Estimation of Viral Population Structure from Amplicon-Based Reads

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Abstract—Accurately estimating the structure of highly diverse viral populations is a challenging task. There are two main impediments to globally reconstructing a population. The first is the presence of sequencing errors in reads. Judiciously differentiating these errors from actual rare variants must be properly handled or the global structure may be ill-defined. Secondly, long conserved regions in the viral genome extend beyond what modern sequencers are capable of producing. As a result, the actual population diversity may be hidden in these targeted regions. We propose VirA, a tool for global reconstruction of a viral population that overcomes these obstacles by combining local error correction and a read-graph approach.

Keywords—next-generation sequencing; viral quasispecies; global reconstruction;

I. Discussion

A quasispecies is a heterogeneous closely-related intra-host viral population. Estimating the structure (i.e., variants and their respective abundance) is categorized into two approaches: local and global reconstruction. Local methods focus on a single targeted PCR region and estimate the diversity. Global reconstruction methods target longer regions and assemble the population.

One approach to sequencing viral populations is to perform PCR for multiple overlapping regions. The resulting pools are then combined and sequenced by technologies such as Roche 454 or Ion Torrent. VirA takes as input the sequenced reads, a reference genome, and the target-specific primers used for PCR. It estimates the viral population by performing the following steps:

1) align the reads to the reference[1]
2) identify each amplicon (i.e., targeted region) and locally reconstruct the population via kGEM[2]
3) construct a read-overlap graph
4) assemble the population using either a Maximum Bandwidth[3] or Multi-commodity Flow method[4].

Preliminary results on 20 simulated HCV E1E2 data show VirA to outperform QuRe in terms of quality of the solution (Figure 1). Each dataset contained 8-12 amplicon (20k reads each) regions spanning 1734bp over 10 variants. All subpopulations conformed to either a uniform or powerlaw (with a = 2.0) distribution. Reads were generated using Grinder 0.5 and contained both substitution/indel errors (uniformly at rate 0.1%) as well as homopolymer errors (according to Balzer model).

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