

Genome analysis

EzMap: a simple pipeline for reproducible analysis of the human virome

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Abstract

Summary: In solid-organ transplant recipients, a delicate balance between immunosuppression and immunocompetence must be achieved, which can be difficult to monitor in real-time. Shotgun sequencing of cell-free DNA (cfDNA) has been recently proposed as a new way to indirectly assess immune function in transplant recipients through analysis of the status of the human virome. To facilitate exploration of the utility of the human virome as an indicator of immune status, and to enable rapid, straightforward analyses by clinicians, we developed a fully automated computational pipeline, EzMap, for performing metagenomic analysis of the human virome. EzMap combines a number of tools to clean, filter, and subtract WGS reads by mapping to a reference human assembly. The relative abundance of each virus present is estimated using a maximum likelihood approach that accounts for genome size, and results are presented with interactive visualizations and taxonomy-based summaries that enable rapid insights. The pipeline is automated to run on both workstations and computing clusters for all steps. EzMap automates an otherwise tedious and time-consuming protocol and aims to facilitate rapid and reproducible insights from cfDNA.

Availability and Implementation: EzMap is freely available at https://github.com/dekoning-lab/ezmap.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Metagenomic studies have largely focused on characterizing the determinants of bacterial and fungal community structures within microbiomes. Although it is not generally apparent that viruses within a human host have sufficient functional interactions to comprise an ecological community, it has become apparent that high-level snapshots of viral abundances can nevertheless be obtained by metagenomic profiling methods and that they can be indicative of host immune status. In particular, cell-free DNA (cfDNA) viromics techniques have been proposed as one way to perform such surveillance of immune function within individual patients (Burnham

et al., 2016; De Vlaminck et al., 2013, 2015; Dinakaran et al., 2014; Ngoi et al., 2016; Rascovan et al., 2016; Young et al., 2015).

To automate exploration of viral activity profiles from cfDNA shotgun sequencing reads, we developed EzMap. EzMap allows the relative abundance of all known viruses in a sample to be determined based solely on DNA sequence. It leverages the power of various publicly available tools and the speed of parallel computation to subtract out human sequences, and then to clean, align, and analyze non-human sequence data. EzMap generates an interactive report in the form of an HTML document containing dynamically generated, interactive figures that enable uncomplicated analysis.

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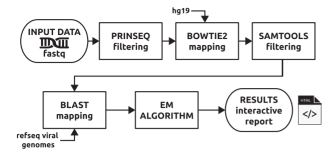


Fig. 1. Flowchart demonstrating the individual steps of the EzMap pipeline. The EM Algorithm step uses the approach of Xia et al. (2011), which we implemented independently

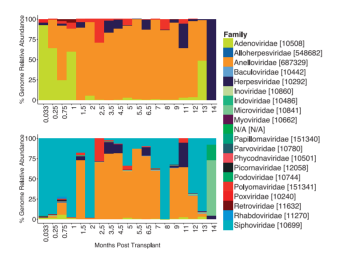


Fig. 2. Genome relative abundance across multiple time-points within the dataset from De Vlaminck *et al.* (2013). Top: six families previously highlighted (De Vlaminck *et al.*, 2013). Bottom: all viral families found (Color version of this figure is available at Bioinformatics online.)

2 Materials and methods

EzMap is partially based on a previously described pipeline (De Vlaminck *et al.*, 2013) (Fig. 1) and was written in Python for SLURM-based computing clusters and multi-core workstation computers. Installation is automated, and requires only the availability of Python 3 and the BioPython library Cock *et al.* (2009). A performance evaluation of the workstation version of EzMap can be found online in the Supplementary Results.

The pipeline workflow is shown in Figure 1 and is fully explained in the Supplementary Methods. Full details of the EM algorithm used (Xia et al., 2011) can also be found there. This important step estimates genome relative abundances, which reflect the relative abundance of each viral genome in the sample. In addition, raw mapping results are output as text files to allow for custom analyses of absolute abundances. The final step of the pipeline generates an interactive report, which can be opened in any modern browser and easily shared. An interactive demo of an EzMap report is available at: http://dekoning-lab.github.io/ezmap/.

3 Discussion

We validated EzMap using the data set from De Vlaminck et al. (2013), in which cfDNA samples from 65 cardiac transplants were sequenced over a prospective time course after transplantation. Our re-analysis was broadly consistent with published findings; some expected variation was observed due to more viral genome sequences being currently available (Fig. 2). One noteworthy discrepancy is the appearance of herpesvirus sequences at time-points from throughout the study (Fig. 2). Interestingly, all of these were identified as HHV-6 viruses, which are known to sometimes be reactivated in transplant recipients (Burnham et al., 2016; Lautenschlager and Razonable, 2012] and can be clinically important. Another discrepancy (Fig. 2, bottom) is the high prevalence of Siphoviridae viruses at nearly all time-points. This appears to be due to the recent sequencing of a number of previously uncharacterized genomes. Interestingly, these viruses have bacteria as their natural hosts, and their dynamics may therefore provide a link between the virome and bacterial microbiome communities.

Author contributions

P.C., S.G. and J.d.K. developed the method, P.C. implemented the method, P.C. and J.d.K. analyzed the data, and P.C., S.G. and J.d.K. wrote the paper.

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Conflict of Interest: none declared.

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