the same species have been indexed with different names. Until addressed, it prevents all information about the same species being brought together. This problem has many sources, such as when new research leads to the relocation of a species to a different genus. For example, a proposal to break up the genus that contains *Drosophila melanogaster* would lead to the species epithet '*melanogaster*' being combined with a different genus name to create a new binomial, *Sophophora melanogaster* [29,30]. Such taxonomic revisions create indisputable homotypic synonyms (*Drosophila melanogaster* and *Sophophora melanogaster* refer to the same species). The names infrastructure must bring together information that was published using either name. A second type of synonymy, heterotypic synonymy, occurs when a taxonomist opines that taxa previously considered distinct are the same. Again, the challenge is to bring information labeled under different names together. The solution to this problem must also manage vernacular names and surrogates.

Most of the alternative names for species come from different ways in which names are represented (Figure 1). Variants are caused by different styles of citing authors, how names are abbreviated, unintended errors, truncations or concatenations. As each string, right or wrong, is associated with one or more usages, all variants must be included within the indexing structure.

Two solutions address the 'many-names-for-one-species' problem. The first standardizes on a 'correct' name and seeks to apply that name universally. This is not viable because the chosen name will be arbitrary when, as is common, there is disagreement about the number of species or how each species should be defined. This solution cannot be applied retrospectively (at least not without the



TRENDS in Ecology & Evolution

Figure 1. Lexical variants of scientific names. A few of the valid alternative spellings of *Cyclotrachelus sodalis*, image from Canadian Biodiversity Information Facility (http://www.cbif.gc.ca/), used with permission.

Opinion

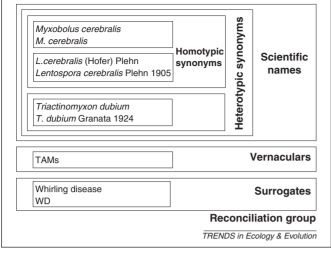


Figure 2. Reconciliation groups. A reconciliation group is an aggregate of all names used to refer to a taxon. It is comprised of one or more scientific names, with or without vernacular names or surrogates. Every name may be written out in one or more ways because the names of genera may or may not be abbreviated, and information about authorities may or may not be included. These lexical variants of names are included in the smallest boxes of the diagram. Homotypic synonyms include names that have the same type material – and they will have the same species element in the binomial name – *Myxobolus cerebralis* and *Lentospora cerebralis* are homotypic synonyms. The heterotypic synonymies are subjective, and emerge from a judgment by one or more taxonomists that *Triactinomyxon dubium* is the same species as was described as *Myxobolus cerebralis*. The vernacular names are non-scientific names that refer to the organisms. Surrogates are terms that also identify the taxon – in this case through the symptoms of the disease.

second solution); is costly to maintain, and does not cover vernacular or surrogate names. The second solution is to link together (reconcile) all known names for a given taxonomic concept (Figure 2). Reconciliation can be applied to any name, and preferred names can be 'flagged' to meet the needs of the first solution. With reconciliation, queries initiated with one name are transformed into actions involving all names.

A second names problem arises when one name is used for more than one taxon. *Bacillus* is a genus of stick insects and of bacteria, *Aotus*, a type of legume and a monkey. This problem risks bringing together information on different organisms, leading to incorrect outcomes. This problem increases as biological research becomes 'bigger' expanding from narrower taxonomic territories to include all taxa. Now the 14% of plant genera that have homonyms elsewhere shifts from an amusing anecdote to a serious problem for data integration [31]. The solution will register homonyms and apply disambiguating devices. Generic names, the most abundant source of homonyms, can be disambiguated with reference to taxonomic context, species names, authorship, or by the included taxa.

Components of GNA

GNA is being developed as a modular structure that can expand and adapt as opportunities and needs emerge. The initial elements (Figure 3) form a virtual layer that integrates information and services from sources (providers) to serve users (consumers).

