Advanced R for Genetic Analysis

Exercises for session 6: XML.

- 1. (a) Write a function to read in the airport locations from the file airportlocation.csv used in session 2, and write out a KML file for a single airport (specified by name or abbreviation) or a set of airports. If you don't have Google Earth installed but do have internet access, you can put your KML file on a website and then supply its URL to Google Maps to see the results.
- (b). Using the Seattle flight data, plot mean arrival delay and mean departure delay for each airport. Use identify() to identify when a point is clicked, and write a KML file for that point.
- 2. Read in the XML file phiSITE767857.xml, which describes promoter sites for a set of bacteriophage viruses (from phisite.org).
- (a) Use xpathApply() to extract the organism name for each site (/phisite/site/organism/name) and the sequence (/phisite/site/sequence)
- 3. Using the phiSITE767857.xml file, extract the sequence for promoter sites that have experimental evidence using xpathApply(). Notes:
- xpathApply() takes a function as its third argument, which is passed each of the XML elements returned by the xpath.
- If site is an element of type "site" then the element site[["evidence"]][["type"]] is the type of evidence (experimental or predicted) for the site
- xmlValue() extracts the actual content from an XML element (eg a string)
- The sequence element for a site is site[["sequence"]]