

Progenetix & GA4GH

A cancer genomics resource built around and driving GA4GH standards

Michael Baudis | SIB - Novartis Symposium | 2021-09-22



1992



Heidelberg

Student of medicine | doctoral thesis in molecular cytogenetics @ DKFZ (Peter Licher) | resident in clinical hematology/oncology | data, clinical studies & cancer systematics

2001



Stanford

Post-doc in hemato-pathology (Michael Cleary) | molecular mechanisms of leukemogenesis | transgenic models | expression arrays | systematic cancer genome data collection | *Progenetix* website

2003



Gainesville

Assistant professor in paediatric haematology | molecular mechanisms of leukemogenesis | focus on bioinformatics for cancer genome data analysis

2006



Aachen

Research group leader in genetics | genomic array analysis for germline alterations | descriptive analysis of copy number aberration patterns in cancer entities

2007



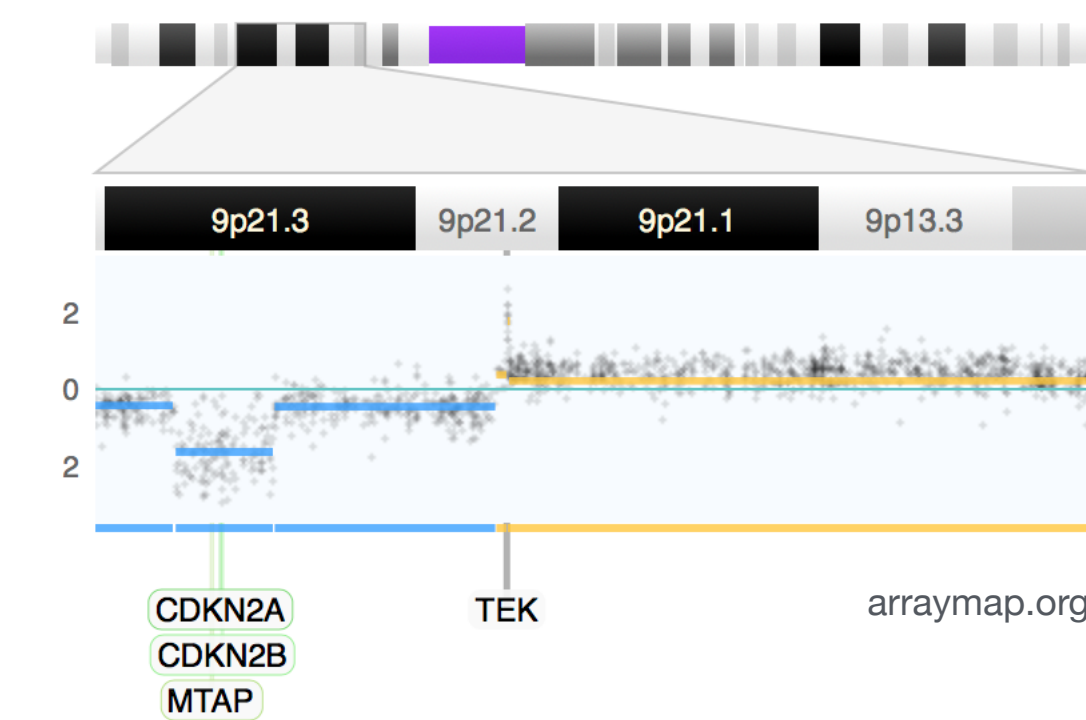
Zürich

Professor of bioinformatics @ IMLS (2015) | systematic assembly of oncogenomic data | databases and software tools | patterns in cancer genomes | *Progenetix* & *arrayMap* resources | GA4GH | SPHN

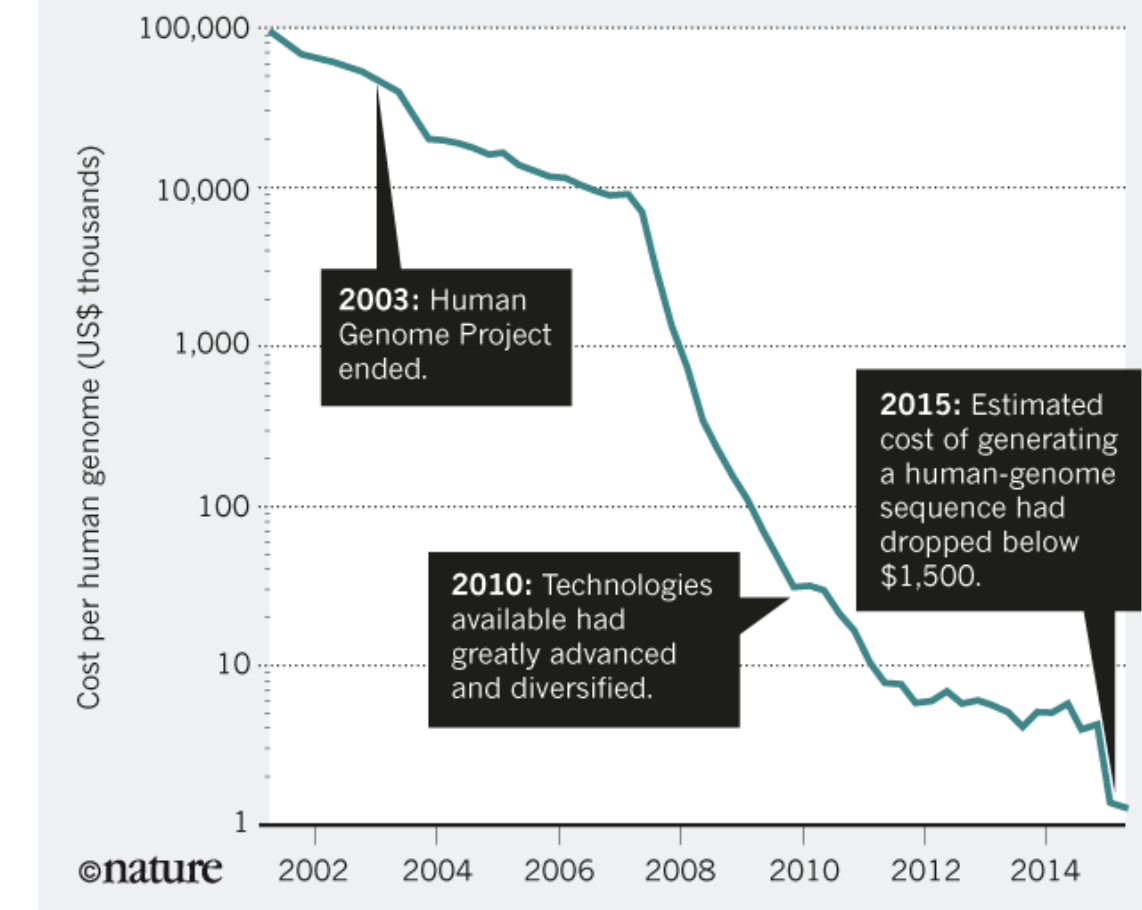


Genome screening at the core of “Personalised Health”

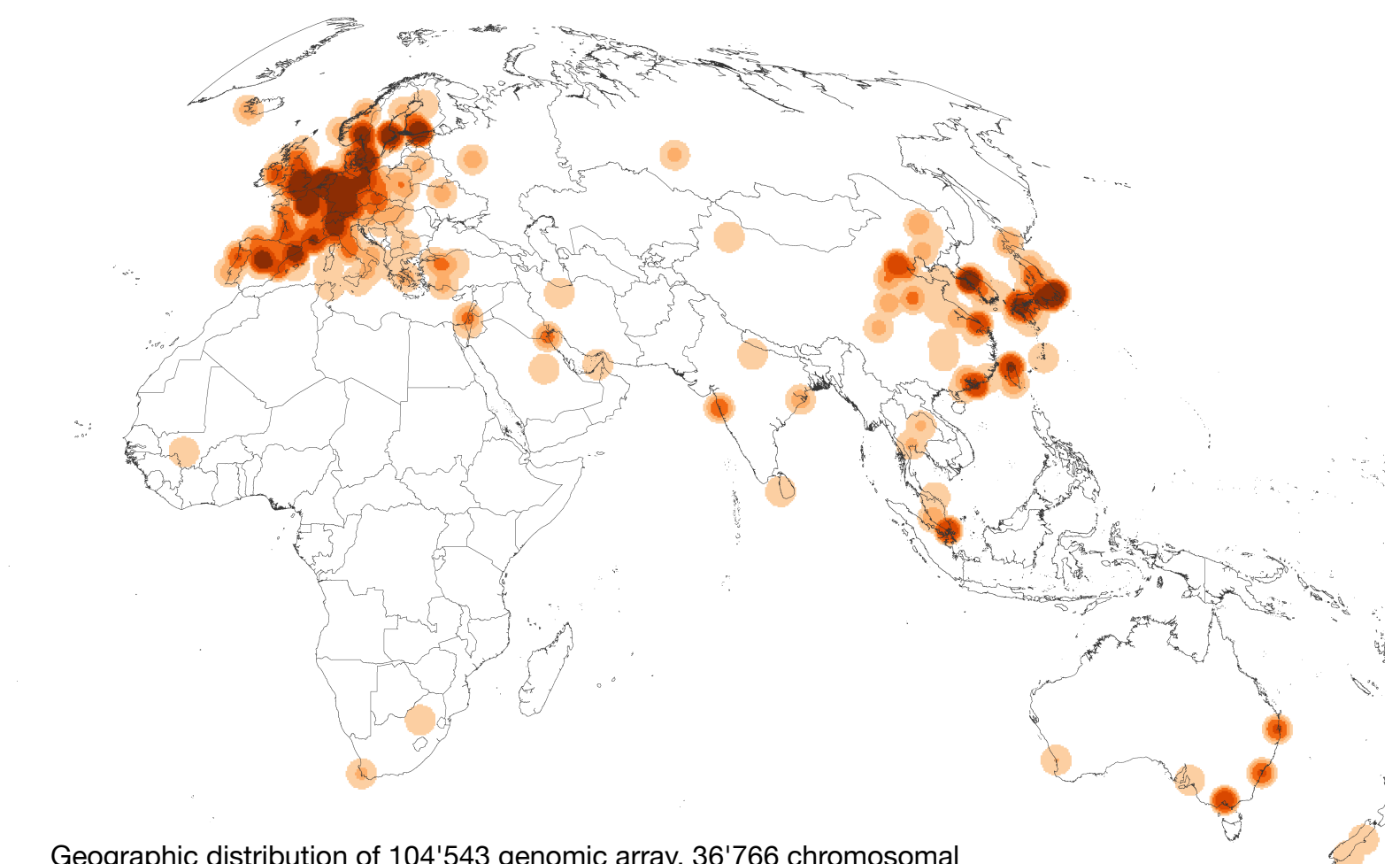
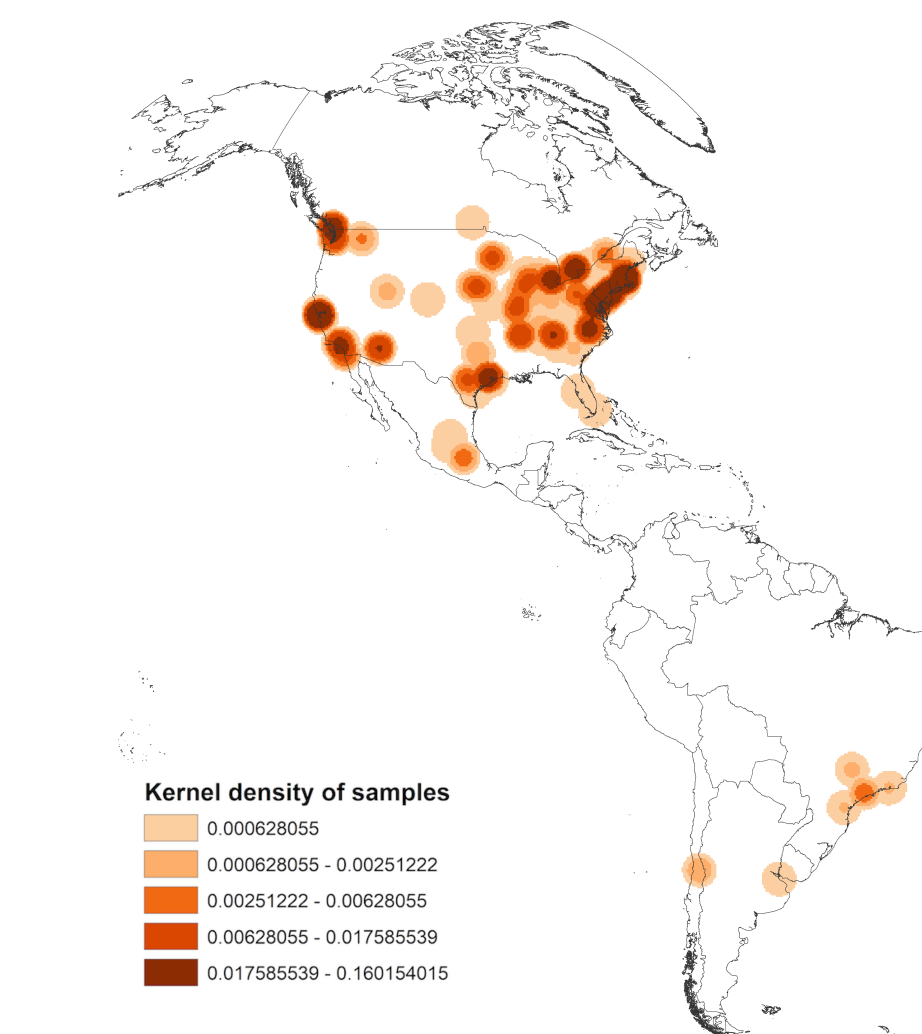
- ▶ **Genome analyses** (including transcriptome, metagenomics) are core technologies for Personalised Health™ applications
- ▶ The unexpectedly large amount of **sequence variants** in human genomes - germline and somatic/cancer - requires huge analysis efforts and creation of **reference repositories**
- ▶ **Standardized data formats** and **exchange protocols** are needed to connect these resources throughout the world, for reciprocal, international **data sharing** and **biocuration** efforts
- ▶ Our work @ UZH:
 - ▶ **cancer genome repositories**
 - ▶ **biocuration**
 - ▶ **protocols & formats**



BETTER, CHEAPER, FASTER
The cost of DNA sequencing has dropped dramatically over the past decade, enabling many more applications.



The future of DNA sequencing. Eric D. Green, Edward M. Rubin & Maynard V. Olson. Nature; 11 October 2017 (News & Views)

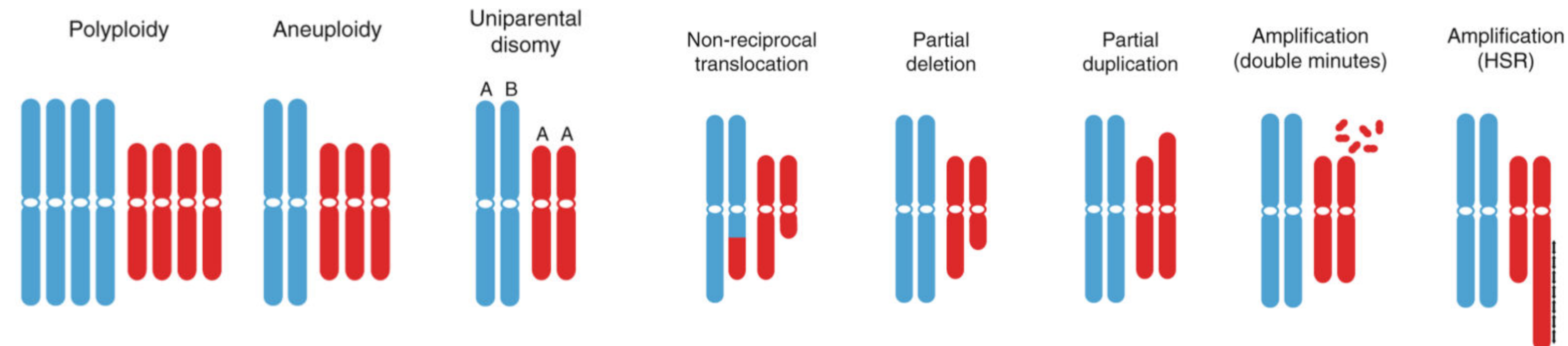


Geographic distribution of 104'543 genomic array, 36'766 chromosomal CGH and 15'409 whole genome/exome based cancer genome datasets

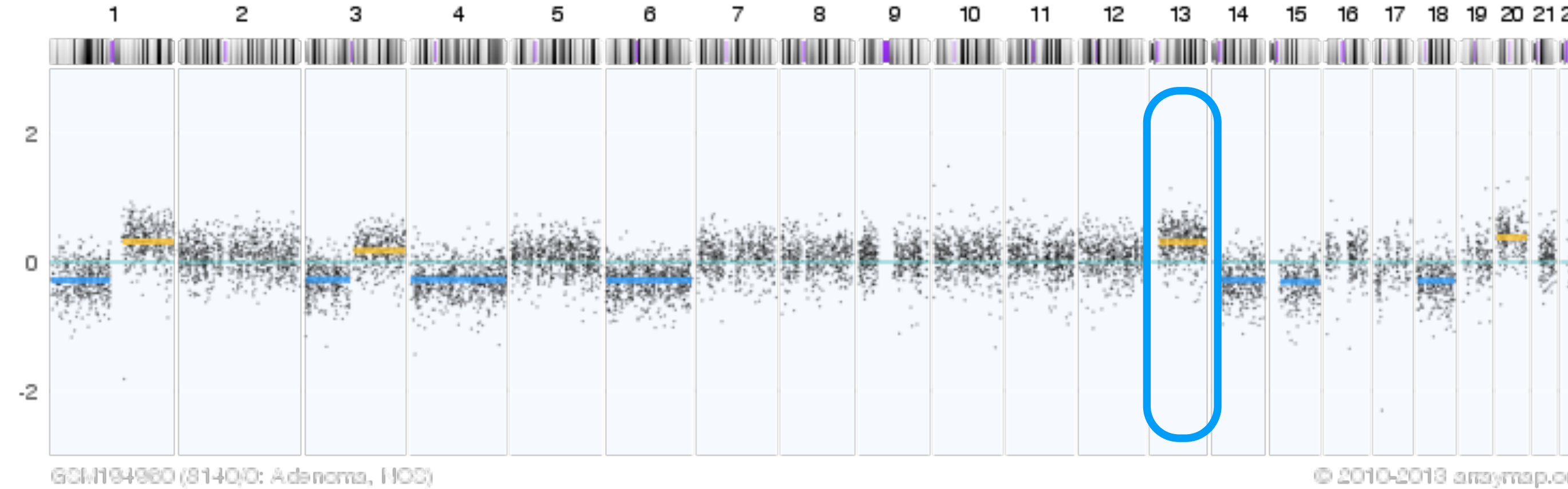
Types of genomic alterations in Cancer

Imbalanced Chromosomal Changes: CNV

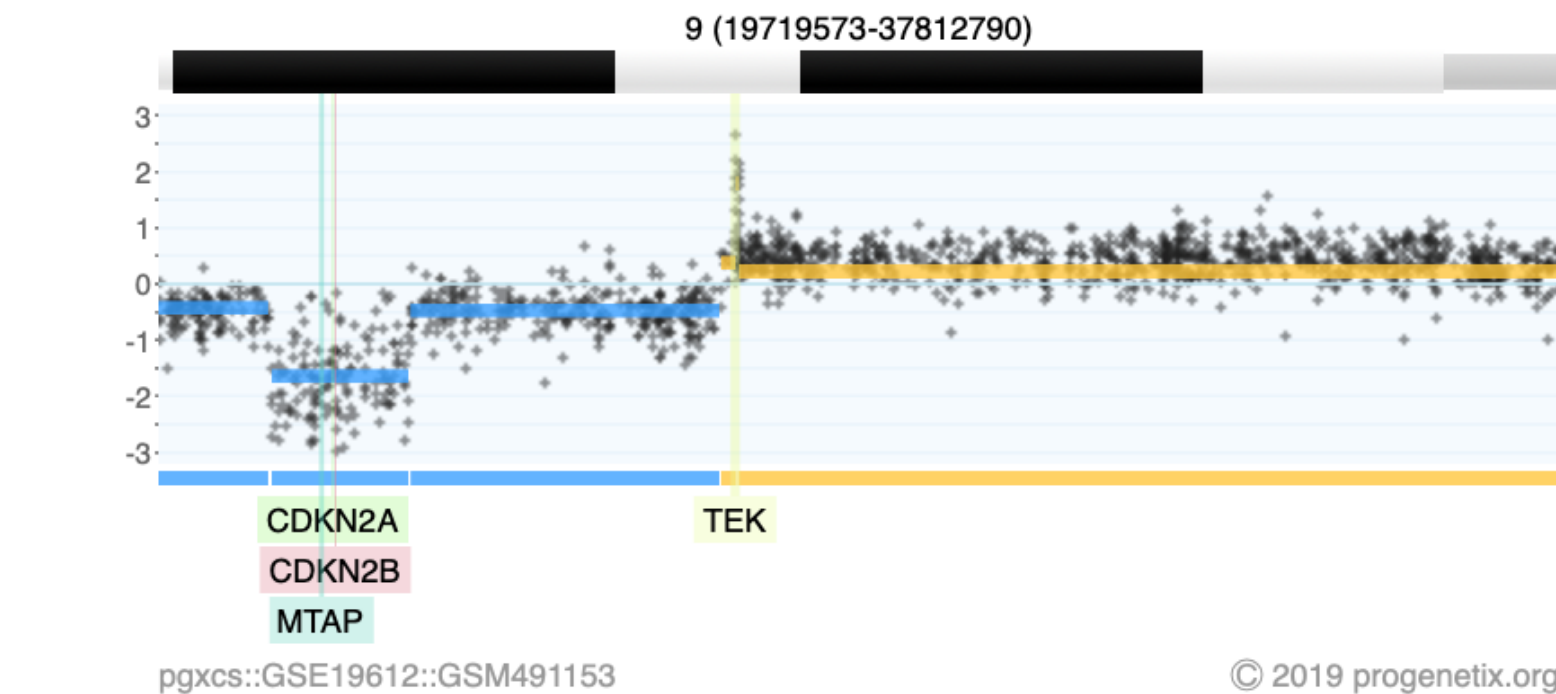
- Point mutations (insertions, deletions, substitutions)
- Chromosomal rearrangements
- **Regional Copy Number Alterations** (losses, gains)
- Epigenetic changes (e.g. DNA methylation abnormalities)



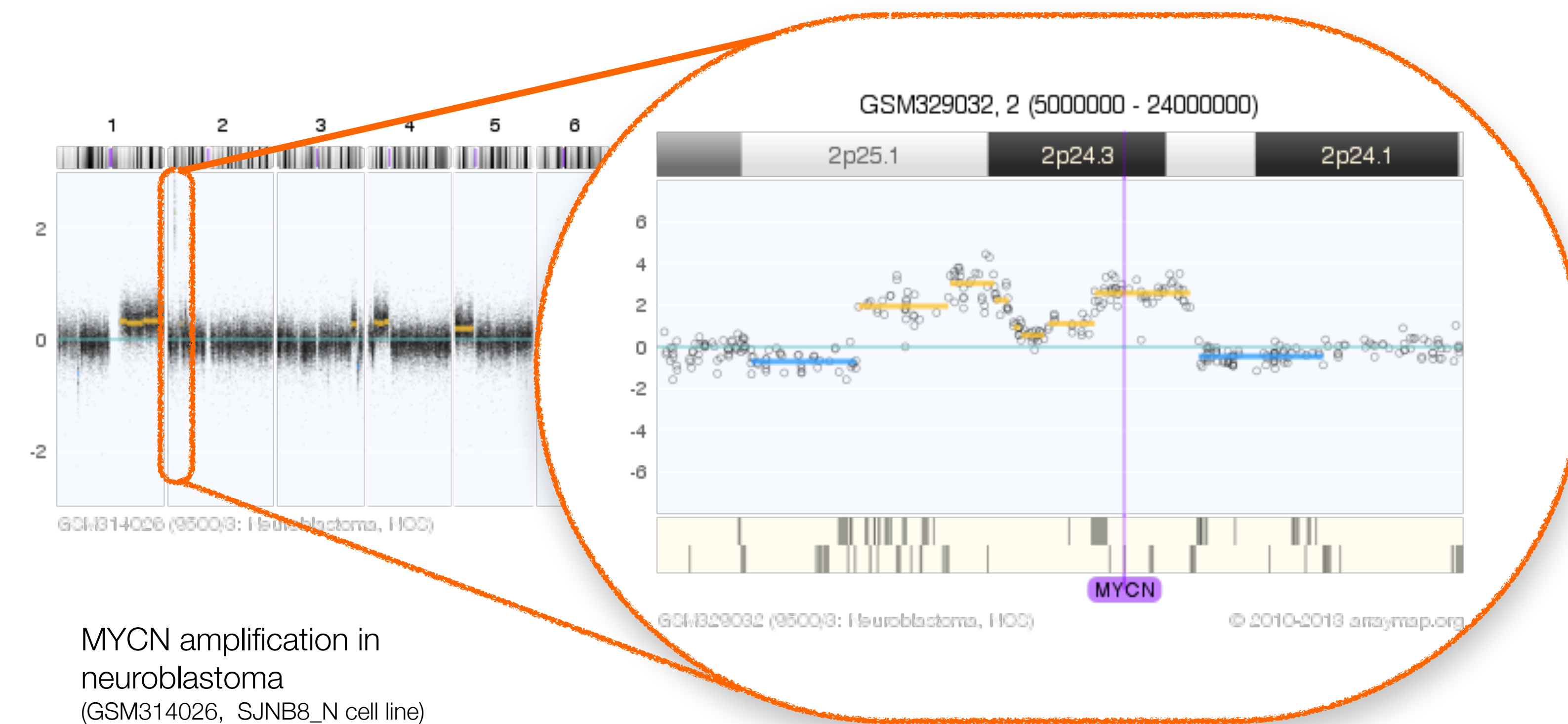
Array-based Detection of Copy Number Variations



