

# ANGSD-wrapper: scripts to streamline and visualize NGS population genetics analysis

Arun Durvasula<sup>1</sup>, Tyler Kent<sup>1</sup>, Siddharth Bhadra-Lobo<sup>1</sup>, Jeffrey Ross-Ibarra<sup>2</sup>

<sup>1</sup>Dept. of Plant Sciences, University of California Davis
<sup>2</sup>Dept. of Plant Sciences, Center for Population Biology, and Genome Center, University of California Davis

#### Introduction

The advent of highly multiplexed sequencing has opened a number of exciting avenues for evolutionary biologists. One of the powerful approaches enabled by inexpensive sequencing is the ability to sequence a large number of individuals, each to relatively low sequencing depth. However, this approach also presents statistical challenges in the analysis of low coverage data. The software ANGSD [1] and related programs [2] were developed to deal with low coverage sequence data. Rather than call genotypes at variable sites, ANGSD performs a number of population genetic analyses on genotype likelihoods, including estimation of the population mutation rate  $\theta$ , the site frequency spectrum, neutrality tests, inbreeding coefficients, and population structure. ANGSD has already been used in several studies to analyze genome sequence data [3] [4]. However, ANGSD requires considerable familiarity with command line tools and remains inaccessible to many biologists that are not from a computational background. Here we present a software package that aids in the preparation of analyses for ANGSD and provides interactive graphing software implemented in R [5] and Shiny [6]. ANGSD-wrapper simplifies multistep analyses such as calculating Tajima's D into a single step. Users supply all the needed information in a single configuration file 1, and after ANGSD has finished calculations, ANGSD-wrapper provides interactive graphing of the results 2.

# Implementation

ANGSD-wrapper is implemented using bash scripts that call ANGSD methods and handle saving intermediate files between the initial data preparation and the final data analysis. Each overall method in ANGSD, such as calculating estimates of  $\theta$ , follows a specific order of program calls. Thus, we have abstracted away the running of each step and provided a set of default values for parameters and instead require the user to supply the data using a configuration file (Figure 1). The user can override the default values of the parameters in the configuration file as well.

Additionally, ANGSD-wrapper contains a powerful graphing application based on R and Shiny (Figure 2). After analysis is done by ANGSD, the user can load the resulting statistics into a web-based application hosted on the user's computer. This application allows the interactive plotting of values such as

estimators of  $\theta$  and Tajima's D as well as the ability to load gene annotations from Ensembl. These features make it easy and intuitive to analyze next generation sequencing data.

