LINK INTEGRATOR
A Link-based Data Integration Architecture

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Abstract: The evolution of the World Wide Web has created a great opportunity for data production and for the construction of public repositories that can be accessed all over the world. However, as our ability to generate new data grows, there is a dramatic increase in the need for its efficient integration and access to all the dispersed data. In specific fields such as biology and biomedicine, data integration challenges are even more complex. The amount of raw data, the possible data associations, the diversity of concepts and data formats, and the demand for information quality assurance are just a few issues that hinder the development of a general proposal and solid solutions. In this article we describe a lightweight information integration architecture that is capable of unifying, in a single access point, several heterogeneous bioinformatics data sources. The model is based on web crawling that automatically collects keywords related with biological concepts that are previously defined in a navigation protocol. This crawling phase allows the construction of a link-based integration mechanism that conducts users to the right source of information, keeping the original interfaces of available information and maintaining the credits of original data providers.

1 INTRODUCTION

World Wide Web and its associated technologies are evolving rapidly as is the ability to develop and deploy customized solutions for users. It is becoming easier over time to create novel applications with attractive interfaces and advanced features. However, this evolution leads to several problems. The major problem is in finding information with certified quality: being extremely easy to deploy new content online, the amount of incorrect and invalid information is growing. This process also leverages content heterogeneity. The same concept can be stored in different media formats or database models and accessed by different kinds of methods. Data integration architectures propose a single unifying framework that can connect to several distinct data sources and offer their content to users through web applications, remote services or any other kind of data exchange API.

Despite the advances in application development and web standards, integrating information is still a challenging task. This difficulty can be mostly seen in life sciences. Biology and biomedicine are using information technologies to solve their problems or share new discoveries. The Human Genome Project – HGP (Collins et al., 1998) – required novel software applications that could help in solving biological problems faster, ease biologists’ everyday tasks, aid in the publication of relevant scientific discoveries and promote researchers’ communication and cooperation. This exponential necessity empowered a new field of research denominated bioinformatics, requiring combined efforts from biologists, statisticians, and software engineers. The success of HGP brought about a new wave of research projects (Adams et al., 1991, Cotton et al., 2008) that resulted in a dramatic increase of information available online.

With the tremendous amount of available data, integration is the most common issue when developing new solutions in the area of bioinformatics. Applications like UniProt (Bairoch et al., 2005) and Ensembl (Hubbard et al., 2002) or warehouses like NCBI (Edgar et al., 2002, Pruitt and Maglott, 2001) aim to integrate biological and biomedical data that is already available. The applied data integration strategies are suitable when the purpose is to cover large data sources that use standard data transfer models. However, when the
data is presented in pure HTML, REST web services, CSV sheets or plain text, the integration process can be quite complex. Along with this problem there is the fact that when small applications are integrated in this environment, the content authorship is lost. This means that the original researcher, who made a great effort to publish his work online, will be hidden behind a small link contained in a simple list among similar links. The architecture described in this paper solves both these problems. We propose an integration model that can easily integrate information by storing its URL and displaying it inside a centralized application. With this, the data integration problem is partially solved and the original application is presented and extended to users.

The following section debates Link Integrator organization and section 3 describe a real world implementation scenario. Finally, section 4 contains some final remarks about our research.

2 LINK INTEGRATOR

To deal with the presented integration issues, one can adopt several distinct strategies for data integration. These approaches differ mostly on the amount and kind of data that is merged in the central database. Different architectures will also generate a different impact on the application performance and efficiency.

Warehouse (Polyzotis et al., 2008, Reddy et al., 2009, Zhu et al., 2008) solutions consist in replicating integrated data sources in a single physical location with a unifying data model. Mediator-based solutions (Haas et al., 2001) rely on a middleware layer for the creation of a proxy between the client and the original integrated servers. Link-based integration (Maglott et al., 2007, Oliveira et al., 2004) strategies simply list URLs linking the original data sources. Arrais work (Arrais et al., 2007) presents a solid analysis on these strategies, their main advantages and disadvantages, resulting in an optimal hybrid solution.

In addition to these strategies, current trends involve developments in the meta-applications area. Mashups (Belleau et al., 2008) and workflows (Oinn et al., 2004) have a growing popularity among data and service integration architectures. When the goal is to offer real-time web-based dynamic integration, with an increased user-control, Lopes’ work (Lopes et al., 2008) presents a valuable solution that can be adapted to several scenarios.

