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Summary: The cBio Cancer Genomics Portal (http://cbioportal.org) is an open-access resource for interactive exploration of multidimensional cancer genomics data sets, currently providing access to data from more than 5,000 tumor samples from 20 cancer studies. The cBio Cancer Genomics Portal significantly lowers the barriers between complex genomic data and cancer researchers who want rapid, intuitive, and high-quality access to molecular profiles and clinical attributes from large-scale cancer genomics projects and empowers researchers to translate these rich data sets into biologic insights and clinical applications. Cancer Discov; 2(5): 401–4. ©2012 AACR.

With the rapidly declining cost of next-generation sequencing, and major national and international efforts, including The Cancer Genome Atlas (TCGA) and the International Cancer Genome Consortium (ICGC) (1), the field of cancer genomics continues to advance at an extraordinarily rapid pace. Data generated by these projects are, however, not easily or directly available to the cancer research community, hindering the translation of genomic data into new biologic insights, drugs, and clinical trials. The cBio Cancer Genomics Portal (http://cbioportal.org), developed at Memorial Sloan-Kettering Cancer Center (MSKCC), was specifically designed to address the unique data integration issues posed by large-scale cancer genomics projects and to make the raw data generated by large-scale cancer genomic projects more easily and directly available to the entire cancer research community (Fig. 1A).

The cBio portal currently contains 5 published data sets (2–5) and 15 provisional TCGA data sets. Provisional TCGA data sets are updated monthly, based on the latest TCGA production runs, and the portal will be continually updated as new TCGA cancer types are added. Published data sets include mutation data, but provisional data sets currently do not. As each cancer type within TCGA is finalized and somatic mutations are validated, mutation data will be released and added to the portal. In addition to mutation data, the portal includes copy number alterations, microarray-based and RNA sequencing-based mRNA expression changes, DNA methylation values, and protein and phosphoprotein levels.

Each data type is stored at the gene level and is then combined with available deidentified clinical data such as overall survival and disease-free survival intervals. The data are then organized as a function of patient and gene, and the portal’s fundamental abstraction is the concept of altered genes; specifically, we classify a gene as altered in a specific patient if it is mutated, homozygously deleted, amplified, or its relative mRNA expression is less than or greater than a user-defined threshold. The notion of altered genes is a powerful simplifying concept that enables users to analyze complex data sets and to develop biologic hypotheses regarding recurrently altered gene sets and biologic pathways.

A key feature of the cBio portal is ease of use. All features of the portal are therefore available through a streamlined 4-step web interface. Specifically, users are guided to select: 1) a cancer study of interest, for example, TCGA Glioblastoma Multiforme (GBM); 2) one or more genomic profiles, for example, mutations and copy number alterations; 3) a patient case set, for example, all “complete” TCGA patients with GBM with mutation, copy number, and mRNA data; and 4) a gene set of interest: users can enter HUGO gene symbols, gene aliases, or Entrez Gene IDs and can enter arbitrary gene sets or pathways of interest. Users also have the option to automatically compute mutual exclusivity and co-occurrence between all pairs of genes. Finally, users have the option of performing cross-cancer queries, a simpler 2-step query, which requires only that users select “All Cancer Studies” and enter a gene set of interest.

For example, to visualize genomic alterations in the retinoblastoma (RB) pathway in the TCGA GBM data, one selects options 1 to 3 as described previously and in step 4 enters: RB1, CDK4, and CDKN2A. Based on the user input, the portal automatically generates a series of reports, each in a separate tab. The first of these reports summarizes genomic data across all patients through a concise graphical summary called an OncoPrint. In this graphical summary, individual genes are represented as rows, individual cases or patients are represented as columns, and glyphs and/or color-coding is used to compactly summarize distinct genomic alterations, including somatic mutations, copy number alterations, and mRNA expression. OncoPrints can be extremely useful for visualizing gene set and pathway...
alterations across a set of cases and for visually identifying trends such as trends in mutual exclusivity or co-occurrence between gene pairs within a gene set. For example, the RB OncoPrint (Fig. 1B) shows that alterations of different members in the RB pathway are mutually exclusive. Statistical tests for co-occurrence/mutual exclusivity are also available in a separate tab.

