

# HTML Table Interpretation by Sibling Page Comparison in the Molecular Biology Domain

Cui Tao and David W. Embley

Department of Computer Science, Brigham Young University, Provo, UT, 84602, USA  
ctao,embley@cs.byu.edu

## Abstract

There are large and growing amount of biological data that reside in different online repositories. Many of these repositories represent their data in tables. In order to automatically understand these online pages, a system that can interpret tables is desired. However, the longstanding problem of automatic table interpretation still illudes us [12]. We offer a solution for the common special case in which so-called sibling pages are available. Sibling pages, which are the pages commonly generated by underlying web databases, are compared to identify and connect non-varying components (category labels) and varying components (data values). We tested our solution on 862 HTML tables. Experimental results show that the system can successfully identify sibling tables, generate structure patterns, interpret different tables using the generated patterns, and automatically adjust the structure patterns as needed.

**Keywords:** Bioinformatics, table interpretation

## 1 Introduction

Catalyzed by world-wide research communities producing publicly available data, the volume of biological data is increasing at a rapid pace. Many online biological data repositories present their information in tables. Tables present information in a simplified and compact way in rows and columns. Data in one row/column usually belongs to the same category or provides values for the same concept. The labels of a row/column describe this category or concept.

Although a table with a simple row and column structure is common, tables can be much more complex. Figure 1 shows an example. The position of table category labels may vary in different tables. Labels commonly appear on the top or left. Occasionally, table designers position labels on the right side of a table. In long tables, labels sometimes appear at the end of a table or in the middle of a table,

every few rows, in order to help a reader find the correspondence between labels and data. Sometimes tables are rearranged to fit the space available. Label-value pairs may appear in multiple columns across the pages or in multiple rows placed below on another down the page. Tables may themselves contain nested tables as does the table in Figure 1. These complexities make automatic table interpretation a challenging task.

## 2 Table Interpretation

To interpret a table is to properly associate table category labels with table data values. Using Figure 1 as an example, we see that *Identification*, *Location*, and *Function* are labels for the large rectangular table. Inside the right cell of the first row is another table with headers *IDs*, *NCBI KOGs*, *Species*, etc. Nested inside of this cell are two tables with labels *CGC name*, *Sequences name*, ... , *Version*, and *Gene Model*, *Status*, ..., and *Amino Acids*. The rest of the information in these tables are data values. Once category labels and data values are found, we want to properly associate them. For example, for the value *F18H3.5*, its associated label should be the sequences of labels *Identification*, *IDs*, and *Sequence name*. We associate one or more sequences of labels with each data value in a table (more, when the table is multi-dimensional). Borrowing notation from Wang [10], the representation of a label-value pair is look like: *Identification.IDs.Sequence name*  $\rightarrow$  *F18H3.5*. The left hand side of the arrow is a sequence of one or more table labels, and the right hand side of the arrow is a data value.

Recent surveys [5, 12] describe the vast amount of research that has been done in table processing and illustrate the challenges of the table interpretation problem. We focus in this paper, however, only HTML tables. A number of HTML table extraction systems use machine learning to recognize tables in web pages (e.g. [3, 11]). Drawbacks of machine learning approaches, however, are that they need training data, and they need to be retrained for tables from different web sites. Other table interpretation systems work based on some simple assumptions and heuris-

## Gene Summary for *cdk-4*

Specify a gene using a gene name ([unc-26](#)), a predicted gene id ([R13A5.9](#)), or a protein ID ([CE02711](#)):

[\[identification\]](#)
[\[location\]](#)
[\[function\]](#)
[\[gene ontology\]](#)
[\[reactome knowledgebase\]](#)
[\[alleles\]](#)
[\[similarities\]](#)
[\[reagents\]](#)
[\[bibliography\]](#)

