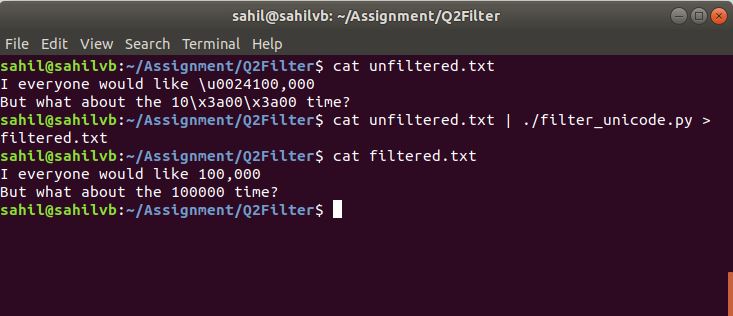
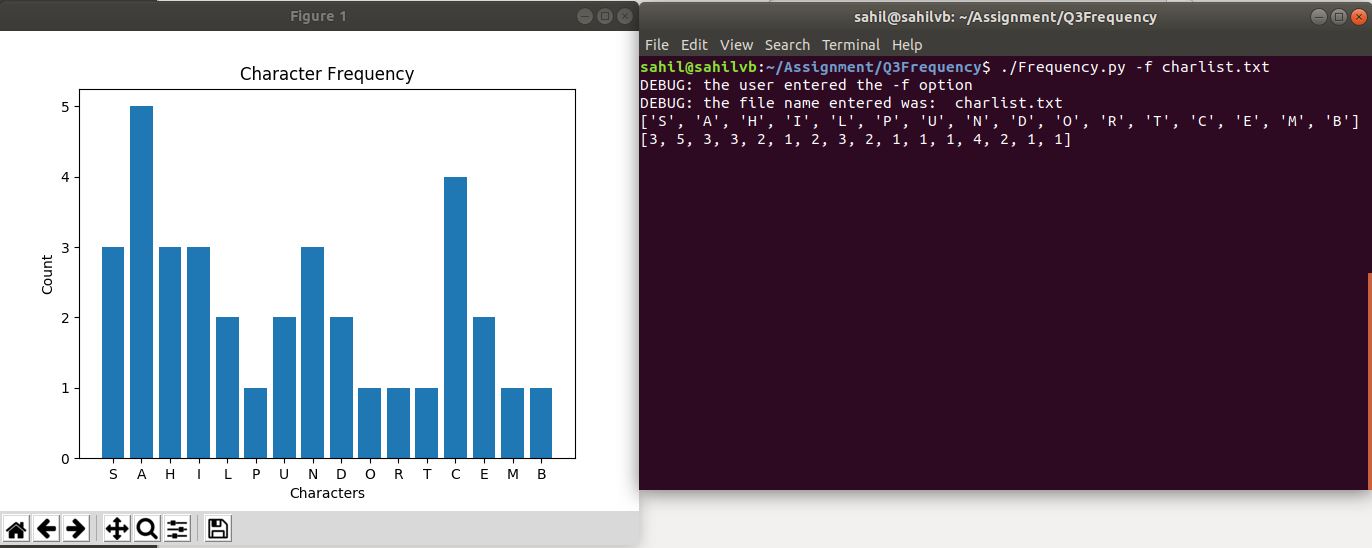
**PYTHON OUTPUTS**