Other reports, each available within a separate tab, include Network Analysis, Correlation Plots, Survival Analysis, Mutation Details, Event Map, Data Download, and Bookmark/E-mail. From these additional reports, we can, for example, observe that many RB1 mutations may have strong functional consequences (Fig. 1C, Mutation Details), as predicted by MutationAssessor.org (6). We can further assess that alterations across a set of cases and for visually identifying trends such as trends in mutual exclusivity or co-occurrence between gene pairs within a gene set. For example, the RB OncoPrint (Fig. 1B) shows that alterations of different members in the RB pathway tend to be mutually exclusive. Statistical tests for co-occurrence/mutual exclusivity are also available in a separate tab.
CDK4 mRNA expression is elevated in amplified cases (Fig. 1D, Plots Tab) and that cases with an RB pathway alteration have worse overall survival than cases without an RB pathway alteration ($P = 0.0513$, log-rank test; Fig. 1E, Survival Tab). Users can also click the Event Map or Data Download reports to copy and paste event information into an external spreadsheet application or click the Bookmark/E-mail tab to share their results with collaborators. Users can also visualize copy number details by choosing to launch a web start version of the Integrative Genomics Viewer [IGV (7)].

The network tab provides interactive analysis and visualization of networks altered in the chosen cancer study. The network consists of pathways and interactions derived from the open-source Pathway Commons Project (8). By default, the network of interest contains all neighbors of all seed genes specified by the user. If more than 50 neighbor nodes exist in the network, all genes are ranked by the frequency of genomic alteration within the specified cancer study and less frequently altered genes are automatically pruned from the network. By default, the portal also automatically overlays multidimensional genomic data onto each node, highlighting the frequency of alteration by mutation and copy number alteration (and optionally mRNA up-/downregulation). This provides an effective means of managing network complexity while automatically highlighting those genes most directly relevant to the cancer type in question. One can also download the full, nonpruned network for more complete visualization and analysis.

For example, we used the portal to identify genomic alterations in the homologous recombination (HR) DNA repair pathway in serous ovarian cancer. BRCA1 and BRCA2 are known to be involved in the HR pathway, but additional defects may also abrogate HR functionality and lead to potential sensitivity to PARP inhibitors (9). To identify potential HR defects in ovarian cancer, we used BRCA1 and BRCA2 as seed nodes for the network view and explored the resulting altered network of interest (Fig. 1F). By this means, we quickly identified alterations in C11orf30/EMSY (6% by amplification, 1.6% by mutation), a known interactor of BRCA2, as a possible alternate means for abrogating HR functionality (9). Users can also filter the network by alteration frequency, as a possible alternate means for abrogating HR functionality (9). To identify potential sensitivity to PARP inhibitors (9). To identify potential sensitivity to PARP inhibitors (9). To identify potential sensitivity to PARP inhibitors (9). To identify potential sensitivity to PARP inhibitors (9).

In summary, the cBio portal facilitates access to cancer genomic data sets for the entire biomedical community. It provides a simple yet flexible interface to integrated data sets, intuitive visualization options, and a programmatic web interface, all of which can aid researchers in translating cancer genomic data into biologic insights and potential clinical applications. By integrating multiple genomic data types and lowering the barrier to access, the portal enables researchers to more easily mine genomic data, test hypotheses regarding genetic alterations in cancer, and place genomic data into the context of prior biologic knowledge. The cBio portal complements existing tools, including the TCGA and ICGC data portals (13), the IGV (7), the UCSC Cancer Genomics Browser (14), and IntOGen (15) by offering a unique focus on analyzing discrete genomic events across integrated data types, ease of use, support for exploratory data analysis, and interactive network analysis.

We anticipate several future directions for the portal. First, we intend to add many additional cancer studies, mostly from the TCGA and ICGC. Based on the current production schedules, we anticipate that the public portal will grow by at least 5 additional tumor types and more than 1,000 tumor samples by the third quarter of 2012. Second, we plan to add several new features, including complete support for miRNA expression, interactive OncoPrints, batch download of complete data sets, summary reports for cancer studies (e.g., frequently mutated genes), and further extensions to the cross-tumor query analysis.

User support for the cBio Cancer Genomics Portal is available via e-mail at: cbioportal@googlegroups.com.
Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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REFERENCES

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