OncoCL-KB, a cancer cell knowledgebase
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THE OncoCL FRAMEWORK

CANCER CELL MESOPHENOTYPE

OncoCL provides a framework for consistent annotation of cancer-associated data allowing queries for how a particular cancer cell type is situated in a pathway of cancer progression. OncoCL constructs a ‘mesophenotype’ to describe the cancer cell reusing other ontologies as much as possible. The CanonicaCell is a normal cell type as represented in CL, the cell type ontology. The CanonicaCell has Mesophenotype comprised of canonical characteristics of that cell type: normal genotype, normal cellular phenotype (such as potency, proliferative capacity, morphology, function) and normal tissue and anatomical location.

IMPLEMENTATION

OncoCL is implemented in Protégé-OWL. We construct five classes: CanonicaCell, CancerCell, Mesophenotype, CancerRelatedGene and CancerHallmark. These classes are embedded in BFO, the Basic Formal Ontology, which facilitates logical consistency checking. Protégé allows browsing of the class hierarchy and definitions of all terms. We include a library of images where appropriate to facilitate annotation (Balloff). Here we show images for the classes ‘adenoma’ and ‘crypt.’

ABSTRACT

OncoCL-KB is a knowledgebase built upon the semantics of our previously developed ontology for describing cancer cell types. OncoCL, OncoCL provides a framework for bringing disparate data together in a structured way, such that a morphologic entity as defined by conventional pathology is ‘anchored’ within cancer progression, associated with its canonical biological counterpart/origin and linked to the molecular genetic abnormalities that characterize it along with the features and properties imparted by the corresponding disrupted cellular pathways, for example, escape from growth regulators or evasion of apoptosis. OncoCL makes use of a number of other molecular and clinical ontologies. The cell type ontology, CL, describes normal cell types and was not designed to capture the pathology of cancer cells. OncoCL builds upon CL as a canonical cell (represented in CL) undergoes oncogenic change, and tumorigenesis with the acquisition of the cancer hallmarks described by Hanahan and Weinberg. Cellular phenotypes are described using the Phenotypic Quality Ontology (PATO) and the Gene Ontology (GO) and cellular location (normal or metastatic) is described using the anatomy ontology, UBERON. OncoCL-KB embeds annotated data sets – including cancer-associated genes and genomic variants, cancer-associated pathways, cancer stem cell markers, and cancer mouse models – in the OncoCL semantic framework that has been checked for logical consistency and valid inference structure. We hope that OncoCL-KB, through the synthesis of complex heterogeneous data related to cancer cells and cancer progression, will provide a resource that will contribute to a better understanding of cancer predisposition, diagnosis, and treatment. This work is supported by NIH with funding through NCI CA155825.

REPRESENTING CANCER PROGRESSION

In OncoCL, cancer progression is represented as shown below: a CanonicaCell undergoes oncogenic change in a CancerRelatedGene thus acquiring a CancerHallmark to become a CancerCell with a change in Mesophenotype. The CancerCell can go on to acquire additional CancerHallmarks due to additional molecular changes in other CancerRelatedGenes to become a different type of CancerCell.

CHARACTERIZING CANCER CELLS IS A CHALLENGE: CSCs

Cancer Stem Cell (CSC) based therapies must distinguish CSCs not only from non-CSC tumor cells but also from normal stem cells. Various CSC specific methods have been studied and seem promising, among them:
- Identification of distinctive CSC surface markers
- Overexpression, mutations, variant isoforms of CSC surface markers
- Evidence of glycosylation of CSC surface marker proteins
- microRNA regulation of stemness genes in CSCs
- The existence of abnormal pre-cancer cell stem cells.

We have extended the OncoCL mesophenotype to capture these data.

CAPTURING CELL SURFACE MARKER DATA

CL describes immune cells in terms of cell surface markers (Diehl et al). The graph shows, for example, that a CD2-positive, CD5-positive, CD44-positive alpha-beta intraepithelial T cell is a type of CD4-positive, alpha-beta intraepithelial T cell which develops from CD4-positive, alpha-beta thymocyte and is part of intestinal epithelium. The cell type under consideration is positive for cell surface markers CD2, CD44, and CD5, which is captured using the ‘has_plasma_membrane_part’ relation to those proteins. We use this procedure in OncoCL.

RELATION ONTOLOGY IS KEY TO LINKING CANCER CELLS TO CELL SURFACE MARKERS

According to good ontology design principles we use only relations in RO, the relation ontology. To describe cancer stem cells we relate them to cell surface markers by means of RO relations: ‘has_plasma_membrane_part’, ‘has_high_plasma_membrane_amount’ and ‘has_low_plasma_membrane_amount’.

BIOMEDICAL ONTOLOGIES PROVIDE THE PIECES

Biomedical ontologies provide a shared understanding of a domain that is human intelligible and computer readable and, consequently, a representational system to support the integration and retrieval of the biological information. In addition, mappings among ontologies enables data integration.

PROTEIN ONTOLOGY, PRO

We use PRO (Natale et al) to describe CSC surface markers. PRO provides a formal classification of protein classes including isoforms, variants and modified forms with protein modifications, such as glycosylation, described by PSI-MOD, the protein modification ontology.

SEQUENCE ONTOLOGY, SO

We use SO (Mungall et al) to describe CSC-specific microRNAs and CSC-specific epigenetic modifications.

CONCLUSIONS

We have augmented OncoCL to better characterize cancer stem cells by allowing the capture of data related to CSC surface markers including distinguishing CSC surface protein modifications, CSC-specific microRNAs and CSC-specific epigenetic modifications. In future work we plan to include descriptions of the tumor microenvironment, characteristics to distinguish EMT state from CSC phenotype, and extend the mesophenotype to include aspects that would distinguish CSCs that arise from normal stem cells from those that arise from other progenitor or differentiated cells.

REFERENCES

Diehl AD et al. (2011) Hematopoietic cell types. Protégé for a macromolecular library. Journal of Biomedical Informatics 44(1)