<b>Identification</b>	<b>IDs:</b>	<table border="1"> <thead> <tr> <th>CGC name</th> <th>Sequence name</th> <th>Other name(s)</th> <th>WB Gene ID</th> <th>Version</th> </tr> </thead> <tbody> <tr> <td><a href="#">cdk-4</a> - (<i>Cyclin-Dependent Kinase family</i>) (via person: <a href="#">Michael Krause</a>)</td> <td><a href="#">F18H3.5</a></td> <td><a href="#">NM_077855</a> (inferred automatically) <a href="#">XO136</a> (inferred automatically)</td> <td><a href="#">WBGene00000406</a></td> <td>1</td> </tr> </tbody> </table>	CGC name	Sequence name	Other name(s)	WB Gene ID	Version	<a href="#">cdk-4</a> - ( <i>Cyclin-Dependent Kinase family</i> ) (via person: <a href="#">Michael Krause</a> )	<a href="#">F18H3.5</a>	<a href="#">NM_077855</a> (inferred automatically) <a href="#">XO136</a> (inferred automatically)	<a href="#">WBGene00000406</a>	1			
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<b>NCBI KOGs*:</b> <a href="#">Protein kinase PCTAIRE and related kinases [KOG0594]</a> <b>Species:</b> <a href="#">Caenorhabditis elegans</a> <b>Other sequence(s):</b> <a href="#">AF083878</a> (Caenorhabditis elegans cyclin-dependent kinase CDK-4 (cdk-4) mRNA, complete cds.) <b>NCBI:</b> [Entrez Genes: <a href="#">15718266</a> ] [ <a href="#">AceView: XO136</a> ] <b>Gene model(s):</b> <table border="1"> <thead> <tr> <th>Gene Model</th> <th>Status</th> <th>Nucleotides (coding/transcript)</th> <th>Protein</th> <th>Amino Acids</th> </tr> </thead> <tbody> <tr> <td><a href="#">F18H3.5a</a> 1, 2</td> <td>confirmed by <a href="#">cDNA(s)</a></td> <td><a href="#">1029/3051 bp</a></td> <td><a href="#">WP:CE18608</a></td> <td><a href="#">342 aa</a></td> </tr> <tr> <td><a href="#">F18H3.5b</a> 1, 2, 3</td> <td>partially confirmed by <a href="#">cDNA(s)</a></td> <td><a href="#">1221/1704 bp</a></td> <td><a href="#">WP:CE28918</a></td> <td><a href="#">406 aa</a></td> </tr> </tbody> </table>	Gene Model	Status	Nucleotides (coding/transcript)	Protein	Amino Acids	<a href="#">F18H3.5a</a> 1, 2	confirmed by <a href="#">cDNA(s)</a>	<a href="#">1029/3051 bp</a>	<a href="#">WP:CE18608</a>	<a href="#">342 aa</a>	<a href="#">F18H3.5b</a> 1, 2, 3	partially confirmed by <a href="#">cDNA(s)</a>	<a href="#">1221/1704 bp</a>	<a href="#">WP:CE28918</a>	<a href="#">406 aa</a>
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**Putative ortholog(s):** [Caenorhabditis briggsae: CBG07433](#) [[syntenic alignment](#)] ([Stein LD et al. best reciprocal blastp match-seg-off](#))

<b>Location</b>	<b>Genetic Position:</b> <a href="#">X:12.69 +/- 0.000 cM</a> [ <a href="#">mapping data</a> ] <b>Genomic Position:</b> <a href="#">X:13518823..13515773 bp</a>
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<b>Function</b>	<b>Mutant Phenotype:</b> [Krause MW] <a href="#">cdk-4</a> is a cyclin dependent kinase related to <a href="#">cdk-4</a> and <a href="#">cdk-6</a> from other organisms. Homozygous <a href="#">cdk-4(gv3)</a> animals usually arrest in L2 due to no, or limited, proliferation of the post-embryonic blast cells. About 3% of animals make it to a late stage of development. Definitions of <a href="#">abbreviations used</a> in the text. <b>RNAi Phenotype(s):</b> <a href="#">Lvl Pvl Unc</a> [For details see: <a href="#">Park M 06 Oct 1999</a> ]
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Figure 1. A sample page from [1].

tics (e.g. [2, 4, 8]). These simple assumptions (labels are either the first row or the first column) are easily broken in complex tables. More sophisticated table interpretation techniques have appeared in recent papers [6, 7, 9]. None of this research makes use of sibling tables, but is complementary to our work and could potentially be used in conjunction with our work in future efforts to improve results for certain cases.

