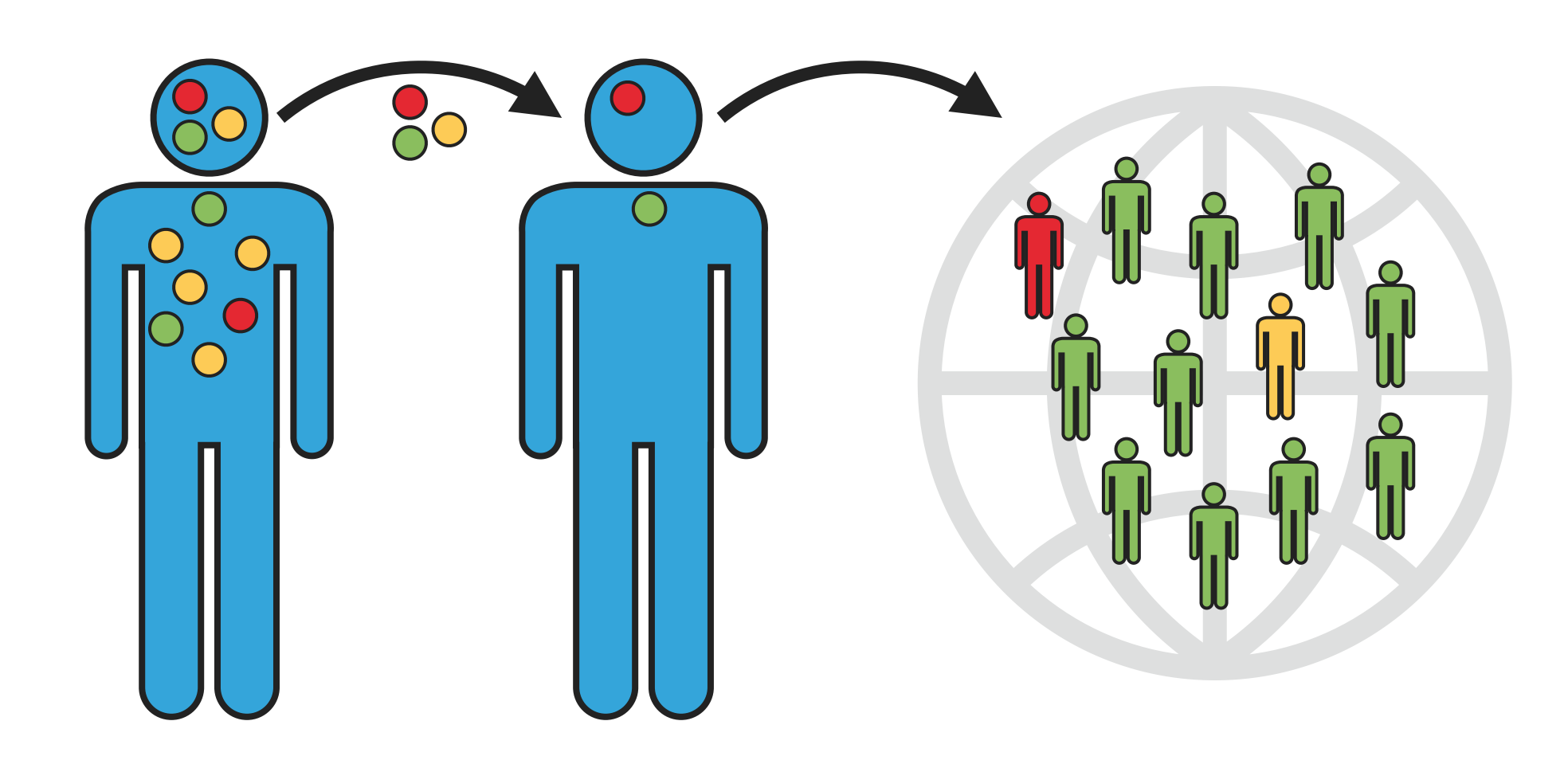
INFLUENZA VIRUS EPIDEMICS ARE LARGELY HOMOGENOUS



How can we most effectively infer phylogenies from genetically homogenous influenza outbreaks?