Gain of chromosome arm 13q in colorectal carcinoma



2-event, homozygous deletion in a Glioblastoma



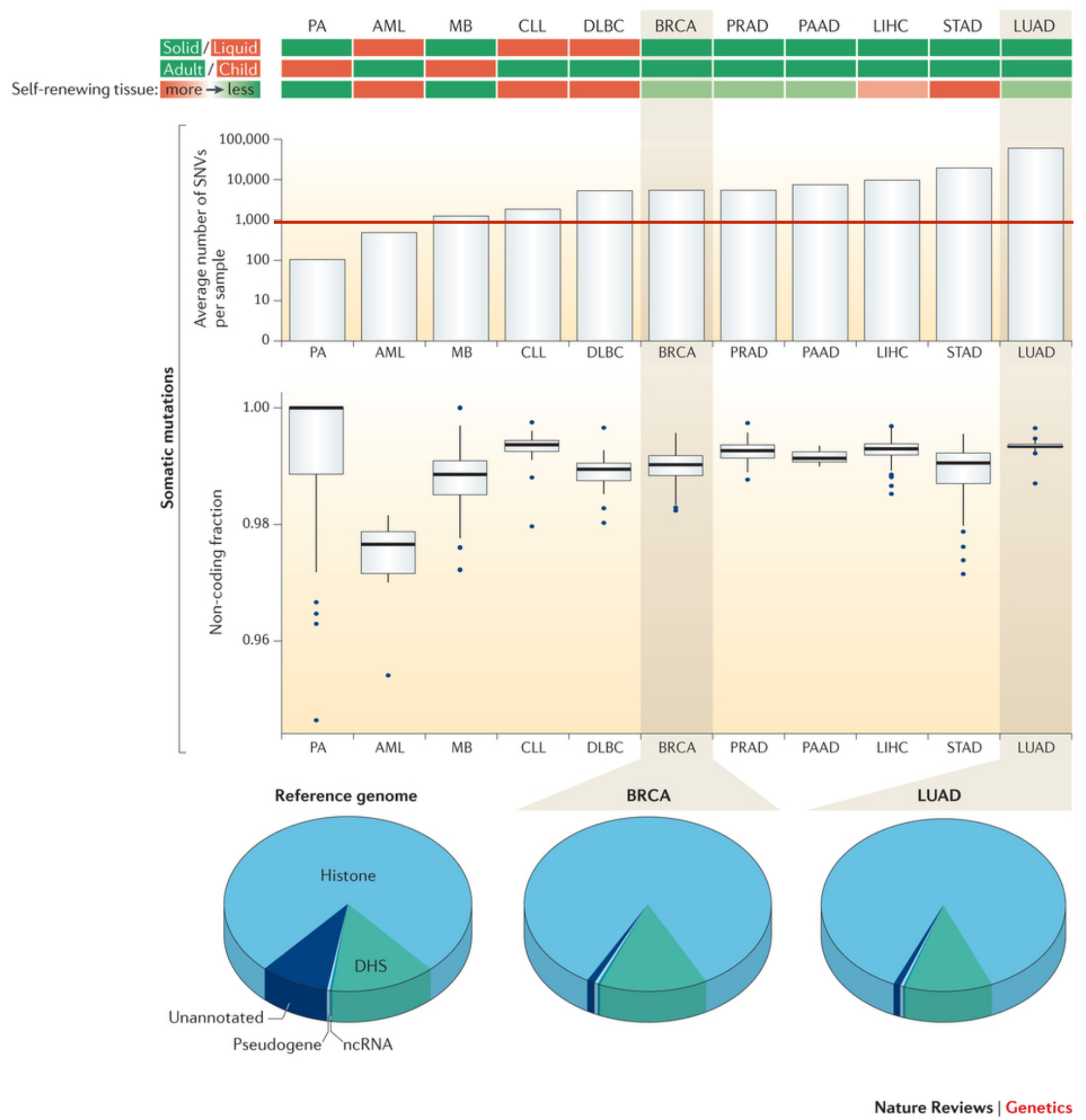
MYCN amplification in neuroblastoma
(GSM314026, SJNB8_N cell line)

low level/high level copy number alterations (CNAs)

arrayMap

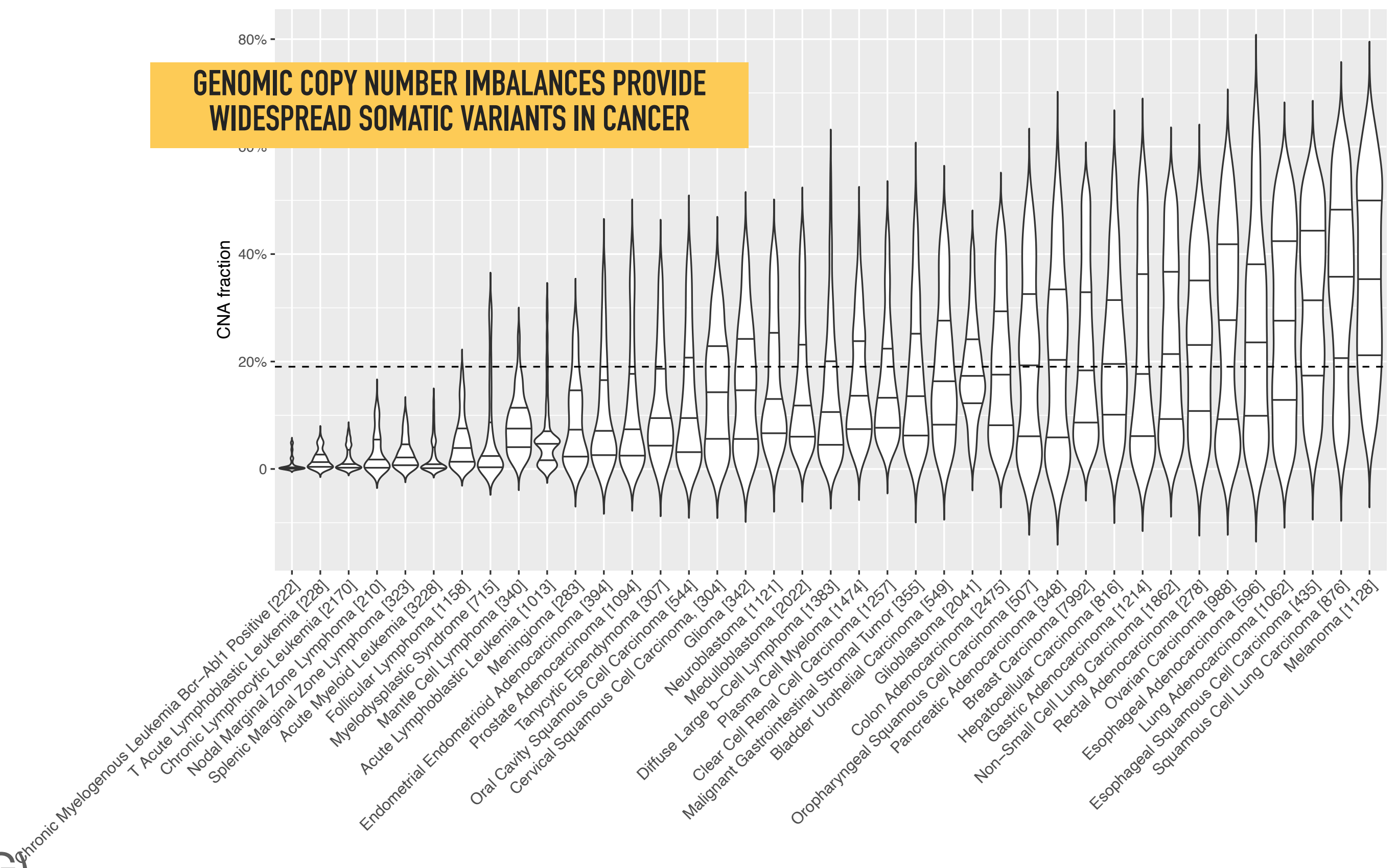


Quantifying Somatic Mutations In Cancer



CANCERS SHOW THOUSANDS OF SINGLE NUCLEOTIDE VARIANTS PER SAMPLE, MOSTLY IN NON-CODING REGIONS

Pan-Cancer Analysis of Whole Genomes (PCAWG) data show widespread mutations in non-coding regions of cancer genomes (Khurana et al., Nat. Rev. Genet. (2016))

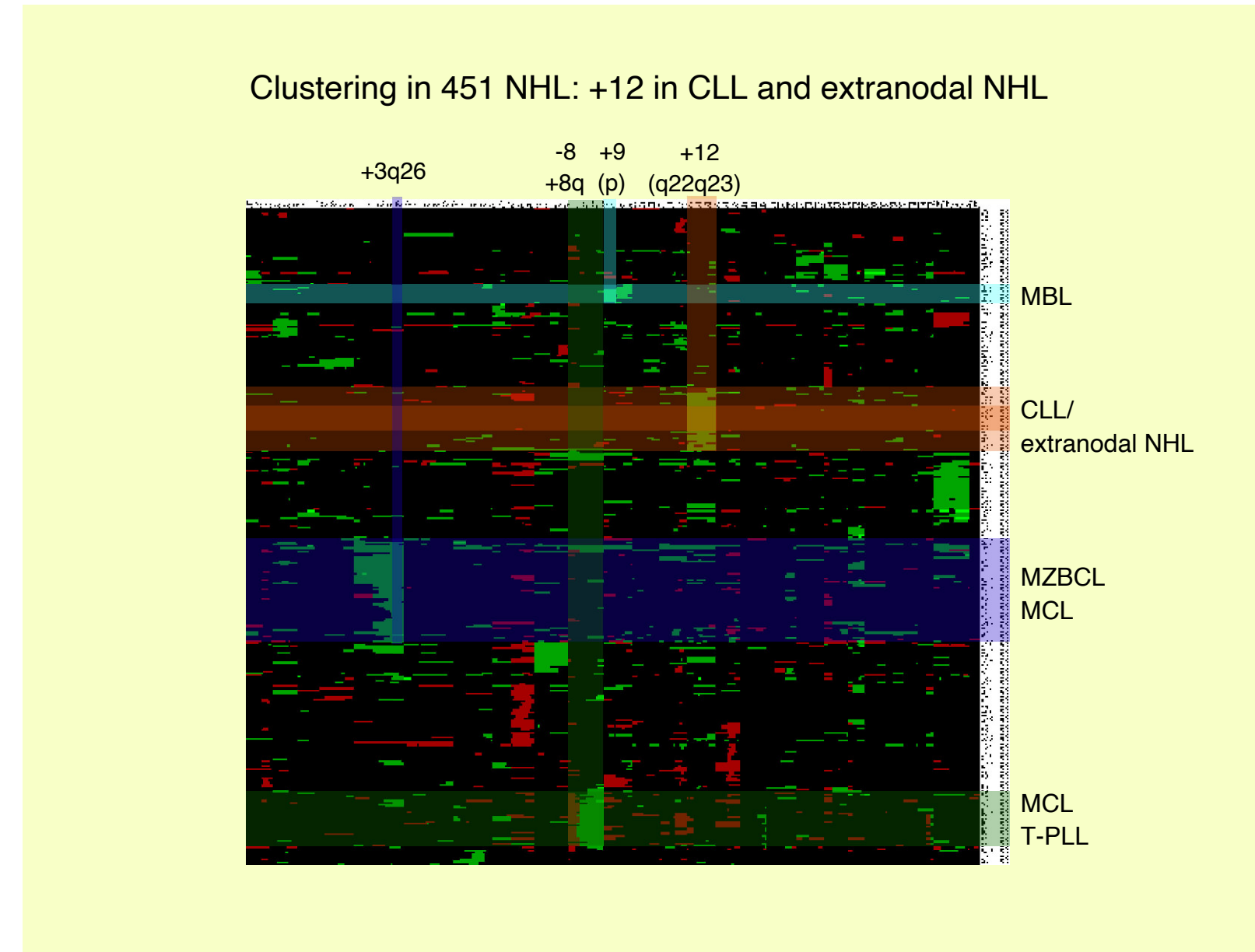
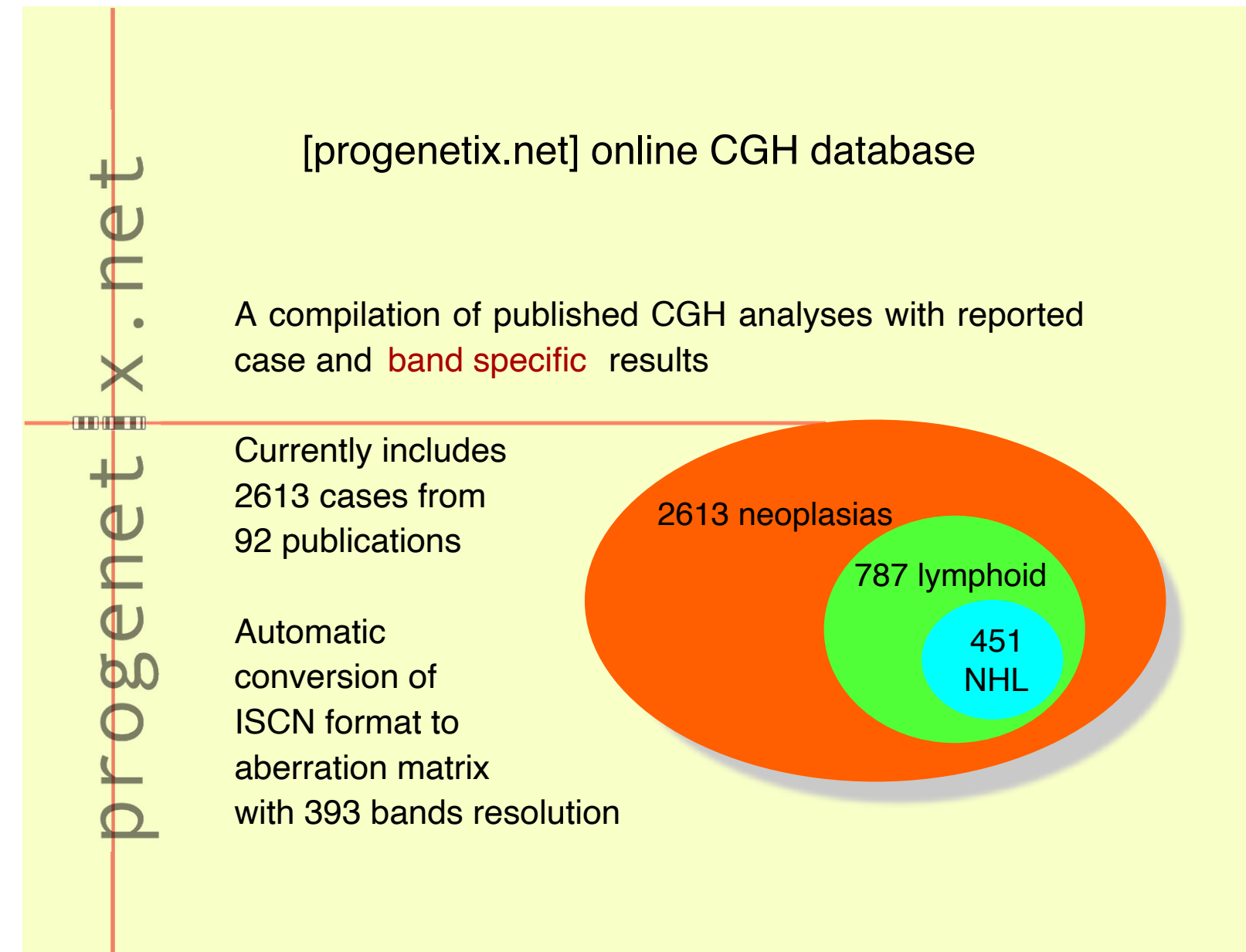


On average ~19% of a cancer genome are in an imbalanced state (more/less than 2 alleles); Original data based on 43654 cancer genomes from progenetix.org

History & Current State...

Origins & trajectory of the Progenetix Resource



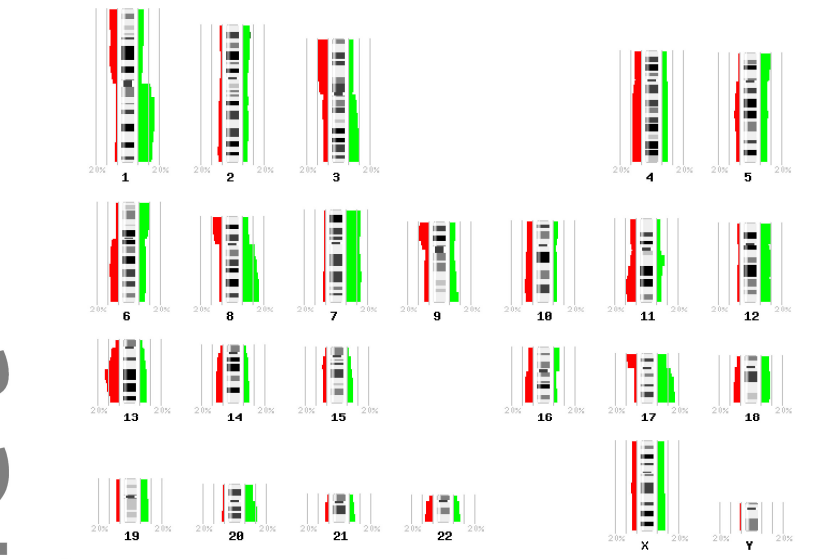


Collection and Transformation of Chromosomal Imbalances in Human Neoplasias for Data Mining Procedures

michael baudis, dept. of pathology, stanford university

Although the deciphering of the human genome has been pushed forward over the last years, little effort has been made to collect and integrate the treasure trove of clinical tumor cases analyzed by molecular-cytogenetic methods into current data schemes. Publicly announced at BCATS 2001, since then [progenetix.net] has been established as the largest public source of chromosomal imbalance data with band-specific resolution. Targets for the use of the data collection may be the description of prediction of oncogene and suppressor gene loci, identification of related loci for pathway creation, and especially the combination of the data with expression array experiments for filtering of relevant genes among the deregulated candidates.

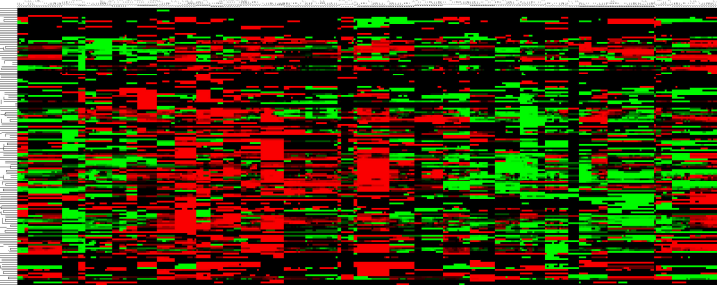
Chromosomal imbalances in 5478 clinical cases from 196 publications
Although not as prominent as in specific subgroups, this large collection shows the non-random distribution of chromosomal gains (green) and losses (red).



Material and Methods Chromosomal aberration data of more than 5478 cases from 196 publications describing results of Comparative Genomic Hybridization (CGH) experiments were collected. Minimal requirements were diagnosis of a malignant or benign neoplasia, analysis of clinical tumor samples and report of the analysis results on a case by case basis, resolved to the level of single chromosomal bands. Data was transformed from the diverse annotation formats to standardized ISCN "rev ish" nomenclature. For the transformation of the non-linear ISCN data to a two-dimensional matrix with code for the aberration status of each chromosomal band per case, a reverse pattern matching algorithm was developed in Perl. Graphical representations and cluster images are generated for all different subsets (Publications, ICD-O-3 entities, meta-groups) and presented on the progenetix.net website.

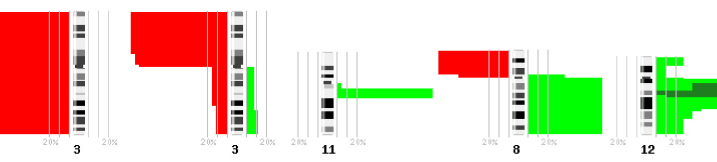


Clustering of the band averages for the different ICD-O entities
Two dimensional clustering groups related disease entities and chromosomal bands with related aberrations.



Results Out of 4896 tumor samples, 3862 (79%) showed chromosomal imbalances by CGH. The average per band probability was 4.5% for a loss (max. 12.9% at 13q1) and 6.5% for a gain (max. 15.6% at 8q23). Differences between neoplastic entities showed in the average frequency and distribution pattern of imbalanced chromosomal regions. Tumor subsets (10 or more cases) with the strongest hot spots for losses were small cell lung carcinomas (ave. 23.3% with max. 96.2% at 3p14p26) and pheochromocytomas (ave. 10.9% with max. 92.7% at 3p); prominent gain maxima were found in pure high grade infiltrating duct carcinomas of the breast (ave. 5.9% with max. 95.7% at 11q13), T-PLL (ave. 4.7% with max. 81.8% for whole 8q) and dedifferentiated liposarcomas (ave. 10.4% with max. 81.8% at 12q13), among others. By cluster analysis, different combinations of chromosomal hot spot regions could be shown to occur in tumors subsummed in the same diagnostic entity; the example of neuroblastomas is shown.

Examples of hotspots of genomic imbalance
SCLC, pheochromocytoma, high grade DCIS, T-PLL, dedifferentiated liposarcoma



Conclusion So far, progenetix.net project was able to:
1. collect a large dataset of genomic aberration data generated through a molecular-cytogenetic screening technique (CGH)
2. develop the software tools to transform those data to a meta format compatible to commonly used genomic interval descriptions
3. produce graphical and numerical output from those data for hot spot detection and statistical analysis.

For future approaches, the data collection will be valuable for filtering data from expression array experiments for relevant genes, and possibly for the description of common and divergent genetic pathways in the oncogenetic process of different tumor entities. The transformed raw data of the progenetix.net collection is available for research purposes over the website.

Distinction of histologically related through their chromosomal aberration pattern
Amplification of the REL locus on 2p16 and gain of 9p(ter) distinguishes primary mediastinal B-cell lymphomas (PMBL, right) from diffuse large cell lymphomas (DLCL, left). The distinction may have clinical implications



Identification of different aberration patterns in Neuroblastoma (289 cases)
N-Myc (2p25) amplification is the hallmark of a subgroup, showing only consistent loss of the terminal portion 1p. Other groups are defined by the loss of 11q, or a "chromosomal instability" phenotype. Gains on 17q are a common feature of all groups. Those patterns may be combined with gene-level information to reconstruct the different pathways leading to malignant transformation.

Progenetix Database in 2003

Text conversion for CNVs

- articles and supplements with **cytoband-based rev ish CGH** results
- sometimes rich, but **unstructured** associated information
- PDFs** readable, but **not well suited for data extraction** (character entities, text flow)

progenetix

TABLE 3. Comparison of Primary Tumors and Metastases by CGH

Case	Gain in common	Loss in common	Primary tumor only	Metastasis only
108		18		
113	7, 8q24-qter, 13q11-qter, 20q11-qter, Xq11-Xter	1p33-pter, 2p21-pter, 4q24-qter, 15q11-q15, 17p11-pter, 18		
LM	12q22-qter, 15q23-qter, 17q11-ter, 20p11-p12, 20q11-ter, 22q11-ter	1p11-p32, 1q24-31, 4, 13q11-pter, 17p11-pter, 18, 20p11-ter	11p11-pter-	12+
145	4q26-q28, 6p11-p13, 8p11-p12, 920q11-qter	1p11-pter, 4q31-qter, 6q11-qter, 8p12-pter, 11, 15q11-qter, 16q11-qter, 17p11-pter, 18, 21q11-qter	13q21-qter+, 20p11-pter-	8q11-qter+, 10-, 6p21-pter-
53	7, 8q11-qter, 9q33-qter, 13q11-qter, 20p11-p12, 20q11-qter	4p13-pter, 4q21-qter, 8p12-pter, 15q14-qter, 18q11-qter, 20p12-pter	5p11-pter-, 5q13-qter-, 14q11-qter-	11+, 16p11-pter+, 17q11-qter+, 19+, 21q11-qter+, 22q11-qter+
147	7, 13q11-qter, 20q11-qter	8p21-pter, 18	4p14-pter-, 4q28-qter+, 8p11-21-, 17q11-q2+, 21q11-qter-	11q22-qter+, 16+, 1p11-33-

TABLE 1. Clinical Data

Case number	Age	Sex	Site	Stage ^a	Grade ^b	Diagnosis of metastatic disease ^c
2	40	M	Transverse colon	IV	3	Synchronous
6	79	M	Ascending colon	IV	2	Synchronous
9	73	M	Transverse colon	II	2	N/A
11	56	M	Rectosigmoid	IV	2	Metachronous
12	70	F	Sigmoid colon	IV	2	Synchronous
13	65	M	Descending colon	II	9	Synchronous
14	60	M	Rectum	III	3	Metachronous
15	51	F	Rectum	III	2	Metachronous
19	63	M	Rectosigmoid Junction	III	2	Synchronous
20	63	M	Rectum	IV	9	Metachronous
21	64	F	Sigmoid colon	IV	2	Synchronous
35	71	M	Rectum	III	9	Metachronous
49	72	M	Cecum	IV	3	Synchronous
53	72	F	Sigmoid colon	IV	2	Synchronous
104	61	M	Sigmoid colon	IV	2	Metachronous
105	58	M	Ascending colon	II	2	Metachronous
107	77	F	Cecum	IV	2	Metachronous
108	53	F	Splenic flexure	IV	2	Synchronous
112	68	M	Rectum	III	3	Synchronous
113	41	M	Splenic flexure	IV	2	Synchronous
114	49	M	Splenic flexure	IV	3	Synchronous
116	73	M	Rectosigmoid	III	9	Metachronous
120	24	F	Descending colon	IV	2	Synchronous
123	62	F	Rectum	III	2	Metachronous
124	42	M	Rectum	IV	9	Synchronous
145	70	M	Rectosigmoid	IV	2	Synchronous
147	86	F	Cecum	IV	2	Synchronous

^aAJCC/UICC staging system (Hutter and Sabin, 1986).^bGrade of primary tumor: 1–3, low, moderate, high grade; 9, grading unknown.^cSynchronous, diagnosis of metastatic disease within 12 months following diagnosis of primary tumor; metachronous, diagnosis of metastatic disease after 12 months or later.