At the core of GNA is a 'Global Names Usage Bank' (GNUB) that can index all published statements about life on Earth. The occurrence of a name on one or more occa-

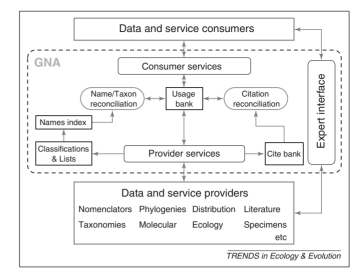


Figure 3. Initial elements of the Global Names Architecture. The virtual layer (dashed) is provided with services to access data and services from providers and to serve the needs of consumers. Given that many providers are also consumers, many interactions will be bidirectional. The central database is the Usage Bank. It indexes and points to occurrence of names in sources. Names, lists and classifications from users are first compiled in a names index and in a classification repository, and citations are captured in CiteBank. As the content comes from many sources, the form and quality will vary. Reconciliation services seek to normalize the content of these databases, contributing consistent, de-duplicated, standardized content to the Usage Bank. An interface allows users to add, comment or correct the underlying infrastructure or annotate records in the databases.

sions within a source constitutes a 'usage'. Usages occur in publications, field notes, databases and classifications, on web pages, specimen labels in museums, and herbarium sheets. Initially, the usage bank will emphasize usages that bear on nomenclature [32,33]. It will interconnect with prospective Web-based registry systems that will be used to formally establish new species instead of continuing the tradition of erecting new species in scientific publications [33]. Through its association with nomenclators, the usage bank will inform the names architecture of correct scientific names and their spellings, will link to taxonomic treatments and specimens to provide insights into synonymies and taxonomic concepts. The first iteration of the usage bank is ZooBank (http://zoobank.org/), the ICZN registry for names of animals [34,35]. Efforts are underway to incorporate nomenclators for fungi.

The names index (GNI) is a simple index of all unique forms of name strings (i.e. correctly and incorrectly spelled scientific names with or without author information, or nomenclatural annotations, or vernacular names, or surrogates for names). The index (http://gni.globalnames.org/) currently includes about 19 000 000 names. The index links to data held by contributors and provides a simple discipline-specific means of linking distributed information (Figure 4, model 'c'). NameLink, a prototype tool (http://labs.eol.org/?q=node/10/), recognizes names in documents and inserts anchors to which links known to GNI or to other digital objects can be attached. The names index is being enhanced with services to reconcile different versions of names and to disambiguate homonyms.

Biologists express their understanding of evolutionary relationships as classifications and trees. Both can be represented as parent–child structures, and are therefore interchangeable means of grouping or navigating data. Many catalogues of species, such as lists of marine species (http://www.marinespecies.org/), place lists within hierarchies of convenience. When the names are extracted to form simple lists, they can quickly filter data sets, instantly converting, for example, an encyclopedia of all life into an encyclopedia of marine life. Hierarchies can communicate insights into evolutionary history, and can be used to infer the distributions of attributes and test phylogenetic hypotheses. By accessing list and hierarchy repositories such as the GBIF Checklist Bank (http://names.gbif.org/), GNA can exploit the informatics and biological value inherent in parent_child structures and lists.

CiteBank (http://citebank.org/) is an open repository for bibliographic citations relating to biodiversity. It fosters collaboration to build definitive reference lists. With content coming from many sources, the styles of citation vary and CiteBank must provide reconciliation services to map variant forms together. CiteBank will include a document submission module to allow sharing of documents while complying with the 'safe harbor' principles of the Digital Millennium Copyright Act (http://www.copyright.gov/ legislation/dmca.pdf). The early version of CiteBank contains bibliographies of the BHL, other digital libraries, publishers, institutional repositories, and contributed bibliographies from specialist groups. CiteBank will have tools, like those in use by BHL, to find names in documents and automatically provide taxonomic indices.

Reconciliation and disambiguation services are being included to overcome the problems that accompany the federation of distributed but non-standardized information. Variously formed names and citations will have to be rendered into standard forms. This is achieved through reconciliation. First generation 'fuzzy' matching algorithms (http://www.cmar.csiro.au/datacentre/ taxamatch.htm) applied to names discover lexical variants and have reduced almost 19 000 000 names to about 6 000 000 reconciliation groups. Fuzzy matching is supplemented with parsing algorithms that reveal that, for example, 'Mycosphaerella eryngii (Fr. ex Duby) Johanson ex Oudem. 1897', 'Mycosphaerella eryngii (Duby) ex Oudem. 1897', and 'Mycosphaerella eryngii (Fr. ex Duby) ex Oudem. 1897', all contain the same canonical binomial, Mycosphaerella *eryngii*, allowing all these strings along with their fuzzilymatched variants to be placed in the same reconciliation group. With time, reconciliation services will bring together homotypic synonyms. Homonym discovery tools that flag homonyms and their children will minimize the risks of linking data on different taxa that have the same name.

