

12th Annual Bioinformatics Open Source Conference BOSC 2011

Day 1 (Friday, July 15, 2011)

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| Time | Title | Speaker or Session Chair |
| 9:00-9:15 | Introduction | Nomi Harris (Co- Chair, BOSC 2011) |
| 9:15-10:15 | Keynote: The role of openness in knowledge- based systems for biomedicine | Larry Hunter |
| 10:15-10:45 | Coffee Break | |
| 10:45-12:30 | Session: Genome Content Management | Chair: Peter Rice |
| 10:45-11:05 | Unipro UGENE: an open source toolkit for complex genome analysis | Konstantin Okonechnikov |
| 11:05-11:25 | Exploring the genome with Dalliance | Thomas Down |
| 11:25-11:45 | InterMine - Using RESTful Webservices for Interoperability | Alex Kalderimis |
| 11:45-11:55 | easyDAS: Automatic creation of DAS servers | Bernat Gel |
| 11:55-12:05 | Enacting Taverna Workflows through Galaxy | Kostas Karasavvas |
| 12:05-12:15 | Mobyle 1.0: new features, new types of services | Hervé Ménager |
| 12:15-12:25 | BioMart 0.8 offers new tools, more interfaces, and increased flexibility through plug-ins | Junjun Zhang |
| 12:25-12:32 | Running Workflows Through Taverna Server | Donal Fellows |
| 12:30-2:00 | Lunch | |
| 1:30-2:30 | Poster Session | |
| 2:30-3:30 | Session: Visualization | Chair: Jan Aerts |
| 2:30-2:50 | Cytoscape 3.0: Architecture for Extension | Michael Smoot |
| 2:50-3:10 | Applying Visual Analytics to Extend the Genome Browser from Visualization Tool to Analysis Tool | Jeremy Goecks |
| 3:10-3:20 | WebApollo: A web-based sequence annotation editor for community annotation | Nomi Harris |
| 3:20-3:30 | The isobar R package: Analysis of quantitative | Florian P. |
| 3.20-3.30 | proteomics data | Breitwieser |
| 3:30-4:00 | | |
| | proteomics data | |
| 3:30-4:00 | proteomics data Coffee Break | Breitwieser Chair: Thomas |
| 3:30-4:00 4:00-5:30 | proteomics data Coffee Break Session: Next-Generation Sequencing Stacks: building and genotyping loci de novo from | Breitwieser Chair: Thomas Down |
| 3:30-4:00 4:00-5:30 4:00-4:20 | proteomics data Coffee Break Session: Next-Generation Sequencing Stacks: building and genotyping loci de novo from short-read sequences Large scale NGS pipelines using the MOLGENIS | Chair: Thomas Down Julian Catchen |

| Time | Title | Speaker or Session Chair |
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| 5:00-5:10 | A Scalable Multicore Implementation of the TEIRESIAS Algorithm | Frank Drews |
| 5:10-5:20 | Biomanycores, open-source parallel code for manycore bioinformatics | Jean-Frédéric Berthelot |
| 5:20-5:30 | GemSIM: General, Error-Model Based Simulator of next-generation sequencing | Kerensa McElroy |
| 5:30-6:30 | Poster Session and BOFs | |
| 7:00 | Optional dinner for BOSC attendees | Location TBA |

Day 2 (Saturday, July 16, 2011)