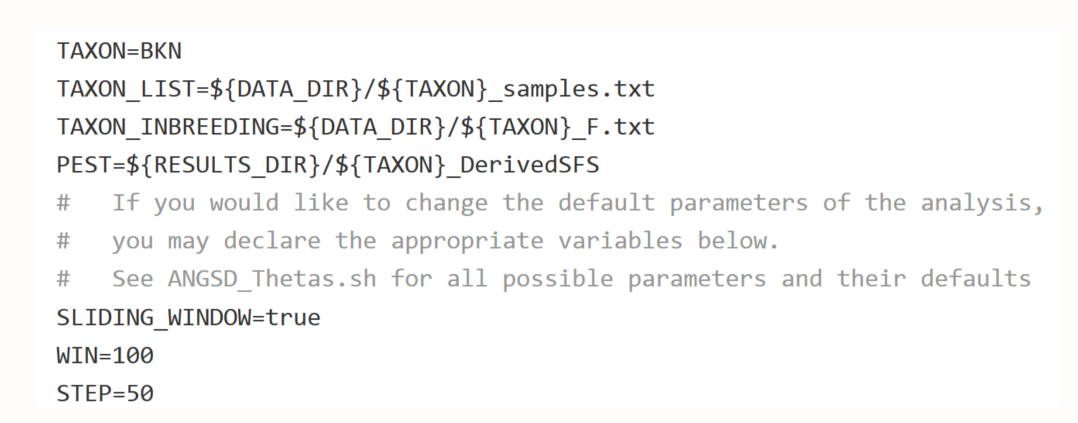


Figure 1: Example configuration file

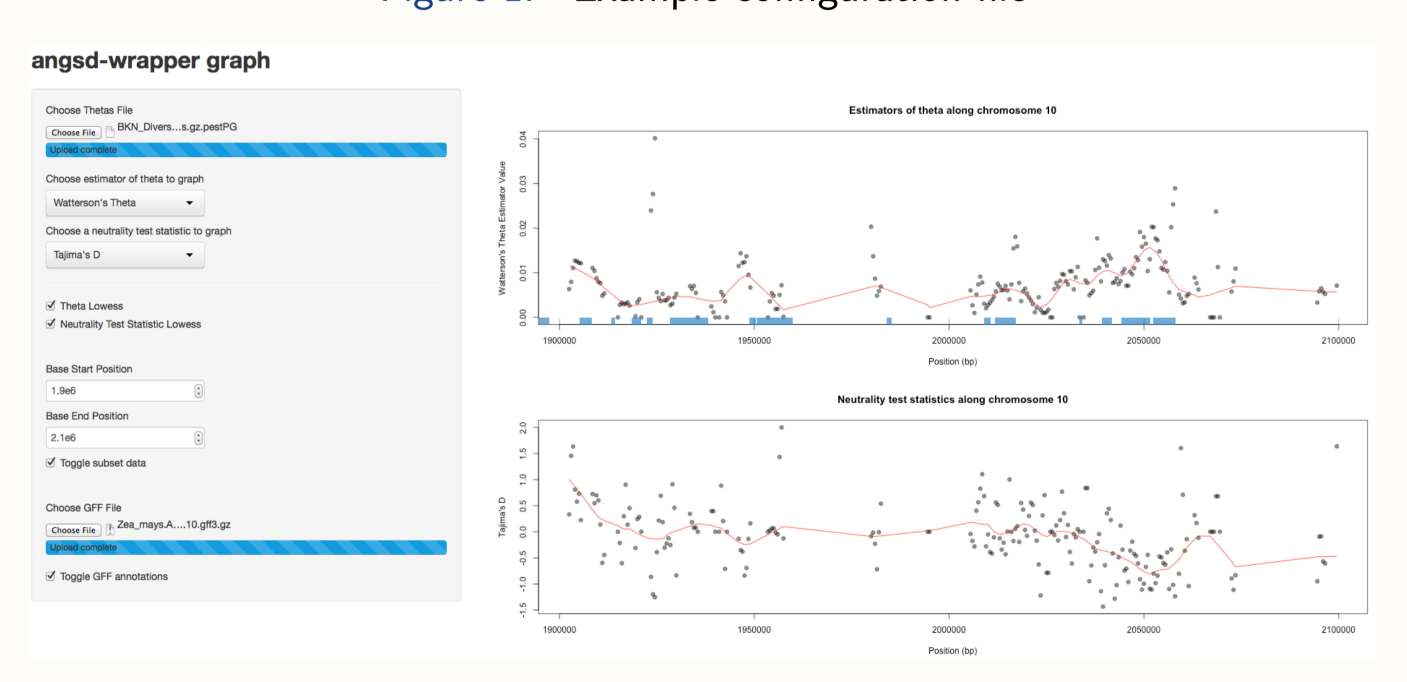


Figure 2: Interactive plotting with Shiny showing a 200kb region along chromosome 10 of *Zea mays*. Blue rectangles are annotated genes.

## **Example Application**

We sequenced 8 genomes of the wild rice *Oryza glumaepatula*, 4 each from populations allopatric and sympatric to cultivated fields of the domesticated *Oryza sativa* ssp. *indica*, in order to investigate evidence of crop-wild introgression. *O. glumaepatula* is a potentially endangered species 1.8 MY diverged from domesticated rice [7] and native to Central and South America [8]. Both species share the AA genome [8] and we have successfully crossed them experimentally, providing reason to believe the possibility of natural hybridization and thus the risk of wild population decline and extinction [9]. Preliminary analysis using ANGSD-wrapper is suggestive of introgression (Figure 3): nucleotide diversity in the sympatric population of *Oryza glumaepatula* is more

strongly correlated with domesticated rice (Spearman's  $\rho = 0.537$ ) than with the allopatric population ( $\rho = XXX$ ). Nucleotide diversity in the allopatric population, in contrast, shows little correlation with diversity in domesticated samples ( $\rho = 0.139$ ).

