BioHDF : Open binary file formats for large scale data management
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The first wave of Next Generation ("Next Gen") sequencing technologies are providing
large numbers of laboratories with "Genome Center" kinds of throughput to make
discoveries and develop new assays never before imagined. However, widespread
adoption of Next Gen will be hindered because current bioinformatics programs do not
scale; they are inefficient in data storage, processing, and memory utilization. The most
popular programs typically copy and recopy data to new files many times during
processing, require that all data be maintained in random access memory (RAM) when
running, and cannot incrementally process data. To overcome these issues, fundamental changes in data management and processing are needed.

Geospiza and The HDF Group are collaborating to develop portable, scalable,
bioinformatics technologies based on HDF5 (Hierarchical Data Format -
http://www.hdfgroup.org). We call these extensible domain-specific data technologies
"BioHDF." BioHDF will implement a data model that supports primary DNA sequence
information (reads, quality values, meta data) and results from sequence assembly and
variation detection algorithms. BioHDF will extend HDF5 data structures and library
routines with new features (indexes, compression, graph layouts) to support the high
performance data storage and computation requirements of Next Gen Sequencing.
BioHDF will include APIs, software tools, and a viewer based on HDFView to enable its
use in the bioinformatics and research communities. Using BioHDF, researchers will be
able to perform de novo sequencing, do resequencing-based SNP discovery, analyze
genotyping data, and export datasets in formats ready for submission to key databases.
As a programming environment, BioHDF can be easily extended to accept data from
new data collection platforms and format data for interchange with many databases.
BioHDF will be delivered to the research community as an open source technology.

In preliminary studies, HDF5’s feasibility for managing large volumes and complex
biological data was tested. The first test case looked at DNA sequencing-based SNP
discovery. Through this study, HDF’s strengths and data organization features (groups,
sets, multidimensional arrays, transformations, linking objects, and general data storage
for other binary data types and images) were evaluated to determine how well these
features would handle SNP data. Other test cases were added to test the ability of HDF
to handle extremely large datasets associated with HapMap data and chromosomal
scale LD (Linkage Disequilibrium) calculations. Data from preliminary studies and new
work with Next Gen sequence data will be presented.