GENES, CHROMOSOMES & CANCER 25:82–90 (1999)

Chromosome Arm 20q Gains and Other Genomic Alterations in Colorectal Cancer Metastatic to Liver, as Analyzed by Comparative Genomic Hybridization and Fluorescence In Situ Hybridization

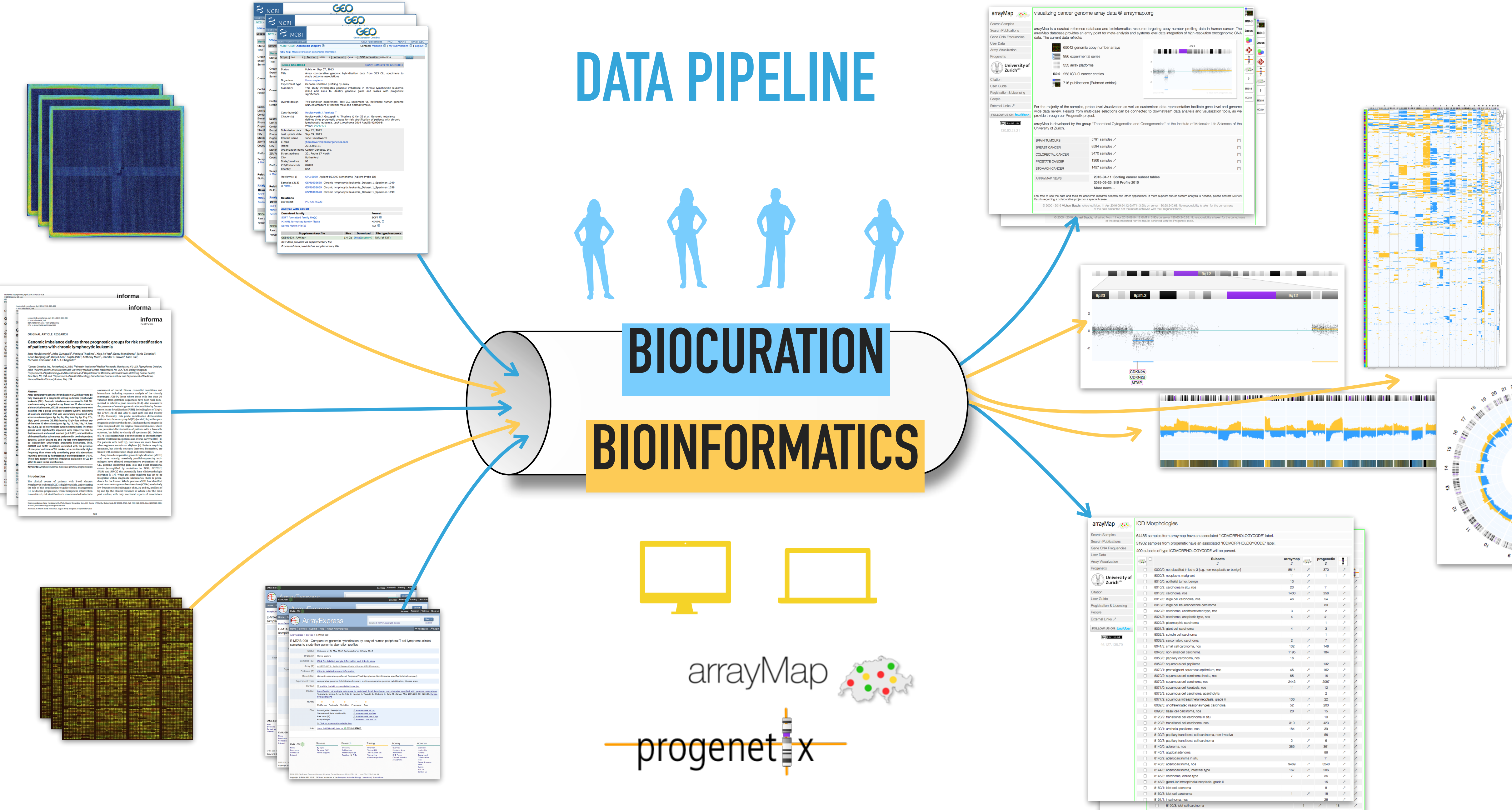
W. Michael Korn,¹ Toru Yasutake,² Wen-Lin Kuo,¹ Robert S. Warren,³ Colin Collins,¹ Masao Tomita,² Joe Gray,¹ and Frederic M. Waidman¹

Progenetix: Data Scopes

Biomedical and procedural "Meta"data types

- Diagnostic classification
 - mapping text-based cancer diagnoses to standard classification systems
- Provenance data
 - store identifier-based pointers
 - geographic attribution (individual, biosample, experiment)
- Clinical information
 - **core set** of typical cancer study values:
 - ➡ stage, grade, followup time, survival status, genomic sex, age at diagnosis
 - balance between annotation effort and expected usability

DATA PIPELINE



Data Curation - Happy RegExing!

Extracting clinical and technical metadata from GEO SOFT file

```
^SAMPLE = GSM174832
!Sample_title = 9194
!Sample_geo_accession = GSM174832
!Sample_status = Public on May 01 2007
!Sample_submission_date = Mar 13 2007
!Sample_last_update_date = Mar 13 2007
!Sample_type = genomic
!Sample_channel_count = 1
!Sample_source_name_ch1 = Bone marrow with 96% blasts
!Sample_organism_ch1 = Homo sapiens
!Sample_taxid_ch1 = 9606
!Sample_characteristics_ch1 = Immunotype: common ALL; Age: 9.2 yrs; Gender: F
!Sample_molecule_ch1 = genomic DNA
!Sample_extract_protocol_ch1 = QiaAmp purification kit (Qiagen)
!Sample_label_ch1 = biotin
!Sample_label_protocol_ch1 = Biotinylated DNA was prepared according to the standard
manual 701930 Rev.3 or 100k assay manual 701684 Rev.3, Affymetrix).
!Sample_hyb_protocol = Hybridizations were performed according to the standard Affy
701930 Rev.3 or 100k assay manual 701684 Rev.3, Affymetrix) using an Affymetrix hybridization
!Sample_scan_protocol = Scanning performed according to the standard Affymetrix protocol
or 100k assay manual 701684 Rev.3, Affymetrix) using an Affymetrix scanner 3000.
!Sample_description = primary ALL diagnosis sample
!Sample_data_processing = copy number detection using CNAG2.0 software (http://www.cnag2.org)
!Sample_platform_id = GPL3718
!Sample_contact_name = Roland,P.,Kuiper
!Sample_contact_email = r.kuiper@antrg.umcn.nl, e.verwiel@antrg.umcn.nl
!Sample_contact_phone = +31243610868
!Sample_contact_fax = +31243668752
!Sample_contact_department = Human Genetics
!Sample_contact_institute = Radboud University Nijmegen Medical Centre
!Sample_contact_address = Geert Grootplein 10
!Sample_contact_city = Nijmegen
!Sample_contact_zip/postal_code = 6525GA
!Sample_contact_country = Netherlands
!Sample_supplementary_file = ftp://ftp.ncbi.nlm.nih.gov/geo/samples/GSM174nnn/GSM174832/suppl/GSM174832.CEL.gz
!Sample_supplementary_file = ftp://ftp.ncbi.nlm.nih.gov/geo/samples/GSM174nnn/GSM174832/suppl/GSM174832.CHP.gz
!Sample_series_id = GSE7255
```

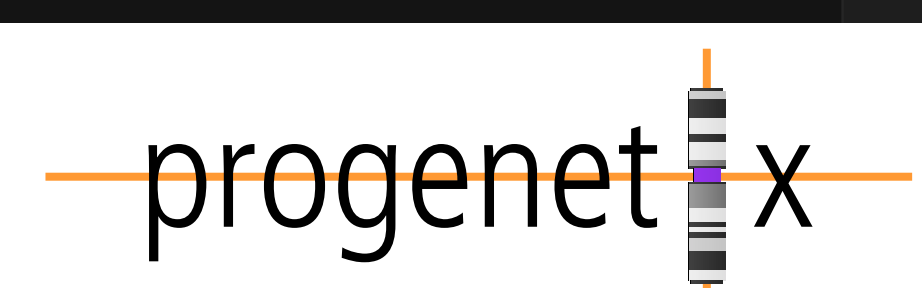
```
foreach (grep { ! /characteristics_ch\d/ } @in) {
    my ($key, $value) = split(' = ', $_);
    $key =~ s/[^w]/_/g;
    if ($key =~ /submission_date/i) {
        $sample->{ YEAR } = $value;
        $sample->{ YEAR } =~ s/^.*?(\d\d\d\d)$/\1/;
    }
}
```

```
$mkey->{ samplekey } = 'AGE';
$mkey->{ matches } = [ qw( age )];

( $mkey->{ retv }, $mkey->{ retk } ) = _grepmeta( $mkey, $meta );

if ($mkey->{ retv } =~ /^(.+?)$/) {
    if ($mkey->{ retv } =~ /month/i) {
        $mkey->{ retk } .= '_months';
        $mkey->{ retv } =~ s/[\^d\.]/ /g;
    }
}

$sample->{ $mkey->{ samplekey } } = _normNumber($mkey->{ retv });
if ($mkey->{ retk } =~ /month/i) { $sample->{ $mkey->{ samplekey } } /= 12 }
if ( $sample->{ $mkey->{ samplekey } } == 0 ) { $sample->{ $mkey->{ samplekey } } = 'NA' }
$sample->{ $mkey->{ samplekey } } = sprintf "%,.2f", $sample->{ $mkey->{ samplekey } };
```



Data Curation - Happy RegExing!

Extracting clinical and technical metadata from GEO SOFT file

```
^SAMPLE = GSM286922
!Sample_title = 481 - mAbID:75320
!Sample_geo_accession = GSM286922
!Sample_status = Public on Sep 04 2008
!Sample_submission_date = May 06 2008
!Sample_last_update_date = Nov 26 2008
!Sample_type = genomic
!Sample_channel_count = 2
!Sample_source_name_ch1 = Normal Lymphocytes
!Sample_organism_ch1 = Homo sapiens
!Sample_taxid_ch1 = 9606
!Sample_characteristics_ch1 = Tissue: lymphocytes
!Sample_molecule_ch1 = genomic DNA
!Sample_extract_protocol_ch1 = Sample DNA Extraction Protocol
!Sample_extract_protocol_ch1 = Other: The DNA was isolated by Qiagen DNe
!Sample_label_ch1 = cy5
!Sample_label_protocol_ch1 = NimbleGen Cy5 Sample Labeling Protocol
!Sample_label_protocol_ch1 = Other: Proprietary protocol information available at http://www.nimblegen.com/technology/index.html
!Sample_source_name_ch2 = 481
!Sample_organism_ch2 = Homo sapiens
!Sample_taxid_ch2 = 9606
!Sample_characteristics_ch2 = Gender: male
!Sample_characteristics_ch2 = Age: 49
!Sample_characteristics_ch2 = Tissue: lymph node
!Sample_characteristics_ch2 = Disease state: Lymphoma
!Sample_characteristics_ch2 = Individual: 481
!Sample_characteristics_ch2 = Clinical info: Submitting diagnosis: DLBCL
!Sample_characteristics_ch2 = Clinical info: Final microarray diagnosis: ABC DLBCL
!Sample_characteristics_ch2 = Clinical info: Follow up status: ALIVE
!Sample_characteristics_ch2 = Clinical info: Follow up years: 10.75
!Sample_characteristics_ch2 = Clinical info: Chemotherapy: CHOP-Like Regimen
!Sample_characteristics_ch2 = Clinical info: ECOG performance status: 2
!Sample_characteristics_ch2 = Clinical info: Stage: 4
!Sample_characteristics_ch2 = Clinical info: LDH ratio: 0.82
!Sample_characteristics_ch2 = Clinical info: Number of extranodal sites: 1
```

Channel 1 is normal -> Cave value swap!

Gender or "chromosomal sex"?

context indicates years, but if it would be a medulloblastoma...

This survival status annotation not known to parser...

```
$mkey->{ samplekey } = 'DEATH';
$mkey->{ matches } = [
    'death',
    'dead ',
    'vital_status',
    'dead_alive',
    'alive_dead',
];
(
    $mkey->{ retv }, $mkey->{ retk } ) = _grepmeta( $mkey, $meta );
if ( $mkey->{ retv } =~ /^(.+?)$/ ) {
    $sample->{ $mkey->{ samplekey } } = _normDeath($mkey->{ retv }) }
```

The Progenetix oncogenomic resource in 2021

Qingyao Huang^{1,2}, Paula Carrio-Cordo^{1,2}, Bo Gao^{1,2}, Rahel Paloots^{1,2} and Michael Baudis^{1,2,*}

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Citation details: Huang, Q., Carrio-Cordo, P., Gao, B. et al. The Progenetix oncogenomic resource in 2021. *Database* (2021) Vol. 2021: article ID baab043; DOI: <https://doi.org/10.1093/database/baab043>

Abstract

In cancer, copy number aberrations (CNAs) represent a type of nearly ubiquitous and frequently extensive structural genome variations. To disentangle the molecular mechanisms underlying tumorigenesis as well as identify and characterize molecular subtypes, the comparative and meta-analysis of large genomic variant collections can be of immense importance. Over the last decades, cancer genomic profiling projects have resulted in a large amount of somatic genome variation profiles, however segregated in a multitude of individual studies and datasets. The Progenetix project, initiated in 2001, curates individual cancer CNA profiles and associated metadata from published oncogenomic studies and data repositories with the aim to empower integrative analyses spanning all different cancer biologies. During the last few years, the fields of genomics and cancer research have seen significant advancement in terms of molecular genetics technology, disease concepts, data standard harmonization as well as data availability, in an increasingly structured and systematic manner. For the Progenetix resource, continuous data integration, curation and maintenance have resulted in the most comprehensive representation of cancer genome CNA profiling data with 138 663 (including 115 357 tumor) copy number variation (CNV) profiles. In this article, we report a 4.5-fold increase in sample number since 2013, improvements in data quality, ontology representation with a CNV landscape summary over 51 distinctive National Cancer Institute Thesaurus cancer terms as well as updates in database schemas, and data access including new web front-end and programmatic data access.

Database URL: progenetix.org

Table 1. Statistics of samples from various data resources

Data source	GEO	ArrayExpress	cBioPortal	TCGA	Total
No. of studies	898	51	38	33	1939
No. of samples	63 568	4351	19 712	22 142	138 663
Tumor	52 090	3887	19 712	11 090	115 357
Normal	11 478	464	0	11 052	23 306
Classifications					
ICD-O (Topography)	100	54	88	157	209
ICD-O (Morphology)	246	908	265	140	491
NCIt	346	148	422	182	788
Collections					
Individuals	63 568	4351	19 712	10 995	127 549
Biosamples	63 568	4351	19 712	22 142	138 663
Callsets ^a	63 568	4351	19 712	22 376	138 930
Variants	5 514 126	118 4170	1 778 096	2 654 065	10 716 093

^aset of variants from one genotyping experiment; ICD-O, International Classification of Diseases for Oncology; NCIt, National Cancer Institute Thesaurus.

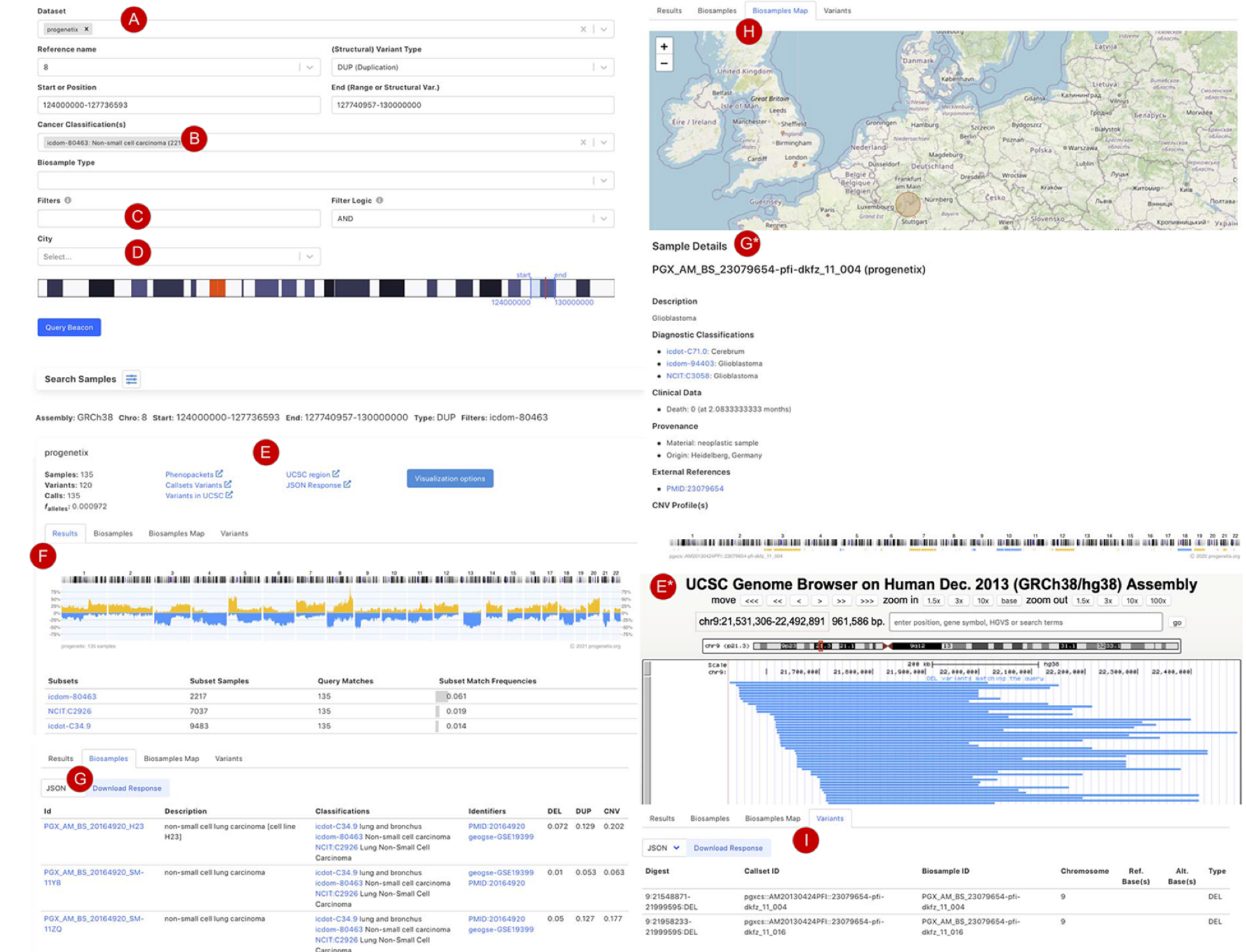


Figure 3. Beacon-style query using fuzzy ranges to identify biosamples with variants matching the CNA range. This example queries for a continuous, focal duplication covering the complete MYC gene's coding region with ≤ 6 Mb in size. A: Filter for dataset; B: filter for cancer classification (NCIt and ICD-O-3 ontology terms available); C: additional filter, e.g. Cellosaurus; D: additional filter for geographic location; E: external link to UCSC browser to view the alignment of matched variants; F: cancer type classification sorted by frequency of the matched biosamples present in the subset; G: list of matched biosamples with description, statistics and reference. More detailed biosample information can be viewed through 'id' link to the sample detail page; H: matched variants with reference to biosamples can be downloaded in json or csv format.

Progenetix in 2021

Cancer Genomics Reference Resource

- >116'000 cancer CNV profiles, mapped to >800 NCIt codes
- majority of data from genomic arrays with ~50% overall from SNP platforms with original data re-processing
- structured diagnostic encodings for NCIt, ICD-O 3, UBERON
- identifier mapping for PMID, GEO, Cellosaurus where appropriate
- core biosample and technical metadata annotations where accessible
- publication database and code mapping services



[Cancer CNV Profiles](#)

[Search Samples](#)

[Studies & Cohorts](#)

[arrayMap](#)

[TCGA Samples](#)

[DIPG Samples](#)

[Gao & Baudis, 2021](#)

[Cancer Cell Lines](#)

[Publication DB](#)

[Services](#)

[NCIt Mappings](#)

[UBERON Mappings](#)

[Upload & Plot](#)

[Download Data](#)

[Beacon⁺](#)

[Progenetix Info](#)

[About Progenetix](#)

[Use Cases](#)

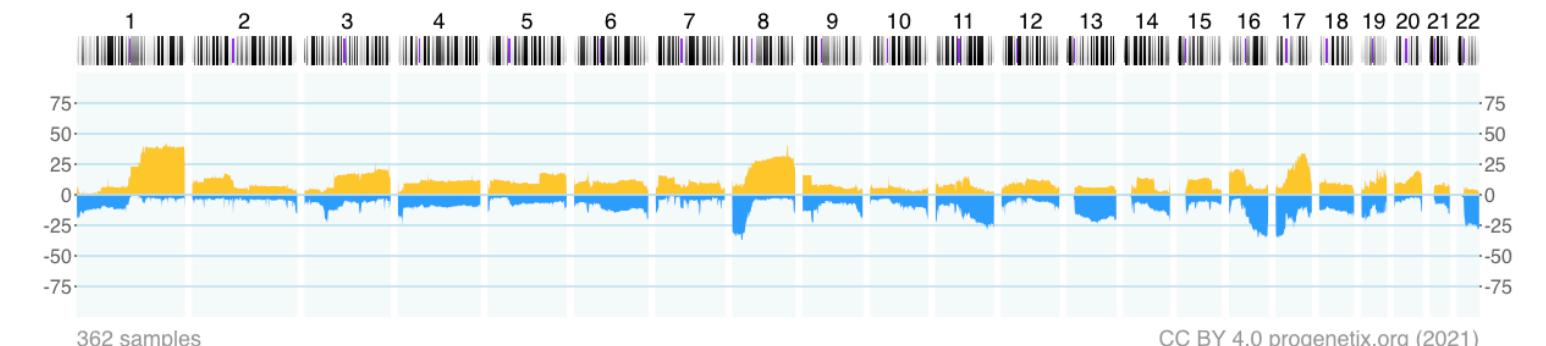
[Documentation](#)

[Baudisgroup @ UZH](#)

[Cancer genome data @ progenetix.org](#)

The Progenetix database provides an overview of mutation data in cancer, with a focus on copy number abnormalities (CNV / CNA), for all types of human malignancies. The data is based on *individual sample data* from currently **139448** samples.

Breast Cancer by AJCC v6 Stage (NCIT:C90513)

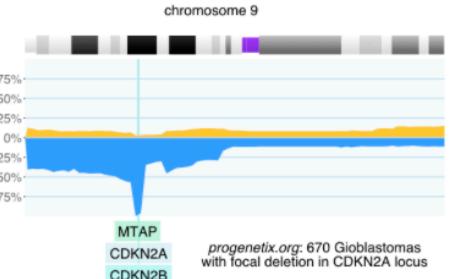


[Download SVG](#) | [Go to NCIT:C90513](#) | [Download CNV Frequencies](#)

Example for aggregated CNV data in 362 samples in Breast Cancer by AJCC v6 Stage. Here the frequency of regional **copy number gains** and **losses** are displayed for all 22 autosomes.

[Progenetix Use Cases](#)

[Local CNV Frequencies](#)



A typical use case on Progenetix is the search for local copy number aberrations - e.g. involving a gene - and the exploration of cancer types with these CNVs. The [[Search Page](#)] provides example use cases for designing queries. Results contain basic statistics as well as visualization and download options.

[Cancer CNV Profiles](#)

The progenetix resource contains data of **810** different cancer types (NCIt neoplasm classification), mapped to a variety of biological and technical categories. Frequency profiles of regional genomic gains and losses for all categories (diagnostic entity, publication, cohort ...) can be accessed through the [[Cancer Types](#)] page with direct visualization and options for sample retrieval and plotting options.

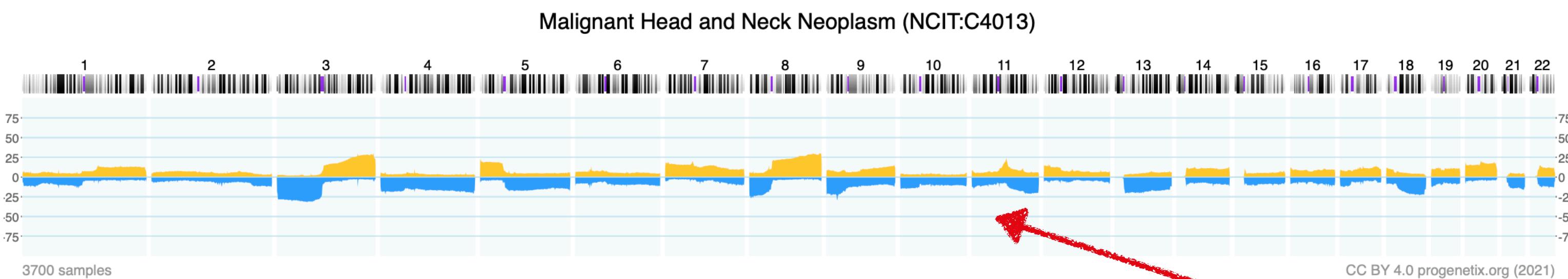
[Cancer Genomics Publications](#)

Through the [[Publications](#)] page Progenetix provides **4025** annotated references to research articles from cancer genome screening experiments (WGS, WES, aCGH, cCGH). The numbers of analyzed samples and possible availability in the Progenetix sample collection are indicated.

