## 1 Reverse complement

In the DNA double helix, two strands twist together and "face" each other. The two strands are reverse-complementary, i.e., reading one strand in reverse order and exchanging each base with its complement gives the other strand. A and T are complementary; C and G are complementary.

For example, given the DNA sequence

AGTAGCAT

the reverse sequence is

TACGATGA

so the reverse complement is

ATGCTACT

- (a) Write a function rComplement (dna) to return the reverse complement of a DNA strand. *Use a loop* to reverse the strand—do not use vectorized code. dna is a vector of characters. Assume that dna contains only the letters 'A', 'T', 'C', and 'G'. If dna is the empty vector return the empty vector.
- (b) Now write a function rCompBulk(mat) to return the reverse complements of a set of DNA strands. mat is a matrix of characters; each row of the matrix represents one strand of DNA (so mat contains only the letters 'A', 'T', 'C', and 'G'). Return a matrix the same size as mat such that the rth row of the returned matrix is the reverse complement of the rth strand of DNA (the rth row of mat). Again use loops—do not use vectorized code.

## 2 Counting a DNA pattern

Write a function countPattern(dna,p) to find out (and return) how many times a pattern p occurs in dna. Assume both parameters to be strings that contain the letters 'A', 'T', 'C', and 'G' only. Note that if p is longer than dna, then p appears in dna zero times. Use the built-in function strcmp to compare two strings. Again, use a loop to solve this problem.

## 3 Counting a DNA pattern—challenge edition!

Rewrite the function countPattern(dna,p) without using the strcmp function.