Challenge Session Speakers

We have selected a range of speakers that, together with the keynote speaker, will cover developments in a broad range of active research topics in modern biological data visualization, from genes to proteins to organisms to populations. The speakers include developers of visualization tools widely used in biology, as well as active users who apply visualization methods to discover new biological knowledge.

Engaging today’s genomics resources

Ting Wang
Assistant Professor at the Genetics Department and Center for Genome Sciences, Washington University in St. Louis

Abstract

Advances in next-generation sequencing platforms have reshaped the landscape of genomic and epigenomic research. Large consortia such as the ENCylopedia of DNA Elements (ENCODE), the Roadmap Epigenomics project and The Cancer Genome Atlas (TCGA) have generated tens of thousands of sequencing-based genome-wide datasets, creating a reference and resource for the scientific community. Small groups of researchers are now able to rapidly obtain huge volumes of genomic data and are in great need of putting their own data in the context of the consortium data for comparison. These data are often accompanied by rich metadata describing the sample and experiment, critical for their interpretation. Fast evolving technologies, such as those interrogating long-range chromatin interactions, are generating new types of data in similarly large volumes. Visualizing, navigating, and interpreting such data in a meaningful way is a daunting challenge. In this presentation I will discuss the present challenges in genomic data visualization and the rapid evolution of Genome Browsers.

Bio

Ting Wang is an assistant professor at the Genetics Department and Center for Genome Sciences at Washington University in St. Louis. He has a PhD in Computational Biology. His research focuses on understanding genetic and epigenetic impact of transposable element on human regulatory networks and their role in human diseases. He develops algorithms for identifying regulatory motifs, and analytical and visualization methodologies to integrate genomic and epigenomic data. He is a co-inventor of the UCSC Cancer Genomics Browser, inventor of the Wash U Epigenome Browser, and a co-investigator of the Epigenome Roadmap Mapping Centers.

Tools for Interactive Visualization and Analysis of Complex Biological Structures

Thomas Ferrin
Professor, Department of Pharmaceutical Chemistry and Department of Bioengineering and Therapeutic Sciences; Director, Resource for Biocomputing, Visualization, and Informatics, University of California, San Francisco

Abstract

As experimental methods to determine the detailed structures of molecules and molecular assemblies continue to rapidly advance, and as databases of modeled macromolecular complexes also swiftly grow in both size and complexity, the challenges of easily visualizing, analyzing, and comparing these structures become increasing difficult. Particularly onerous are models that encompass scales that range from the atomic to cellular and data that is dynamical in nature. Yet the ambitious goal of creating an accurate spatiotemporal model of the cell and all of its molecular processes, with maximal accuracy, resolution, and completeness, is nevertheless crucial to a thorough understanding of biological function and the molecular basis of disease. Because of the complexity of this data, interactive 3D visualization tools greatly facilitate analysis tasks by enabling researchers to explore, interpret, and manipulate their data. This talk will highlight the challenges associated with visualization of macromolecules at multiple spatial resolutions and illustrate some of the existing tools used to address this problem domain. Also to be discussed are the challenges that tool builders must face if their tools are to be adopted by the larger scientific community.

Bio

Tom originally hails from Minneapolis, where he first became interested in computing while writing codes for a Control Data Corp. model 6600 computer, the first machine to be called a “supercomputer.” In the early 1970’s, while working as a bioengineer, he became enamored with the application of minicomputers such as the Digital Equipment Corp. PDP-8 and Data General Nova for real-time control and analysis of scientific experiments. As the data collected from these experiments became larger and more complex, he turned to interactive computer graphics as a means of displaying and analyzing the data, and ultimately gaining insight into the underlying biophysical processes at play. Thus began a 40+ year career at the University of California, San Francisco (UCSF) focused on developing tools for the interactive visualization and analysis of molecular structure at scales ranging from the atomic to the supramolecu-
lar. In addition to tool development, Tom’s lab has a long history of documenting, distributing, and providing training for these tools. Their UCSF Chimera package is used worldwide and has been cited in more than 5,400 journal articles. In recent years his research group has also contributed tools that emphasize integrating diverse types of biomolecular data, including atomic-resolution coordinates, density maps, sequences, annotations, and network representations of protein similarity, binding interactions, and biological pathways.