

# General

All the configuration settings are in [config.js](#) and tools.js file.

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## config.js

### CellBase Configuration

The host and version of CellBase are configured in the section below

#### CellBase

```
var cellbase = {
  hosts: ["bioinfodev.hpc.cam.ac.uk/cellbase-4.5.0-beta"],
  version: "v4"
};
```

### OpenCGA Configuration

The below section in [config.js](#) allows to do the following:

- OpenCGA Host and version can be configured.
- There is a section called projects which allows to browse public projects without any login.
- Cookies section is useful when the application expects users to login and their cookie names that store the actual user name and session ID can be configured here.

#### OpenCGA

```
var opencga = {
  host: "bioinfodev.hpc.cam.ac.uk/opencga-1.0.0-rc3",
  version: "v1",
  projects: [
    {
      name: "Project A",
      alias: "proj_a",
      studies : [
        {
          name: "Study 1",
          alias: "std_1"
        }
      ]
    }
  ],
  cookies: {
    userName: "iva_userId",
    sessionId: "iva_sid"
  }
};
```

### Visual Settings

Consequence Type, Population frequency and Protein Substitution Scores which can be found in [config.js](#) are enabled to have colors in web application. They can be modified to your needs.

## Consequence Type

```
var consequenceTypes = {
    // This is the impact color. It allows to customise both the impact
    categories and desired colors
    color: {
        high: "red",
        moderate: "orange",
        low: "blue",
        modifier: "green"
    },
    /* 'Title' is optional. if there is no title provided then 'name' is
    going to be used.
    There are two more optional properties - 'checked' and 'color'. They
    can be set to display them default in web application.
    Similarly 'description' is optional as well.
    */
    categories: [
        {
            id: "",
            name: "",
            title: "Intergenic",
            description: "",
            terms: [
                {
                    id: "SO:0001631",
                    name: "upstream_gene_variant",
                    title: "upstream gene variant",
                    description: "A sequence variant located 5' of a gene",
                    impact: "modifier"
                },
                .
                .
                .
                .
            ]
        },
        .
        .
        .
    ]
};
```

### Population Frequency

```
var populationFrequencies = {
  color: {
    veryRare: "red",
    rare: "yellow",
    average: "orange",
    common: "blue"
  },
  studies: [
    {
      id: "1kG_phase3",
      title: "1000 Genomes",
      populations: [
        {
          id: "ALL",
          title: "All populations [ALL]",
          active: true
        },
        {
          id: "EUR",
          title: "European [EUR]"
        },
        .
        .
        .
        .
        .
      ]
    },
    .
    .
    .
  ],
};

};
```

### Protein Substitution Score

```
var proteinSubstitutionScores = {
  // This is to show the predictions in respective colors
  sift: {
    deleterious: "red",
    tolerated: "green"
  },
  polyphen: {
    probablyDamaging: "red",
    possiblyDamaging: "orange",
    benign: "green",
    unknown: "black"
  }
};
```

*tools.js*