

3<sup>rd</sup> transPLANT Training Workshop - October 2014  
Exploiting and understanding Solanaceous genomes

# Mining integrated data sources using LAILAPS

**Matthias Lange**  
**IPK**



# Mining integrated data sources using LAILAPS

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[www.IPK-Gatersleben.de](http://www.IPK-Gatersleben.de)

## IPK - Leibniz Institute of Plant Genetics and Crop Plant Research



- over 70 years tradition
- "Magdeburger Börde" very fruitful soil
- source of the breeding industry in Germany
- total staff: 550
- scientists: 200  
(10 Bioinformaticians)
- 30 research groups

- Bioinformatic Research Topics
- data management & information retrieval
  - sequence-, network- & image-analysis



Instructor:  
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## Aims

- Training of Novel Technology Platform
- Get Feedback and Requirements for Sustainable Service

## Scope

- Basic Concepts of Information Retrieval
- LAILAPS Modules for Integrative IR
- Trait driven IR
- Personalized Ranking
- Embed into Custom Web Sites

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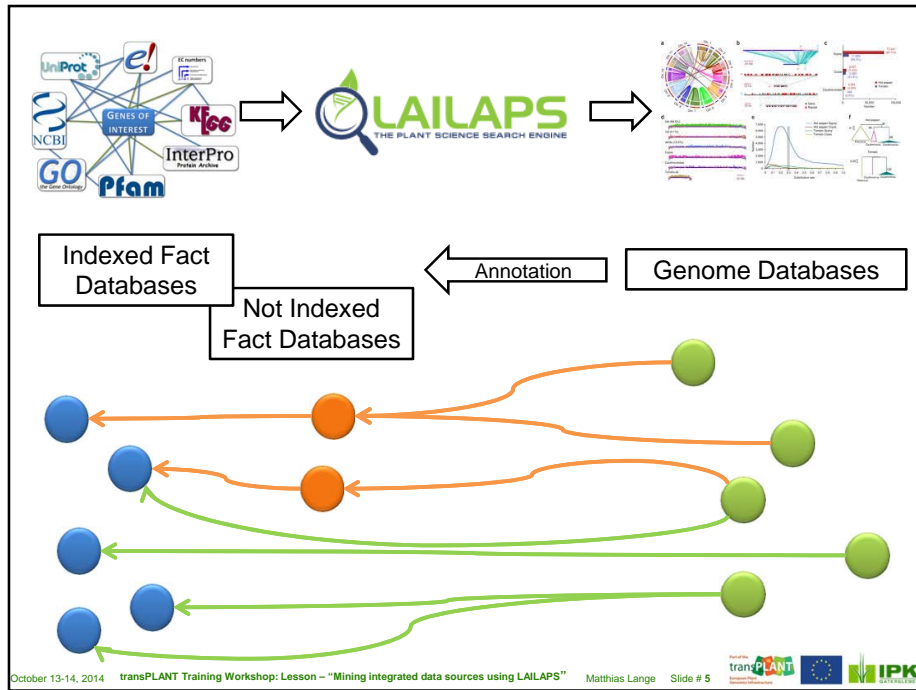
Life Science Application for Information Retrieval and Lightweight API for Portable Search Engines

- Integrative search over distributed data resources
- Information retrieval corpus over genome knowledge bases
- Link trait to genome and -omics resources
- Full text search and personalized relevance ranking



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The screenshot shows the LAILAPS web interface with a sidebar and a main content area. The sidebar includes links for "LAILAPS", "Indexed Databases", "Referenced Genomic Resources", "LAILAPS full screen", "Demo Screen", "CLIENTS", "Portlet", "Drupal Plugin", "PUBLICATIONS", "Paper", "Talks", "Poster", and "Built by Maven".

The main content area is titled "Ontologies" and lists the following:
 

- Trait Ontology
- Controlled vocabulary to describe each trait as a distinguishable feature, characteristic, quality or phenomenon of a developing or mature individual.
- Gramineae Taxonomy Ontology
- Primarily derived from NCBI Taxonomy, this taxonomy ontology focuses on the Poaceae (Gramineae) family of plant taxonomy only.
- NCBI Taxonomy
- The Taxonomy Database is a curated classification and nomenclature for all of the organisms in the public sequence databases.
- Plant Ontology (PO)
- Controlled vocabulary (ontology) that describes plant anatomy and morphology and state of development for all plants.
- Gene Ontology (GO)
- The Gene Ontology project is a major bioinformatics initiative with the aim of standardizing the representation of gene and gene product attributes across species and databases.

The content area also includes sections for "Protein Sequence Databases" (listing Pfam, UnIProtKB/Swiss-Prot, UniProtKB/UniMML, and Protein Database (PDB)) and "Plant Diversity Resources" (listing Taxonomic Allium Reference Collection, Garlic and Shallot Core Collection, and Genbank Information System of the IPK Gatersleben).

A large diagonal watermark reading "Indexed Fact Databases" is overlaid across the middle of the page.

At the bottom of the slide, there is a footer with the date "October 13-14, 2014", the workshop title "transPLANT Training Workshop: Lesson – ‘Mining integrated data sources using LAILAPS’", the name "Matthias Lange", and "Slide # 6". Logos for transPLANT, the European Union, and IPK are also present.

Linked Genome Databases

The screenshot shows a web-based application titled "LAILAPS - Linked Databases". On the left, there's a sidebar with links like "Referenced Genomic Resources", "LAILAPS full screen", "Demo", "Screenshot", "CLDITIS", "Portlet", "Drupal Plugin", "PUBLICATIONS", "Paper", "Talks", and "Poster". The main content area is titled "Genomes" and lists several databases:

- EnsemblPlants
- Ensembl
- GnpGis
- PlantsDB
- BARLEX
- POLAPGEN DB
- BMRF
- MetaCrop
- OPTIMAS-DW

Below these are sections for "Observations of Phenotypic Traits", "Protein Function Prediction", and "Systems Biology". A "Built by maven" logo is visible at the bottom left. At the bottom right are logos for "Part of the transPLANT European Project", the European Union, and IPK.

## Information Retrieval

- “... Information retrieval is the activity of obtaining information resources relevant to an information need from a collection of information resources ...”

The diagram is a triangle divided into three horizontal sections. The top section is labeled "Decision" at the apex and "Synthesis" in the middle. The middle section is labeled "Summary" at the top, "Analysis" in the middle, and "Organize" at the bottom. The bottom section is labeled "Collect" at the bottom left, "Information" in the middle, and "Facts" at the bottom right. The word "Knowledge" is written vertically in the center of the triangle.