Progenetix

Cancer Type CNA Data

- hierarchical aggregation of cancer samples
- pre-computed CNA frequencies for fast overview
- sample retrieval for custom grouping, visualization



Cancer CNV Profiles

Search Samples

Studies & Cohorts

arrayMap

TCGA Samples

DIPG Samples

Gao & Baudis, 2021

Cancer Cell Lines

Publication DB

Services

NCIt Mappings

UBERON Mappings

Upload & Plot

Download Data

Beacon⁺

Progenetix Info

Cancer Types

The cancer samples in Progenetix are mapped to several classification systems. For each of the classes, aggregated data is available by clicking the code. Additionally, a selection of the corresponding samples can be initiated by clicking the sample number or selecting one or more classes through the checkboxes.

Sample selection follows a hierarchical system in which samples matching the child terms of a selected class are included in the response.

Cancer Classification: NCIT Cancer Core

Filter subsets ... Hierarchy Depth: 2 levels

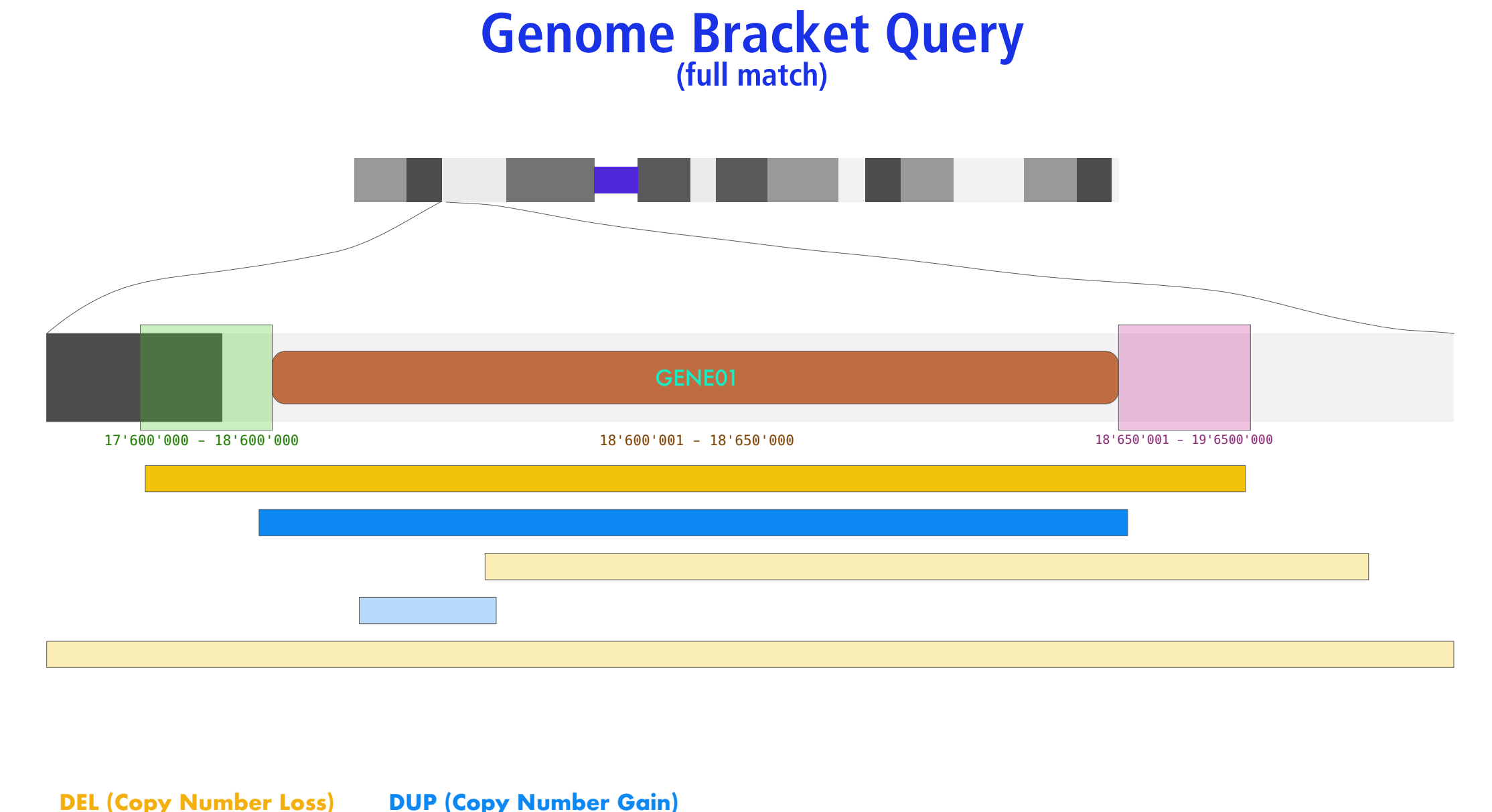
No Selection

- ▼ NCIT:C3262: Neoplasm (116232 samples)
 - ▼ NCIT:C3263: Neoplasm by Site (110429 samples)
 - NCIT:C156482: Genitourinary System Neoplasm (16534 samples)
 - NCIT:C2910: Breast Neoplasm (15556 samples)
 - NCIT:C27939: Lobular Neoplasia (92 samples)
 - NCIT:C36083: Intraductal Breast Neoplasm (275 samples)
 - NCIT:C27942: Intraductal Proliferative Lesion of the Breast (270 samples)
 - NCIT:C2924: Ductal Breast Carcinoma In Situ (270 samples)
 - NCIT:C36090: Intraductal Papillary Breast Neoplasm (5 samples)
 - NCIT:C40405: Breast Fibroepithelial Neoplasm (41 samples)
 - NCIT:C40406: Breast Soft Tissue Neoplasm (3 samples)
 - NCIT:C5206: Papillary Breast Neoplasm (15 samples)
 - NCIT:C9335: Malignant Breast Neoplasm (15528 samples)
 - NCIT:C3010: Endocrine Neoplasm (3521 samples)
 - NCIT:C3030: Eye Neoplasm (280 samples)
 - NCIT:C3052: Digestive System Neoplasm (15285 samples)
 - ▼ NCIT:C3077: Head and Neck Neoplasm (3961 samples)
 - NCIT:C3260: Neck Neoplasm (2565 samples)
 - NCIT:C3361: Salivary Gland Neoplasm (274 samples)
 - NCIT:C3375: Skull Neoplasm (161 samples)
 - NCIT:C4013: Malignant Head and Neck Neoplasm (3700 samples)
 - NCIT:C133187: Mucosal Melanoma of the Head and Neck (13 samples)
 - NCIT:C155790: Malignant Skull Neoplasm (17 samples)
 - NCIT:C164198: Head and Neck Sarcoma (1 sample)
 - NCIT:C170467: Metastatic Malignant Head and Neck Neoplasm (15 samples)

Progenetix in 2021

Variant and Metadata for Sample Discovery

- positional queries for genomic variants using the GA4GH Beacon protocol
- metadata queries (diagnoses, identifiers, clinical classes ...) using Beacon "filters"



progenetix

Cancer CNV Profiles

Search Samples

Studies & Cohorts

- arrayMap
- TCGA Samples
- DIPG Samples
- Gao & Baudis, 2021
- Cancer Cell Lines

Publication DB

Services

- NCIt Mappings
- UBERON Mappings

Upload & Plot

Download Data

Beacon+

Progenetix Info

- About Progenetix
- Use Cases
- Documentation
- Baudisgroup @ UZH

Search Samples

CDKN2A Deletion Example MYC Duplication TP53 Del. in Cell Lines K-562 Cell Line

Gene Spans Cytoband(s)

This example shows the query for CNV deletion variants overlapping the CDKN2A gene's coding region with at least a single base, but limited to "highly focal" hits (here i.e. $\leq \sim 1\text{Mbp}$ in size). The query can be modified e.g. through changing the position parameters or diagnosis.

Gene Symbol i
Select...

Chromosome i (Structural) Variant Type i
9 DEL (Deletion)

Start or Position i End (Range or Structural Var.) i
21500001-21975098 21967753-22500000

Minimum Variant Length i Maximal Variant Length i

Reference ID(s) i
Select...

Cancer Classification(s) i Clinical Classes i
NCIT:C3058: Glioblastoma (4375) X Select...

Genotypic Sex i Biosample Type i
Select... Select...

Filters i Filter Logic i
exact AND

Filter Precision i
exact

City i

Chromosome 9 i
21500001-21975098 21967753-22500000

Query Database

Progenetix in 2021

Query Results and Variant Frequencies

- genomic variant + metadata queries provide relative result counts / frequencies for mapped entities (NCIt, ICD-O ...)
- disease-specific CNA event scores
- representation of genome -wide CNA frequency profiles / context
- link-outs to download options, subset visualization, sample exploration ...



progenetix

Search Samples Modify Query

Cancer CNV Profiles

Search Samples

Studies & Cohorts

- arrayMap
- TCGA Samples
- DIPG Samples
- Gao & Baudis, 2021
- Cancer Cell Lines

Publication DB

Services

- NCIt Mappings
- UBERON Mappings

Upload & Plot

Download Data

Beacon⁺

Progenetix Info

- About Progenetix
- Use Cases
- Documentation
- Baudisgroup @ UZH

Assembly: GRCh38 Chro: 9 Start: 21500001-21975098 End: 21967753-22500000

Type: DEL Filters: NCIT:C3058

progenetix

Samples: 678 Variants: 297 Calls: 687

Found Variants (.pgxseg) All Sample Variants (.json) All Sample Variants (.pgxseg) Show Variants in UCSC

UCSC region JSON Response Visualization options

Results Biosamples Biosamples Map Variants

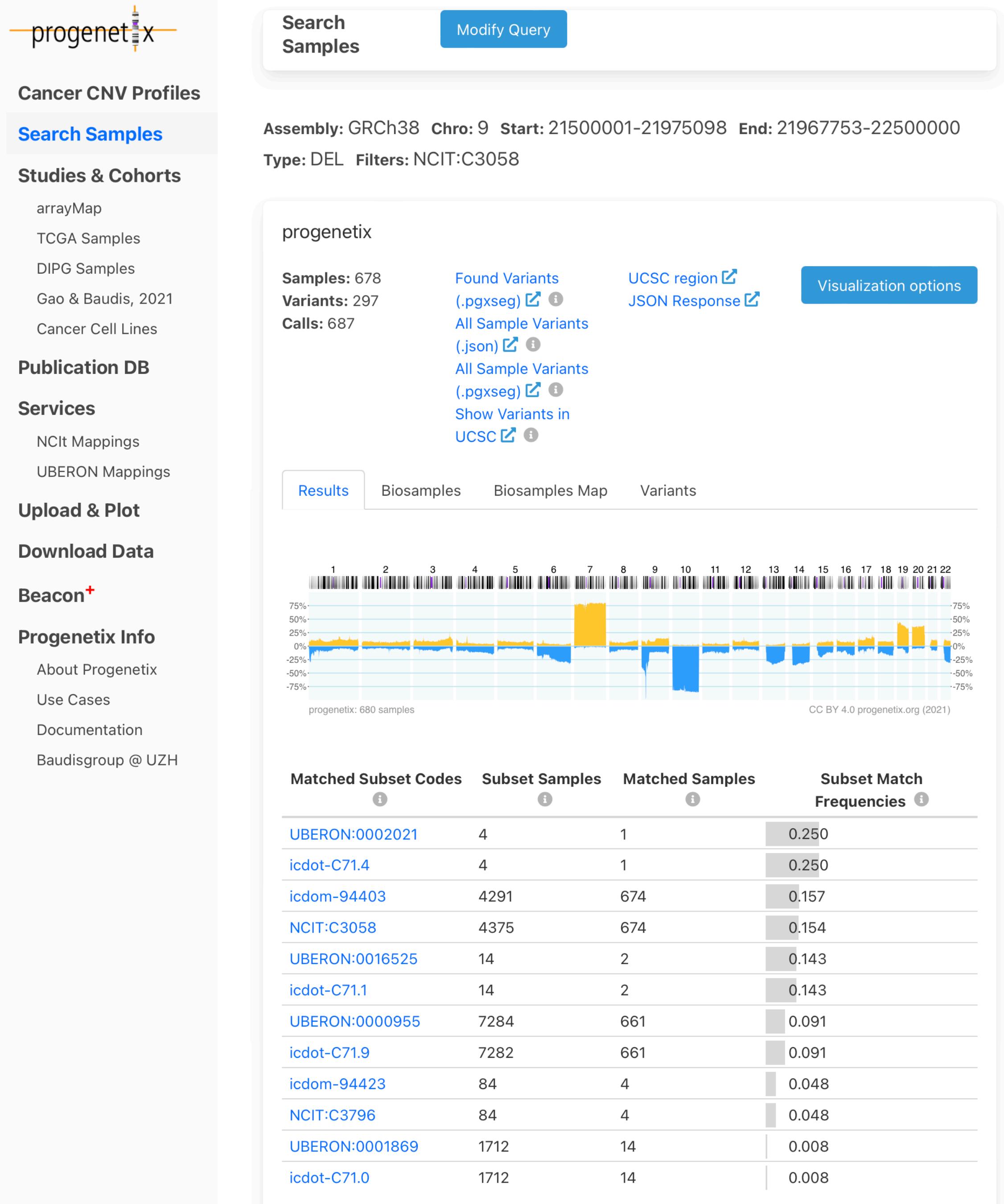
A horizontal plot showing CNA event scores across chromosomes 1 through 22. The y-axis ranges from -75% to 75%. A prominent yellow bar is visible on chromosome 7, indicating a high-frequency event. Below the plot, the text "progenetix: 680 samples" is displayed.

CC BY 4.0 progenetix.org (2021)

Matched Subset Codes	Subset Samples	Matched Samples	Subset Match Frequencies
UBERON:0002021	4	1	0.250
icdot-C71.4	4	1	0.250
icdom-94403	4291	674	0.157
NCIT:C3058	4375	674	0.154
UBERON:0016525	14	2	0.143
icdot-C71.1	14	2	0.143
UBERON:0000955	7284	661	0.091
icdot-C71.9	7282	661	0.091
icdom-94423	84	4	0.048
NCIT:C3796	84	4	0.048
UBERON:0001869	1712	14	0.008
icdot-C71.0	1712	14	0.008

Progenetix in 2021

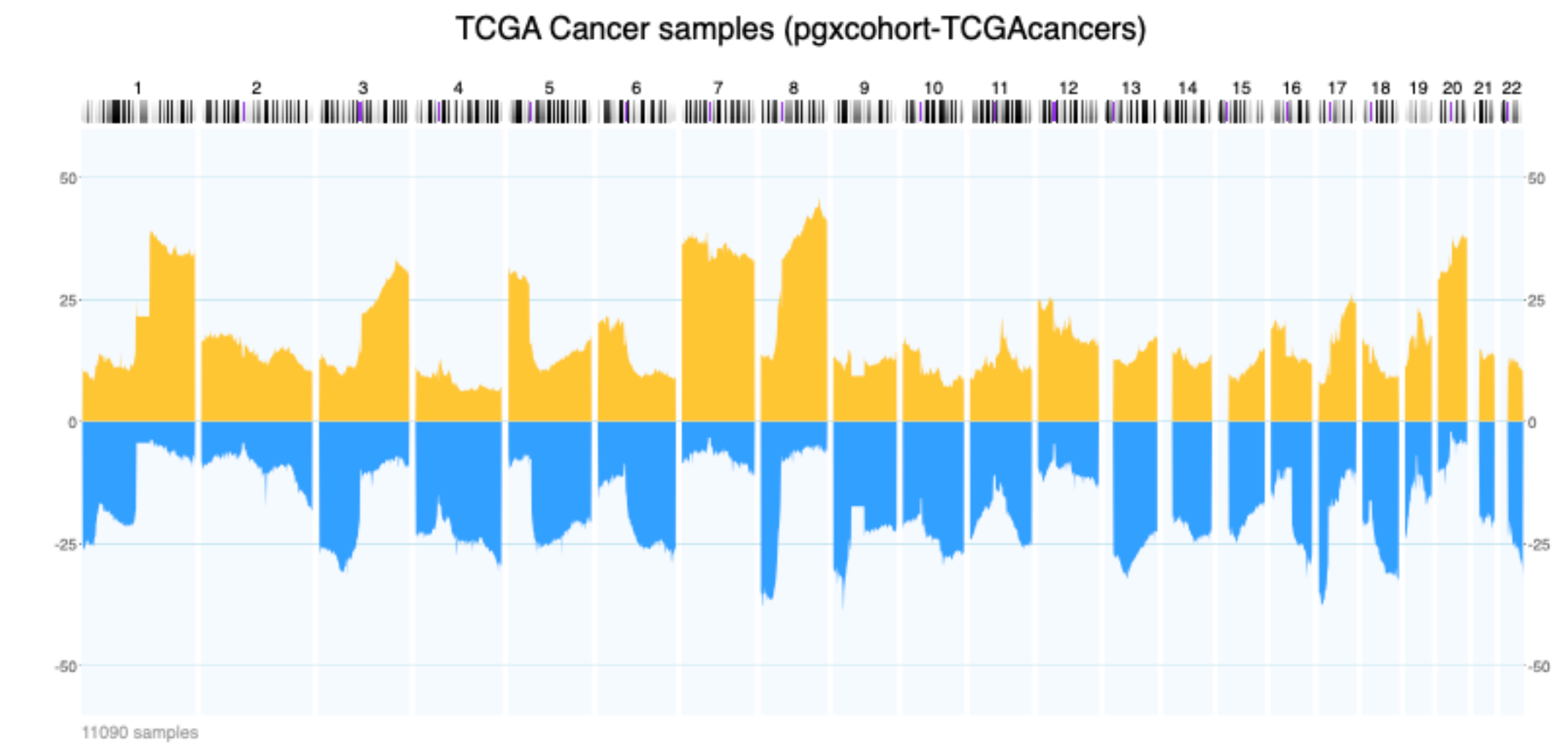
Query Results and Variant Frequencies



Progenetix API

Data & Plots

- "all" of the data can be accessed using API calls
- segmented CNV data in .pgxsg (columnar) and JSON format
- histograms for disease, study or cohort from precomputed frequencies - live generated as SVG for embedding with plot options



[https://progenetix.org/cgi/PGX/cgi/collationPlots.cgi?
datasetIds=progenetix&id=pgxcohort-TCGAcancers&-
size_plotimage_w_px=800&-size_plotarea_h_px=300&-
value_plot_y_max=60](https://progenetix.org/cgi/PGX/cgi/collationPlots.cgi?datasetIds=progenetix&id=pgxcohort-TCGAcancers&-size_plotimage_w_px=800&-size_plotarea_h_px=300&-value_plot_y_max=60)

Progenetix

Services, Documentation...

- services e.g. for disease code translation (NCIt
<=> ICD-O; UBERON ...)
- API & documentation "progressing" ...

progenetix

Services: Ontologymaps (NCIt)

The **ontologymaps** service provides equivalence mapping between ICD-O and other classification systems, notably NCIt and UBERON. It makes use of the sample-level mappings for NCIT and ICD-O 3 codes developed for the individual samples in the Progenetix collection.

NCIT and ICD-O 3

While NCIT treats diseases as **histologic** and **topographic** described entities (e.g. **NCIT:C7700: Ovarian adenocarcinoma**), these two components are represented separately in ICD-O, through the **Morphology** and **Topography** coding arms (e.g. here **8140/3 + C56.9**).

More documentation with focus on the API functionality can be found on the [documentation pages](#).

The data of all mappings can be retrieved through this API call: [{JSON}](#)

Code Selection

gio|

NCIT:C3058: Glioblastoma
NCIT:C3288: Oligodendrogloma
NCIT:C4326: Anaplastic Oligodendrogloma
NCIT:C3903: Mixed Glioma
NCIT:C3059: Glioma
NCIT:C3796: Gliosarcoma
NCIT:C4822: Malignant Glioma
NCIT:C3308: Paraganglioma
NCIT:C4831A: Renal Paraganglioma

About Progenetix

News
Cancer CNV Profiles
Search Samples
arrayMap
Studies & Cohorts
Publication DB
Services
Upload & Plot
Documentation
Beacon+
Baudisgroup @ UZH

NCIthesaurus

Progenetix :: Info

Structural Cancer Genomics Resource Documentation and Example Pages

News
About...
API
Documentation
Publications
Progenetix Home

Related Sites

Progentix Data
Baudisgroup @ UZH
Beacon+
SchemaBlocks {S}[B]
Beacon Project

Github Projects

baudisgroup
progenetix
ELIXIR Beacon

Tags

API Beacon BeaconPlus
BeaconV2 GA4GH Perl Python
TCGA article bycon code
documentation identifiers licensing
ontologies prefixes schemas
services statistics tools website

Welcome to the *Progenetix* documentation pages

The **Progenetix Resource Documentation** provides information and links related to the **Progenetix** cancer genome resource and the related **Progenetix code repositories** contains projects, such as data conversion scripts, ontology mappings and code for the **Beacon+** project.

Progenetix Website Code Repositories

- **Progenetix Source Code**
- **Related Projects**

Latest News

Progenetix File Formats

Standard Progenetix Segment Files [.pgxseg](#)

Progenetix uses a variation of a standard tab-separated columnar text file such as produced by array or sequencing CNV software, with an optional metadata header for e.g. plot or grouping instructions.

@mbaudis 2021-02-22: [more ...](#)

Beacon+ and Progenetix Queries by Gene Symbol

We have introduced a simple option to search directly by Gene Symbol, which will match to *any* genomic variant with partial overlap to the specified gene. This works by expanding the Gene Symbol (e.g. **TP53**, **CDKN2A** ...) into a range query for its genomic coordinates (maximum CDR).

Such queries - which would e.g. return all whole-chromosome CNV events covering the gene of interest, too - should be narrowed by providing e.g. **Variant Type** and **Maximum Size** (e.g. 2000000) values.

@mbaudis 2021-02-22: [more ...](#)

Gene Symbol
MYC
MYCBP (1:38864669-38873304)
MYCBPAP (17:56508545-50531427)
MYCL (1:39897371-39901887)
MYCN (2:15940586-15946096)

The Progenetix oncogenomic resource in 2021

Qingyao Huang, Paula Carrio Cordero, Bo Gao, Rahel Paloots, Michael Baudis

bioRxiv. doi: <https://doi.org/10.1101/2021.02.15.428237>

This article provides an overview of recent changes and additions to the Progenetix database and the services provided through the resource.

2021-02-15: [more ...](#)

Diffuse Intrinsic Pontine Glioma (DIPG) cohort

Diffuse Intrinsic Pontine Glioma (DIPG) is a highly aggressive tumor type that originates from glial cells in the pons area of the brainstem, which controls vital functions including breathing, blood pressure and heart rate. DIPG occurs frequently in the early childhood and has a 5-year survival rate below 1 percent. Progenetix has now incorporated the DIPG cohort, consisting of 1067 individuals from 18 publications. The measured data include copy number variation as well as (in part) point mutations on relevant genes, e.g. TP53, NF1, ATRX, TERT promoter.

@qingyao 2021-02-15: [more ...](#)

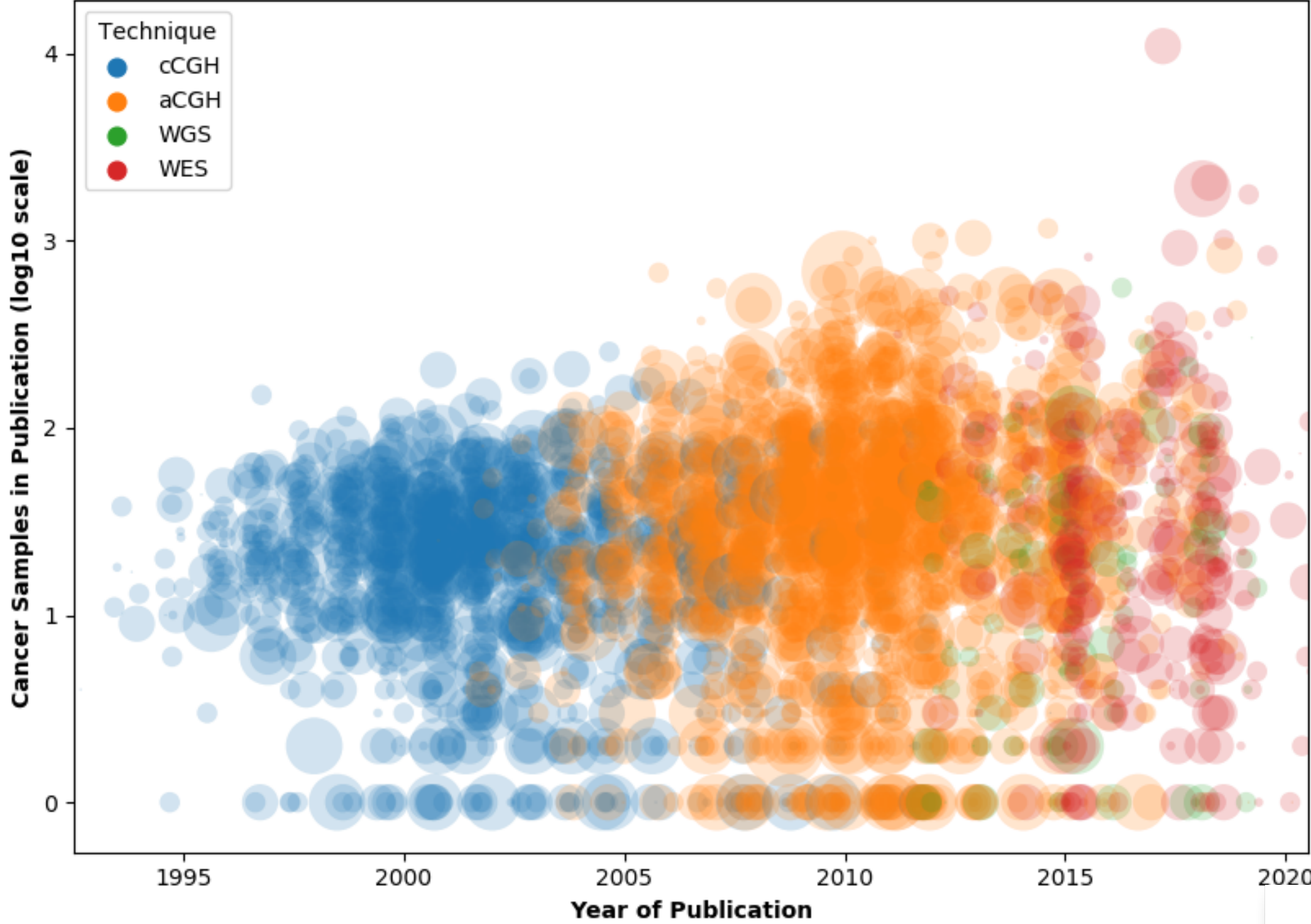
arrayMap is Back

After some months of dormancy, the **arrayMap** resource has been relaunched through integration with the new **Progenetix** site. All of the original arrayMap data has now been integrated into Progenetix, and of today the [arraymap.org](#) domain maps to a standard Progenetix search page, where only data samples with existing source data (e.g. probe specific array files) will be presented.