**Q2. Filtering the Text**

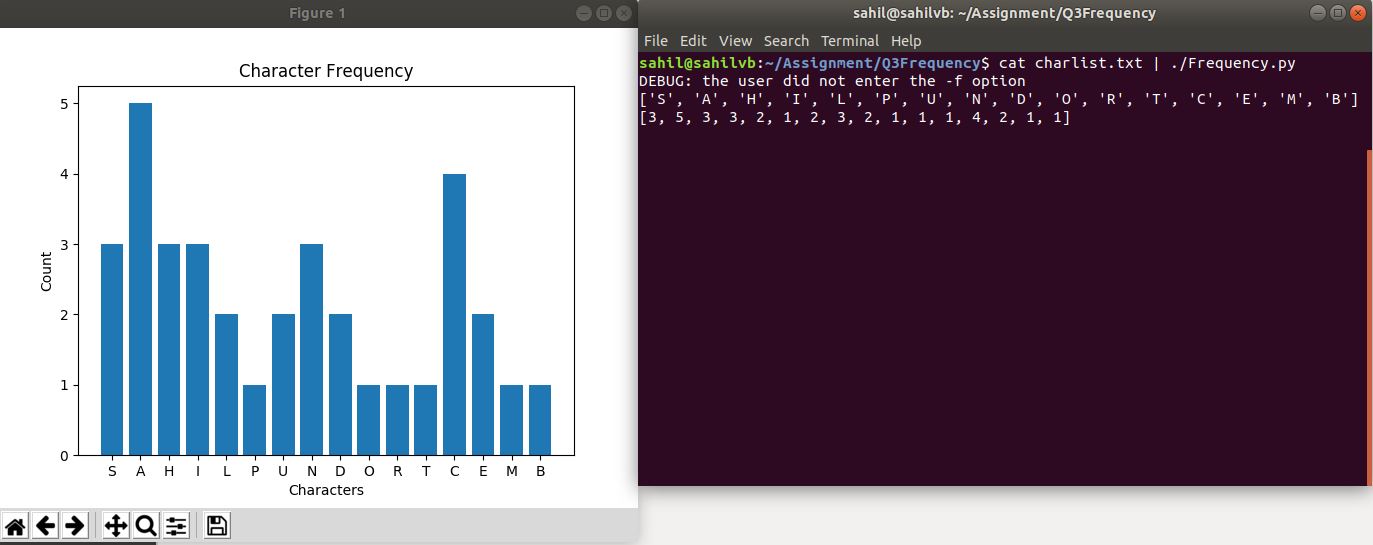


**Q3. Frequency Count**

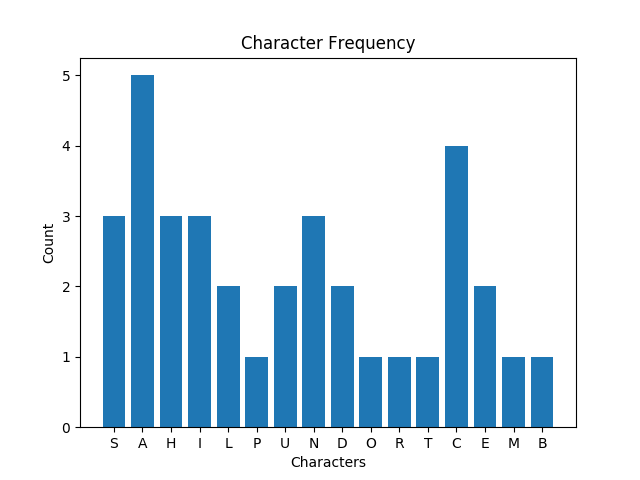
1. **Using FILE (-f command)**

****

1. **Using CAT File Command**