Volker Lepczynski 2011

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# Information Retrieval Components

## 1. Linguistic

- text / data decomposition
- language processing
- synonyms, homonyms



## 2. Indexing

- efficient search in content

## 3. Presentation

- intuitive user interface
- related entries („page like this“)
- query suggestion („did you mean“)



## 4. Relevance estimation

- feature extraction
- ranking functions
- pertinence (subjective user relevance profiles)



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## Crawling



- Extraction
  - flat files, SQL, REST
- Transformation
  - parse, encoding
- Loading
  - excerpts, keywords, synonyms

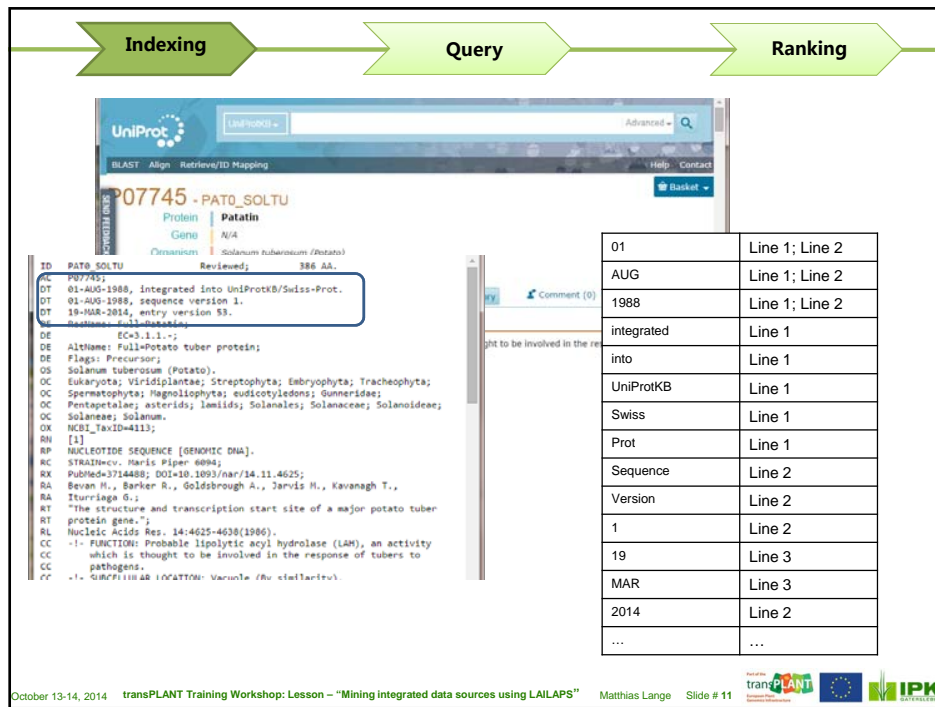
## Indexing

- compile/split into documents
- tokenization
  - words, substances, IDs
- Inverted Index
  - unique tokens -> documents

ID	Term	Document position
1	blue	1, 3, 3, 2
2	butterfly	1, 2, 3, 5
3	butterfly	1, 4
4	butterfly	1
5	forged	2, 5
6	great	2, 1, 2
7	honey	1, 5
8	needs	3, 8
9	search	3, 11
10	sky	2
11	search	3, 10
12	sky	2, 6, 3, 3
13	wind	2, 14

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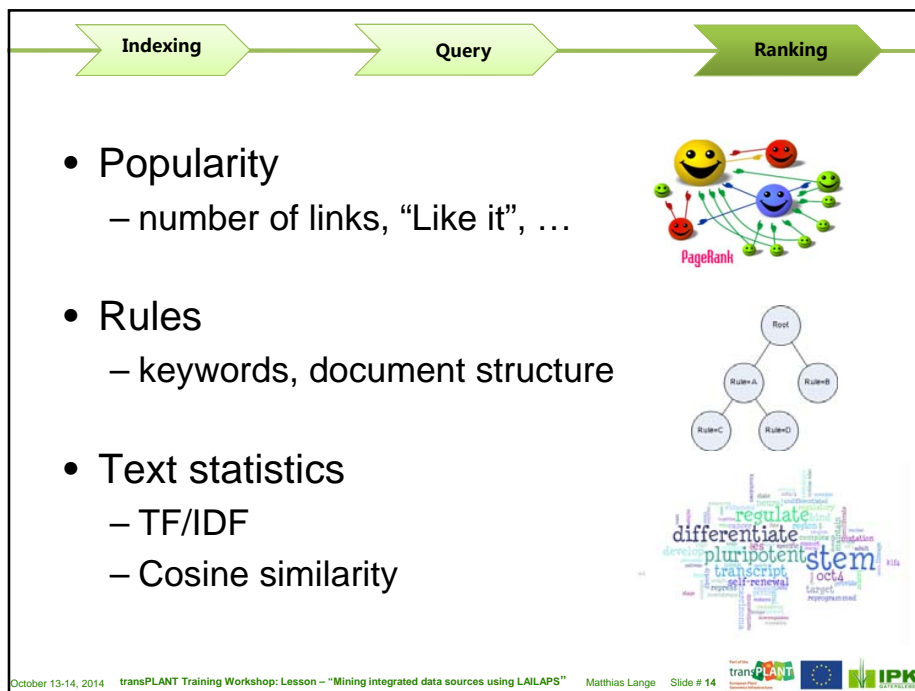
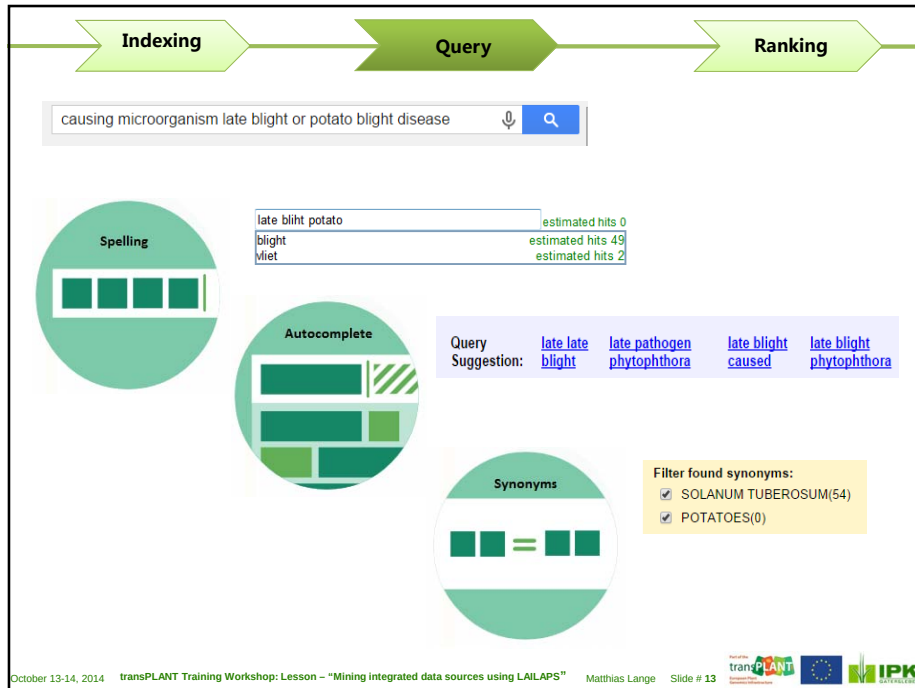


## Exercise I – Inverse Text Index

- ETL of UniProt Documents
  - each participant separately:
    - select one document from /exercise\_1/uniprot\_document.txt (paste URL into Web browser)
    - create reverse text index of first CC and OS line
    - order index alphabetically
- find matching documents
  - all participants:
    - how much documents match to late blight potato?

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The flowchart consists of three green chevron-shaped boxes arranged horizontally. The first box is labeled "Indexing", the second is labeled "Query", and the third is labeled "Ranking".

- term frequency x inverse document frequency (TF-IDF)

TF('example', D2) = 3  
IDF ('example') =  $\log(2/1) = 0.30$   
TF-IDF =  $3 * \log 2 = 0.9$

TF('this', D2) = 1  
IDF ('this') =  $\log(2/2) = 0$   
TF-IDF =  $1 * \log 0 = 0$

1. Document 2 is specific for "example"
2. Document 2 is not specific for "this"

Document 1	
Term	Term Count
this	1
is	1
a	2
sample	1

Document 2	
Term	Term Count
this	1
is	1
another	2
example	3

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## Exercise II – TF-IDF ranking

- compute the most relevant UniProt entry for combination of search terms “blight” “tomato”
  - use documents D of Exercise I
  - count number matching documents
  - count matches per document
  - compute  $TF-IDF(\text{blight}, D) * TF-IDF(\text{tomato}, D)$
  - which is the most relevant UniProt entry?