@mbaudis 2021-02-06: [more ...](#)



Number of tumor samples for each publication across the years



Map of the geographic distribution (by first author affiliation) of the 104'543 genomic array, 36'766 chromosomal CGH and 15'409 whole genome/exome based cancer genome datasets.

The numbers are derived from the 3'240 publications registered in the Progenetix database.



Progenetix Publication Collection

The current page lists articles describing whole genome screening (WGS, WES, aCGH, cCGH) experiments in cancer, registered in the Progenetix publication collection. For each publication the table indicates the numbers of samples analysed with a given technology and if sample profiles are available in Progenetix.

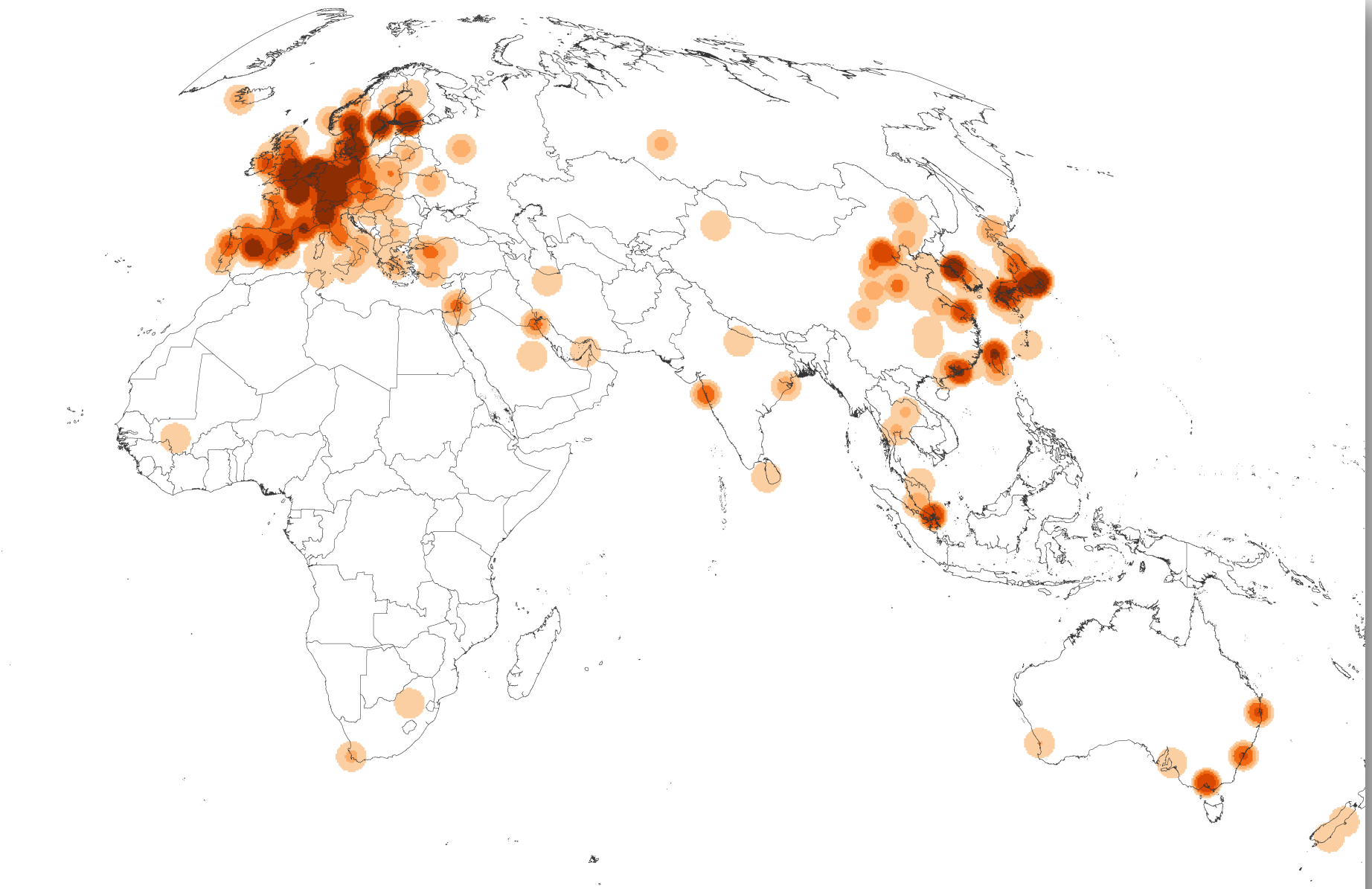
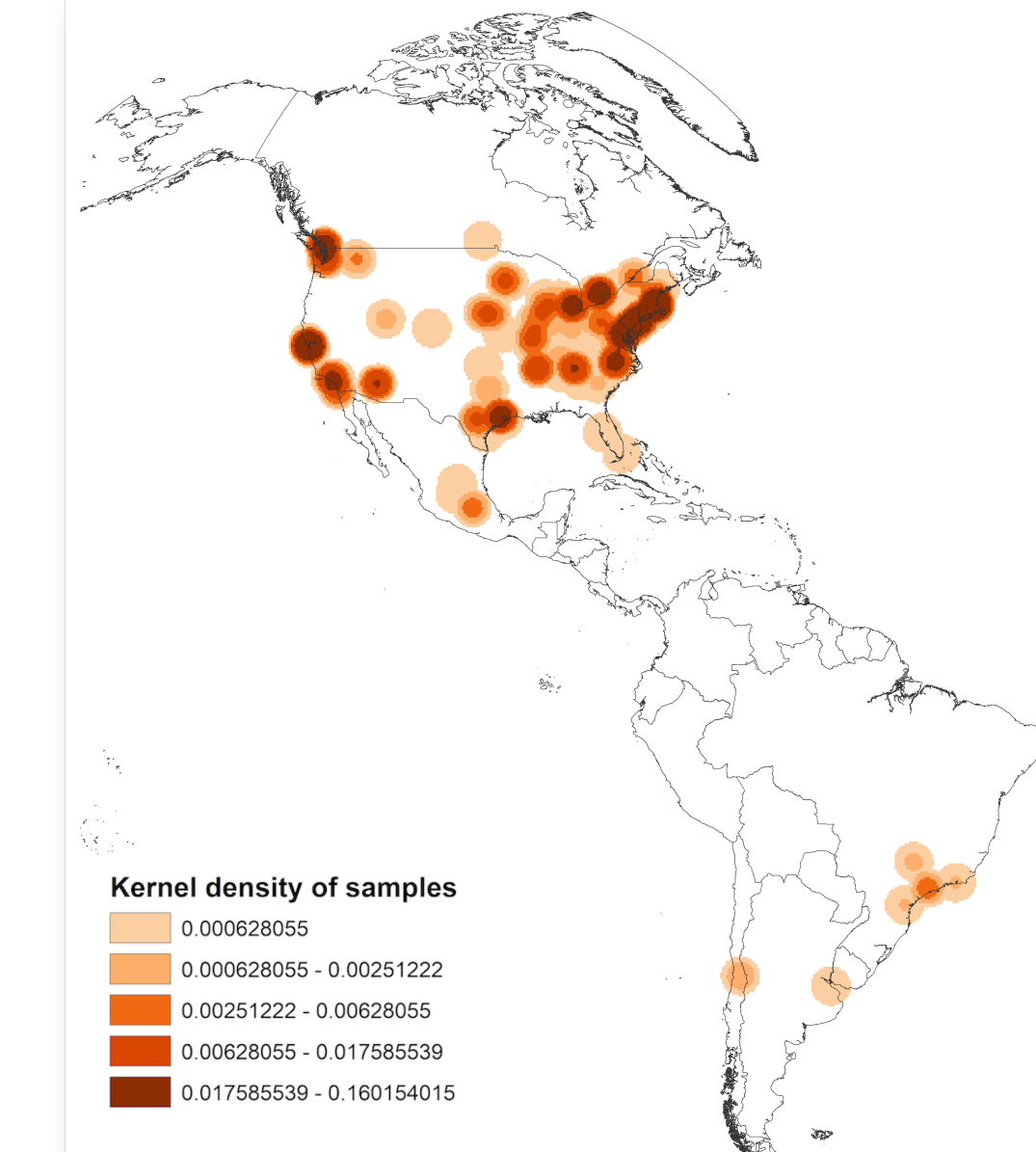
Please [contact us](#) to alert us about additional articles you are aware of. The inclusion criteria are described in the documentation [🔗](#).

Filter [i](#) City [i](#)

 Type to search... [▼](#)

Publications (3324)

id i ▾	Publication	cCGH	aCGH	WES	WGS	pgx
PMID:34103027	Peng G, Chai H, Ji W, Lu Y, Wu S et al. (2021) Correlating genomic copy number alterations with clinicopathologic findings in 75 cases of ... BMC Med Genomics	0	79	0	0	0
PMID:34059130	Tsui DWY, Cheng ML, Shady M, Yang JL et al. (2021) Tumor fraction-guided cell-free DNA profiling in metastatic solid tumor patients. ... J Clin Oncol	0	0	5	113	0



Progenetix and GA4GH Beacon

Implementation driven development of a GA4GH standard



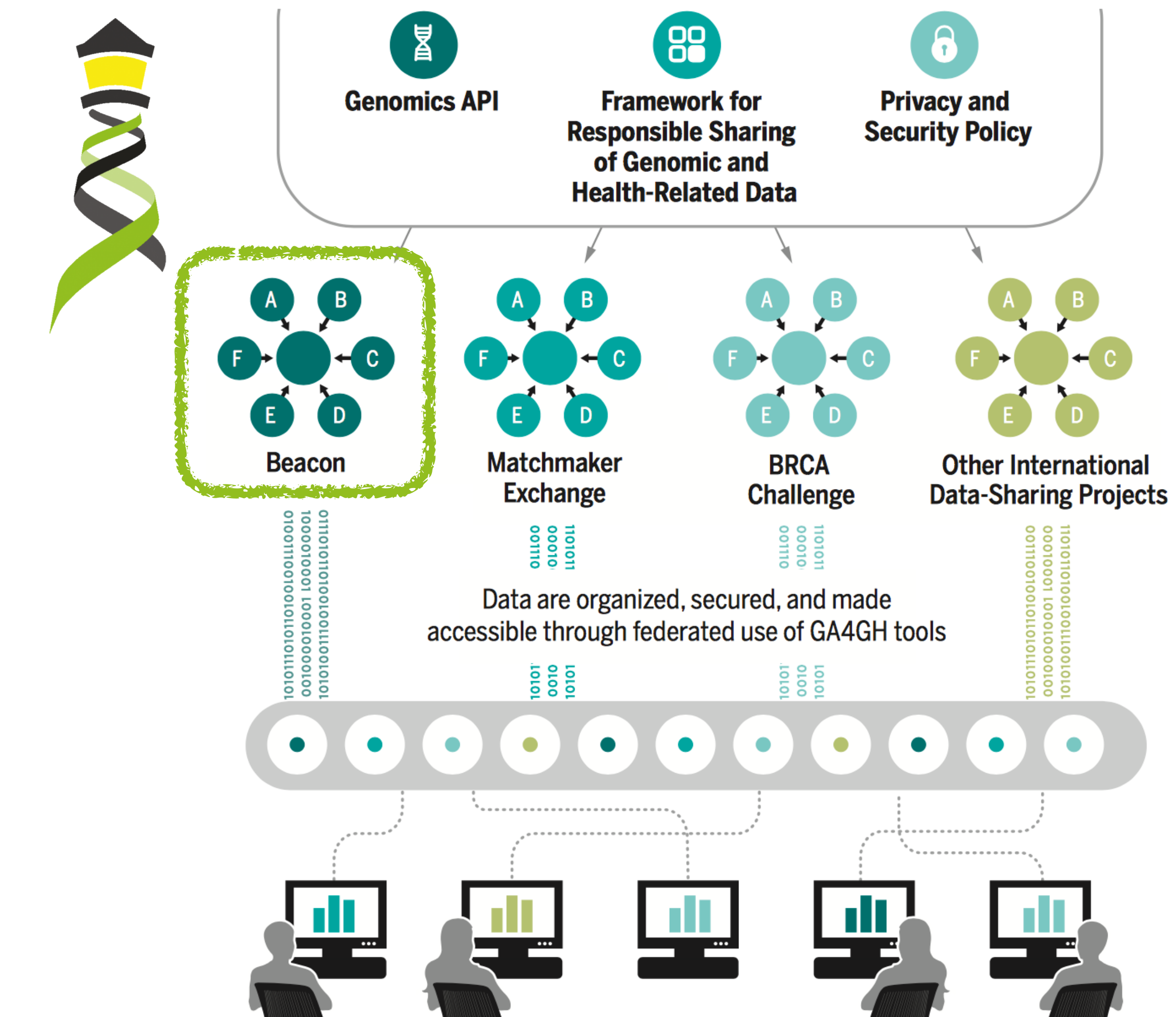


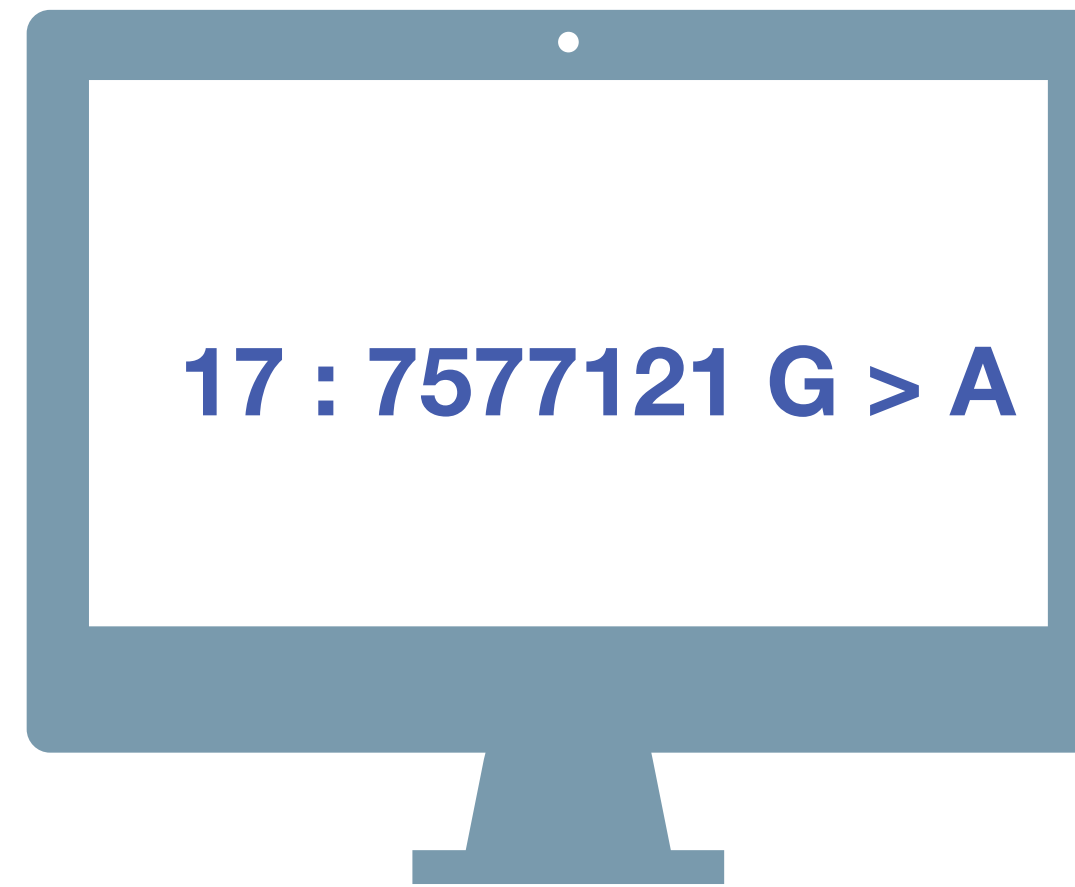
GENOMICS

A federated ecosystem for sharing genomic, clinical data

Silos of genome data collection are being transformed into seamlessly connected, independent systems

A federated data ecosystem. To share genomic data globally, this approach furthers medical research without requiring compatible data sets or compromising patient identity.





Beacon

A **Beacon** answers a query for a specific genome variant against individual or aggregate genome collections

YES | NO | \0

ELIXIR Beacon Project

- Driver project on GA4GH roadmap
- aligns with Discovery Work Stream
- strong impact on GA4GH developments as a concrete, funded project

The screenshot shows two cards. The left card is titled 'Driver Projects' and contains text about real-world genomic data initiatives. The right card is titled 'ELIXIR Beacon' and provides links to its implementation studies and champions.

Driver Projects
GA4GH Driver Projects are real-world genomic data initiatives that help guide our development efforts and pilot our tools. Stakeholders around the globe advocate, mandate, implement, and use our frameworks and standards in their local contexts.

ELIXIR Beacon
<https://www.elixir-europe.org/about/implementation-studies/beacons>

Europe
Champions: Jordi Rambla, Juha Tornroos, Gary Saunders

v1.1 and roadmap

- structural variations** (DUP, DEL) in addition to SNV
- ... more structural queries (translocations/fusions...)
- Beacon queries as entry for **data handover** (outside Beacon protocol)
- layered authentication system using **ELIXIR AAI**
- v2** **filters** for phenotypic & technical metadata
- v2** Extended quantitative responses
- Ubiquitous **deployment** (e.g. throughout ELIXIR network)



Beacon+ by Progenetix

From Beacon Query to Explorative Analyses of CNV Patterns

- Since 2016 the Progenetix resource has been used to model options for Beacon development
 - 138334 individual samples from 698 cancer types
- The consistent use of hierarchical diagnostic codes allows the use of Beacon "filters" for histopathological/clinically scoped queries
- Beacon's handover protocols can be utilized for data retrieval and, well, handing over to additional services, e.g.
 - downloads
 - visualization
 - use of external services (UCSC browser display...)



Search Samples

CNV Request Allele Request Range Query All Fields

CNV Example

This query type is for copy number queries ("variantCNVrequest"), e.g. using fuzzy ranges for start and end positions to capture a set of similar variants.

Dataset
progenetix

Cohorts

Genome Assembly GRCh38 / hg38

Gene Symbol

Reference name 9 **(Structural) Variant Type** DEL

Start or Position 19000001-21975098 **End (Range or Structural Var.)** 21967753-24000000

Minimum Variant Length **Maximal Variant Length**

Cancer Classification(s)

Filters

City

Query Database

Beacon v2 Requests

POSTing Queries

- Beacon v2 supports a mix of dedicated endpoints with REST paths
- POST requests using JSON query documents
- final syntax for core parameters still in testing stages

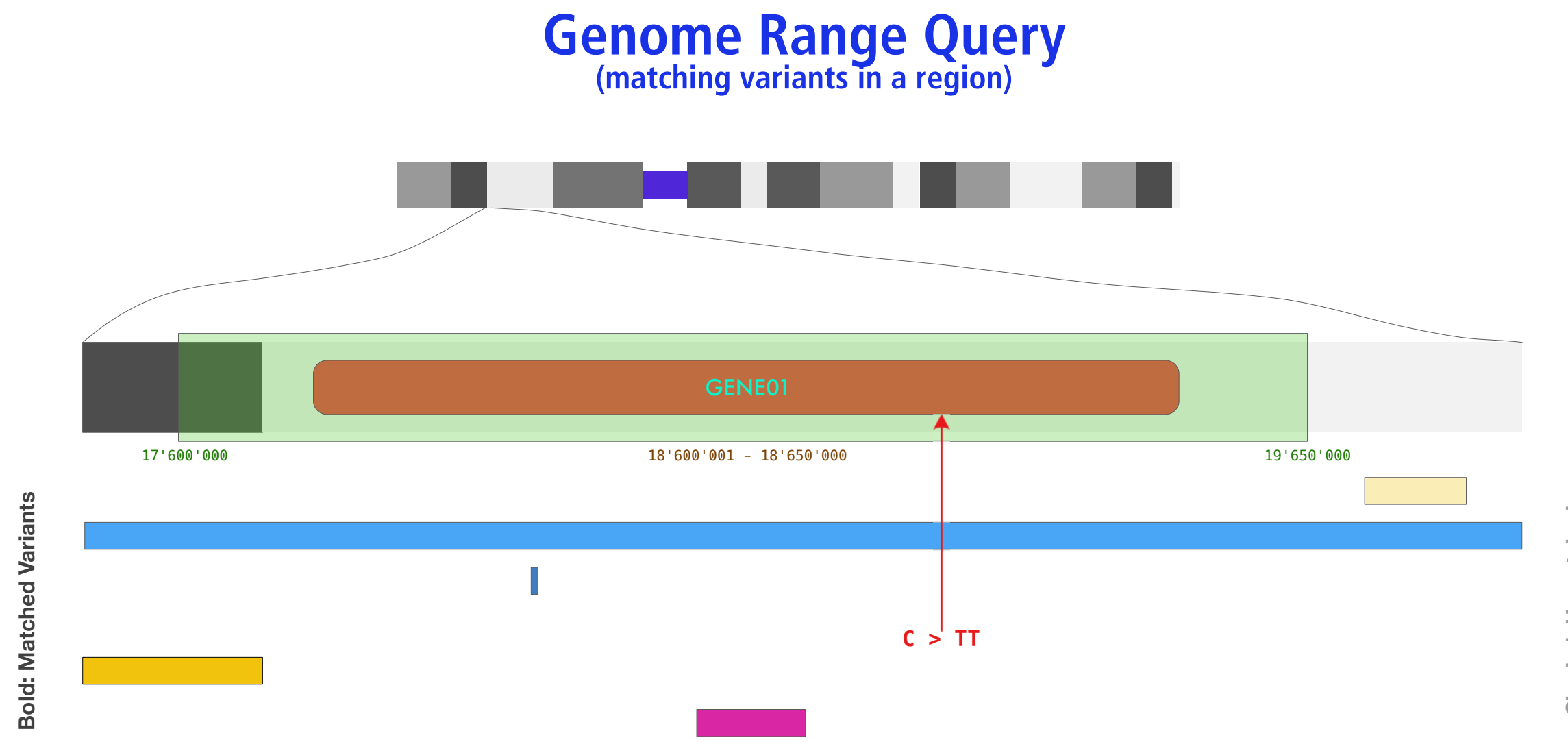
```
{  
  "$schema": "beaconRequestBody.json",  
  "meta": {  
    "apiVersion": "2.0",  
    "requestedSchemas": [  
      {  
        "entityType": "individual",  
        "schema": "https://progenetix.org/services/schemas/Phenopacket/"  
      }  
    ],  
    "query": {  
      "requestParameters": {  
        "datasets": {  
          "datasetIds": ["progenetix"]  
        }  
      },  
      "filterLogic": "OR"  
    },  
    "pagination": {  
      "skip": 0,  
      "limit": 10  
    },  
    "filters": [  
      { "id": "NCIT:C4536" },  
      { "id": "NCIT:C95597" },  
      { "id": "NCIT:C7712" }  
    ]  
  }  
}
```



