

DESIGN OF A STANDOFF OBJECT-ORIENTED MARKUP LANGUAGE (SOOML) FOR ANNOTATING BIOMEDICAL LITERATURE

Jing Ding and Daniel Berleant

Department of Electrical & Computer Engineering, Iowa State University, Ames, IA, USA

Keywords: text mining, markup language, bioinformatics

Abstract: With the rapid growth of electronically available scientific literature, text mining is attracting increasing attention. While numerous algorithms, tools, and systems have been developed for extracting information from text, little effort has been focused on how to mark up the information. We present the design of a standoff, object-oriented markup language (called SOOML), which is simple, expressive, flexible, and extensible, satisfying the demanding needs of biomedical text mining.

1 INTRODUCTION

With the rapid growth of electronically available biomedical literature, information extraction from unrestricted text is attracting increasing attention. While numerous algorithms and tools have been developed for extracting the information, little effort has been focused on markup methodology, *i.e.*, how to annotate and manage the extracted information. Among the most intuitive solutions are inline XML¹ markup, as in the GENIA corpus (Kim *et al.*, 2003) or the Medstract Gold Standards Corpora². Although straightforward, inline markup is not sufficient for dealing with complex annotation structures, nor does it cope well with other issues such as copyright and annotation reuse, *etc.* We propose a standoff object-oriented markup language (SOOML) and present its design. The design is based on software engineering principles and intended to meet the challenging needs of text mining annotations.

In section 2, we analyze the requirements of text mining annotations and review existing markup methodologies. Section 3 describes the concepts and mechanisms in the design of SOOML. A brief discussion of how the design meets the requirements is presented in section 4.

¹ <http://www.w3.org/XML>

² <http://medstract.org/gold-standards.html>

2 MOTIVATION OF THE WORK

Annotation structures in bioinformatics text mining can be complex, as illustrated in Fig. 1. In the figure, we wish to annotate three entities, *protein kinase C alpha isoform*, *protein kinase C beta isoform* and *Akt*. The surface strings of the first two entities not only contain gaps, but also overlap with each other. There are also two events in the sentence fragment. The events and the entities have hierarchical relationships. For instance, *entity[3]* plays a role in *event[a]*; and *event[a]* itself plays a role in *event[b]*. This complexity poses demanding requirements for a markup language.

2.1 Annotation requirements

Expressiveness: Complex annotation structures (gaps, overlaps, and hierarchies, *etc.*) can be expressed easily and intuitively.

Resolution: In many text-mining applications, it is important to show where the extracted information is in the original text. The required resolution varies.

Reusability: The language must be able to support annotation reuse—*i.e.*, annotating annotated text.

Independence of availability: The availability of annotations should not be restricted by the copyrights on the original documents.

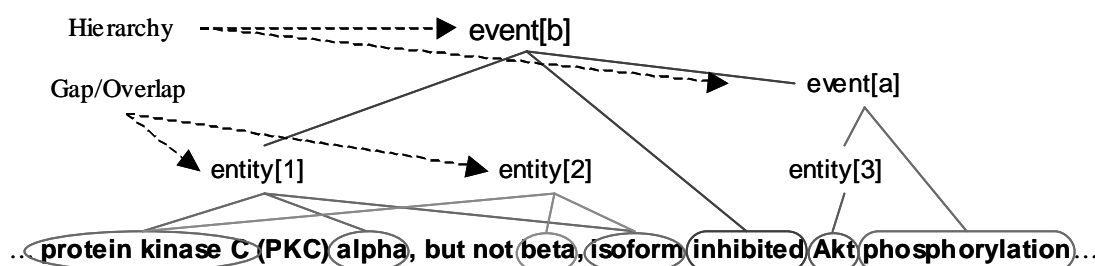


Figure 1: Sample annotations in biomedical domain.

Flexibility: The language should adapt well to various text formats and annotation needs.

Efficiency: It should be time and space efficient in storing, retrieving and processing annotations.

Extensibility: The language should be easily extensible to incorporate biomedical ontologies (e.g., UMLS³, and Gene Ontology⁴).

2.2 Existing works

There are some initiatives to develop “standard” annotated corpora in the bioinformatics field, notably the GENIA corpus and the Medstract Gold Standards Corpora. Both annotate MEDLINE⁵ abstracts using in-line XML tags. In-line markup has some limitations:

- Gapped, overlapped or hierarchical structures cannot be easily expressed.
- Availability of the annotations may be restricted by the copyright of the original document.
- In-line markup “contaminates” the original documents. The “contamination” complicates pipelined processing, because later stages cannot see directly the original text.

A solution to these limitations is standoff markup (annotations stored separately from the original documents). With the standoff approach, the question arises as to how to map an annotation back to its original text. The annotation graph and W3C’s XPointer framework address the issue.