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## Connect to the LAILAPS Server

The screenshot shows the LAILAPS homepage with a search bar at the top containing the query "Productive". Below the search bar, there is a list of search results. The first result is highlighted in green and contains the URL <http://lailaps.ipk-gatersleben.de>. Other results include "Application: <http://pgrc.ipk-gatersleben.de/lailaps>". The page features the LAILAPS logo and the text "Welcome to LAILAPS" and "integrative search engine for plant genomics data". Logos for transPLANT and IPK Gatersleben are visible in the top right corner.

## LAILAPS Modules – Home Page

The screenshot shows the LAILAPS search interface. A search bar at the top contains the query "late blight potato tomato". Below the search bar, a speech bubble points to the search results area, listing features like "conjunctive keywords: late blight", "phrase search: \"late blight\"", and "Wildcard search: plant\*". Another speech bubble points to the search results table, listing "estimated hits for entire query", "index and link statistics", and "spelling correction", "word before cursor", and "estimated hits". A third speech bubble points to the search bar itself, listing "login roles: guest (predefined personal ranking profile)", "manager (personal ranking)", and "admin (control, ranking, configuration, user)". The page includes the LAILAPS logo and the text "Integrated Search in Genomics Resources" and "trans-National Infrastructure for Plant Genomic Science". Logos for transPLANT and IPK Gatersleben are visible in the top right corner.

## LAILAPS Modules – Result Page

The screenshot shows the LAILAPS Result Page. At the top, there's a search bar with the query "late blight potato disease". Below it, a sidebar contains several filter options:

- suggested queries
- click to search
- download result
- select indexed facts
- select linked genome databases
- direct or indirect annotation filter
- synonym filter

The main content area shows search results for three entries:

- 090329\_RecName\_Full-Putative late blight resistance protein homolog**: Results 1-8 from 19 displayed (Ranked: 231 FilterResults). This entry has a green link to a found fact.
- 090328\_RecName\_Full-Putative late blight resistance protein homolog RIB-19**: Results 1-8 from 19 displayed (Ranked: 231 FilterResults). This entry has an orange link to a found fact.
- 090328\_RecName\_Full-Putative late blight resistance protein homolog RIB-18**: Results 1-8 from 19 displayed (Ranked: 231 FilterResults). This entry has an orange link to a found fact.

Annotations in genome databases are highlighted in green (direct) and orange (indirect).

## LAILAPS Modules – Fact Browser

The screenshot shows the LAILAPS Fact Browser. On the left, there's a sidebar for rating the record:

- original record from fact database
- rate relevance (used for training)
- predicted score (by page interaction)

The main content area shows a detailed record for a resistance protein:

- Function: Confers resistance to late blight (Phytophthora infestans) races carrying the avirulence gene Avr1b. Resistance proteins guard the plant against pathogens that contain an appropriate avirulence protein via an indirect interaction with this avirulence protein. That triggers a defense system including the hypersensitive response, which restricts the pathogen growth.
- GO - Molecular function: ADP binding, ATP binding, Nucleotide binding.
- GO - Biological process: plant-type hypersensitive response, Plant defense.
- Keywords: Biological process, Hypersensitive response, Plant defense.
- Keywords: Ligand, ATP-binding, Nucleotide-binding.
- Names & Taxonomy: Protein name: Putative late blight resistance protein homolog RIB-13; Gene name: RIB-13.

On the right, there's a panel for related facts and linked annotations in genome databases:

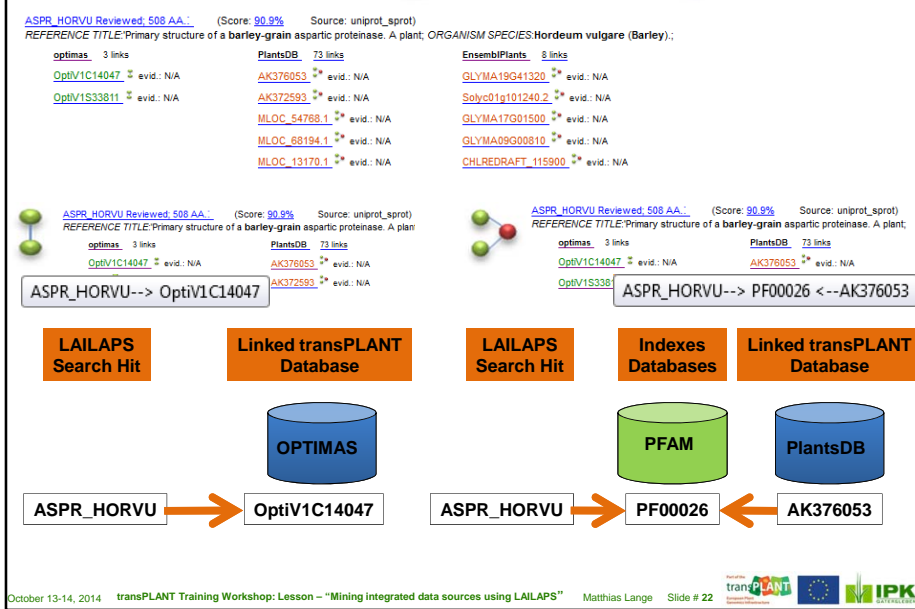
- related facts
- linked annotations in genome databases (same as result page)

## LAILAPS Modules – Related Facts & Cross Ref.

The screenshot shows two LAILAPS search results side-by-side. Both results are for the same gene, **scaffold\_803427.1**, which is a positive uncharacterized protein. The left result is from EnsemblPlants and the right is from PGSC0003DMG40200457B. Each result includes a reference title, gene ID, species, and location. Below each result is a detailed view of the gene's characteristics and its relationship to other genes and databases.

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## LAILAPS: Direct and Indirect Links



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## LAILAPS Modules – Download Search Result

The screenshot shows a search results page for 'late blight potato disease'. A large arrow points down to the table below.

A	B	C	D	E	F	G	H	I
relevance_score	annotation_identifier	annotation_source	annotation_abstract	annotation_detail_URL	annotated_resource_URL	annotation_evidence	annotation_link_type	annotation_link_graph
1	Q8L320;	uniprot_sprot	DESCRIPTION:ReName: Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_3630.1">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_3630.1</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_3630.1
2	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_31.1">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_31.1</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_31.1
3	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_3287.1">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_3287.1</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_3287.1
4	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_8142.1">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_8142.1</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_8142.1
5	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_8057.1">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_8057.1</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_8057.1
6	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_79783.3">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_79783.3</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_79783.3
7	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_78491.4">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_78491.4</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_78491.4
8	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_78474.3">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_78474.3</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_78474.3
9	Q8L320;	uniprot_sprot	Full/putative late blight resistance protein homolog RIB-13; COMMENTS OR NOTES:- DESCRIPTION:ReName:	<a href="http://www.uniprot.org/uniprot/Q8L320">http://www.uniprot.org/uniprot/Q8L320</a>	<a href="http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_77918.2">http://mpis-helmholtz-muenchen.de/plant/barley/ga/reports/ppl/entryElement.jsp?gene=MLOC_77918.2</a>	1.0	indirect link	Q8L320->PF00931<-MLOC_77918.2

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## Connect to the LAILAPS Training Server

The screenshot shows the LAILAPS search application interface. A green box highlights the 'Training:' section.