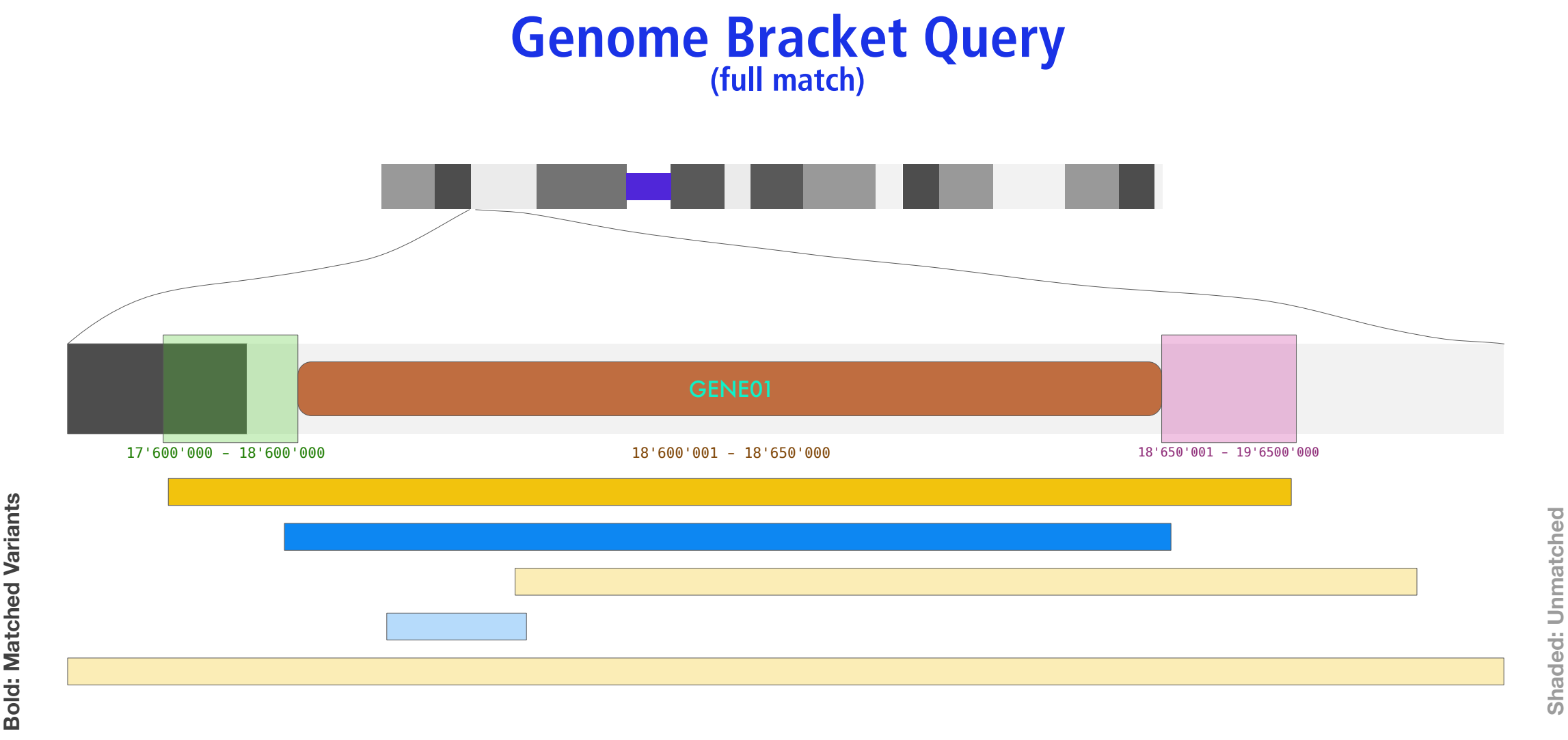


Beacon v2: Extended Variant Queries

Range and Bracket queries enable positional wildcards and fuzziness



- Genome Range Queries provide a way to "fish" for variants overlapping an indicated region, e.g. the CDR of a gene of interest
- Additional parameters (e.g. variant type, reference or alternate bases) limit the scope of the responses
- new Beacon v2 size parameters to limit structural variants (e.g. "focal" CNVs)



- Genome Bracket Queries allow to search for structural variants with start and end positions falling into defined sequence ranges
- allows to query any contiguous genomic variant (and in principle also can step in for range queries)
- typical use case is e.g. the query for variants such as duplications covering the whole CDR of a gene, while limiting the allowed start or end regions

Beacon v2 Filters

Example: Use of hierarchical classification systems (here NCI ICD neoplasm core)

- Beacon v2 "filters" assumes inclusion of child terms when using hierarchical classifications
 - ➡ implicit *OR* with otherwise assumed *AND*
 - implementation of hierarchical annotations overcomes some limitations of "fuzzy" disease annotations



Beacon+ specific: Multiple term selection with OR logic

<input checked="" type="checkbox"/>	> NCIT:C4914: Skin Carcinoma	213
<input type="checkbox"/>	> NCIT:C4475: Dermal Neoplasm	109
<input checked="" type="checkbox"/>	> NCIT:C45240: Cutaneous Hematopoietic and Lymphoid Cell Neoplasm	310

Filters: NCIT:C4914, NCIT:C4819, NCIT:C9231, NCIT:C2921, NCIT:C45240, NCIT:C6858, NCIT:C3467, NCIT:C45340, NCIT:C7195, NCIT:C3246, NCIT:C7217



progenetix

Variants: 0 *f*alleles: 0 Callsets Variants ↗ UCSC region ↗ Legacy Interface ↗ [Show JSON Response](#)

Results **Biosamples**

Id	Description	Classifications	Identifiers	DEL	DUP	CNV
PGX_AM_BS_MCC01	Merkel cell carcinoma	icdot-C44.9 Skin, NOS icdom-82473 Merkel cell carcinoma NCIT:C9231 Merkel Cell Carcinoma	PMID:9537255	0.116	0.104	0.22
PGX_AM_BS_MCC02	Merkel cell carcinoma	icdot-C44.9 Skin, NOS icdom-82473 Merkel cell carcinoma NCIT:C9231 Merkel Cell Carcinoma	PMID:9537255	0.154	0.056	0.21
PGX_AM_BS_MCC03	Merkel cell carcinoma	icdot-C44.9 Skin, NOS icdom-82473 Merkel cell carcinoma NCIT:C9231 Merkel Cell Carcinoma	PMID:9537255	0.137	0.21	0.347
PGX_AM_BS_MCC04	Merkel cell carcinoma	icdot-C44.9 Skin, NOS icdom-82473 Merkel cell carcinoma NCIT:C9231 Merkel Cell Carcinoma	PMID:9537255	0.158	0.056	0.214
PGX_AM_BS_MCC05	Merkel cell carcinoma	icdot-C44.9 Skin, NOS icdom-82473 Merkel cell carcinoma NCIT:C9231 Merkel Cell Carcinoma	PMID:9537255	0.107	0.327	0.434

Standardized Data

Data re-use depends on standardized, machine-readable metadata

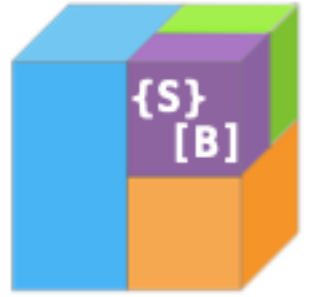
- Multiple international initiatives (ELIXIR, GA4GH, MONARCH...) and resource providers (EBI, NCBI ...) work on the generation and implementation of data annotation standards
- emerging / established principles are the use of hierarchical coding systems where individual codes are represented as CURIEs
- other formats for non-categorical annotations based on international standards, e.g.
 - ISO (ISO 8601 time & period, ISO 3166 country codes ...)
 - IETF (GeoJSON ...)
 - W3C (CURIE ...)
- these standards become pervasive throughout GA4GH's ecosystem (e.g. Phenopackets ...)

```
"data_use_conditions" : {  
    "label" : "no restriction",  
    "id" : "DUO:0000004"  
},  
  
"provenance" : {  
    "material" : {  
        "type" : {  
            "id" : "EFO:0009656",  
            "label" : "neoplastic sample"  
        }  
    },  
    "geo" : {  
        "label" : "Zurich, Switzerland",  
        "precision" : "city",  
        "city" : "Zurich",  
        "country" : "Switzerland",  
        "latitude" : 47.37,  
        "longitude" : 8.55,  
        "geojson" : {  
            "type" : "Point",  
            "coordinates" : [  
                8.55,  
                47.37  
            ]  
        },  
        "IS0-3166-alpha3" : "CHE"  
    },  
    {  
        "age": "P25Y3M2D"  
    }  
}
```

GA4GH {S}[B] SchemaBlocks

Standardized formats and data schemas for developing an "Internet of Genomics"

- “cross-workstreams, cross-drivers” initiative to document GA4GH object **standards** and **prototypes** launched in December 2018
- documentation and implementation examples provided by GA4GH members
- not a rigid, complete data schema
- object **vocabulary** and **semantics** for a large range of developments
- ▶ **Beacon** as contributor and user
- ▶ 2021: going forward through integration with GA4GH TASC efforts, towards "standards library"



Biosample sb-phenopackets ↗	
{S}[B] Status [i]	implemented
Provenance	◦ Phenopackets
Used by	◦ Phenopackets
Contributors	◦ GA4GH Data Working Group ◦ Jules Jacobsen ◦ Peter Robinson ◦ Michael Baudis ◦ Melanie Courtot ◦ Isuru Liyanage
Source (v1.0.0)	◦ raw source [JSON] ◦ Github
Attributes	
Type:	object
Description:	A Biosample refers to a unit of biological material from which the genomic DNA, RNA, proteins) for molecular analyses (e.g. sequencing, array hybridization, spectrometry) are extracted. Examples would be a tissue biopsy, a single cell from a culture or single cell gel fraction from a gradient centrifugation. Several instances (e.g. technical replicates) or types of experiments (e.g. genome experiments) may refer to the same Biosample.
FHIR mapping:	Specimen.
Properties	
Property	Type
ageOfIndividualAtCollection	https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/Age.json [SRC] [HTML]
ageRangeOfIndividualAtCollection	https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/AgeRange.json [SRC] [HTML]
description	string
diagnosticMarkers	array of https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/OntologyClass.json [SRC] [HTML]
histologicalDiagnosis	https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/OntologyClass.json [SRC] [HTML]
htsFiles	array of https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/HtsFile.json [SRC] [HTML]
id	string
individualId	string
isControlSample	boolean
phenotypicFeature	array of https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/PhenotypicFeature.json [SRC] [HTML]
procedure	https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/Procedure.json [SRC] [HTML]
sampledTissue	https://schemablocks.org/schemas/sb-phenopackets/v1.0.0/Tissue.json [SRC] [HTML]
HtsFile sb-phenopackets ↗	
{S}[B] Status [i]	implemented
Provenance	◦ Phenopackets
Used by	◦ Phenopackets
Contributors	◦ Jules Jacobsen ◦ Peter Robinson
Source (v1.0.0)	◦ raw source [JSON] ◦ Github
Attributes	
Type:	object
Description:	A file in one of the HTS formats (https://samtools.github.io/hts-specs)
Properties	
Property	Type
description	string
genomeAssembly	string
htsFormat	
individualToSampleIdentifiers	object
uri	string

schemablocks.org

Beacon v2 Paths

Progenetix utilizes Beacon v2 REST paths

- Beacon v2 paths are used in the Beacon specification to scope query and delivery
- Progenetix uses a default `/biosamples/` + query path for its front end queries, and then collection specific methods for data retrieval (see next)
- current implementation addresses a core subset of all options, and evaluates some still moving targets
 - variants_interpretations
 - variant instances versus prototypes
 - ...



Base `/biosamples`

`/biosamples/` + query

- `/biosamples/?filters=cellosaurus:CVCL_0004`
 - this example retrieves all biosamples having an annotation for the Cellosaurus CVCL_0004 identifier (K562)

`/biosamples/{id}/`

- `/biosamples/pgxbs-kftva5c9/`
 - retrieval of a single biosample

`/biosamples/{id}/variants/` & `/biosamples/{id}/variants_in_sample/`

- `/biosamples/pgxbs-kftva5c9/variants/`
- `/biosamples/pgxbs-kftva5c9/variants_in_sample/`
 - retrieval of all variants from a single biosample
 - currently - and especially since for a mostly CNV containing resource - `variants` means "variant instances" (or as in the early v2 draft `variantsInSample`)

Base `/variants`

There is currently (April 2021) still some discussion about the implementation and naming of the different types of genomic variant endpoints. Since the Progenetix collections follow a "variant observations" principle all variant requests are directed against the local `variants` collection.

If using `g_variants` or `variants_in_sample`, those will be treated as aliases.

`/variants/` + query

- `/variants/?assemblyId=GRCh38&referenceName=17&variantType=DEL&filterLogic=AND&start=7500000&start=7676592&end=7669607&end=7800000`
 - This is an example for a Beacon "Bracket Query" which will return focal deletions in the TP53 locus (by position).

`/variants/{id}/` or `/variants_in_sample/{id}/` or `/g_variants/{id}/`

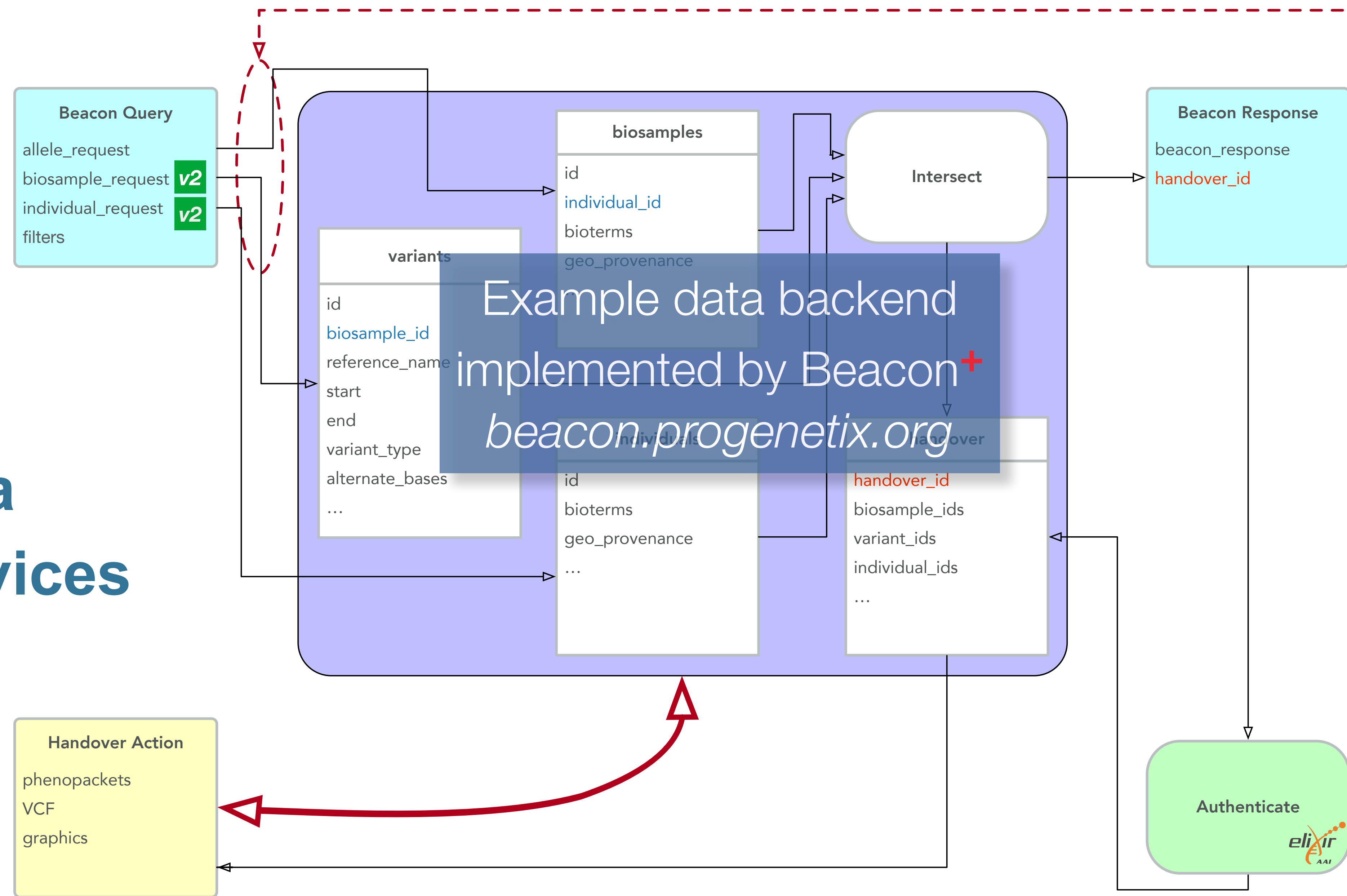
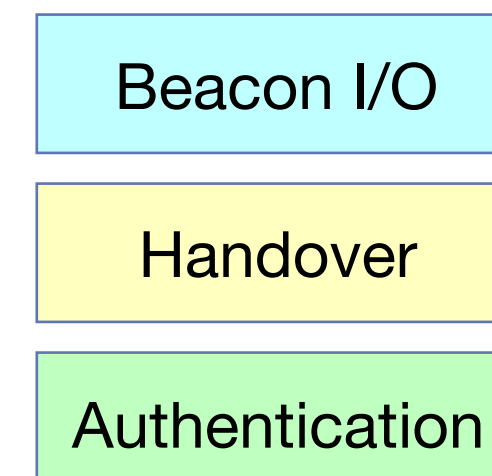
- `/variants/5f5a35586b8c1d6d377b77f6/`
- `/variants_in_sample/5f5a35586b8c1d6d377b77f6/`

`/variants/{id}/biosamples/` & `variants_in_sample/{id}/biosamples/`

- `/variants/5f5a35586b8c1d6d377b77f6/biosamples/`
- `/variants_in_sample/5f5a35586b8c1d6d377b77f6/biosamples/`

Beacon & Handover

Beacons v1.1
supports data
delivery services





Onboarding

Demonstrating Compliance

- onboarding server run by CRG (EGA team)
- registering the URI of a server's map document will initiate traversal and testing of services
- blueprint for Beacon service registries
- to be used as demonstrator in GA4GH approval process for the Spring 2022 session

The screenshot shows a grid of service registries being tested against the Beacon API specification. Each row contains the service name, its logo, and a summary of its compliance status across various API endpoints.

Service	Configuration	analysis	dataset	interactor	variantInSample	BeaconMap	biosample	genomicVariant	run	variantAnnotation	EntryTypes	cohort	individual	variantAnnotation
Centre Nacional Analisis Genomica (CNAG-CRG)	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗
Progenetix	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗
baudisgroup at UZH and SIB	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗
European Genome-Phenome Archive (EGA)	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗
bioRxiv	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗
bioRxiv Genomics	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗	✗

Legend at the bottom:

- Green checkmark: Matches the Spec
- Red X: Not Match the Spec
- Grey question mark: Not Implemented

ga4gh-beacon / beacon-framework-v2 Public

Code Issues 18 Pull requests 2 Discussions Actions Wiki Security Insights Settings

main 7 branches 0 tags Go to file Add file Code About

jrambla Merge pull request #51 from ga4gh-beacon/configuration-typos-fixes ...

common	de-lining \n
configuration	speling in configuration -> filteringTermsSchema
requests	de-lining \n
responses	de-lining \n
.gitignore	Initial commit
LICENSE	Initial commit
README.md	Adding naming conventions to readme
endpoints.json	de-lining \n

README.md

beacon-framework-v2

Beacon Framework version 2

Introduction

The GA4GH Beacon specification is composed by two parts:

- the Beacon Framework (in *this* repo)
- the Beacon Model (in the [Models repo](#))

The Beacon Framework is the part that describes the overall structure of the API

progenetix / bycon Public

Code Issues Pull requests 1 Actions Projects Wiki Security Insights Settings

master 3 branches 0 tags Go to file Add file Code About

mbaudis Update README.md 5064e89 11 seconds ago 519 commits

beaconServer	datatables, genesRefresher	6 days ago
byconeer	datatables, genesRefresher	6 days ago
config	datatables, genesRefresher	6 days ago
lib	intervalFrequencies service & some library shuffling	5 months ago
schemas	datatables, genesRefresher	6 days ago
services	genespan method for gene request size reduction	2 days ago
remnants	biocharacteristics removal; shuffling of beaconsv2 references...	21 days ago
.gitignore	biocharacteristics removal; shuffling of beaconsv2 references...	21 days ago
LICENSE	Create LICENSE	12 months ago
README.md	Update README.md	11 seconds ago
__init__.py	intervalFrequencies service & some library shuffling	5 months ago
requirements.txt	add non-interactive mode	16 months ago

README.md

License CC0 1.0

Bycon - a Python-based environment for the Beacon v2 genomics API

The `bycon` project - at least at its current stage - is a mix of Progenetix (i.e. GA4GH object model derived, MongoDB implemented) - data management, and the implementation of middleware & server for the Beacon API.

More information about the current status of the package can be found in the inline documentation which is also [presented in an accessible format](#) on the [Progenetix website](#).

About

Bycon - A Python Based Beacon API (beacon-project.io) implementation leveraging the Progenetix (progenetix.org) data model

Readme

CC0-1.0 License

Releases

No releases published [Create a new release](#)

Packages

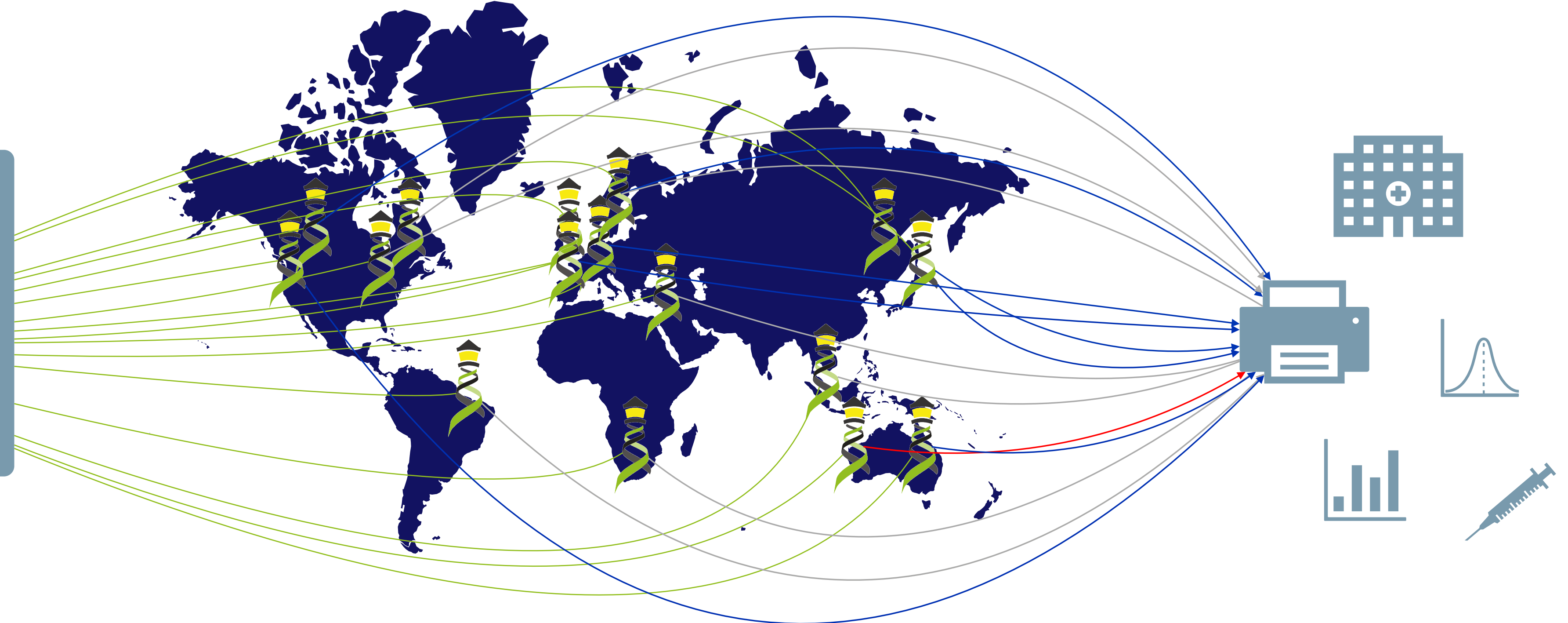
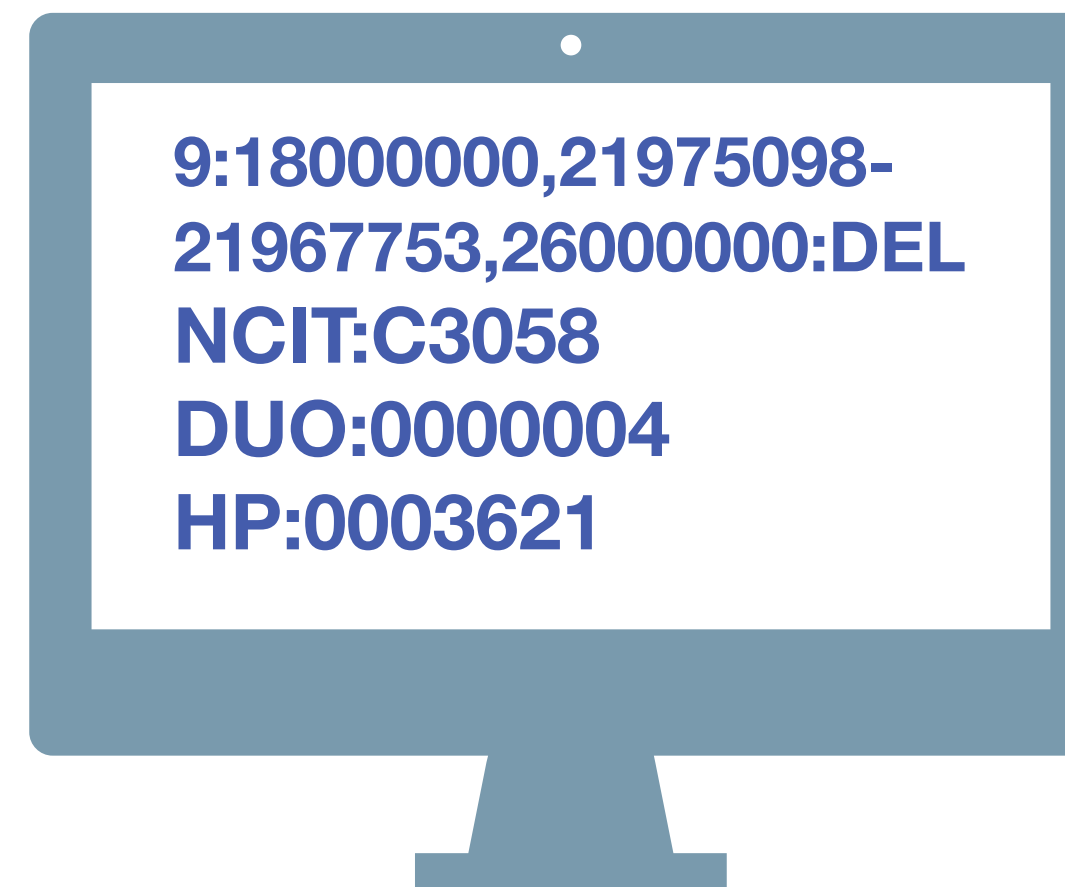
No packages published [Publish your first package](#)

