Recall the definitions: The set of RNA strands S is defined (recursively) by:

Basis Step:  $A \in S, C \in S, U \in S, G \in S$ Recursive Step: If  $s \in S$  and  $b \in B$ , then  $sb \in S$ 

where sb is string concatenation. The function rnalen that computes the length of RNA strands in S is defined recursively by:

 $\begin{array}{lll} & & rnalen: S & \rightarrow \mathbb{Z}^+ \\ \text{Basis Step:} & \text{If } b \in B \text{ then} & rnalen(b) & = 1 \\ \text{Recursive Step:} & \text{If } s \in S \text{ and } b \in B \text{, then} & rnalen(sb) & = 1 + rnalen(s) \end{array}$ 

The function basecount that computes the number of a given base b appearing in a RNA strand s is defined recursively by:

 $\text{Basis Step:} \quad \text{If } b_1 \in B, b_2 \in B \\ \text{Basic Step:} \quad \text{If } b_1 \in B, b_2 \in B \\ \text{Recursive Step:} \quad \text{If } s \in S, b_1 \in B, b_2 \in B \\ \text{Basecount}(\ (b_1, b_2)\ ) \quad = \begin{cases} 1 & \text{when } b_1 = b_2 \\ 0 & \text{when } b_1 \neq b_2 \end{cases} \\ \text{Basecount}(\ (s, b_2)\ ) & \text{when } b_1 = b_2 \\ basecount(\ (s, b_2)\ ) & \text{when } b_1 = b_2 \\ basecount(\ (s, b_2)\ ) & \text{when } b_1 \neq b_2 \end{cases}$