**Welcome to LAILAPS**  
integrative search engine for plant genomics data

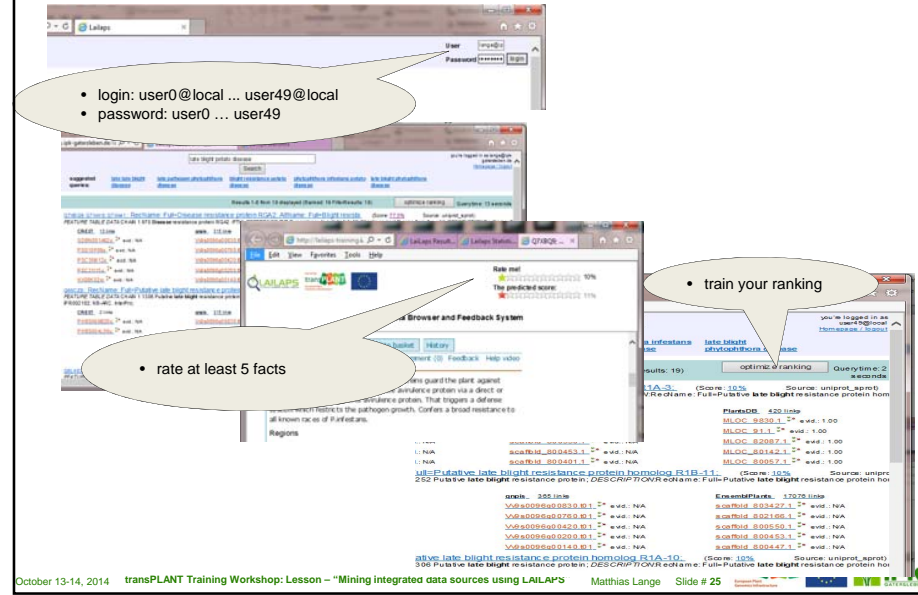
**Training:**

Web site: <http://lailaps-training.ipk-gatersleben.de>  
Application: <http://lailaps-training.ipk-gatersleben.de/lailaps/>

Integrated Search in Genomics Resources  
trans-National Infrastructure for Plant Genomic Science

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## LAILAPS Modules – Personalized Ranking



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## Exercise III – IR using LAILAPS

- plant material: find material in German germplasm collection
  - **material from IPK genbank information system:**  
potato phureja
  - **material from IPK genbank information system by characterisation:**  
garlic white stripes
  - **geo location of genbank donors:**  
tomato netherlands
- traits: download all transcript positions at EnsemblPlant genomes with direct annotation to:
  - potato late blight disease
  - potato starch content
  - potato virus resistance
- metabolic activity: find metabolic facts and for starch synthesis in potato
  - which pathways are involved (use BioModels and MetaCrop database)
- query own traits

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Slide # 26



# Portlet Embedding

Plugin LAILAPS search as service into own Web sites

The screenshot shows the MetaCrop interface. On the left, there's a sidebar with links like 'Overview', 'About', and 'Login'. The main content area has tabs for 'Home', 'Pathways', 'Databases', 'Substances', and 'Gen'. A central panel titled 'Choose organism' displays small images of various plants and their names: Hordeum vulgare, Triticum aestivum, Oryza sativa, Zea mays, Solanum tuberosum, Brassica napus, Beta vulgaris, Arabidopsis thaliana, and Medicago truncatula. To the right, there's a 'Trax Search' section with a search bar containing 'QAILAPS transPLANT' and a 'Start BLAST Search' button. The top right corner features the IPK Gatersleben logo.

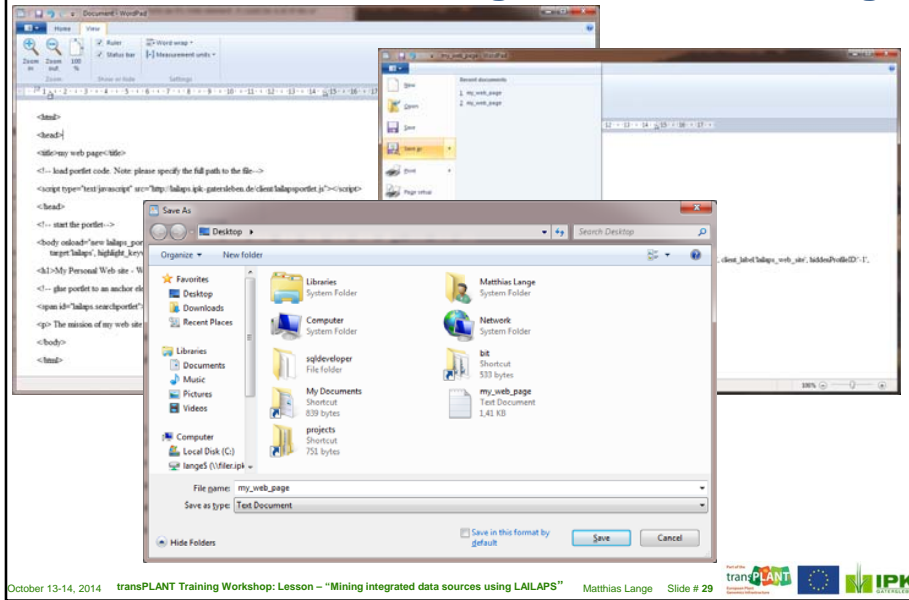
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## Embed LAILAPS – Copy HTML Template

The screenshot shows two browser windows. The left window is titled 'LAILAPS Portlet' and contains instructions on how to embed LAILAPS into a web site using a JavaScript library. It includes a 'Copy' button and a 'Paste' input field. The right window is titled 'Parameter' and shows the detailed JavaScript code for embedding. The code includes parameters like 'parentID', 'url', 'client\_label', 'highlight\_keyword\_search', and 'database\_ids'. Below the code is a 'Sample HTML client site' section with a large block of sample HTML code.

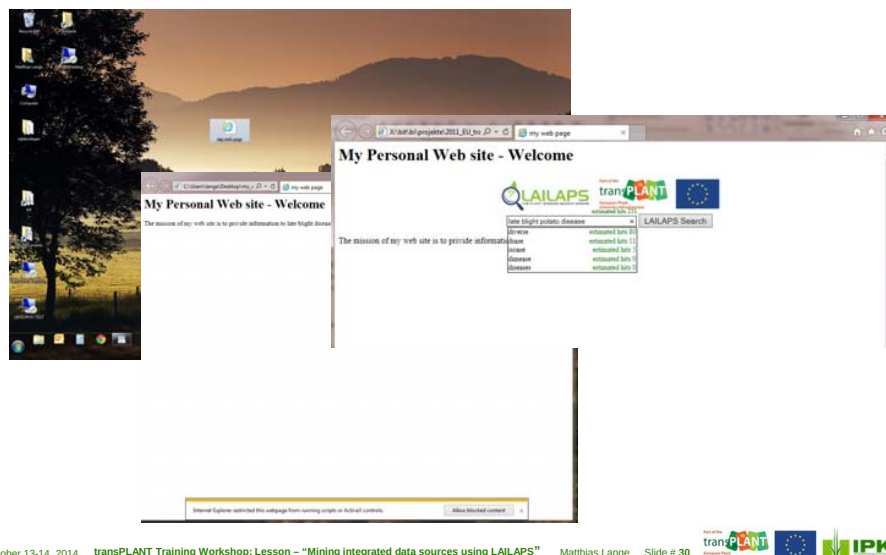
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## Embed LAILAPS – Design Custom Web Page



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## Embed LAILAPS – Open in Web Browser



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## Embed LAILAPS – Test It

The screenshot shows a web browser window with two tabs. The left tab is titled 'My Personal Web site - Welcome' and displays a simple website with a logo, some text, and a sidebar with a search bar. The right tab is titled 'LAILaps Result Browser' and shows the LAILAPS search results for the query 'late blight potato disease'. The results list several entries, each with a title, source, and abstract. The LAILAPS interface includes filters for databases, sequencing methods, and search terms.