### 3 Sibling Page Comparison

If we have another page, such as the one in Figure 2, that has the same structure as the one in Figure 1, the system maybe able to obtain enough information about the structure to make automatic interpretation possible. Molecular biology web resources usually generate output pages after receiving a user query by placing the results into a predefined page structure. Thus, pages from the same web site are usually structured in the same way. We call pages that are from the same web site and have similar structures *sibling pages*. The two pages in Figures 1 and 2 are a pair of sibling pages. They have the same basic structure, with the same top banners that appear in all the pages from this web

site, with the same table title (*Gene Summary for some particular gene*), and a table that contains information about the gene. Corresponding tables in sibling pages are called *sibling tables*. If we compare the two large tables in the main part of the sibling pages, we can see that the first columns of each table are exactly the same. If we look at the cells under the *Identification* label in the two tables, both contain another table with two columns. In both cases, the first column contains identical labels *IDs*, *NCBI KOGs*, ..., *Putative orthology(s)*. Further, the tables under *Identification.IDs* also have identical header rows. The data rows, however, vary considerably. General speaking, we can look for commonalities to find labels and look for variations to find values.

Given that we can find most of the label and data cells in this way, our next task is to infer the general structure pattern of the web site and of the individual tables embedded within pages of the web site. For each table, we first locate the positions of values and labels. For example, consider the two nested tables in Figures 1 and 2 that start with *CGC name*. The top rows of the tables on the two pages are identical while the two second rows vary considerably. We thus determine that the first row is a row of labels and the second row is a row of values. Depending on the posi-

## Gene Summary for *dyb-1*

Specify a gene using a gene name ([unc-26](#)), a predicted gene id ([R13A5.9](#)), or a protein ID ([CE02711](#)):

[\[identification\]](#) [\[location\]](#) [\[function\]](#) [\[gene ontology\]](#) [\[reactome knowledgebase\]](#) [\[alleles\]](#) [\[similarities\]](#) [\[reagents\]](#) [\[bibliography\]](#)

<b>Identification</b>	<b>IDs:</b>	<u><a href="#">CGC name</a></u>	<u><a href="#">Sequence name</a></u>	<u><a href="#">Other name(s)</a></u>	<u><a href="#">WB Gene ID</a></u>	<u><a href="#">Version</a></u>
		<i>dyb-1</i> - ( <i>DystroBrevin homolog</i> ) (via person: <a href="#">Laurent Segalat</a> )	<a href="#">F47G6.1</a>	<a href="#">NM_058459</a> (inferred automatically) <a href="#">1B963</a> (inferred automatically)	<a href="#">WBGene00001115</a>	1
	<b>Concise Description:</b>	The <i>dyb-1</i> gene encodes a homolog of mammalian alpha-dystrobrevin (DTNA, OMIM:601239), mutation of which can lead to left ventricular noncompaction with congenital heart defects. [ <a href="#">details</a> ]				
	<b>NCBI KOGs*:</b>	Beta-dystrobrevin [ <a href="#">KOG4301</a> ]				
<b>Species:</b>	<i>Caenorhabditis elegans</i>					
<b>NCBI:</b>	[ <a href="#">Entrez Genes: 14670171</a> ] [ <a href="#">AceView: 1B963</a> ]					
<b>Gene model(s):</b>	<u><a href="#">Gene Model</a></u>	<u><a href="#">Status</a></u>	<u><a href="#">Nucleotides (coding/transcript)</a></u>	<u><a href="#">Protein</a></u>	<u><a href="#">Swissprot</a></u>	<u><a href="#">Amino Acids</a></u>
	<a href="#">F47G6.1</a> 1, 2	confirmed by cDNA(s)	<a href="#">1773/7391 bp</a>	<a href="#">WP_CE26812</a>	<a href="#">DTN1 CAEEL</a>	<a href="#">590 aa</a>
<b>Putative ortholog(s):</b>	<i>Caenorhabditis briggsae</i> : <a href="#">CBG22285</a> [ <a href="#">syntenic alignment</a> ] (Stein LD et al. ; best reciprocal blastp match-seg-off)					
<b>Location</b>	<b>Genetic Position:</b> <a href="#">I:15.38 +/- 0.361 cM</a> [ <a href="#">mapping data</a> ]					
	<b>Genomic Position:</b> <a href="#">I:1483084..1490474 bp</a>					
<b>Function</b>	<b>Mutant Phenotype:</b>	Definitions of <a href="#">abbreviations used</a> in the text.				
	<b>RNAi Phenotype(s):</b>	<a href="#">WT</a> [For details see: <a href="#">Ahringer JA 16 Nov 2000</a> ]				

Figure 2. A second sample page from [1].

tion of labels and values, we try to match the table with a pre-defined structure pattern templates or a combination of several pattern templates. Once we found a match, we generated a structure pattern for the set of sibling tables, which records the location of the table in each sibling page, the position of labels and values, and how the labels and values associate to each other.

