



**HELMHOLTZ
ZENTRUM FÜR
INFEKTIONSFORSCHUNG**

**This is an by copyright after embargo allowed publisher's PDF of an article
published in**

**Bartsch, A., Bunk, B., Haddad, I., Klein, J., Münch,
R., Johl, T., Kärst, U., Jänsch, L., Jahn, D., Retter,
I.**

**GeneReporter-sequence-based document retrieval and
annotation**

**(2011) Bioinformatics, 27 (7), art. no. btr047, pp.
1034-1035.**

GeneReporter—sequence-based document retrieval and annotation

Annekathrin Bartsch¹, Boyke Bunk¹, Isam Haddad¹, Johannes Klein¹, Richard Münch¹, Thorsten Johl², Uwe Kärst², Lothar Jänsch², Dieter Jahn¹ and Ida Retter^{1,*}

¹Institute for Microbiology, Technische Universität Braunschweig, Spielmannstr. 7, 38106 Braunschweig and ²Cellular Proteomics Group, Helmholtz Centre for Infection Research, Inhoffenstr. 7, 38124 Braunschweig, Germany

Associate Editor: Jonathan Wren

ABSTRACT

Summary: GeneReporter is a web tool that reports functional information and relevant literature on a protein-coding sequence of interest. Its purpose is to support both manual genome annotation and document retrieval. PubMed references corresponding to a sequence are detected by the extraction of query words from UniProt entries of homologous sequences. Data on protein families, domains, potential cofactors, structure, function, cellular localization, metabolic contribution and corresponding DNA binding sites complement the information on a given gene product of interest.

Availability and implementation: GeneReporter is available at <http://www.genereporter.tu-bs.de>. The web site integrates databases and analysis tools as SOAP-based web services from the EBI (European Bioinformatics Institute) and NCBI (National Center for Biotechnology Information).

Contact: i.retter@tu-bs.de; ida.retter@helmholtz-hzi.de

Supplementary information: Supplementary data are available at *Bioinformatics* online.

Received on October 3, 2010; revised on December 1, 2010; accepted on January 9, 2011

1 INTRODUCTION

In face of next-generation sequencing and high-throughput analyses, the link between obtained data and existing knowledge is crucial. Automatic annotation pipelines provide useful evidence of potential functions for genes and proteins, but in a last essential step, the scientist must manually evaluate the available information. Usually, the necessary evidence is derived from scientific publications, databases and *in silico* predictions. Thus, tools that provide a combination of all of these relevant data for a gene or protein of interest are of high practical impact. In this context, GeneReporter offers a customizable workflow for the integrated application of protein sequence analysis and document retrieval.

A large number of diverse text-mining tools exist that provide different strategies and interfaces to satisfy the extensive data-mining demands in biomedical sciences (Krallinger *et al.*, 2010). GeneReporter identifies citations related to a gene or protein sequence of interest. The UniProt annotations of homologous sequences are used to derive keywords such as gene names, synonyms and species. These keywords provide the query terms for a subsequent literature search in PubMed (Sayers *et al.*, 2010). In

*To whom correspondence should be addressed.

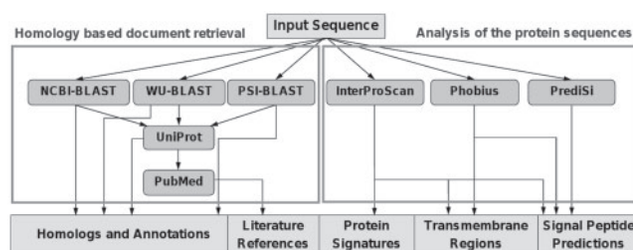


Fig. 1. Workflow of the GeneReporter analysis process. Arrows indicate data transfer and processing. Input and output is depicted as rectangles, web services are depicted as rounded rectangles.

this way, GeneReporter extends and replaces MineBlast (Dieterich *et al.*, 2005), a similar tool which is discontinued. In comparison with other tools that connect literature to sequence information, like quickLit (Gilchrist *et al.*, 2008) and Metis (Mitchell *et al.*, 2005), GeneReporter is characterized by highly customizable query options, the integration of InterPro and the direct access to the original EBI and NCBI databases.

2 REQUESTING LITERATURE AND SEQUENCE ANALYSIS

The user can enter up to 10 nt or protein sequences to submit a query on the GeneReporter web site. Two different types of analyses are provided: (i) *homology-based document retrieval* searches information on homologous sequences from the UniProt Knowledgebase (UniProt Consortium, 2010) and citations from PubMed. (ii) *Analysis of the protein sequences* requests protein annotations from InterPro (Hunter *et al.*, 2009), Phobius (Käll *et al.*, 2007) and PrediSi (Hiller *et al.*, 2004). The complete workflow is depicted in Figure 1. An example for an application is given as Supplementary Material.

Using homology-based document retrieval, the first step is a BLAST search in UniProtKB, where the user can select the desired algorithm. NCBI-BLAST (Altschul *et al.*, 1997) and WU-BLAST (Lopez *et al.*, 2003) result in a different ranking of homology matches, and therefore yield different query word extractions from the respective UniProtKB entries. PSI-BLAST (Altschul *et al.*, 1997) is the most sensitive algorithm and beneficial for sequences that fail to result in significant hits with the other algorithms. Either Swiss-Prot or the complete UniProtKB can be chosen as BLAST

target database. The UniProtKB entries of the resulting BLAST hits are parsed for gene names, synonyms and species names, which are used as query terms for the subsequent PubMed request. This literature search can be further specified, e.g. by additional query terms and years of publication. The option ‘organism-specific search’ adds the respective species name to the PubMed search string. Query and result options and the construction of the PubMed queries are described in detail in the Supplementary Material.