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## Exercise IV – LAILAPS Portlet

- Create an custom HTML file and include LAILAPS search for PubMed Cross references

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## Lessons Learned

- Concepts of IR to interpret search engine results
- Smart querying
- Apply LAILAPS as integrated IR environment for plant genomics
- Train the LAILAPS Ranking System
- Embed LAILAPS into custom Web sites



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## Acknowledgements

<http://lailaps.ipk-gatersleben.de>

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- Daniel Arend (IPK DPPN)
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- Uwe Scholz (IPK BIT)
- Danuta Schüler (BIT LIMS)
- Stephan Weise (IPK Genbank)

<http://edal.ipk-gatersleben.de>

HelmholtzZentrum münchen  
German Research Center for Environmental Health



PLANT RESEARCH INTERNATIONAL  
WAGENINGEN UR



TGAC

The Genome Analysis Centre™



DPPN partners:

HelmholtzZentrum münchen  
Deutsches Forschungszentrum für Gesundheit und Umwelt



transPLANT: European Commission within its 7<sup>th</sup> Framework Program, under the thematic area “Infrastructures”, contract number 283496.

Federal Ministry  
of Education  
and Research



DPPN: German Ministry of Education and Research (project identification number: 031A053).

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# The LAILAPS Search Engine: A Feature Model for Relevance Ranking in Life Science Databases

**Matthias Lange, Karl Spies, Christian Colmsee, Steffen Flemming, Matthias Klapperstück and Uwe Scholz**

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## Summary

Efficient and effective information retrieval in life sciences is one of the most pressing challenge in bioinformatics. The incredible growth of life science databases to a vast network of interconnected information systems is to the same extent a big challenge and a great chance for life science research. The knowledge found in the Web, in particular in life-science databases, are a valuable major resource. In order to bring it to the scientist desktop, it is essential to have well performing search engines. Thereby, not the response time nor the number of results is important. The most crucial factor for millions of query results is the relevance ranking.

In this paper, we present a feature model for relevance ranking in life science databases and its implementation in the LAILAPS search engine. Motivated by the observation of user behavior during their inspection of search engine result, we condensed a set of 9 relevance discriminating features. These features are intuitively used by scientists, who briefly screen database entries for potential relevance. The features are both sufficient to estimate the potential relevance, and efficiently quantifiable.

The derivation of a relevance prediction function that computes the relevance from this features constitutes a regression problem. To solve this problem, we used artificial neural networks that have been trained with a reference set of relevant database entries for 19 protein queries.

Supporting a flexible text index and a simple data import format, this concepts are implemented in the LAILAPS search engine. It can easily be used both as search engine for comprehensive integrated life science databases and for small in-house project databases. LAILAPS is publicly available for SWISSPROT data at

<http://lailaps.ipk-gatersleben.de>

## 1 Introduction

“Finding information in the WWW is not much of a challenge. Just head for Google or Entrez and get the related web page or database entry.” This issue can be heard frequently talking to biologists, who search information about a certain biological object [1]. However, finding reliable information about the function of a protein or seeking the protein that is involved in a certain activity in the cell cycle, is much more challenging. One has the choice of about 1,100 life science databases and billions of database records [2]. Even if one reduces the number of databases and records using database integration systems and powerful query systems, there

are still too many results for a simple query like "arginase" - an enzyme involved in the urea cycle. As shown in Figure 1, one gets 6739 hits in NCBI Entrez Protein databases, 2610 in Uniprot, and 568 in KEGG GENES (data from December 2009).



**Figure 1: Search Engines in Life Science – the screenshots show examples of popular search engines, to investigate protein function: Google, Entrez, UniProt and KEGG.**

Intuitively, the first choice are web search engines. Web site ranking techniques order query hits by relevance. But, trying to apply ranking methods that were developed to rank natural language text or WWW-sites to life science content and databases is questionable [3]. For example, the top ranked Google hit for "arginase" is a Wikipedia page. This is because the page is referenced by a high number of web-pages or Google assigned a manual defined priority rank. Here, the hypothesis is: *A high hyperlink in-degree of a page means high popularity and high popularity means high relevance.*

In order to find scientific relevant database entries, scientists need strong scientific evidence in relation to the specific research field. A dentist has other relevance criteria than a plant biologist or a patent agent. The intuitive and commonly used way at the scientist's desktop is query refinement. Criteria like who published, in which journal, for which organism, evidence scores, surrounding keywords etc. matter. Even complete search guides, e.g. for dentists were published [4].

Other ranking algorithms use Term Frequency - Inverse Document Frequency (TF-IDF) as ranking criteria. Apache-Lucene<sup>1</sup> is a popular implementation of this concept and is frequently

<sup>1</sup><http://lucene.apache.org>

used in bioinformatics, like LuceGene from the GMOD project [5], which is used for the EBI ‘google’ like search frontend EBeye. The TF-IDF approach works well, but misses the semantic context between the database entries and the query.

Another approach is the probabilistic relevancy ranking [6], whereby probabilistic values for the relevance of database fields and word combinations have to be predefined. In combination with a user feedback system, the probabilistic approach shows promising ranking performance [7].

Semantic search engines use methods from natural language processing and dictionaries to predict the semantic most similar database entries. Such conceptual search strategies, implemented in GoPubMed [8] or ProMiner [9], are frequently used algorithms in text mining projects.

The combination and abstraction of the mentioned relevance indicators motivated the development of a feature model for life science databases, a user feedback system and a machine learning ranking engine. In the next sections we present the LAILAPS system as method for relevance ranking in life science databases [10]. In particular, we present the used ranking concept, the used feature model, its implementation as neural network and ranking benchmarks.

## 2 LAILAPS Method for Relevance Ranking

Nearly every search engine, including LAILAPS, incorporates a scoring or ranking function to calculate a relevance for an entry. The central LAILAPS hypothesis is that the relevance score is context-dependent and the absolute rank position can be determined by sorting the relative scores. We apply information theory and postulate that the relevance of a database entry depends on two factors: its *content* and its *interpretation* by the search engine user. For the first factor we found that the relevance decision is based on a small number of core properties of the content. These core properties are used to deduce a feature model that expresses all important properties of a database entry as feature vector. The factor user interpretation is realized by LAILAPS in form of a feedback system and hand curated reference sets of relevant rated database entries. Both factors are used for the scoring algorithm. The algorithm uses artificial neural networks as method to estimate the relevance score of a database entry based on these factors.

### 2.1 LAILAPS Feature Model

A relevance scoring function for life science databases is highly dependent on the underlying data. In contrast to full text data like PubMed or even traditional web sites, a database entry in a life science database is

1. structured and split into blocks (e.g. attributes, entities),
2. enriched by metadata (e.g. name of an attribute, quality information),
3. a compressed excerpt of a fact and
4. a mix of scalar, pure values and natural language text (e.g. stoichiometric biochemical reaction vs. function description).