The annotation graph (AG) is an abstraction of existing standoff annotation formats used by linguistic databases for textual and audio signals (Bird and Liberman, 1999). The signals are viewed as one-dimensional streams. An AG is a directed graph with nodes representing time points or character positions in a stream and with edges representing the text between two positions or audio content between two time points. Annotations are labeled on the edges. Conceptually, AG is equivalent to inline XML, with nodes corresponding to start and end

tags, and edges the text enclosed in tag pairs. Hence, AG has the same limited expressive power as inline XML.

The XPointer framework⁶ is a W3C standard for identifying text fragments in XML files. Two mechanisms (XPath⁷ and string matching) are used in XPointer for different resolutions. XPath is for identifying text nodes (text fragments, such as titles or abstracts in MEDLINE files). The string-matching mechanism is for locating exact words within a node. Together, they provide the resolution required by SOOML. However, XPointer has drawbacks. First, XPointer expressions can be long and complex. For example, the XPointer expression for entity[1] in Fig. 1 would be something like this:

```
xpointer(
  string-range(//MedlineCitation[PMID=1234]//AbstractText,
    'protein kinase C') |
  string-range(//MedlineCitation[PMID=1234]//AbstractText,
    'alpha') |
  string-range(//MedlineCitation[PMID=1234]//AbstractText,
    'isoform'))
```

More than 200 characters are needed to describe the original string of 29 characters. Another drawback is that ambiguity may arise if the matching string occurs multiple times in a text node. To avoid the ambiguity, the expression has to be longer and more complex. This complexity and inefficiency make XPointer not an ideal match to the requirements.

3 DESIGN

Since none of the existing systems mentioned above fully meets the requirements, we therefore designed a standoff object-oriented markup language (SOOML), which defines one concept (monad) and four mechanisms (object orientation, standoff markup, annotation mapping, and annotation inclusion). Please keep in mind that SOOML is not another new member of the ever-growing xxML fam-

³ <http://www.nlm.nih.gov/research/umls/>

⁴ <http://www.geneontology.org/>

⁵ <http://www.ncbi.nlm.nih.gov/entrez/query.fcgi>

⁶ <http://www.w3.org/TR/xptr-framework/>

⁷ <http://www.w3.org/TR/xpath>

Token: ...protein kinase C (PKC) alpha, but not beta, isoform inhibited Akt phosphorylation...
Monad: ... 12 13 14 15 16 17 18 19 20 21 22 23 ...

Figure 2: Monads tokenized at white space.

ily, such as GPML (the schema used in the GENIA corpus) or SBML (Systems Biology Markup Language) (Hucka *et al.*, 2003). A typical xxML defines a set of XML tags (a small ontology) for a specific purpose, usually focusing on WHAT is marked up. SOOML, on the other hand, is more interested in WHERE is the markup. Another important difference is the relationship with XML. While those xxMLs are all extensions to XML, SOOML has no direct connection with XML. It is a set of concepts and mechanisms, which can be implemented in XML syntax, C/Java-style syntax, or any other custom-made syntax. For simplicity, we will use Java-style syntax to show examples in the following sections.

3.1 The monad concept

The monad concept as used here is a variant of the concept with the same name in the text database field (Doedens, 1994). A stream of text is broken down into atomic units (tokens) at specified delimiting characters (*e.g.*, the space character). Integer indexes are assigned to each token in the order of text-flow. A monad is a token plus its assigned index. Delimiters may be counted as monads or discarded. If no delimiter is specified, each character is a monad. Fig. 2 shows a list of monads tokenized at the space character with the delimiters discarded. The **context** of a monad is the underlying text stream (the **text context**) along with the delimiting character set and whether delimiters are included or discarded (the **delimiting context**). Thus, given a context, an integer is unambiguously mapped to a text token, and vice versa.

3.2 Mechanisms

Object orientation. All annotations are instances of various classes. All classes are directly or indirectly derived from the base class annotation (*Inheritance*). An annotation object may contain other objects as components (*composition*) and is responsible for storing the extracted information (*data abstraction and encapsulation*).

Standoff. A collection of various objects is stored in an annotation file separate from the original document. The objects are identified by their class types and unique IDs.

Annotation mapping. Annotation mapping is achieved at three levels: document, text node and within a node.

1. An annotation is mapped to its original document using W3C's URI (Uniform Resource Identifier) addressing⁸. For example:

```
http://www.iastate.edu/~berleant/med35.xml
ftp://www.pub.iastate.edu/users/index.htm
file://c:\My Documents\manuscript\SOOML.txt
```

2. Within an XML document, text nodes are addressed using XPath. For example, the following XPath expressions map to the title and the abstract, respectively, of the MEDLINE citation whose PMID is 12345, in a given MEDLINE XML file:

```
//MedlineCitation[PMID=12345]//ArticleTitle
//MedlineCitation[PMID=12345]//AbstractText
```

