



Genome analysis

Coral: a web-based visual analysis tool for creating and characterizing cohorts

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Abstract

Summary: A main task in computational cancer analysis is the identification of patient subgroups (i.e., cohorts) based on metadata attributes (patient stratification) or genomic markers of response (biomarkers). Coral is a web-based cohort analysis tool that is designed to support this task: Users can interactively create and refine cohorts, which can then be compared, characterized, and inspected down to the level of single items. We visualize the evolution of cohorts and also provide intuitive access to prevalence information. Coral can be utilized to explore any type of cohort and sample set. Here, we focus on the analysis of genomic data from cancer patients and therefore included in the public version data from the AACR Project GENIE, The Cancer Genome Atlas, and the Cell Line Encyclopedia.

Availability and Implementation: Coral is publicly available at <https://coral.caleydoapp.org>. The source code is released at <https://github.com/Caleydo/coral>.

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Supplementary information: Supplementary data are available at *Bioinformatics* online.

1 Introduction

A frequent goal in cancer research is the characterization of patient or sample cohorts based on experimental and metadata. A common approach is to define multiple cohorts based on shared characteristics, for instance, by splitting the samples based on their response to a treatment. One then aims to identify shared properties, find differences, and explore cohort characteristics in detail. Interactive visualization tools can assist users in creating, refining, and comparing cohorts based on attributes from high-dimensional space. Current tools, however, do not allow comparison or refinement of more than two cohorts, compare attributes only individually, or do not show the relationships between the cohorts created.

To address this deficiency, we present *Coral*, a web-based tool for interactively creating and characterizing multiple cohorts (Fig. 1). Coral allows users to keep track of the cohort evolution by showing which attributes have been used to create them and how the cohorts are related to each other. Additionally, analysts can use tailored visualizations to characterize the cohorts and explore them in detail. Coral's database contains mutation data from the AACR Project GENIE (AACR Project GENIE Consortium, 2017), with mRNA expression, DNA copy number, and mutation data from The Cancer Genome Atlas (TCGA;

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<https://cancergenome.nih.gov>) and the Cell Line Encyclopedia (CCLE; <https://portals.broadinstitute.org/ccle>), and two depletion screen datasets. Data pre-processing steps are described in the Supplementary Notes.

We demonstrate the utility of our tool by reproducing findings from a recently published article about *KRAS*^{G12C} somatic mutations (Nassar *et al.*, 2021) and using a case study exploring *KRAS* mutations in TCGA data (see Supplementary Notes, Figs. S3–S17, and Video).

2 Software Description

Coral is a client-server web-application that utilizes a Postgres database and is implemented in TypeScript and Python. The Coral interface follows an overview+detail approach in which two components are shown in a horizontal split-screen setup.

The **Cohort Evolution View** (upper panel) presents the evolution of all cohorts as a graph. Operations and cohorts are encoded as nodes connected by edges to represent the analysis flow (see Fig. 1A). The first cohort includes all items of the loaded dataset and is created automatically (Fig. 1B). When the user selects a cohort in the graph, it is assigned a color that is used consistently in all visualizations. Selected cohorts are loaded into the **Action View** (lower panel), which allows users

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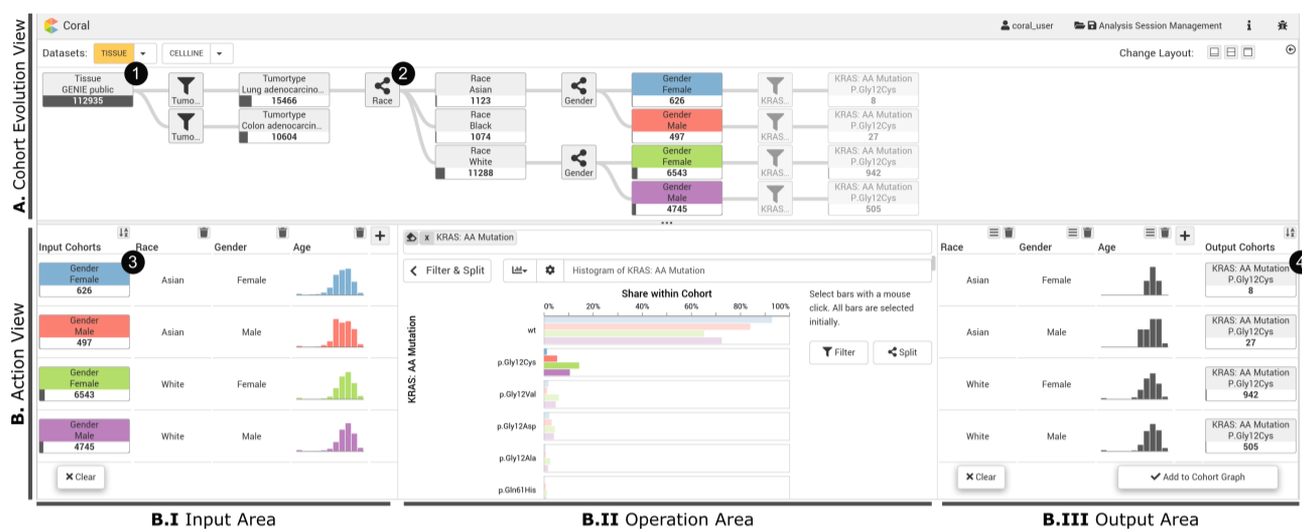