| Time | Title | Speaker or Session Chair |
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| 8:45-8:50 | Announcements | Nomi Harris and Peter Rice |
| 8:50-9:50 | Keynote: Into the Wonderful | Matt Wood |
| 9:50-10:15 | Securing and sharing bioinformatics in the cloud | Richard Holland |
| 10:15-10:45 | Coffee Break | |
| 10:45-12:30 | Session: Cloud Computing | Chair: Brad Chapman |
| 10:45-11:05 | Mygene.info: Gene Annotation as a Service - GAaaS | Chunlei Wu |
| 11:05-11:25 | Cloud BioLinux: open source, fully-customizable bioinformatics computing on the cloud for the genomics community and beyond | Konstantinos Krampis |
| 11:25-11:35 | OBIWEE : an open source bioinformatics cloud environment | Jonathan Piat |
| 11:35-11:45 | SeqWare: Analyzing Whole Human Genome Sequence Data on Amazon's Cloud | Brian O'Connor |
| 11:45-12:05 | Sequencescape - a cloud enabled Laboratory Information Management Systems (LIMS) for second and third generation sequencing | Lars Jorgensen |
| 12:05-12:15 | Enabling NGS Analysis with(out) the Infrastructure | Enis Afgan |
| 12:15-12:25 | Hadoop-BAM: A Library for Genomic Data Processing | Aleksi Kallio |
| 12:30-2:00 | Lunch | |
| 1:00-2:00 | Poster Session | |
| 2:00-3:30 | Session: Semantic Web and Misc. Open Source Projects | Chair: Peter Cock |
| 2:00-2:10 | SADI for GMOD: Bringing Model Organism Data onto the Semantic Web | Ben Vandervalk |
| 2:10-2:20 | Scufl2: Because a workflow is more than its definition | Stian Soiland-Reyes |
| 2:20-2:30 | OntoCAT - an integrated programming toolkit for common ontology application tasks | Tomasz Adamusiak |
| 2:30-2:50 | Debian Med: individuals' expertise and their sharing of package build instructions | Steffen Möller |

| Time | Title | Speaker or Session Chair |
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| 2:50-3:10 | The BALL project: The Biochemical Algorithms Library (BALL) for Rapid Application Development in Structural Bioinformatics and its graphical user interface BALLView | Andreas Hildebrandt |
| 3:10-3:20 | Biopython Project Update | Peter Cock |
| 3:20-3:27 | What's new with GMOD | Scott Cain |
| 3:27-3:34 | Exploring human variation data with Clojure | Brad Chapman |
| 3:30-4:00 | Coffee Break | |
| 4:00-4:30 | Session: Misc. Open Source Projects | Chair: Jim Procter |
| 4:00-4:10 | EMBOSS: New developments and extended data access | Peter Rice |
| 4:10-4:20 | G-language Project: the last 10 years and beyond | Kazuharu Arakawa |
| 4:20-4:30 | A Framework for Bioinformatics on the Microsoft Platform | Simon Mercer |
| 4:30-5:20 | Panel: Multi-Institution Collaboration | Moderator: Brad Chapman Panelists: Richard Holland, Hilmar Lapp, Jean Peccoud, Peter Rice |
| 5:20-5:30 | Presentation of awards | Nomi Harris |
| 5:30-6:30 | BOFs | |

Last-minute schedule updates will be posted at http://www.open-bio.org/wiki/BOSC_2011_Schedule

Posters

| No. | Poster Title and Author(s) |
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| 1 | Unipro UGENE: an open source toolkit for complex genome analysis (Okonechnikov |
| | et al) |
| 2 | Exploring the genome with Dalliance (Down, Hubbard) |
| 3 | InterMine - Using RESTful Webservices for Interoperability (Kalderimis et al.) |
| 4 | easyDAS: Automatic creation of DAS servers (Gel et al.) |
| 5 | Enacting Taverna Workflows through Galaxy (Karasavvas et al.) |
| 6 | Mobyle 1.0: new features, new types of services (Menager et al.) |
| 7 | BioMart 0.8 offers new tools, more interfaces, and increased flexibility through plug- |
| | ins (Zhang et al.) |
| 8 | Running Workflows Through Taverna Server (Fellows et al.) |
| 9 | Cytoscape 3.0: Architecture for Extension (Smoot) |
| 10 | Applying Visual Analytics to Extend the Genome Browser from Visualization Tool to |
| | Analysis Tool (Goecks et al.) |
| 11 | The isobar R package: Analysis of quantitative proteomics data (Breitwieser, Colinge) |
| 12 | Large scale NGS pipelines using the MOLGENIS platform: processing the Genome of |
| | the Netherlands (Byelas et al.) |
| 13 | Bio-NGS: BioRuby plugin to conduct programmable workflows for Next Generation |
| | Sequencing data (Bonnal et al.) |
| 14 | Goby framework: native support in GSNAP, BWA and IGV 2.0 (Dorff et al.) |
| 15 | Biomanycores, open-source parallel code for many-core bioinformatics (Giraud et al.) |