These properties lead to the conclusion that classical ranking methods miss important properties and are suboptimal for life science database entries. Consequently, we had to define our own set of features that combines traditional and life science database specific ones. Motivated by the observation of user behavior during search engine result inspection, we introduced a set of 9 features  $F$ , presented in table 1. These features are intuitively used by scientists, who briefly screen database entries for potential relevance. The features are both sufficient to estimate the potential relevance, and efficiently quantifiable.

	feature	description
$F_1$	attribute	in which attribute the query term was found
$F_2$	database	in which database the found entry is included
$F_3$	frequency	the frequency of all query terms in the entry and attribute
$F_4$	cooccurrence	express how close and in which order the query terms were found
$F_5$	keyword	gives information, if good or bad keyword are present near to the query terms
$F_6$	organism	to which organism the database entry relates to
$F_7$	sequence length	the length of the sequence described by the database entry
$F_8$	text position	which portion of the attribute is covered by the query term
$F_9$	synonym	gives information if the hit was produced by an automatic synonym expansion

**Table 1: Overview of the LAILAPS feature set**

## 2.2 LAILAPS Relevance Ranking

The starting point of a ranking process is the query. Here we define a query as a set of AND-linked terms. Each term is a combination of alpha-numeric characters. Terms can be grouped into phrases, by quotation marks. We restrict the logical join of terms and phrases to a conjunctive form.

$$Q = t_1 \wedge t_2 \wedge \dots \wedge t_n \mid t_n \in \text{term or phrase} \quad (1)$$

The result of each query is a set  $R$  of database entries, which include all query terms or phrases. For these database entries, all attribute hits and related text positions are extracted:

$$\text{find}(Q) \rightarrow \{R\} \mid R = (t, d, \{(a, p)\}) \quad (2)$$

Where,  $t$  is the query term;  $d$  is the database entry; and  $\{(a, p)\}$  is a set of attribute-position pairs.

Based on the mentioned features, we define 9 feature functions  $find_f$ , that compute for each element  $r$  of the query result set  $R$  a scalar score  $\omega$  as element in the final feature vector  $\vec{\Omega} = (\omega_1, \dots, \omega_9)$ :

$$score_f(r) \rightarrow \omega \mid f \in F_1, \dots, F_9 \quad (3)$$

The final step is a ranking function, which computes a relevance score  $\tau$  from this 9-dimensional feature vector  $\vec{\Omega}$ :

$$relevance(\vec{\Omega}) \rightarrow \tau \quad (4)$$

Because this function constitutes a regression problem, we chose a machine learning approach to estimate this function. For a given vector of quantified features of a particular database entry, a continuous relevance score must be predicted. The focus of the relevance function is to automatically learn to recognize complex patterns from the feature vectors and make intelligent relevance predictions. Hence, we choose a machine learning approach to estimate the function  $relevance(\vec{\Omega})$ .

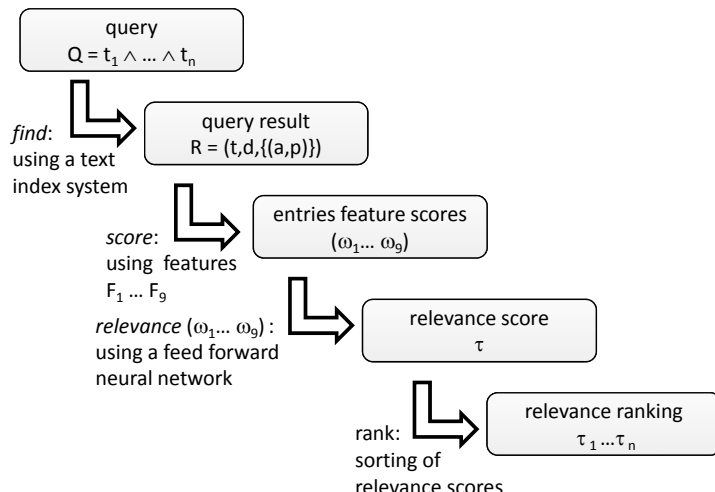
After we have predicted relevance scores, they can be ordered such as

$$\tau_1 < \tau_2 \Rightarrow \tau_1 \text{ is less relevant than } \tau_2 \quad (5)$$

The final relevance ranking is the order of all relevance scores:

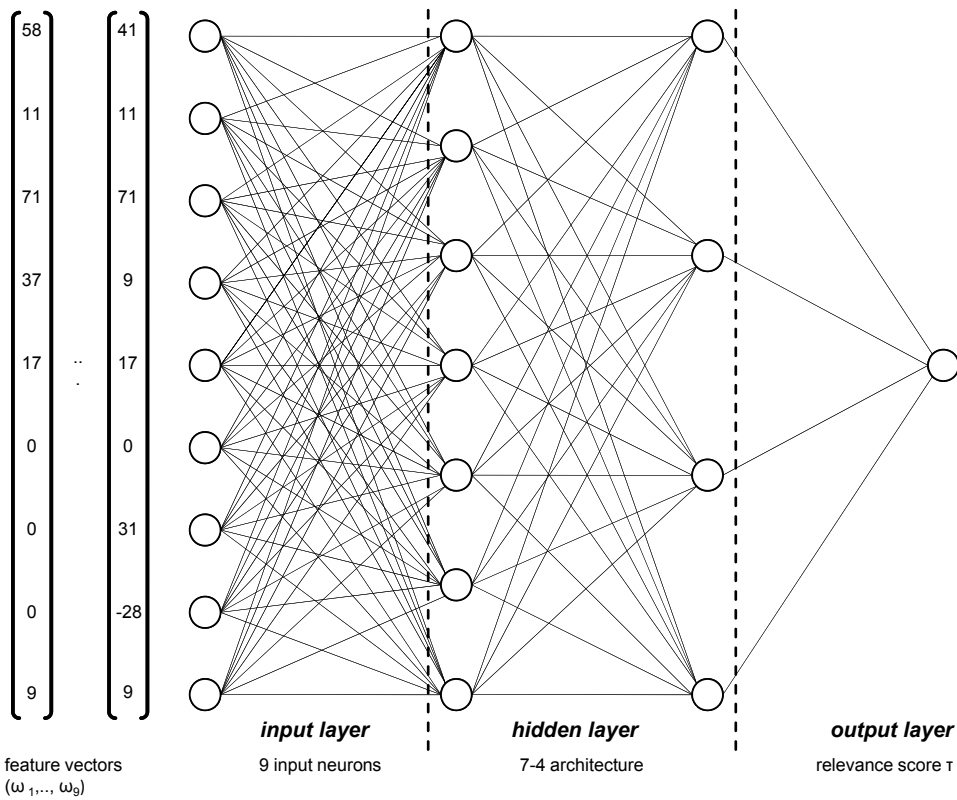
$$(\tau_1, \dots, \tau_n) \mid \forall \tau : \tau_{n-1} < \tau_n \quad (6)$$

As shown in figure 2 those ranking tasks have to be executed for each query as sequential workflow.



**Figure 2: The LAILAPS Ranking Workflow**

As mentioned before, the function  $relevance$  constitutes a regression problem. We have to map a vector of 9 features to a scalar, continuous relevance score. In particular, supervised methods perform well for text mining systems [11]. We found that artificial neural networks show best



**Figure 3: neural network for relevance ranking – the used neural network predict for a vector  $\vec{\omega}$  of 9 feature values the relevance of the database entry.**

performance for our regression problem [12]. Using a set of reference data, we trained a feed-forward neural network with 9 neurons at the input and 7-4 neuron architecture in the hidden layer (see Figure 3). In order to train the network, we split up the training data into 80% for training and 20% for testing and used 500 training epochs. These parameters were estimated by minimizing the mean squared error  $\epsilon$  over the training set:

$$\epsilon = \frac{1}{n} \sum_{i=1}^n (\tau'_i - \tau_i)^2 \mid n = \text{size of training set}; \tau'_i = \text{manual score}; \tau_i = \text{predicted score} \quad (7)$$

In each epoch the change of  $\epsilon$  was checked. After 500 epochs, no significant decrease was found and the final mean square error was 0.33 [12].