Fig. 1: Using Coral to reproduce $KRAS^{G12C}$ mutation findings by Nassar *et al.* (2021). The *Cohort Evolution View* (A) shows that the user starts by filtering the AACR Project GENIE data to include only patients with either non-small cell lung cancer (NSCLC) or colorectal cancer. The user then splits the NSCLC cohort by race, and subsequently the resulting cohorts *Asian* and *White* by gender. The resulting cohorts are selected and highlighted (■, ■, ■, ■) across all views. The *Input Area* (B.I) shows the cohorts' distributions for race, gender, and age. Using *View, Filter & Split* in the *Operation Area* (B.II), the user subsequently assesses the four cohorts with respect to their frequencies of various $KRAS$ mutations and selects the samples that contain $KRAS^{G12C}$. By applying the *Filter* operation, the user generates the corresponding four output cohorts displayed in the *Output Area* (B.III).

to perform cohort characterization and creation operations. New cohorts created by these operations are also added to the graph, which results in a circular workflow within the tool (see Supplementary Notes, Fig. S1). The Action View is divided into three areas: the *Input Area*, the *Operation Area*, and the optional *Output Area*, which is shown if the operation results in new cohorts (Fig. 1B). The *Operation Area* provides the following operations that can be applied to the input cohorts.

View. In addition to the visualizations in the *Input Area* (Fig. 1B.I), which provide context to the input cohorts, the *View* operation is the main route to exploring the dataset and investigating how the values of one or more attributes are distributed across cohorts. The visualization is based on the number and type of attributes, as explained in Supplementary Table S1.

Filter & Split. The *Filter & Split* operation is used to create cohorts from the loaded dataset. Fig. 1B shows four input cohorts being filtered using the categorical attribute *KRAS mutation status*. Users can adjust the selection, as illustrated in Fig. 1B.II, where, for each input cohort, the user selected all samples with a $KRAS^{G12C}$ mutation. In contrast, the *Split* operation can be used to divide a cohort into multiple sub-cohorts, as shown in Fig. 1B.II, where the NSCLC cohort is split based on *Race*.

Compare. Additionally to comparing cohorts using the visualizations in the *Input Area* and the *View* operation, Coral offers a *Compare* operation to test for statistically significant differences. Supplementary Figs. S6–S7 show that the NSCLC and the colorectal cancer cohorts in Fig. 1 differ significantly in their age and gender distributions. The compare operation integrates the TourDino view (Eckelt *et al.*, 2019) to confirm visual patterns. Cohorts are tested pairwise for differences. The test results are summarized in a matrix showing each p-value. The applied test, its score, and a visualization are shown on demand.

Prevalence. Prevalence is the proportion of items with a certain characteristic in a cohort. In Fig. 1, the proportion of patients with a $KRAS^{G12C}$ mutation (*sample cohort* ④) among female Asian patients with NSCLC (*reference cohort* ③) is 8/626, which equals a prevalence of about 1.3%. Coral provides a dedicated analysis view to assess prevalence estimates (see Supplementary Fig. S13). After selecting the *sample cohort* with items that have the characteristic of interest, the user can flexibly define the *reference cohort* by applying or skipping *Filter &*

Split operations used to create the *sample cohort*. The cohorts' sizes and the resulting prevalences are then displayed in a bar chart.

Inspect Items. To identify outliers or to assess single data points, users need to move from the aggregated level down to the level of individual items. We use Taggle (Furmanova *et al.*, 2020), a tabular visualization technique in which the items of the cohorts are displayed. Users can select attributes, whose data is displayed in the table, and sort, filter, and group the data (see Supplementary Fig. S16).

3 Conclusion

Coral is an analysis tool for interactively creating cohorts based on a range of attributes, and visually represents the evolution of cohorts in an overview graph. The cohorts can be characterized, either by size or by a rich set of experimental and metadata. Users are able to gain insights into the cohort prevalences and also to drill down to the level of single-item information. Technically and conceptually, Coral can be applied to fields beyond cancer genomics. It can cater to the needs of a broad range of users, including bioinformaticians, biologists, and researchers in other disciplines.

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