For further analysis of the protein sequence, GeneReporter submits a query to InterProScan that matches the sequence against InterPro. This database comprises predictive signatures that assign protein families, various domains and functional sites for a protein of interest. The input sequence can also be analysed by Phobius and PrediSi, which search for putative transmembrane regions and signal peptides.

To assure long-term up-to-date datasets and analysis tools, GeneReporter utilizes standardized web services from the EBI (Goujon *et al.*, 2010), the NCBI (Sayers *et al.*, 2010) and our institute. The processing time of a query strongly depends on these services. The web service providers bind the access of their services to certain rules in order to avoid overload and abuse of their resources. To match these rules, a local queuing system monitors and limits the number of simultaneous queries. Details on cut-offs and limits are provided as Supplementary Material.

3 RESULTS

The results are summarized on an overview page. For each query sequence, this page provides a link to a detailed view of the obtained data for the requested services. The result overview page can be bookmarked and results can be retrieved from this URL for at least 24 h. For further analysis, results can be downloaded as Excel or tab-delimited text files.

The detailed view provides one result tab for each requested service. The BLAST result tab shows homologous protein sequences. It is complemented with annotations from the UniProt database, e.g. organism name and GO terms, in order to facilitate their evaluation. The PubMed result tab shows gene-related citations ordered by the respective PubMed queries. Query words that were matched within title and abstract are marked in bold. For each query word combination, the link ‘This query in PubMed’ performs the corresponding query on the PubMed web site. This allows the manual modification and specification of the automatically generated queries with all the sophisticated features of the PubMed search interface. Furthermore, GeneReporter provides citations from UniProt entries of the BLAST hit sequences. In general, these references comprise the key papers on the respective gene or protein. Figure 2 shows the PubMed tab of an example search for a hypothetical protein from *Pseudomonas aeruginosa* C3719.

The output from InterPro, Phobius and PrediSi requests is given in additional tabs. The InterProScan and Phobius output includes graphical visualizations of signature matches and transmembrane regions within the proteins of interest.

Sequence: PACG_03312
View references from UniProt

search again with this sequence
back to overview
back to top
This query in PubMed

Gene name: PA2G_04443, Transcriptional regulator Dnr
Species name: Pseudomonas aeruginosa 2192

PubMed ID	Authors	Journal	Date	Abstract
19472662	Nicolletta Cavallaro...	Microbiology (Reading, England), 155(Pt 9):2638-44	2009 Sep	The transcription factor DNR from <i>Pseudomonas aeruginosa</i> specifically represses nitric oxide and hemX for the activation of a latent promoter in <i>Escherichia coli</i> .
19420222	Garcia Gardina...	Journal of molecular biology, 378(5):1003-15	2008 May	NO sensors in <i>Pseudomonas aeruginosa</i> : structure of the transcriptional regulator DNR.
15932158	Hiroshi Arai...	Journal of bacteriology, 187(12):3960-9	2005 Jun	Transcriptional regulation of the flavohemoglobin gene for aerobic nitric oxide detoxification by the second nitric oxide-responsive regulator of <i>Pseudomonas aeruginosa</i> .

Fig. 2. Screenshot of the homology-based document retrieval result. Query sequence in this example: UniProt AcNo A3KZR4.

ACKNOWLEDGEMENTS

We would like to thank the EBI and NCBI for providing web service access to their tools and databases. We would also like to thank Max Schobert for intensive testing and discussions and Patrick Jäkel for the layout of the GeneReporter logo.

Funding: Grants of the German Federal Ministry of Education and Research (BMBF) for the European transnational research initiative on ‘Systems Biology of Microorganisms’, SysMO (Psysmo) (grant number 0313980D); the Volkswagen Foundation (I/81448).

Conflict of Interest: none declared.

REFERENCES

- Altschul,S.F. *et al.* (1997) Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic Acids Res.*, **25**, 3389–3402.
- Dieterich,G. *et al.* (2005) MineBlast: a literature presentation service supporting protein annotation by data mining of BLAST results. *Bioinformatics*, **21**, 3450–3451.
- Gilchrist,M.J. *et al.* (2008) Evading the annotation bottleneck: using sequence similarity to search non-sequence gene data. *BMC Bioinformatics*, **9**, 442.
- Goujon,M. *et al.* (2010) A new bioinformatics analysis tools framework at EMBL-EBI. *Nucleic Acids Res.*, **38**, W695–W699.
- Hiller,K. *et al.* (2004) PrediSi: prediction of signal peptides and their cleavage positions. *Nucleic Acids Res.*, **32**, W375–W379.
- Hunter,S. *et al.* (2009) InterPro: the integrative protein signature database. *Nucleic Acids Res.*, **37**, D211–D215.
- Lopez,R. *et al.* (2003) WU-Blast2 server at the European Bioinformatics Institute. *Nucleic Acids Res.*, **31**, 3795–3798.
- Käll,L. *et al.* (2007) Advantages of combined transmembrane topology and signal peptide prediction—the Phobius web server. *Nucleic Acids Res.*, **35**, W429–W432.
- Krallinger,M. *et al.* (2010) Analysis of biological processes and diseases using text mining approaches. *Methods Mol. Biol.*, **593**, 341–382.
- Mitchell,A.L. *et al.* (2005) METIS: multiple extraction techniques for informative sentences. *Bioinformatics*, **21**, 4196–4197.
- Sayers,E.W. *et al.* (2010) Database resources of the National Center for Biotechnology Information. *Nucleic Acids Res.*, **38**, D5–D16.
- UniProt Consortium (2010) The Universal Protein Resource (UniProt) in 2010. *Nucleic Acids Res.*, **38**, D142–D148.