3. Using the monad concept, mapping to a location within a node is trivial. It is done by simply listing the monads of an annotation. For example, given the context in Fig. 2, some of the annotations in Fig. 1 may look like the following:

```
context.file = <file address>;
context.node = <node address>;
context.delimiters = " ";
context.includeDelimiters = no;
entity[1] = {12, 13, 14, 16, 20};
entity[2] = {12, 13, 14, 19, 20};
entity[3] = {22};
event[a] = { actor1 = entity[3],
            action = {23},
            actor2 = null};
event[b] = { actor1 = entity[1],
            action = {21},
            actor2 = event[a]};
```

Annotation inclusion. An annotation file can use annotation objects defined in other annotation files. Suppose the entity objects in Fig. 1 are defined in a file named *entities.xml*. Then other files can include *entities.xml* and reference the entities by their names, prefixed with a unique identifier string assigned to *entities.xml*. For example:

```
#include "entities.xml" prefix "a"
...
event[a] = { actor1 = a:entity[3],
            action = {23},
            actor2 = null};
event[b] = { actor1 = a:entity[1],
            action = {21},
            actor2 = event[a]};
```

4 FULFILLING THE REQUIREMENTS

The four mechanisms work closely together to meet all of the text-mining annotation requirements.

Expressiveness: The requirement of expressive power has two aspects—the need to express complex hierarchical structures and the need to express arbitrarily distributed surface strings. For the former, the object-oriented mechanism enables modeling of complex hierarchical structures, similar to the use of object-oriented programming languages in modeling software application environments. For the latter, the monad-based mapping mechanism enables listing the tokens anywhere in a document.

Resolution of annotation mapping: Monad-based mapping also enables annotation mapping at various resolutions. Monads are the atomic units in SOOML. Their sizes, hence the mapping resolutions, are determined by the set of delimiting characters. For example, “insulin-induced” can be treated as a single monad (an adjective) in linguistic part-of-speech tagging, so it need not be tokenized at the character “-.” However, this resolution is not enough for protein name recognition. Therefore, it should be split into two monads (“-” being a delimiter). The finest resolution is single character mapping (using “null” delimiting character).

Reusability: The inclusion mechanism enables annotation reuse, facilitating modular design of complex text-mining systems in accordance with software engineering principles. It is not necessary to design a single powerful super-module to extract the information all at once. Specialized modules can target particular aspects of a complicated task and create annotations on top of each other. The standoff mechanism leaves the original documents unchanged, thereby avoiding interference among different modules and/or applications.

Independence of availability: The standoff mechanism separates annotations from the original documents, and the monad-based mapping mechanism avoids copying any contents from the original text. Therefore, the availability of the annotations is independent of the originals.

Flexibility: SOOML’s mechanisms enable annotating files of various formats in a consistent way (“one shoe fits all”). First, the standoff mechanism separates annotations from the original documents; therefore, the formats and the organization of the annotations are not restricted by those of the original documents. In contrast, in-line markup methods have to follow the formats of the original texts. Second, although SOOML’s mapping mechanisms (es-

pecially node-level mapping using XPath) are designed for XML-based original documents, they can be extended easily to any document with well-defined fields or sections, because they are conceptually equivalent to the nodes in XML documents. The worst case is for those files without any apparent internal structures. SOOML can still treat such a file as a single large text node, and monadize it from the first token to the last.

Efficiency: The monad-based mapping mechanism is space efficient. Instead of copying the content from the original documents, it uses monads (equivalent to pointers) pointing to the sources. For example, it takes only five integers to mark up entity[1] in Fig. 1, while XPointer needs over 200 characters. This makes SOOML an ideal format for annotation storage and exchange, as well as for serving as an intermediate data-flow format among the modules/applications.

The monad-based mapping mechanism also greatly reduces the complexity of annotation processing. First, gapped and overlapped annotations are handled in exactly the same way as continuous and non-overlapped ones. Second, monad-based mapping does not create any ambiguities, which are inevitable in string matching-based processing.

Extensibility: Because it is object oriented, SOOML can be integrated readily with other ontologies. Ontologies typically already have well-defined hierarchical structures. All we need to do is define the main ontology entries as subclasses of the annotation class (or one of its subclasses). The rest of the ontology is automatically included in the hierarchy.

In conclusion, we presented here the design of a standoff object-oriented markup language (SOOML), which provides an expressive, efficient, flexible and extensible framework for text annotation in bioinformatics – as well as other similar applications.

REFERENCES

- Bird, S. and Liberman, M. (1999) A Formal Framework for Linguistic Annotation. *Technical Report MS-CIS-99-01, Department of Computer and Information Science, University of Pennsylvania.*
- Doedens, C.-J. (1994) in *Text Databases. One Database Model and Several Retrieval Languages.* Amsterdam and Atlanta, GA.
- Hucka, M., *et al.* (2003) The Systems Biology Markup Language (SBML): A Medium for Representation and Exchange of Biochemical Network Models. *Bioinformatics* 19: 524-531.
- Kim, J.D., Ohta, T., Tateisi, Y., and Tsujii, J. (2003) GENIA Corpus - A Semantically Annotated Corpus for Bio-textmining. *Bioinformatics* 19: i180-i182.

⁸ <http://www.w3.org/Addressing>