florian leitner data & machine learning architect

contact	experience

Address Line 2015-now Data Cataltyics, S.L. Madrid, Spain Data & machine learning architect (International consultant) E-28005 Madrid Architecting & operating (Jenkins, Ansible) low-latency, high-performance language processing and machine learning solutions for large scale (B+) and "in-+34 6XX XXX XXX finte" data on streaming platforms (Spark, Flink, clients' in-house systems; into Cassandra, Aurora, MySQL, Postgres, etc.). Integrating machine learning lileitner@ braries (TensorFlow, Chainer, DyNet, VowpalWabbit, FastText, MLlib, Factorie, datacatalytics.com SMILE, CoreNLP, NLP4J, SpaCy, SciKit Learn, etc.). Architecting the curation of finance/economy-related texts for supervised learning and training clients' languages analyst teams to do the annotations. Teaching & hiring. 2014-2015 **Technical University of Madrid (UPM)** Madrid, Spain German: native NeuroNLP researcher (Juan de la Cierva research fellow) Training neurobiology-specific NER taggers (Wapiti, Factorie, word2vec). Min-English: master ing the graph of molecular interactions between neuro-entities using deep parsing (GENIA parser), extracting semantic triplets from (2M+) full-text documents Spanish: proficient (Spark, Theano). Working towards a graph-based entity grounding system (OrientDB). Statistical analyses of citation data (R, Postgres). Teaching. 2013-2014 National Cancer Research Centre (CNIO) Madrid, Spain computing BioNLP researcher (Post-doc) Extracting the mammalian transcriptome from all (22M+) abstracts in medicine text mining and & biology. Detecting abstracts containing mammalian gene regulation informadeep learning tion (SciKit Learn, SPECIES, GENIA tagger). Grounding mentions of genes and transcription factors by developing a rule-based gene symbol mapper. Shallow natural language parsing (Wapiti, CRFSuite) sentences containing genes and factors for tranprocessing scription event extraction (LibSVM). Leading a team of scientists to conduct the curation of mammalian transcriptome data. Teaching. distributed and cloud computing 2006-2013 **National Cancer Research Centre (CNIO)** Madrid, Spain Bioinformatician (PhD student) data warehousing Steering BioCreative (BC) community challenges: task design, orchestrating and engineering data curation, deep system analyses, conference organization, and systematic evaluations (R) of participating systems. Lead architect of the BioCrestatistical and ative Meta-Server, "BCMS", a geographically distributed, streaming NLP and numerical analyses text mining platform (TwistedPython, Postgres), in coopwith a dozen intlie-

(GNU) POSIX systems

C++/Python, Java/Scala (R, SQL, C/CUDA)

software development

and operations on

2003-2006 Institute for Molecular Pathology (IMP)

Data architect (MSc student)

Architecting a molecular [sequence] resource (Postgres) to analyze posttranslational modifications. Creating web interfaces for sequence meta-data via a DSL to bootstrap a web framework (LISP, Struts). Distributed indexing of the resources and designing an information retrieval interface (Lucene, Struts).

search teams. Designing an application to maintain up-to-date, local mirrors

of PubMed, UniProt, and GenBank (Postgres). Mining abstracts for protein-

protein interaction (PPI) information using the BCMS API. Developing the chal-

Vienna, Austria

lenges' web interface and team management framework (Django, Postgres).

profile

Data curation, statistical modeling, and software engineering of intelligent business solutions. Development of domain-specific language understanding and information extraction technology. Architecting & operating distributed data warehouses and online processing frameworks ("DataOps"). Sectors with specialty expertise: Biotech/pharma and finance/banking.

education

2012-now	Continuous professional education Deep learning, distributed data processing, scalable machine learning,	MOOCs IT Sec certifications, etc.
2012	PhD Bioinformatics, with distinction Online assessment of protein interaction information extraction system	Autonomous University of Madrid (UAM), Spain S.
2006	MSc Molecular Biology, with distinction Integration of biological databases and mining for post-translational mo	Vienna University, Austria odifications.