The scale of the challenge to manage billions of data objects about millions of species arising on thousands of Web-sites can be addressed through algorithms and by promoting information exchange with machine-readable standards and protocols. Yet, the properties of the species include a myriad of idiosyncrasies and are defined by complex interactions that defy rule-based analysis and organization [36]. The automated processes will not serve biology perfectly. The names architecture compensates with an interface that allows experts to identify gaps, correct errors, disambiguate homonyms and help build reconciliation groups. Elements of the interface will allow names to be added, edited, 'deleted' or commented on; other functions will enable editing, merger or division of reconciliation groups, as well as the integration of vernacular and surrogate names. Flagging tools can be used to annotate names and their relationships, and finally, classification tools will allow users to build or improve classifications.

The big new biology needs to be readied to participate in newer trends of data integration, such as semantic data linking (Figures 4 and 5). As biologists digitize data and make them available through web services, they have relied on search engines and hyperlinks to make content discoverable and to draw attention to related data. More automated data federation has been made possible through the adoption of web services, data standards, universally unique identifiers and atomization of content. Now common keywords can foment a rich digital world of linked-data able to generate unsuspected insights [37]. A little time spent with Google Earth reveals how information generated for quite different purposes can be integrated using common denominators, such as georeferences, to deliver rich new services. The resulting semantic web has an almost anarchic quality, but it has enormous potential ([38], http://richard.cyganiak.de/2007/10/lod/). Semantic data-linking can be improved with services that manage

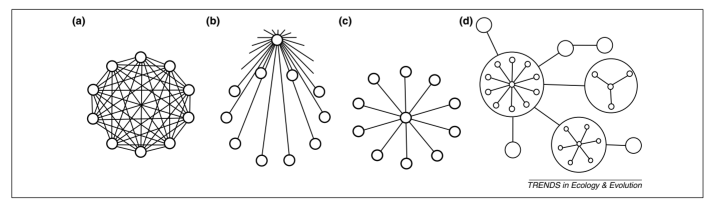


Figure 4. Models of interconnections among information at Web-sites. The oldest system (a) relies on hyperlinks between pages at each site. Such systems are costly to establish and maintain. Remote indexing of sites (b) is exemplified by search engines that allow users to visit one site and from it gain access to many other sites; the approach is very efficient but, as a 'one-size-fits-all' solution, may fail to accommodate the nuances of a discipline. A hub and spoke model (c) relies on specialist knowledge and agreed communications protocols to show content from a distributed array of sites through a unified specialist portal; structures like this for biology are made possible through a names-based infrastructure. The future will likely witness integrated linked systems (d) that flexibly interconnect differing arrays of single Web-sites, data in semantic data clouds, and indexed systems. The appeal is that such systems can evolve into increasingly complex and customizable structures as more indexing and management services become available.

