This document serves as a quick-start guide and reference card for installing and using pong. Please refer to the software manual, which can be downloaded from the Ramachandran Lab website, for comprehensive information.

## Installing pong

pong has been tested on Mac OS X (10.8–10.11), Linux (Ubuntu 15.04, Linux Mint 17.2), and Windows 7.

pong is hosted on PyPI and can thus be easily installed with pip. In order to run pong, you need Python 2.7 and a modern web browser (e.g. Chrome, Firefox, Safari). pong is not compatible with Internet Explorer.

To install pong, run:

```bash
pip install pong
```

### Linux-specific tips

On some Linux systems, the install command may fail due to a "permission denied" error. In that case, try running `sudo pip install pong` instead. You will be prompted to enter in your computer login password. If that doesn't work either, try running the command as the super user:

```bash
su # you will be prompted to enter an administrator password
pip install pong
exit
```

### Mac-specific tips

Note that the Apple system default Python cannot run pong. Python 2.7 and pip can be installed manually, but we find that setup is easiest if you have Homebrew and Homebrew-installed Python. Using Homebrew, pong's dependencies, and subsequently pong itself, can be installed [ or updated ] as follows:

```bash
brew install python     [ brew upgrade python ]
pip install pip         [ pip install --upgrade pip ]
pip install pong        [ pip install --upgrade pong ]
```
Running pong

pong is executed through the command line. To see a complete list of options, run:

```
pong -h
```

**Note to Windows users:** On some Windows systems, pong may not be installed as an executable, and thus cannot be run using this command. Instead, replace `pong` with `python \path\to\pong`.

We have provided an example dataset on which you can run pong (download available at the Ramachandran Lab website). Using data from the 1000 Genomes Project Phase3 (2,426 individuals), we performed 8 runs of ADMIXTURE at each value of $K$ from $K=2$ to $K=8$. To analyze the example dataset with pong, navigate to the unzipped example directory and run:

```
pong -m pong_filemap -n pop_order_expandednames.txt -i ind2pop.txt
```

Information regarding pong’s application to the input data will be displayed to your terminal window. After its algorithms complete, pong initializes a web server on `localhost:4000` (you can change the port on which pong operates with the command line option `--port`). Once you navigate to `localhost:4000` on your web browser, pong will detect a new browser connection and begin rendering the visualization.

Running pong on your own data

pong’s setup was designed to be low-hassle. This section details the required and optional input files that pong handles.

**Clustering output**

pong accepts clustering output files, or $Q$ matrices, from a variety of clustering inference programs. The command line option `--c`, or `--ignore_cols`, tells pong how many leading columns of each row of the input $Q$ matrix files to skip before parsing individual membership coefficients. For example, use `--c 5` for some versions of STRUCTURE indiv output to ignore covariate metadata. In the case that no leading columns should be skipped (e.g. for ADMIXTURE output), it is not necessary to provide `--c 0` because the default value of `--c` is 0.

**A few related notes:**

- Don’t worry about trailing columns; they will not be parsed (after `--c` leading columns are ignored, only the first $K$ columns are used).
- pong parses $Q$ matrix files as whitespace-delimited by default. If, for some reason, this is not the
case with your input data, you can specify any column delimiter with the option `--col_delim`
(e.g. `--col_delim ','` for CSV files, or `--col_delim "\t"` for tab-delimited files).

- While pong does handle leading/trailing columns in Q matrices, it does not handle
  leading/trailing rows. These rows must be stripped prior to analysis with pong (i.e. for a dataset
  with $n$ samples, every Q matrix file should have $n$ lines).

**filemap file**

A bit of information about the input Q matrices must be provided, in the form of a filemap. A filemap is
a three-column, tab-delimited file. Each line contains the following information about a single Q
matrix:

1. A unique ID (e.g. `iter5_K=7`; this must contain at least one letter, and cannot contain a
   hashmark `&#35`. Integers, spaces, and other special characters are accepted.)
2. The $K$ value
3. The Q matrix file path, relative to the filemap. Thus, if the filemap is in the same directory as the
   Q matrix files, this is just the name of the Q matrix file.

It is important that the columns of your filemap are in the right order and that the file is tab-delimited.
Use the filemap provided with the example dataset for reference. Use the command line option `--m`,
or `--filemap`, to pass pong the path to your your filemap.

**Population labels (optional)**

**ind2pop data**

It is common for Q matrices to be annotated with population metadata. In many datasets, individuals
are assigned a population code/label/number; we refer to this as ind2pop data. If provided with this
information, pong’s visualization will sort individuals by population, partition populations with black
lines, and sort individuals within each population by their membership in the population’s major
cluster.* Other covariates, such as continent or language spoken, can also be used as ind2pop in place
of population metadata.

You can provide ind2pop data with the command line option `--i`, or `--ind2pop`. The argument to
this option can be either of the following:

- An integer, representing the Q matrix column number that contains ind2pop data. For example,
  use `--i 4` for standard STRUCTURE output.
- The path to an ind2pop file, where line $i$ of this file contains the ind2pop data (i.e. population
code/label/number) for the individual represented by line $i$ of the Q matrix files.

Note that the population codes/labels/numbers provided as ind2pop data should not contain any
whitespace.
Population order and detailed names

If you provide *ind2pop* data, you may provide an additional file specifying the desired order in which to display the populations (as the argument to the command line option `-n`). This file should have one population code/label/number per line; the top-to-bottom order of the file corresponds with the left-to-right display order of the populations.

You may want to more descriptive population names to be displayed (e.g. if the input *ind2pop* data are numbers, cryptic three-letter codes, etc.); in this case, you can add a second column (tab-delimited) to the population order file containing population names. Space characters are allowed in these names. See the file *pop_order_expandednames.txt* in the example dataset for reference.

Custom colors for visualization (optional)

*pong* provides default colors (unless $K_{max} > 26$), but it is possible to provide *pong* with a file containing a set of colors to use for visualization (using the command line option `-l`). This file must contain at least $K_{max}$ colors, with one color per line. Because *pong*’s visualization is web-based, colors can be provided in any format that is accepted by CSS (e.g. `#ff0000`, `rgb(255,0,0)`, `red`).

Additional resources

The example dataset, software manual, and other relevant materials can be found at the Ramachandran Lab website. For discussions and feedback on *pong*, refer to the *pong* Google Group.

Contact Aaron Behr or Sohini Ramachandran for more information.

*We have found that this sorting operation makes it much easier to understand the distribution of cluster membership within each population. Note that *pong* performs this sorting operation on a single $Q$ matrix (the bottom-most plot in the main visualization, which is the representative run of the major mode of the highest $K$-value) and propagates that order through all the other visualized $Q$ matrices, such that the ordering of individuals is consistent across all plots.*