Stat 261, lab 5

The data for this lab are in the file MLLT3_small.vcf. These are data on genomic variants in a gene called MLLT3. The file includes many meta-data lines that start with ##, and then a header for the data columns that starts with #. The separator for the data columns is the tab character.

- 1. Use the appropriate read_* function to read these data into R, skipping the meta-data lines.
- 2. Use the spec() function to print a list of the column specifications. Copy this and make changes so that the REF and ALT variables are read in as factors.
- 3. Rename the first column CHROM. Hint: Beware the #-sign in the column name. This is what the text calls a "nonsyntactic" name. How do you refer to such a name?