The crucial step for the neural network training is a set of true positive, manual evaluated relevance rankings. Our industrial and academic partners provided a set of plant metabolic queries with 1089 manually relevance ranked database records (see Table 2). This reference ranking list was separated into three confidence classes: high, medium and low. With these data, a neural network for plant metabolism was trained.

### 2.3 LAILAPS Implementation

LAILAPS was developed as a 3-tier web application using Apache Tapestry<sup>2</sup> as web application-framework, ORACLE as database backend and a JAVA-implemented ranking logic featured by

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<sup>2</sup><http://tapestry.apache.org>

Query Text	Size	Category Split-Up (hi/me/lo)
industrial use case 1	20	6 / 4 / 10
”pinene synthase”	18	10 / 3 / 5
industrial use case 2	39	8 / 13 / 18
industrial use case 3	64	14 / 32 / 18
”gamma tocopherol methyltransferase”	38	21 / 9 / 8
”ent-kaurene synthase”	65	17 / 38 / 10
”chlorophyll synthase”	77	17 / 54 / 6
industrial use case 4	134	35 / 68 / 31
”cinnamyl-alcohol dehydrogenase”	214	45 / 36 / 133
industrial use case 5	17	3 / 4 / 10
”dihydrokaempferol 4-reductase”	65	9 / 29 / 27
”l-ascorbate peroxidase”	100	69 / 12 / 19
”morphine 6-dehydrogenase”	35	2 / 15 / 18
”zeaxanthin epoxidase”	51	21 / 2 / 28
”squalene monooxygenase”	84	24 / 30 / 30
”acetoacetyl-coa synthetase”	68	14 / 36 / 18

**Table 2:** Overview of the training data set.

the Java Object Oriented Neural Engine (JOONE)<sup>3</sup>. The backend database stores the loaded life science databases in an entity-attribute-value (EAV) [13] adapted database schema. This flexible concept enables the import of RFC-compatible CSV-formatted<sup>4</sup> exports from life science databases, whereas each row comprise a database record and its columns the fields.

For the imported databases, an inverse text index is computed and synonyms are loaded. In the public available system, we provide protein synonyms extracted from UNIPROT/SWISSPROT [14] and BREND [15]. The ranking logic

1. computes all matched positions per database entry,
2. extracts for each database entry an n-dimensional feature vector with 9 basis feature classes and
3. predicts a relevance probability using user specific neural networks, which maps the feature vectors to the users specific relevance score.

Since the ranking profile is computed in context of the authenticated user, a valid user login is recommended. Otherwise, a default context with a pre-trained neural network is used.

The relevance ordered query hits are assigned to the ranking profile and rendered by the middleware into a number of web pages (see Figure 4). The embedded feedback system provides a tool for the user to rate the relevance of a particular database entry. The ratings are stored in the database backend and used to accumulate user training data. The LAILAPS feedback system is

<sup>3</sup><http://www.jooneworld.com/>

<sup>4</sup><http://tools.ietf.org/html/rfc4180>

transparently embedded into the result browser. By opening the database detail browser, AJAX<sup>5</sup> code is injected into the original data HTML presentation, which is for example, provided by the SRS@EBI data retrieval system [16]. This code collects the user rating of the database entry and tracks user interactions. This feedback is used to enrich the original training data in the related ranking profile.

### 3 Results and Discussion

Searching scientific databases effectively necessitates the use of contemporary software to locate desired and meaningful information according to the users scientific or project priorities. However, the combination of relevance ranking and life science data retrieval is still missing in life science information systems. LAILAPS fills this gap and provides a search engine for integrated or single instance life science databases in combination with an efficient ranking system.

We have evaluated the relevance prediction using the standard measures for precision  $Pr$ , recall  $Re$  and  $F_1$  score:

$$Pr = \frac{TP}{TP + FP}; \quad Re = \frac{TP}{TP + FN}; \quad F_1 = \frac{2 * Pr * Re}{Pr + Re}$$

As mentioned, we have used a curated reference set of plant metabolic queries. In order to decide whether a database entry has been correctly ranked or not, we do not consider its concrete ranking position. Because of combinatorial explosion, for human curators it is nearly impossible to find a correct relevance order among hundreds of database entries. Rather, the knowledge quality of a certain database entry is crucial. Consequently, the database entries were classified into three confidence classes: "HIGH", "MEDIUM" and "LOW". The "HIGH"-class comprises the top entries with proven and reliable knowledge. The class "MEDIUM" includes all those, that could be interesting but are uncertain. The class "LOW" includes all data, that has insufficient knowledge value or bad quality indicators. For each query, those classes form sub sets of a query result  $R$  such as  $R = R_H \cup R_M \cup R_L$ , whereas  $R_H \cap R_M \cap R_L = \emptyset$ . Each set forms a continuous window in the list of results. E.g., for a query result of 100 database entries, the window for  $R_H$  ranges from position 1-20,  $R_M$  from position 21-80, and  $R_L$  from 81-100.

In order to evaluate the LAILAPS rankings, we have to compare a list of relevance ordered database entries with the confidence classes of the reference set. We can't compare absolute ranking positions, because the elements in the confidence classes have no order. But we can say in which range (window) of rank positions a LAILAPS ranked entry should be, to fall into a certain confidence class. This consideration is used to define the true positives (TP), false positives (FP) and false negatives (FN):

For example, for one use case the curators sorted 20 entries into the class "HIGH". 18 of the top 20 LAILAPS ranked result are in the window  $R_H$ . In this case, the precision is  $\frac{18}{18+2} = 0.9$ .

Because all database entries of the reference set were found by LAILAPS, the recall is  $Re = 100\%$ . This is because the text indexing is the basis for matching query terms. The text index uses a tokenizer, that decomposes text into words. LAILAPS use the same text decomposition

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<sup>5</sup>Asynchronous JavaScript and XML

error type	LAILAPS benchmark semantics
true positive (TP)	the rank position is in the same window as in the reference set
false positive (FP)	the rank position is in a different window as in the reference set
false negative (FN)	the database entry was not found by LAILAPS

**Table 3: Definition of evaluation error types.**

rules and the same databases as the reference retrieval systems. Furthermore, no synonym expansion of query terms was used. Thus, the hits of the reference systems could be reproduced by LAILAPS and no false negative hits exit.

The overall benchmarking result of 16 queries is shown in Figure 5. The average recall, precision and  $F_1$  values are shown in Table 4.

confidence class	precision	recall	$F_1$
”HIGH”	62%	100%	76
”MEDIUM” $\cup$ ”HIGH”	81%	100%	90

**Table 4: Evaluation of LAILAPS relevance ranking results.**

In average we achieve a better precision than existing search engines [3]. Training and benchmark data for the non-industrial use cases are available by request to the authors.

The training data were collected in the application scenario of plant research and queries for protein functions. In order to bring LAILAPS to a broad community, we provide a public installation of the LAILAPS search engine. This will help to improve performance and to include more user domain specific ranking profiles. Because of limited database and project resources, the public, non-commercial version is restricted to SWISSPROT data. A comprehensive set of databases is available for registered users on request to the authors.