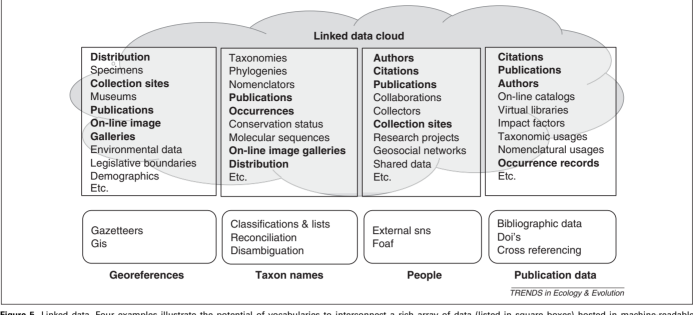


Figure 5. Linked data. Four examples illustrate the potential of vocabularies to interconnect a rich array of data (listed in square boxes) hosted in machine-readable environments, and provide the basis of cross-walks among knowledge areas (bolded terms). Georeferencing, publications and social networks (people) are already enhanced by services (rounded boxes), such as FOAF (Friend of a friend) that can be used to describe or build networks (e.g. http://network.nature.com/) or through technologies which interconnect the scientific literature (such as serviced by CrossRef, http://www.crossref.org/, with almost half a billion links). Other links may identify where individuals have collected specimens and, being attached to location, provide for potential cross-walks to other data linked by georeferences such as maps, images, habitat type, environmental conditions, local institutions and experts. Names of organisms, enhanced with reconciliation and disambiguation services, open up almost any data relating to organisms. As an illustration, a 16S sequence of a rare spider in GenBank, through latitude-longitude metadata, allows access to maps showing where in Argentina this organism was found, integrates the result with distributional data from the Global Biodiversity Information Facility that show the known world distribution, lists all publications in the Biodiversity Heritage Library that refer to the organism, all publications by the data depositor and of her colleagues, her obituary, and a description of the species emphasizing venomousness caused by a toxin that blocks sodium channels and has an LD50 of 0.0005 mg/kg.

discipline-specific data, metadata and ontologies. Datalinking for biology will benefit from rich services associated with taxonomic names, such as those that address the names problems (Figure 5). To fulfill this role, the GNA will emphasize web services that broadcast and collate new knowledge in forms that are readily understood by other machines.

And where is all this heading?

One reflection of the big new biology will be a biologically informed Internet. Users of search engines will find all information about a species irrespective of which name was used; no longer will biologists need to unpack nomenclatural history, but can expect systems to know that much of the information about *Pneumocystis pneumoniae* can be found under the name *Pneumocystis carinii*. We can expect electronic documents to be automatically brought up to date in matters nomenclatural and taxonomic, and for names in documents viewed with browsers to automatically link to other resources of our choosing.

The first beneficiaries of GNA will be the communities from which its architects and engineers are drawn. Nomenclaturalists will have access to on-line reference information cross-linked to searchable page images from on-line virtual libraries. Taxonomists will be able to check on all previously used names and will not create new homonyms. They will register new species quickly and easily, linking them to descriptions on-line without the delays associated with representing knowledge with ink on paper. Taxonomists will adapt on-line classifications to suit their own needs, and the parent-child statements they create will be captured and drawn together to assemble an editable and

tious treatment and the quality of data will improve. The capacity of this 'crowd sourcing' to be creative as well as

complementary data.

critical was powerfully demonstrated with Open Mapping that produced the most useful maps in the immediate aftermath of the recent Haitian earthquake (http://haiti. openstreetmap.nl/). With a virtual data commons, data become part of a dialog, and we can expect more tools to allow users to annotate data, or for nature lovers to confirm, deny or track the spread of invasive species or to register biological responses to climate change. Connections among previously unassociated data will provide a fertile pasture to nourish new hybrid scientists who combine biology and computer sciences. From them and those working at the boundaries of the different subdisciplines of biology we can expect a flush of new services, analytical tools and visualizations to reveal trends, patterns and dis-

dynamic catalogue of all life. Ecologists will find services to

ensure that they identify components of their ecosystems

correctly, and text-editing programs will prompt authors

with the correct names for their objects of study. 'Normal-

izing' names-services will correct names in databases and

data-linking projects will use common identifiers to merge

applications. Biologists of the future, assisted by GNA, can

expect services to keep them abreast of new information

about clades or taxa of interest. Users will have access to

bigger and broader arrays of data, with valuable datasets

identified with automated pointers that inform us that, for

example, other ecologists and molecular biologists who

used this data set also used those other data sets. Through

their availability, suspicious data can be flagged for cau-

The potential of data-linking is evident from mapping

Opinion

continuities in data. They will take an unfamiliar, distant view of the knowledge landscape that is biology to reveal patterns not evident from reductionist approaches, and will direct our attention to features of the underlying biology that deserve study. As a reinvention of comparative biology, such tools will become the 'Macroscope' [39,40] able to extract new insights from the big new biology.

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