Contributors 4

mbaudis Michael Baudis
sofiapfund Sofia
qingyao
KyleGao Bo Gao

Languages

Python 99.9% Shell 0.1%

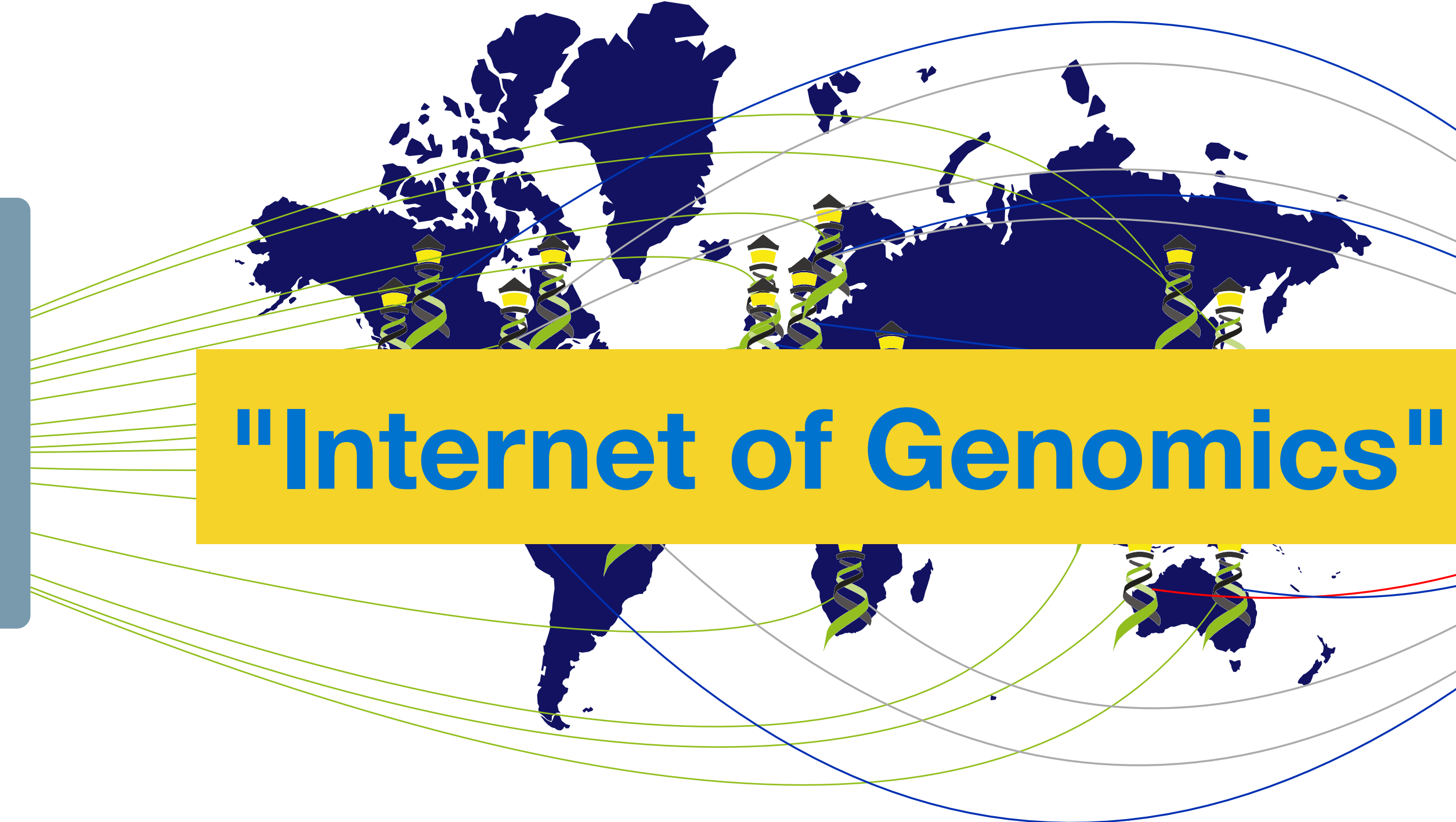
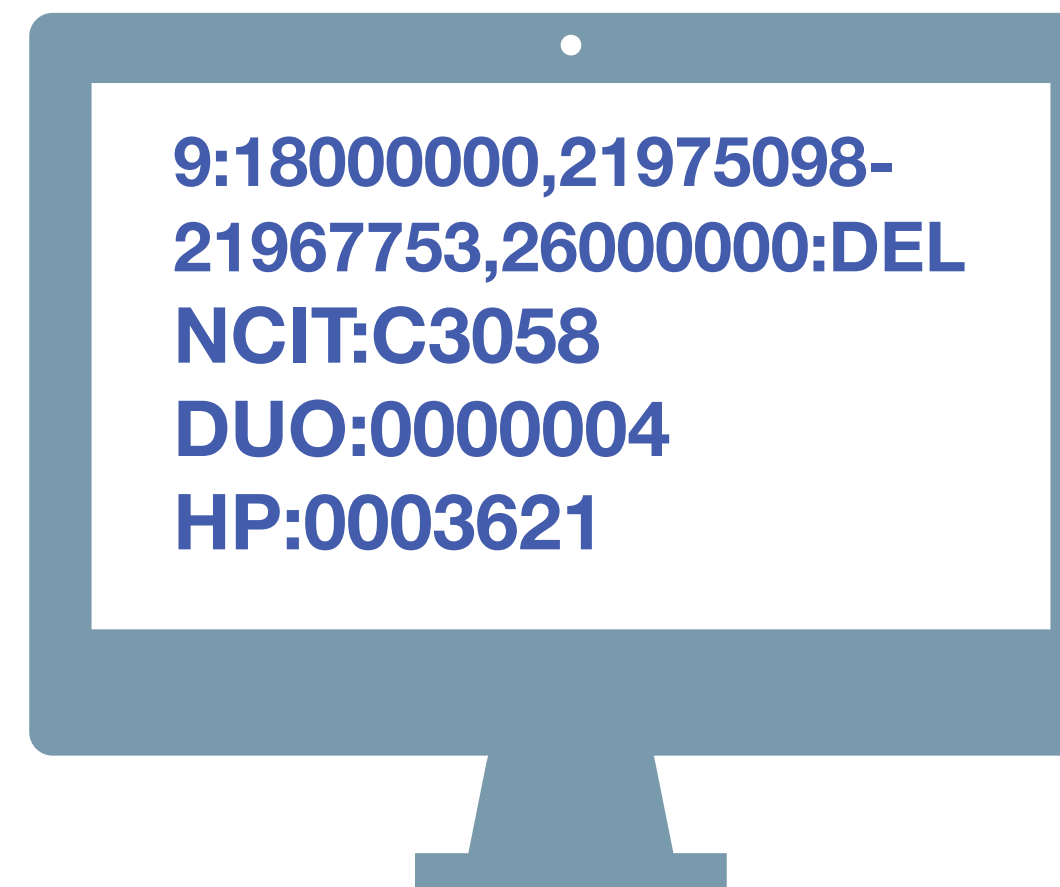


Have you seen deletions in this region on chromosome 9 in Glioblastomas from a juvenile patient, in a dataset with unrestricted access?

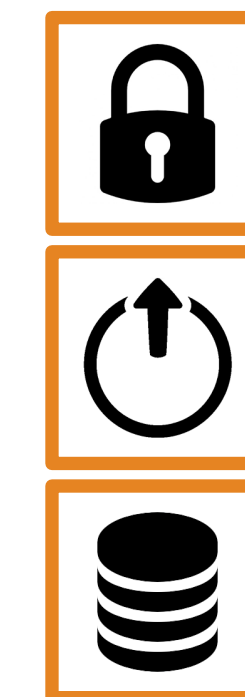


Beacon v2 API

The Beacon API v2 proposal opens the way for the design of a simple but powerful "genomics API".



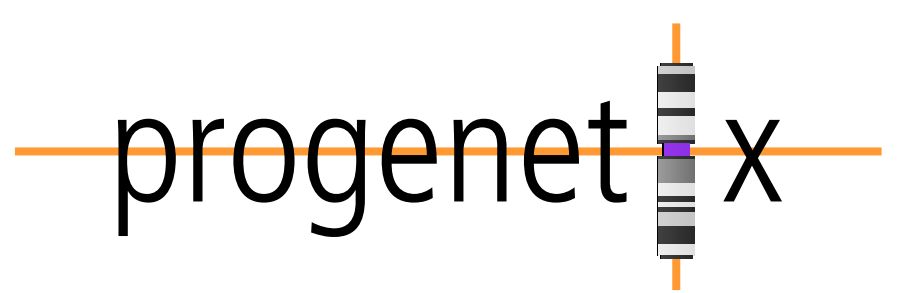
Have you seen deletions in this region on chromosome 9 in Glioblastomas from a juvenile patient, in a dataset with unrestricted access?



Beacon v2 API

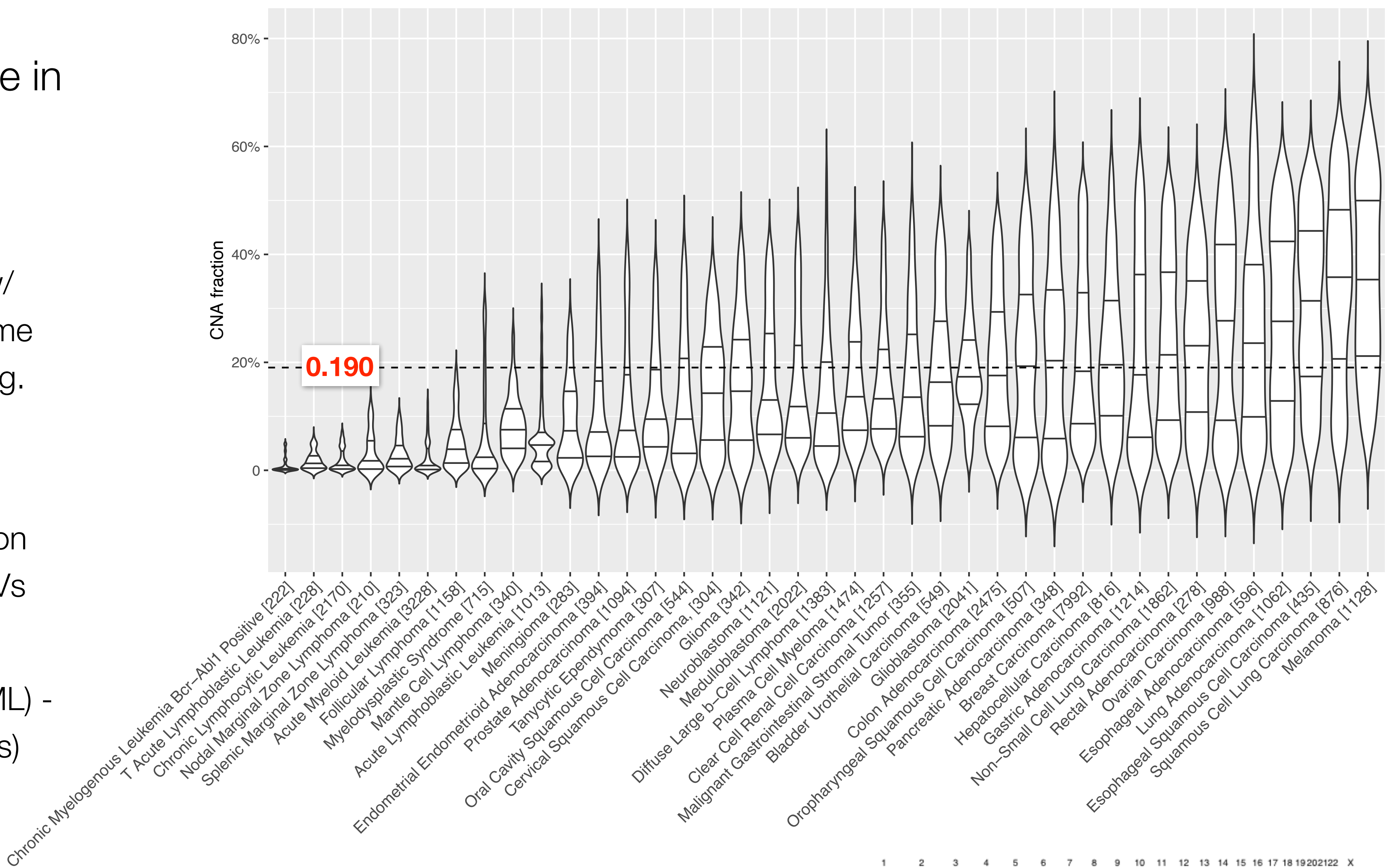
The Beacon API v2 proposal opens the way for the design of a simple but powerful "genomics API".

Progenetix Data Use Cases

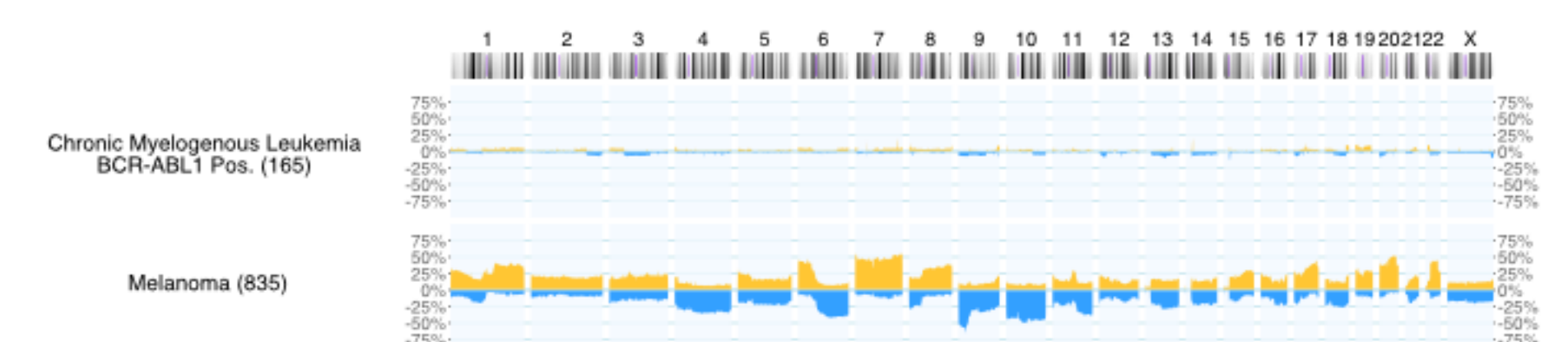


Genome CNV coverage in Cancer Classes

- 43654 out of 93640 CNV profiles; filtered for entities w/ >200 samples (removed some entities w/ high CNV rate, e.g. sarcoma subtypes)
- Single-sample CNV profiles were assessed for the fraction of the genome showing CNVs (relative gains, losses)
- range of medians 0.001 (CML) - 0.358 (malignant melanomas)



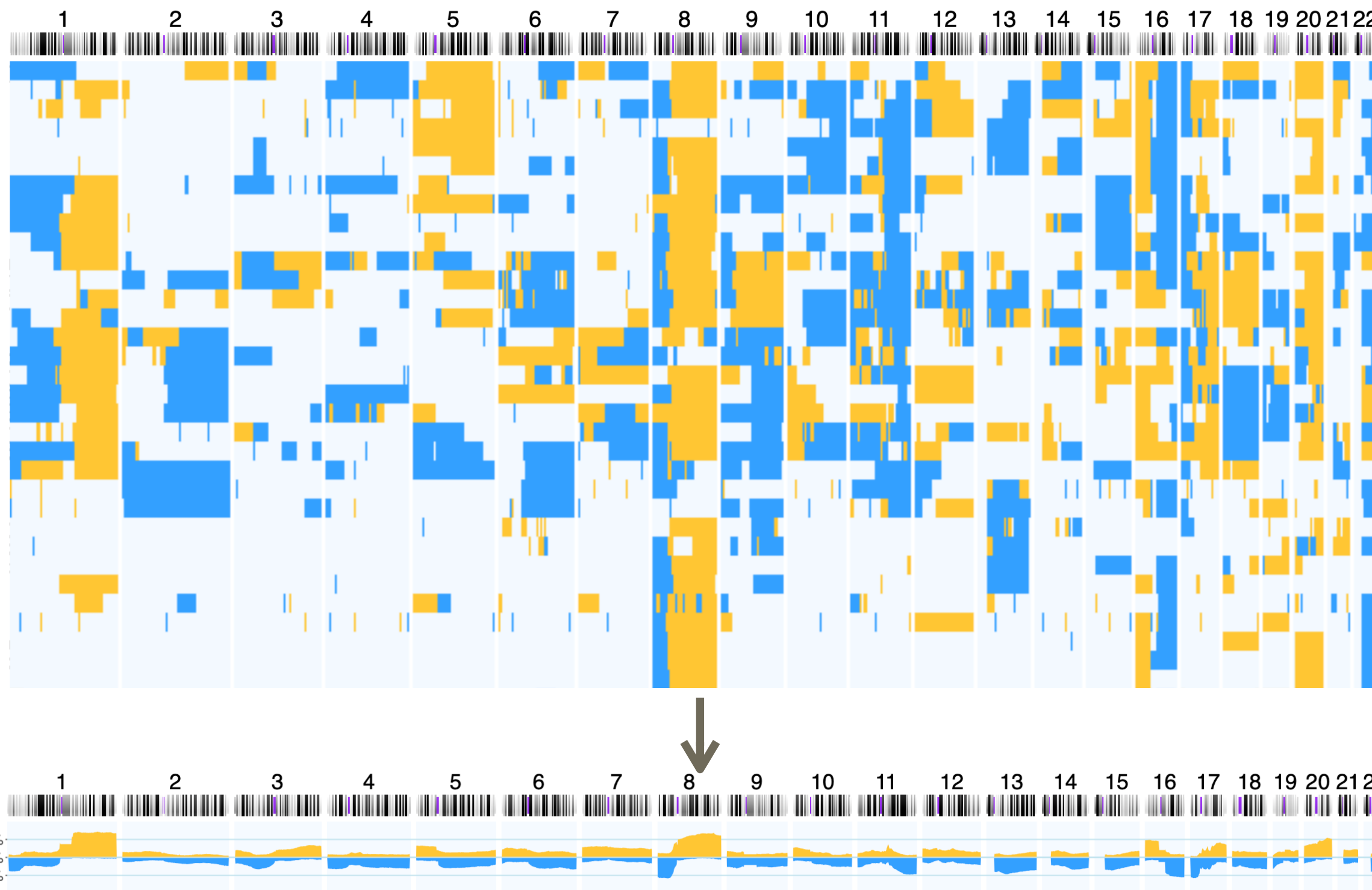
Lowest / Highest CNV fractions =>



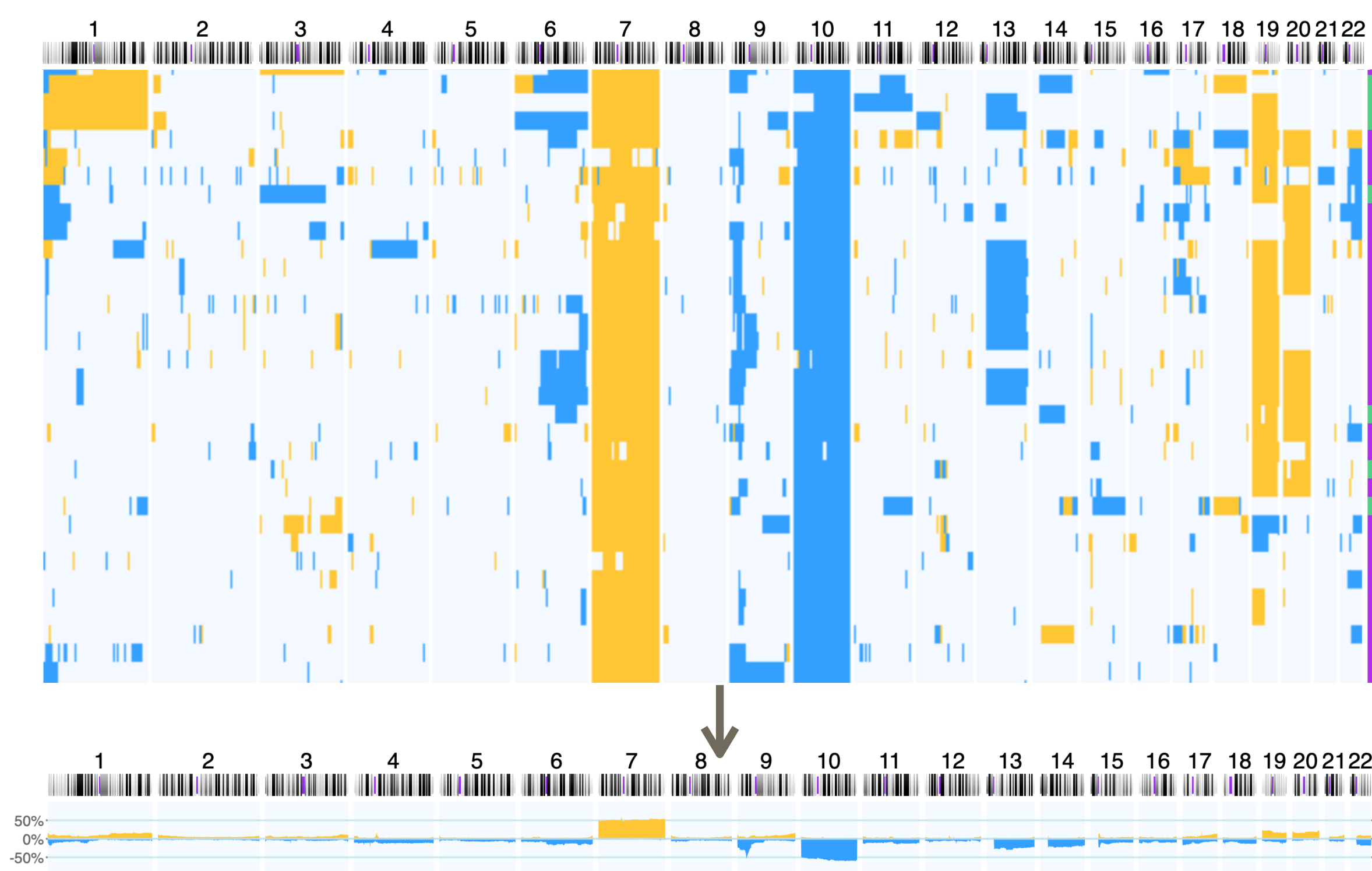
Drivers? Passengers? Markers?