LAILAPS combines a clean, powerful and easy to use human computer interface with a machine learning based, context sensitive ranking system. It comprises a search engine with a self trained neural network ranking system, which brings a new quality and determinism into the scientific knowledge exploration.

## 4 Conclusion

In this paper, we presented a feature model for relevance ranking in life science databases. The presented concept is to learn ranking pattern from the ranking behaviour of scientist, who make use of data retrieval systems. We combined user ranking profiles, a reference set of relevant database entries for 19 protein queries as training set to a ranking model. Queries are formulated as simple keyword lists and will be expanded by synonyms. The model is used to extract per database entry a feature vector. Using supervised machine learning approach, we were able to predict a user specific relevance score per feature vector. We found a feed forward neural network as best performing method for the regression problem of ranking in life science databases.

Supporting a flexible text index and a simple data import format, this concepts are implemented in the LAILAPS search engine. It can easily be used both as search engine for comprehensive

integrated life science databases and for small in-house project databases. Using expert knowledge as training data for a predefined neural network or using users own relevance training sets, a reliable relevance ranking of database hits has been implemented. LAILAPS is public available for SWISSPROT data at <http://lailaps.ipk-gatersleben.de>

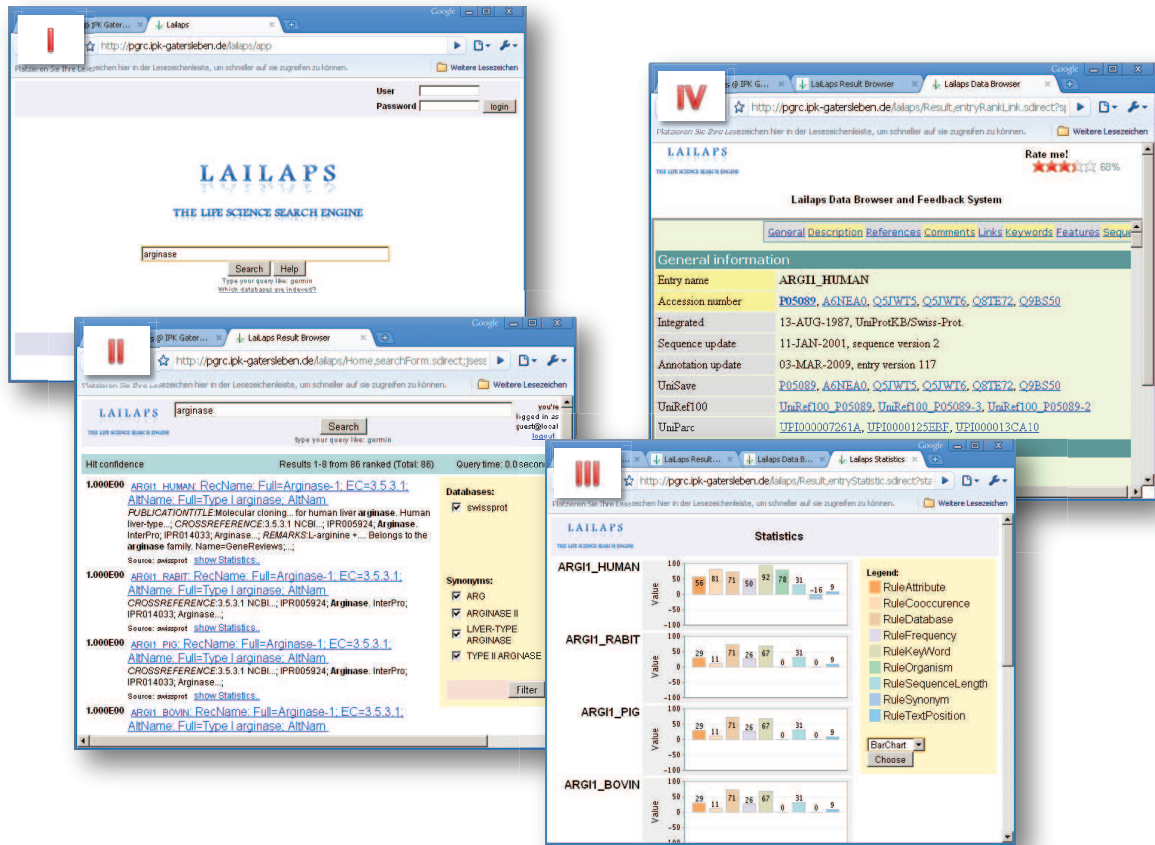
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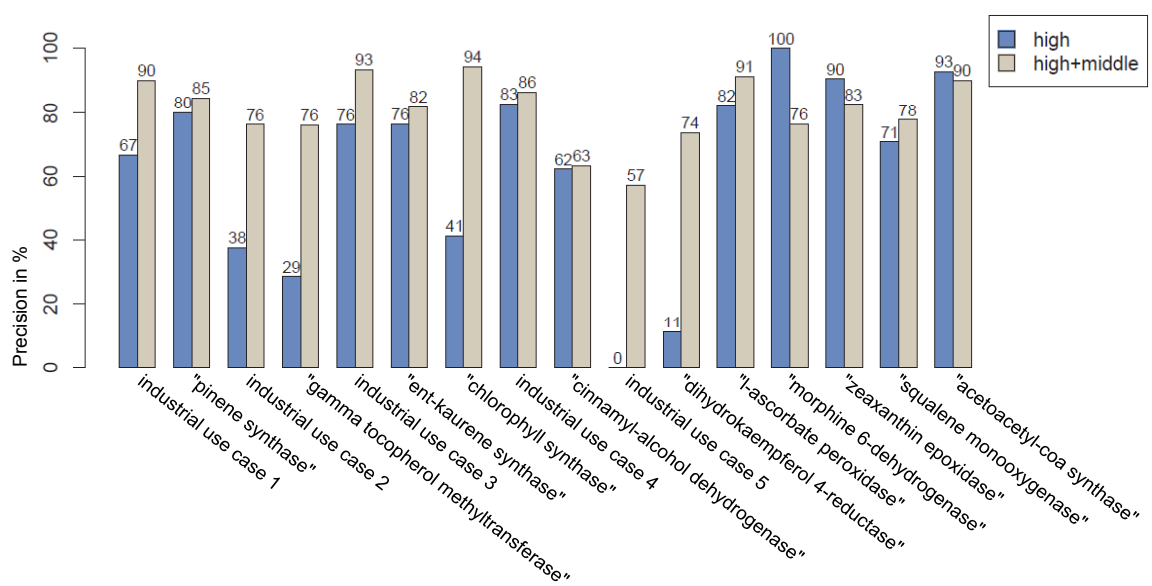
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**Figure 4: LAILAPS view of an example query session – The four browser windows represent the common query workflow of LAILAPS.** Start screen with an optional login to load a customized ranking profile (I). The query might be specified as a single word, combination of words delimited by a whitespace or quoted phrases. The ranked query result is presented as a list of relevance sorted database accessions (II), with a short hit description, the evidence value, the hyper link to the original data source and a link to the scoring statistics in window (III). The exploration of hits is supported by a detail browser and feedback system (IV). The original data is extracted from the original source and displayed. The user have to be aware that LAILAPS is querying the original data from the SRS system, which installed at the EBI (<http://srs.ebi.ac.uk>). Thus, some outdated accession may result in broken links. In this page, the user might rank the relevance of the hit for later training of the user ranking profile.



**Figure 5: LAILAPS recall for use case queries**