| 16 | GemSIM: General, Error-Model Based Simulator of next-generation sequencing (McElroy et al.) | |
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| 17 | Mygene.info: Gene Annotation as a Service - GAaaS (Wu, Su) | |
| 18 | Enabling NGS Analysis with(out) the Infrastructure (Afgan et al.) | |
| 19 | Hadoop-BAM: A Library for Genomic Data Processing (Niemenmaa et al.) | |
| 20 | SADI for GMOD: Bringing Model Organism Data onto the Semantic Web (Vandervalk | |
| | et al.) | |
| 21 | Scufl2: because a workflow is more than its definition (Sioland-Reves et al.) | |
| 22 | OntoCAT - an integrated programming toolkit for common ontology application tasks (Adamusiak et al.) | |
| 23 | Debian Med: individuals' expertize and their sharing of package build instructions (Moeller et al.) | |
| 24 | The BALL project: The Biochemical Algorithms Library (BALL) for Rapid Application Development in Structural Bioinformatics and its graphical user interface BALLView (Hildebrandt et al.) | |
| 25 | What's new with GMOD (Cain et al.) | |
| 26 | EMBOSS: New developments and extended data access (Rice et al.) | |
| 27 | A tool kit for pre-processing genome annotations in Generic Feature Format (GFF) (Sreedharan) | |
| 28 | ADDAPTS: A Data-Driven Automated Pipeline and Tracking System (Naravan et al.) | |
| 29 | BALLView: A versatile molecular visualization and modeling tool (Nickels et al.) | |
| 30 | Beyond Web Services: Calling External Programs from within Taverna (Williams et al.) | |
| 31 | BioMart combines the power of the semantic web with the speed of relational database querying (Baran et al.) | |
| 32 | BioSharing: standards, policies and communication in bioscience (Sansone et al.) | |
| 33 | BioXSD: the XML Schema for everyday bioinformatics (Kalas et al.) | |
| 34 | Cloudgene – A graphical MapReduce interface for cloud computing (Forer et al.) | |
| 35 | Combining RapidMiner operators with bioinformatics services - a powerful combination (Jupp et al.) | |
| 36 | Distributed Web Services for Bioinformatics: Multiple Sequence Alignment (Troshin et al.) | |
| 37 | Endrov - an open source framework for image processing and analysis (Henriksson, Burglin) | |
| 38 | fastapl a utility for processing fasta format data (Horton) | |
| 39 | FLAME: An Open-Source Agent-Based Modelling Platform for High Performance | |
| 40 | Computing of Biological Systems (Bicak et al.) GeneNomenclatureUtils: Tools for annotating genes and comparing gene lists with | |
| 40 | community resources (Croning, Grant) | |
| 41 | GenomeTools - a versatile and efficient bioinformatics toolkit (Gremme et al.) | |
| 42 | InterMine - An Open Source Data Warehouse and Query interface (Hu et al.) | |
| 43 | Jalview 2: Transitions (Procter et al.) | |
| 44 | jsDAS and DASTY3, enabling DAS protein visualization (Garcia et al.) | |
| 45 | KNIME-CADDSuite – integration of the computer aided drug design suite | |
| 1 | (CADDSuite) into the Konstanz Information Miner (KNIME) (Roettig et al.) | |
| 46 | (CADDSuite) into the Konstanz Information Miner (KNIME) (Roettig et al.) OncoPortal: the power of BioMart for oncology research (Wong-Erasmus et al.) | |
| 46 47 | | |
| | OncoPortal: the power of BioMart for oncology research (Wong-Erasmus et al.) | |
| 47 | OncoPortal: the power of BioMart for oncology research (Wong-Erasmus et al.) SEQCRAWLER – A cloud ready indexing platform (Sallou) | |

Ballot for Audience Favorite Talk

Please vote for one talk and give this form to Nomi Harris or Peter Rice (BOSC Co-Chairs) before 4:30pm on the second day. (Note that this ballot is two pages!)

The winner will be announced in the closing session at 5:20 on the second day.

| Best Talk? | Title | Speaker |
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