We have chosen to design a solution based on link integration scenarios. Our choice is mostly due to the fact that the biological content available online is dynamically changing and evolving swiftly.

2.1 Architecture

Link Integrator architecture metaphor relies on typical three-layer architecture. The proposed architecture divides the application structure: data access, application logic and presentation layer.

The data access layer is completely independent from our application. It represents the external data sources that are accessed by originally integrated applications. The application logic layer is crucial in our system: externally it deals with the communications with the integrated applications; internally it deals with the application execution cycle. The presentation layer is where any user in his web browser can access a single unifying interface, designed to be attractive and fulfilling high quality usability patterns.

It is important to highlight that our system does not have any kind of communication or data exchange with the original data sources. This type of activity would breach the initial system requirements, transforming the integration model from link-based to mediator-based.

2.2 Execution

Link Integrator is composed of a map, containing navigational information that the crawler will read to configure its processing cycle; a topic driven web crawler capable of parsing and processing the URLs it gathers and a relational database to store the information gathered by the crawler.

In this system, the most important part is the web crawler. The map contains mostly URL addresses and regular expressions that are used by the crawler to retrieve, parse and process web documents. The crawling results are triplets containing an association between: an Entity, which is the generic category where the found URLs will fit; a Class that represents the container where the link belongs; a link to the Application, the URL, which is the main result of the crawling cycle. Both the Entities and the Classes are defined in the map; the crawler only finishes the triplet by associating the URL with the Class and with the Entity.

The system operation cycle – Figure 1 – is simple. Initially, the system reads a predefined navigation map containing Entities, Classes, URLs and regular expressions to find other URLs. The crawler will then parse and process the web pages generating sets of results – URL-Class-Entity triplets – that will be stored in containers. These containers
can be serialized in XML and passed to another application or, as we do, stored in a relational database. After this crawling and retrieving phase, the application engine generates, from the data stored in the relational database, a tree that is presented to the user in the application’s main workspace.

3 DISEASECARD

DiseaseCard is a link-based integration application with the main purpose of connecting and presenting disease related information spread throughout the Internet. It was first launched in 2005 and has grown to a mature state where up to 2000 rare diseases stored in the Online Mendelian Inheritance in Man (Hamosh et al., 2005) database are covered. When DiseaseCard was created, web applications where mostly based on static HTML layouts to display content. Along with Internet evolution, web application complexity has also evolved: Web2.0 dynamic applications and web service responses cannot be parsed by traditional web crawlers.

Locus specific databases contain information that is as important for researchers as the information already gathered in DiseaseCard. These databases store information about genetic variations and relating this information with other genetics marks can offer newer insights on rare disease studies. Joining DiseaseCard paradigm with amount of information contained in LSDBs we have the perfect test bed for Link Integrator. Adding LSDBs to DiseaseCard will increase the added value it represents in the biomedical community and will validate the goals we set when developing our system. The new on development version of DiseaseCard – Figure 2 – will be supported by the Link Integrator engine that can join information from several LSDBs applications like LOVD (Fokkema et al., 2005) or UMD (Al and Junien, 2000). The current portal is available at www.diseasecard.org.

4 CONCLUSIONS

Taking into account the amount of data available after the Human Genome Project, the area of life sciences is probably the discipline most affected by information quantity, diversity and heterogeneity. Recently, an area of growing importance is related to the human variome. Variation studies often result in
new web applications denominated locus specific databases and they usually contain information about sequence variations among individuals for a particular gene. In addition, content ownership and its growing importance is gaining relevance. Despite the fact that for regular end-users, access to scientific content is easier when provided by a centralized service, researchers who want to publish their work are almost obliged to create their own applications if they want to keep the authorship of their work visible.

The described architecture and application intend to overcome these problems with three key features for both users and researchers. First, integration is based on simple Internet URLs that are parsed and processed to gather the most significant information. This means that developers will not have to make any changes to the application core and that we are able to integrate any URL-accessible content. Secondly, the original applications will be shown inside our application. Thus, the content owners will not be shown as a link but as part of a complete application. Finally, external applications can be extended inside our system: information exchanges, text-mining and other user customization features can be developed to enhance the original applications.

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