Although we look for commonalities to find labels and look for variations to find data values, we must be careful about being too strict. Sometimes there are additional or missing label-value pairs. The two tables beginning with *Gene Model* Figures 1 and 2 do not share exactly the same structure. The table in Figure 1 has five columns and three rows, while the table in Figure 2 has six columns and two rows. Although they are not exactly the same, we can still identify the structure pattern by comparing them. The top rows in the two tables are very similar. Observe that the table in Figure 2 only has an additional *Swissprot* column inserted between the *Protein* and *Amino Acids* columns. Although the labels for the two tables are not identical, we can still tell that they are table headers.

In addition to discovering the structure pattern for a web site, we can also dynamically adjust the pattern if the system encounters a table that varies from the pattern. There are two ways to adjust a structure pattern. (1) We can adjust the location of a table. If the table in the recorded location

does not match with the pattern, we check for tables in the neighboring positions and see if we can find a match. If so, we add the new position in the pattern as an alternative location where we could possibly locate the sibling table in another sibling page. (2) We can adjust the labels. If there is an additional or missing label, the system can change the pattern by either adding the new label and marking it optional or marking the missing label optional. For example, if we had not seen the extra *Swissprot* column in our initial pair of sibling pages, the system can add *Swissprot* as a new label and mark it as optional.

## 4 Experimental Results

We collected 100 sibling pages from 10 different web sites in the molecular biology domain for a total of 862 HTML tables. Among these tables, the system falsely classified three pairs of layout tables as data tables. The system, however, successfully eliminated these false sibling pairs during pattern generation because it was unable to find a matching pattern. No false patterns were generated. The system was able to recognize 28 of 29 structure patterns. The system missed one pattern because the table contained too many empty cells. If we had considered empty cells as mismatches, the system would have correctly recognize this pattern. As the system processed additional sibling pages,

it found 5 additional sibling tables and correctly interpreted all but one of them. The failure was caused by labels that varied across sibling tables causing them in some cases to look like values. There were 5 location adjustments and 12 label adjustments—all of them correct. One table was interpreted only partially correctly because the system considered the irrelevant information *To Top* as a header.

The time for the pattern generation given a pair of sibling pages consists of: (1) the time to read and parse the two pages and locate all the HTML tables, (2) the time for sibling table comparisons, and (3) the time to select from pre-defined structure templates and generate the pattern. The complexity of parsing and locating HTML tables is  $O(n)$ , where  $n$  is the number of HTML tags. To find the best match for each HTML table, the time complexity is  $O(km^2)$ , where  $k$  is the number of HTML tables in one sibling page and  $m$  is the number of nodes in each table DOM tree. The time complexity for finding the correct pattern for each matched sibling table is  $O(pl)$ , where  $p$  is the number of pattern templates and  $l$  is the number of leaf nodes (cells) in the HTML table. If there is pattern combination involved, this complexity increases multiplicatively. The actual time needed for the pattern generation for a pair of sibling pages is, on average, below or about one second, but reached a maximum of 15 seconds for a complicated web site where pages have more than with more than 20 tables on a Pentium 4 computer running at 3.2 GHz.

The time for table interpretations for a single sibling web page when no adjustment are necessary consists of: (1) the time for parsing a web page and locating each table and (2) the time for matching located table with a pattern. The complexity of parsing and locating HTML tables is  $O(n)$ , where  $n$  is the number of HTML tags. The complexity of matching a located table with the corresponding pattern is  $O(pl)$ , where  $p$  is the number of level path possibilities leading to a possible sibling table and  $l$  is the number of leaf nodes (cells) in the HTML table. The time to do adjustments range from the time to do a simple label adjustment, which is constant, to the time required to re-evaluate all sibling tables, which is the same as the time for initial pattern generation. Overall, the actual time needed for interpreting tables in one page is, on average, below one second, but reached a maximum of 19 seconds for a complicated web page with several tables and several adjustments.

## 5 Conclusion

Many online biological repositories present their information in tables with complicated structures. In this paper, we introduced a system that can successfully interpret these tables automatically. Our system works based on sibling page comparison. By comparing sibling pages from the same site, we are able to find the location of table head-

ers and data entries, and further we are able to infer the general pattern for all pages from the same site.

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