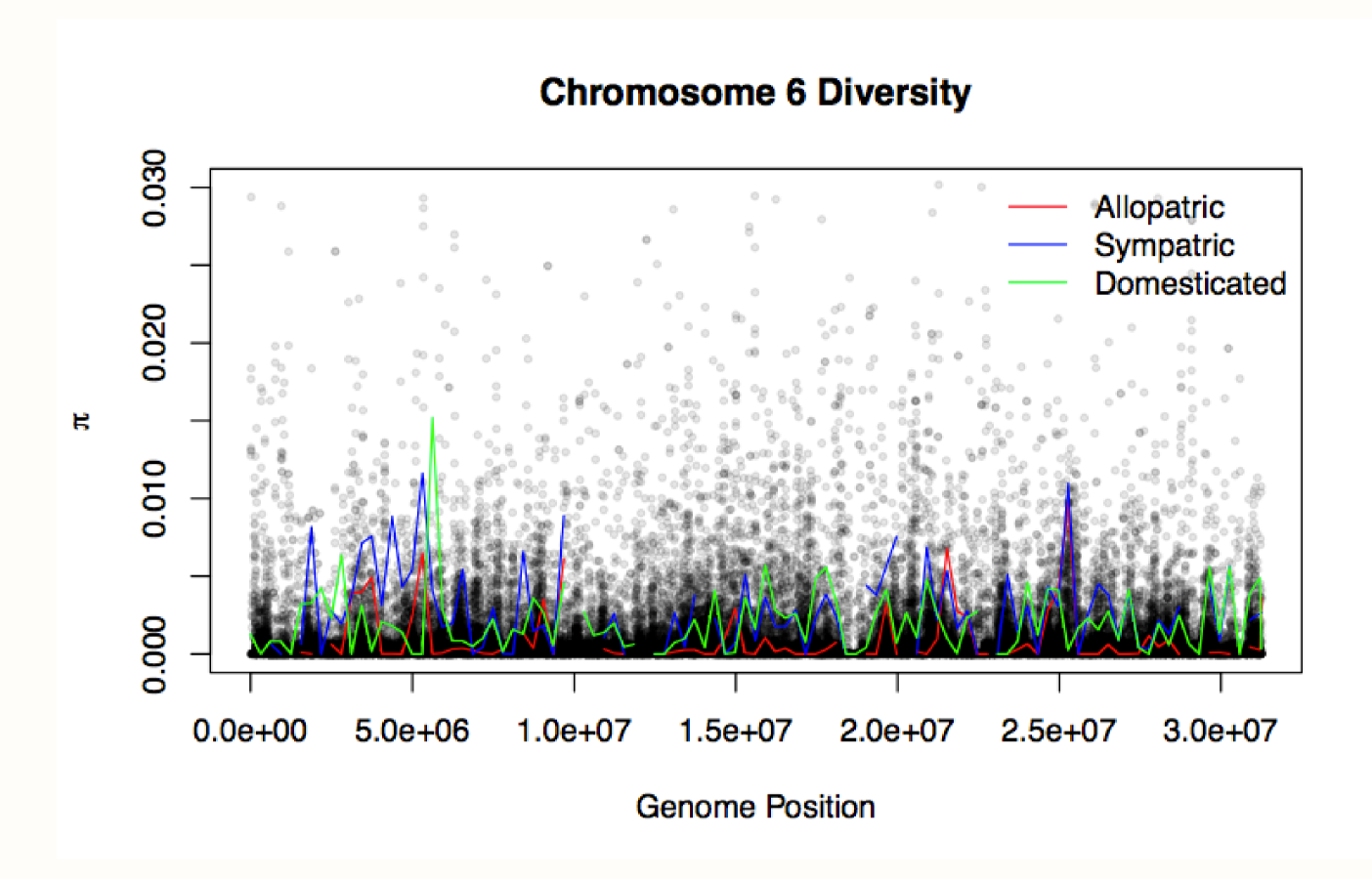


Figure 3: Nucleotide diversity  $(\pi)$  along chromosome 6 of *Oryza glumaepatula* and *Oryza sativa* 

### References

- [1] Korneliussen et al. (2014). BMC Bioinformatics. 15:356
- [2] Fumagalli, M., F. G. Vieira, T. Linderoth and R. Nielsen (2014). Bioinformatics 30(10): 1486-1487.
- [3] Crawford, J., M. M. Riehle, W. M. Guelbeogo, A. Gneme, N. f. Sagnon, K. D. Vernick, R. Nielsen and B. P. Lazzaro (2014). bioRxiv.
- [4] Lohmueller, et al. (2013). American Journal of Human Genetics, 93(6), 1072–1086.
- [5] R Core Team (2014). R Foundation for Statistical Computing, Vienna, Austria http://www.R-project.org/
- [6] Rstudio, Inc. (2013). http://www.rstudio.com/shiny/
- [7] Zhang, Q.-J., et al. (2014). PNAS 111(46): E4954-E4962.
- [8] Vaughan DA, Morishima H, and Kadowaki K (2003). Current Opinion in Plant Biology 6:139-146.
- [9] Rhymer JM and Simberloff D (1996). Annu Rev Ecol Syst 27:83-109.