Disentangling CNA Patterns

Ductal Breast Carcinoma



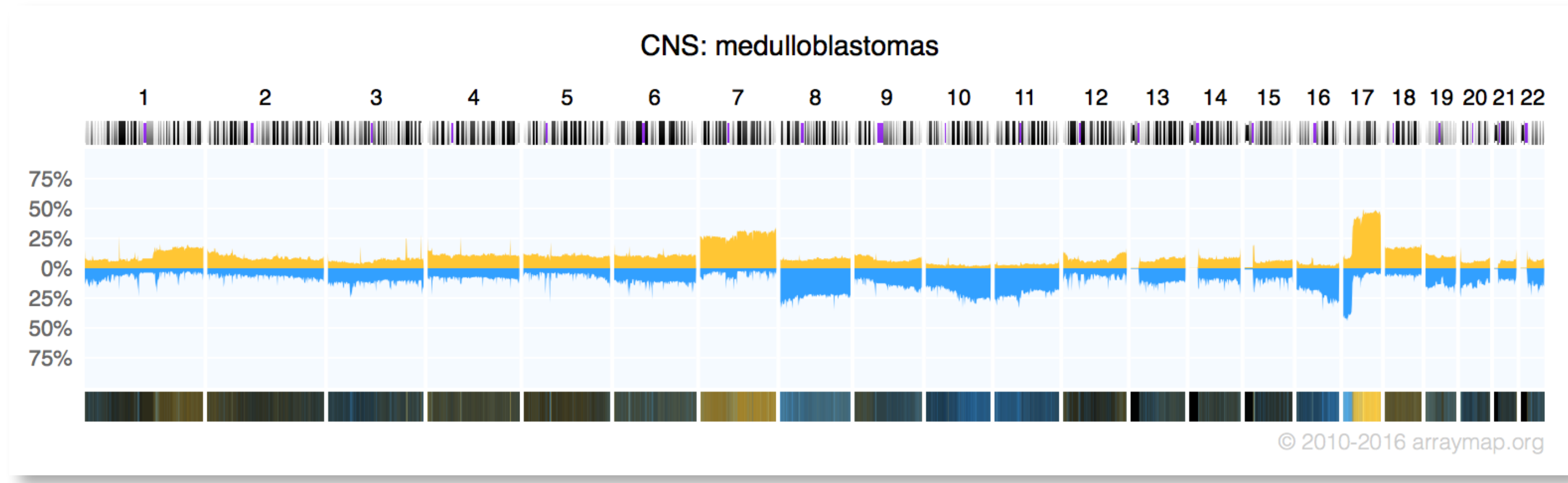
Glioblastoma



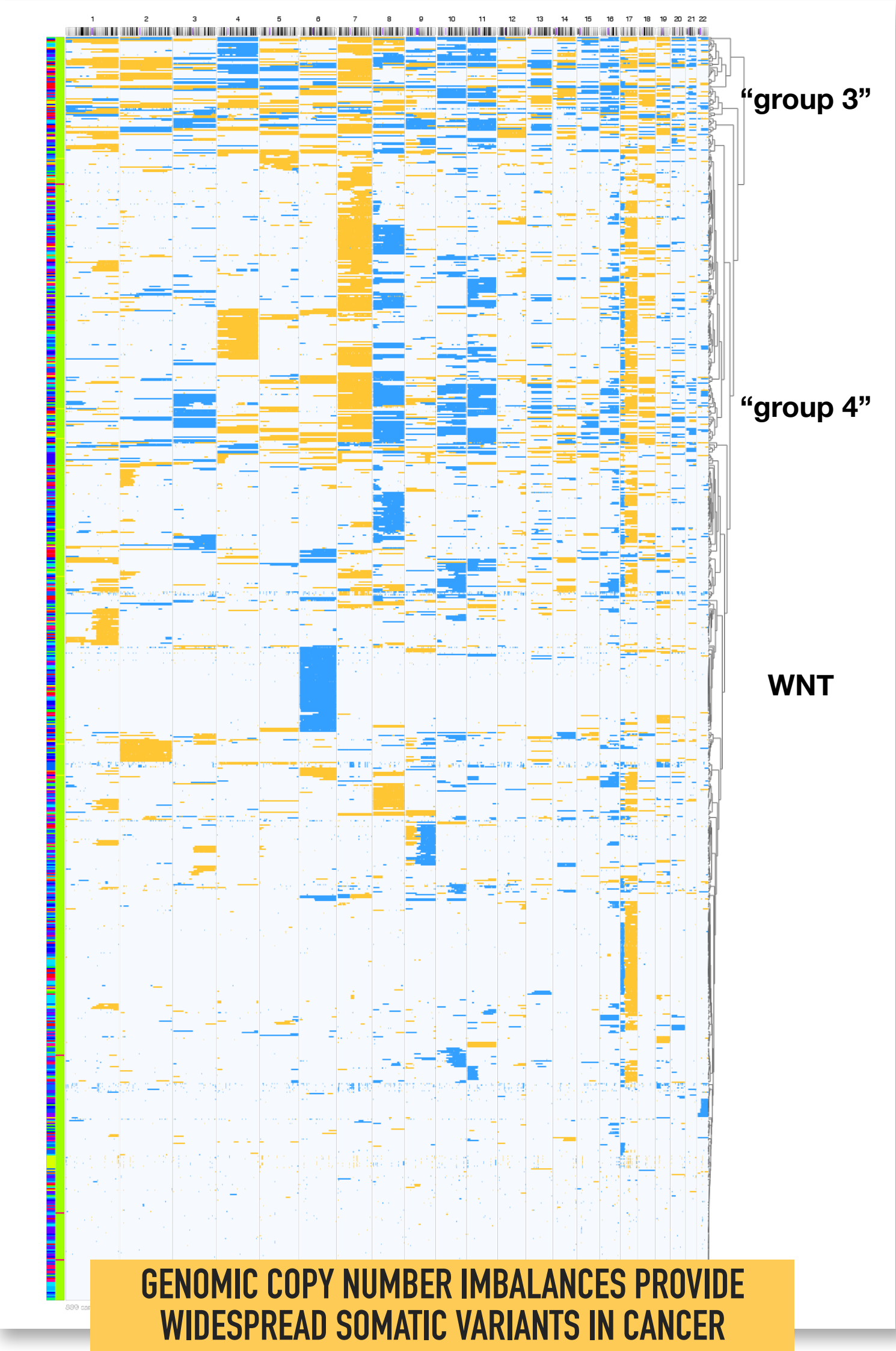
Somatic CNVs In Cancer

Recurrent mutation patterns

How can those patterns be used for classification and determination of biological mechanisms?



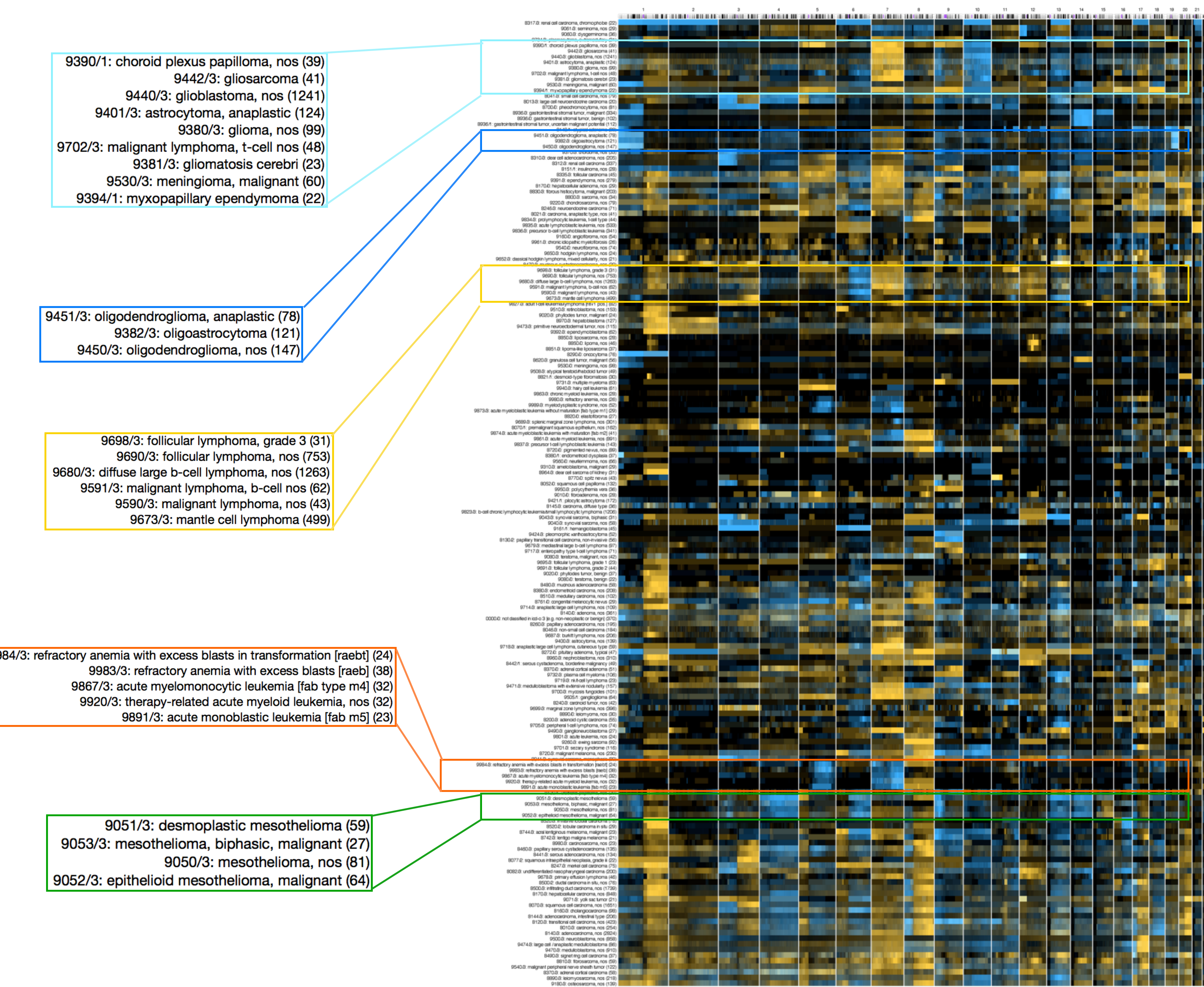
A genomic copy number histogram for malignant medulloblastomas, the most frequent type of pediatric brain tumors, displaying regions of genomic duplications and deletions. These can be decomposed into individual tumor profiles which segregate into several clusters of related mutation patterns with functional relevance and clinical correlation.



Somatic Mutations In Cancer: Patterns

Making the case for genomic classifications

Some related cancer entities show similar copy number profiles



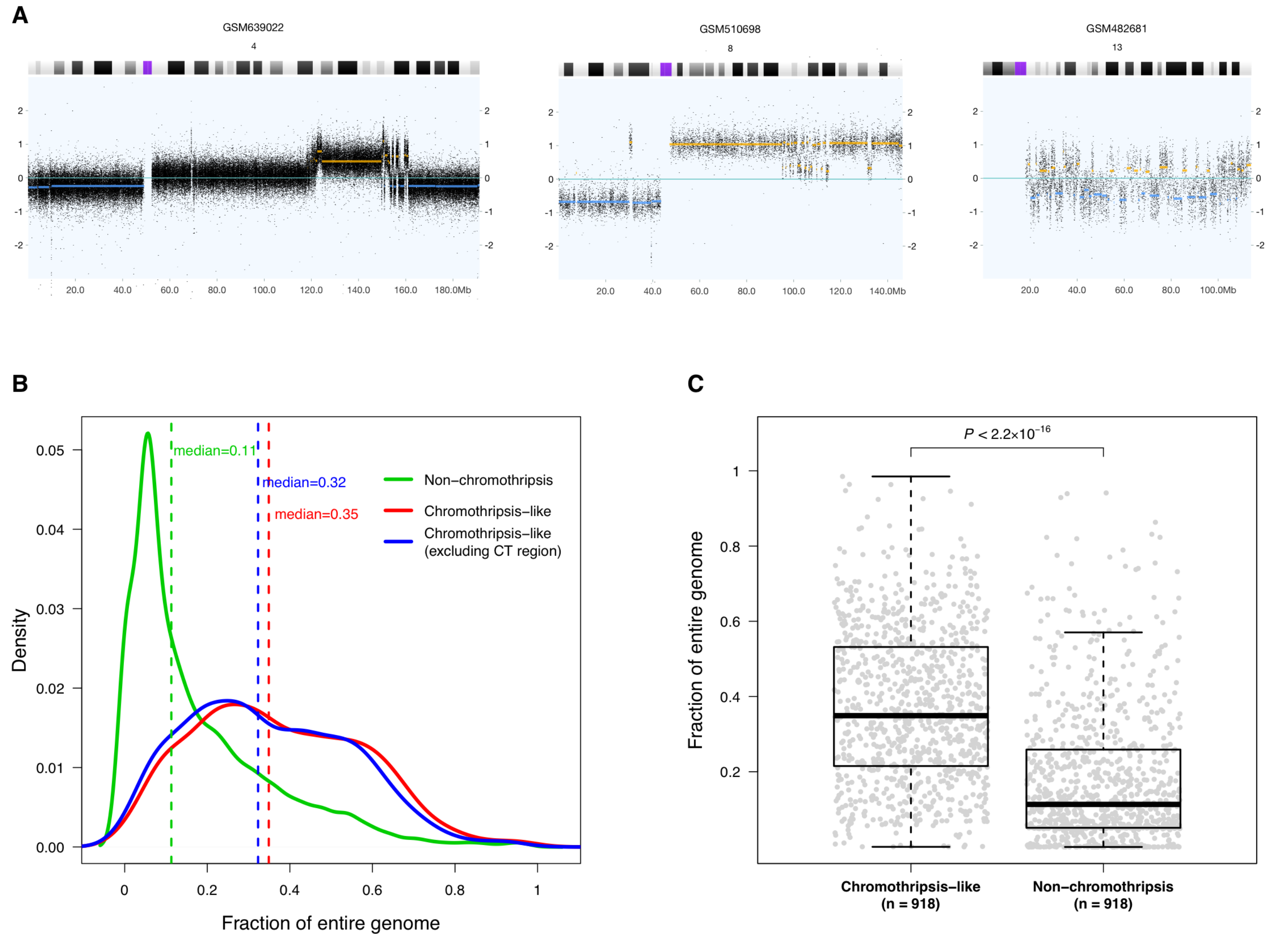
RESEARCH ARTICLE

Open Access

Chromothripsy-like patterns (CTLP)

Chromothripsy-like patterns are recurring but heterogeneously distributed features in a survey of 22,347 cancer genome screens

Haoyang Cai^{1,2}, Nitin Kumar^{1,2}, Homayoun C Bagheri³, Christian von Mering^{1,2}, Mark D Robinson^{1,2*} and Michael Baudis^{1,2*}



Input

22347 arrays
(18458 cases)

Output

1566 chromosomes
(1028 arrays)

Population stratification in cancer samples based on SNP array data

- Despite extensive somatic mutations of cancer profiling data, consistency between germline and cancer samples reached 97% and 92% for 5 and 26 populations
- Comparison of our benchmarked results with self-reported meta-data estimated a matching rate between 88 % to 92%.
- Ethnicity labels indicated in meta-data are vague compared to the standardized output from our tool

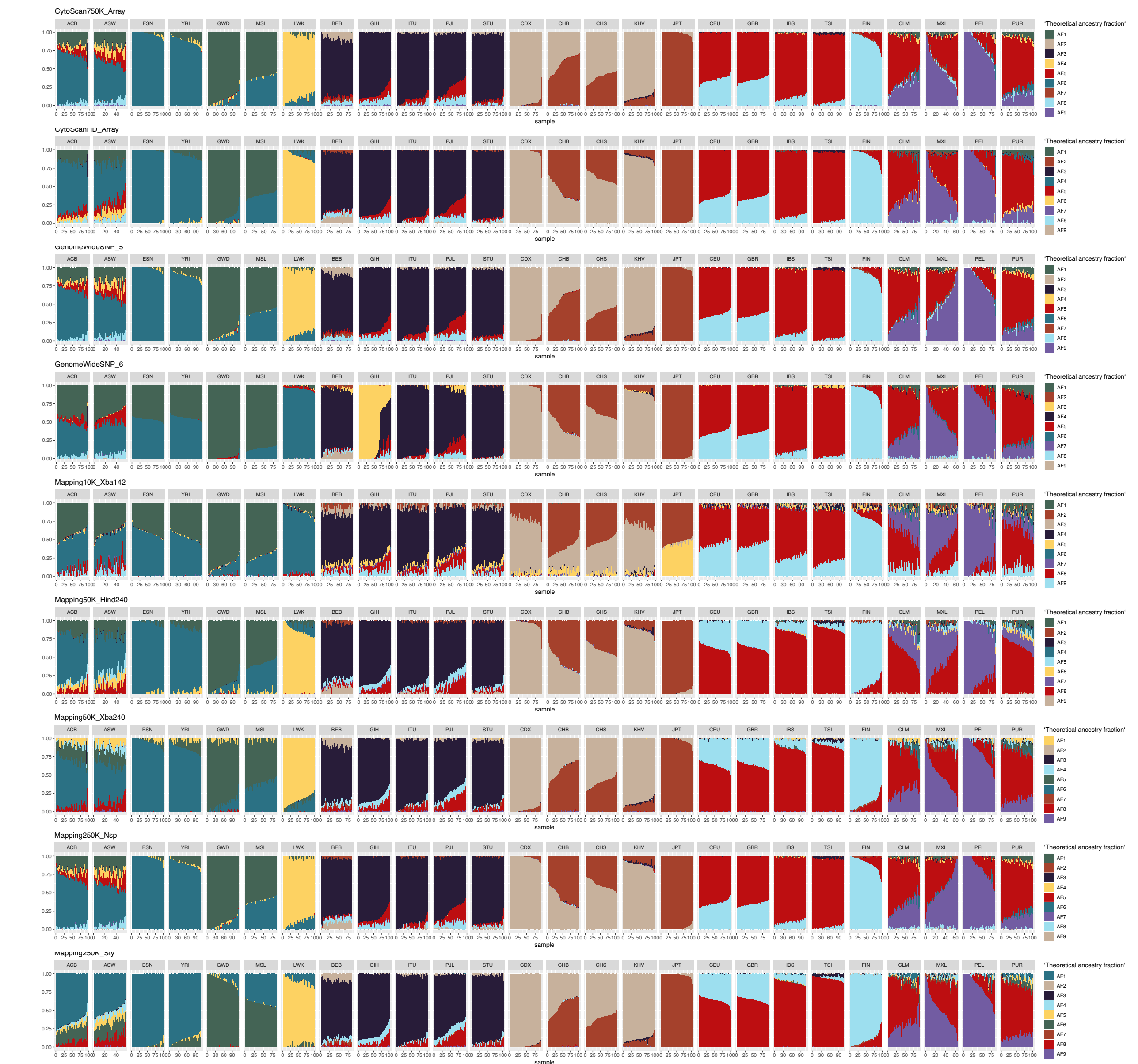
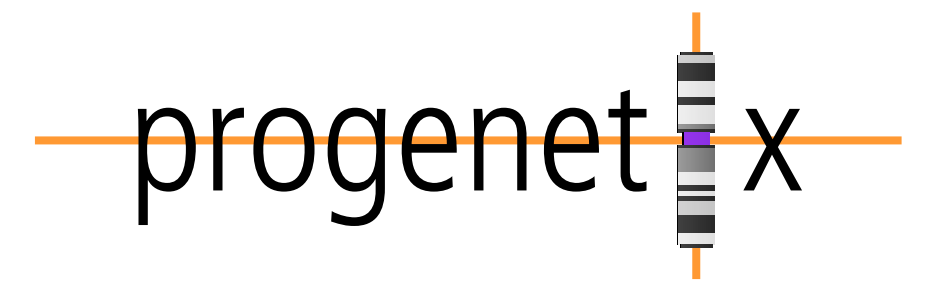


Figure S1 The fraction or contribution of theoretical ancestors ($k=9$) in reference individuals from 1000 Genomes Project with regard to nine SNP array platforms. The x-axis are individual samples, grouped by their respective population. Groups belonging to the same continent/superpopulation are placed neighboring to each other: AFR (1-7), SAS (8-12), EAS (13-17), EUR (18-22), AMR (23-26).

Progenetix Contributes to ELIXIR hCNV



hCNV Implementation Studies 2021-2023 No. 2



Beacon and beyond – Implementation-driven standards and protocols for CNV discovery and data exchange

- reinforce work on priority areas established in the current hCNV Implementation Study
- extend collaborations with the Rare Diseases and Galaxy Communities, EJP-RD and GA4GH
- Expected outcomes
 - ▶ shared CNV resources testing advanced versions of the Beacon protocol
 - ▶ integration of GA4GH standards such as Phenopackets in such resources
 - ▶ tools for data ingestion and export for standard formats (e.g. VCF, Phenopackets) and CNV-specific improvements of such standards
 - ▶ ELIXIR AAI demo on clinical and research hCNV resources
 - ▶ demonstration of Galaxy pipeline adoption for real-world hCNV data analysis projects
- connecting to international partners, e.g. Cancer Genomics Consortium (U.S.)

- ▶ WP1 - hCNV community reference resources
- ▶ WP2 - hCNV Resources and Beacon
- ▶ WP3 - Galaxy Community Intersection and Data Exchange
- ▶ WP4 - Workflows and Tools for hCNV Data Exchange Procedures
- ▶ WP5 - Training and dissemination



hCNV Implementation Studies 2021-2023 No. 2



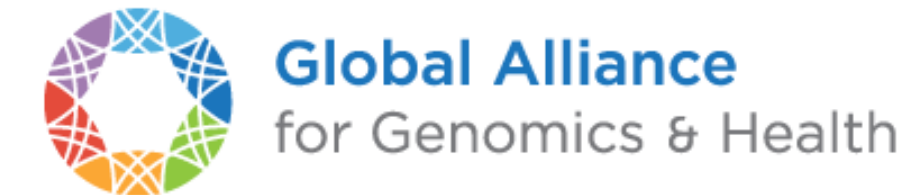
Beacon and beyond – Implementation-driven standards and protocols for CNV discovery and data exchange

The screenshot shows a web interface for the Progenetix database. At the top, it says "Cancer genome data @ progenetix.org". Below that, a text block states: "The Progenetix database provides an overview of mutation data in cancer, with a focus on copy number abnormalities (CNV / CNA), for all types of human malignancies. The data is based on *individual sample data* from currently 139448 samples." A chart titled "Sezary syndrome (icdom-97013)" displays CNV frequencies across 22 chromosomes for 166 samples. The chart has a y-axis ranging from -75 to 75. Below the chart, there are links to "Download SVG", "Go to icdom-97013", and "Download CNV Frequencies". A note below the chart says: "Example for aggregated CNV data in 166 samples in Sezary syndrome. Here the frequency of regional **copy number gains** and **losses** are displayed for all 22 autosomes." At the bottom, there is a dark header with "BANCCO" and navigation links for "Accueil", "Statistique", "Contact", and "Inscription". A login form with fields for "Email" and "Mot de passe" is shown. Below the header, a call-to-action button says "Importer vos données et partager les avec les partenaires du réseau" and "BANCCO - Banque Nationale de CNV Constitutionnelles".

"Galaxify", "Beaconize" & "Phenopack"
Progenetix & RD-CNVdb prototypes



- ▶ WP1 - hCNV community reference resources
- ▶ WP2 - hCNV Resources and Beacon
- ▶ WP3 - Galaxy Community Intersection and Data Exchange
- ▶ WP4 - Workflows and Tools for hCNV Data Exchange Procedures
- ▶ WP5 - Training and dissemination





GA4GH Genome Beacons

A Driver Project of the Global Alliance for Genomics and Health GA4GH and supported through ELIXIR

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[Specification & Roadmap](#)

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[beacon-network.org](#)

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[GA4GH::SchemaBlocks](#)

[GA4GH::Discovery](#)

[Github Projects](#)

[Beacon API and Tools](#)

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[website](#)

Beacon Protocol for Genomic Data Sharing

Beacons provide discovery services for genomic data using the Beacon API developed by the [Global Alliance for Genomics and Health \(GA4GH\)](#). The [Beacon protocol](#) itself standard for genomics data discovery. It provides a framework for public web service against genomic data collections, for instance from population based or disease specific repositories.



Baudisgroup @ UZH

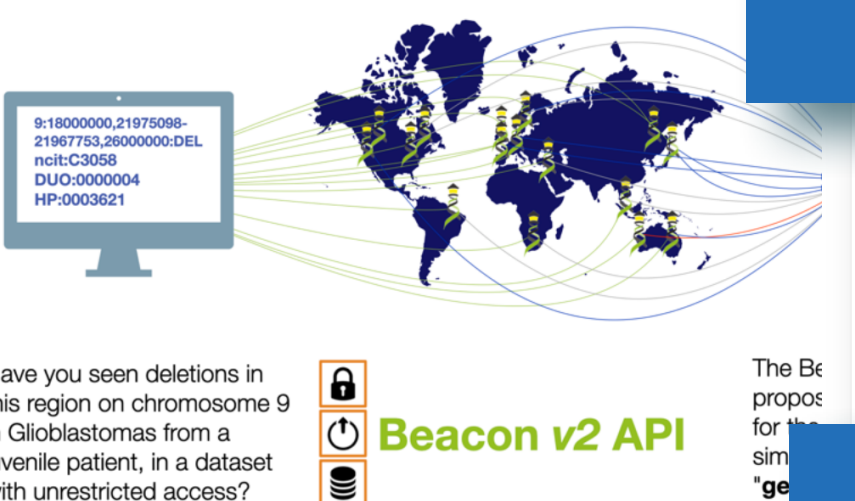
Michael Baudis
(Paula Carrio Cordo)
(Bo Gao)
Qingyao Huang
Sofia Pfund
Rahel Paloots
Hangjia Zhao

Pierre-Henri Toussaint

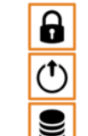
The original Beacon protocol had been designed to be:

- **Simple:** focus on robustness and easy implementation
 - **Federated:** maintained by individual organizations and assembled into networks
 - **General-purpose:** used to report on any variant collection
 - **Aggregative:** provide a boolean (or quantitative) answer about the observed variants
 - **Privacy protecting:** queries do not return information about single individuals
- Sites offering *beacons* can scale through aggregation *Beacon Networks*, which aggregate queries among a potentially large number of international *beacons* and assemble them into a single response. Since 2015 the development of the Beacon protocol has been led by [ELIXIR](#) in close collaboration with international participants. Recent versions of the *Beacon* protocol have expanded its scope:
- providing a framework for other types of genome variation data (i.e. rare variants)
 - allowing for data delivery using *handover* protocol, e.g. to link with clinical environments and allow for data delivery and visualisation services

Beacon v2 - Towards Flexible Use and Clinical Applications



Have you seen deletions in this region on chromosome 9 in Glioblastomas from a juvenile patient, in a dataset with unrestricted access?



Beacon v2 API

beacon-project.io

Reference name	(Structural) Variant
9	DEL (Deletion)
Start or Position	End (Range or Start)
21000001-21975098	21967753-23000000

{S}[B] and GA4GH
Melanie Courtot
Helen Parkinson
many more ...



progenetix.org

Beacon API Leads

Jordi Rambla
Anthony Brooks
Juha Törnroos

Discovery WS

Michael Baudis (Beacon)
Marc Fiume (Networks)

ELIXIR

Gary Saunders
David Lloyd
Serena Scollen
Dylan Spalding

Beacon Team CRG

Laureen Fromont
Babita Singh
Sabela de la Torre Pernas

...

Beacon v2 Scouts

Tim Beck
Joaquin Dopazo
Veronique Geoffroy
Jean Muller
David Salgado
Alex Wagner

...

github.com/ga4gh-beacon/





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Zurich^{UZH}



Global Alliance
for Genomics & Health



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beacon-project.io
schemablocks.org