accomplishments

2015	Company Founder of my own consultancy, with U.S. market coverage	Data Catalytics, S.L.
2013	Special Award - Premio Extraordinario for my PhD thesis work	Autonomous University of Madrid (UAM)
2009	Main Conference Organizer three-day conference with nearly 100 participants	BioCreative, CNIO, Madrid
2009	Services Cup 3 rd place behind competitors from SAP and IBM	IEEE Services-I World Conference

communication skills

2015	Invited Talk The BioCreative challenges: Critical assessmen	Jornada de Tecnología y Salud, Real Academia de Ingeniería, Madrid, Spain of biomedical information extraction.	
2014	Invited Talk Institute for Computational Linguistics (UZH), Zurich, Switzerland From the BioCreative challenges to transcription regulation network extraction.		
2014	Progress Report Retracing transcription regulation networks.	National Cancer Research Centre (CNIO), Madrid, Spain	
2013	Presentation BioLINK @ Intelligent Systems for Molecular Biology (ISMB), Berlin, Germany Mining cis-regulatory transcription networks from literature.		
2013	Panel Host Integration across genomics and medicine.	BioLINK @ Intelligent Systems for Molecular Biology (ISMB), Vienna, Austria	
2011	Invited Talk Annotating scientific manuscripts and the role of text-mining. FEBS Journal Board Meeting, Cambridge, UK		
2010	Progress Report Making biological information accessible.	National Cancer Research Centre (CNIO), Madrid, Spain	
2010	Presentation Intelligent Systems for Molecular Biology (ISMB), Stockholm, Sweden Comparative community assessments for applied biomedical text mining.		
2009	Presentation The BioCreative Meta-Server platform.	Semantic Mining in Biomedicine (SMBM), Turku, Finland	
2008–2013	Presentations Various presentations throughout the BC II-IV v	BioCreative, Madrid, Spain and Bethesda, MD, USA workshops (see conference proceedings).	

publications

journal articles

Data publications correlate with citation impact

Leitner, F, C Bielza, SL Hill, and P Larrañaga

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Salgado, D, M Krallinger, M Depaule, E Drula, A Tendulkar, F Leitner, A Valencia, and C Marcelle

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The Protein-Protein Interaction tasks of BioCreative III: ranking of articles and linking bio-ontology concepts to full text

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Text Mining for Drugs and Chemical Compounds: Methods, Tools and Applications

Vazquez, M., M Krallinger, F Leitner, and A Valencia

Molecular Informatics 30.6-7 (2011) pp. 506-519. 2011

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The FEBS Letters SDA corpus.

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Krallinger, M, A Morgan, L Smith, F Leitner, L Tanabe, J Wilbur, L Hirschman, and A Valencia

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Krallinger, Martin, Florian Leitner, Carlos Rodriguez-Penagos, and Alfonso Valencia

Genome Biology 9 Suppl 2 (2008) S4. 2008

Introducing meta-services for biomedical information extraction.

Leitner, F, M Krallinger, C Rodriguez-Penagos, J Hakenberg, ..., C Sander, and A Valencia

Genome Biology 9 Suppl 2 (2008) S6. 2008

A text-mining perspective on the requirements for electronically annotated abstracts.

Leitner, Florian and Alfonso Valencia

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Leitner, Florian, Martin Krallinger, and Valencia Alfonso

Encyclopedia of Systems Biology pp. 106-110, 2013, Springer New York

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Methods in Molecular Biology pp. 341-382, 2010, Humana Press

Biological Knowledge Extraction

Leitner, Florian, Robert Hoffmann, and Alfonso Valencia

Bioinformatics for Systems Biology pp. 413-433, 2009, Humana Press

conference proceedings

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Overview of the chemical compound and drug name recognition (CHEMDNER) task

Krallinger, Martin, Florian Leitner, Obdulia Rabal, Miguel Vazquez, Julen Oyarzabal, and Alfonso Valencia

BioCreative IV Challenge Evaluation Workshop, 2013

Mining cis-Regulatory Transcription Networks from Literature

Leitner, Florian, Martin Krallinger, Sushil Tripathi, Martin Kuiper, Astrid Lægreid, and Alfonso Valencia

Proceedings of the ISMB BioLINK SIG, 2013

Results of the BioCreative III Interaction Method Task

Krallinger, Martin, Miguel Vazquez, Florian Leitner, and Alfonso Valencia

Proceedings of the BioCreative III Challenge Evaluation Workshop, 2010

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Mardis, Scott, Florian Leitner, and Lynette Hirschman

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Starlinger, Johann, Florian Leitner, Alfonso Valencia, and Ulf Leser

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Assessment of the second BioCreative PPI task: automatic extraction of protein-protein interactions

Krallinger, Martin, Florian Leitner, and Alfonso Valencia

Proceedings of the second BioCreative Challenge Evaluation Workshop, 2007

Exploiting text mining of biological annotations to extract protein interactions for the Semantic web

Krallinger, Martin, Carlos Rodriguez-Penagos, Florian Leitner, and Alfonso Valencia

Seventh International Workshop on Network Tools and Applications in Biology, 2007