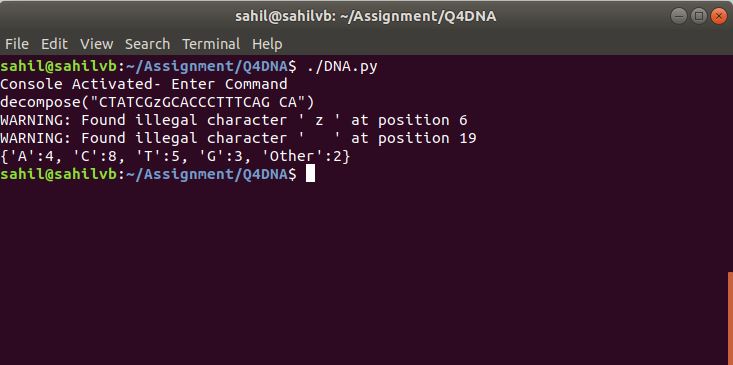
****

1. **Output**

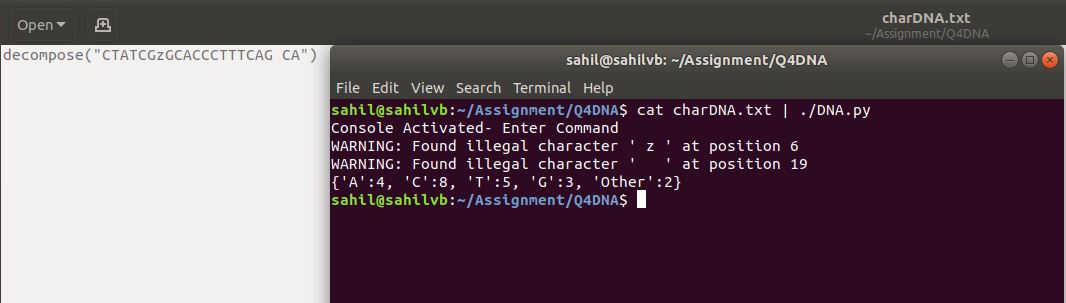
****

**Q4. DNA Decompose**

1. **Using Input on BASH Terminal**

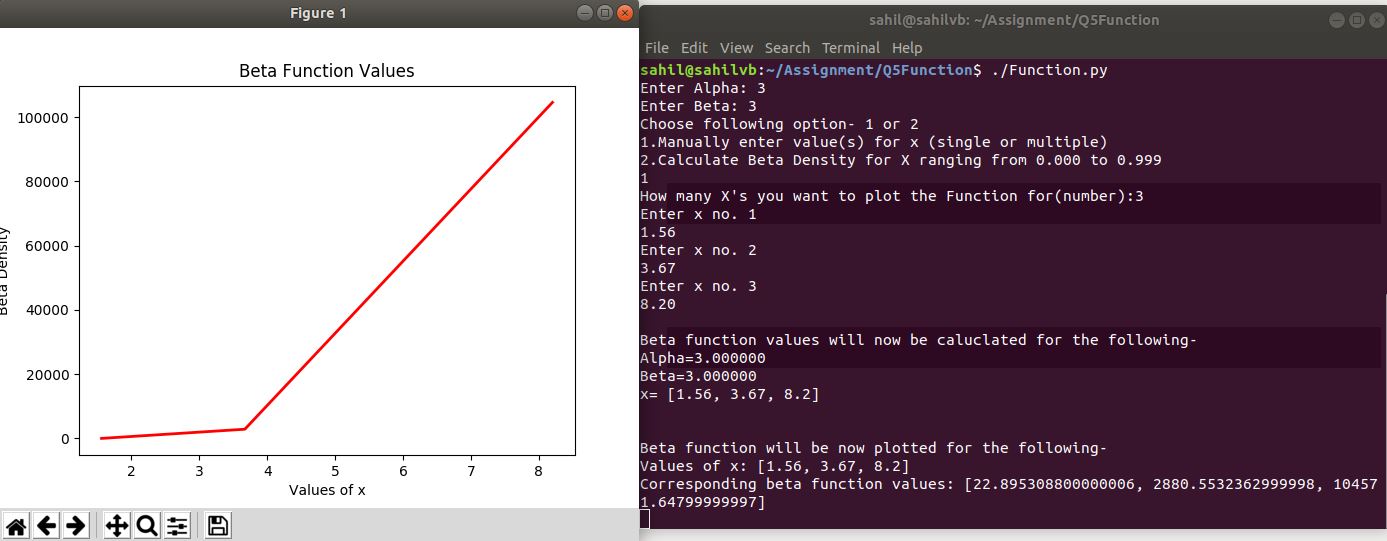
****

1. **Using CAT File command**

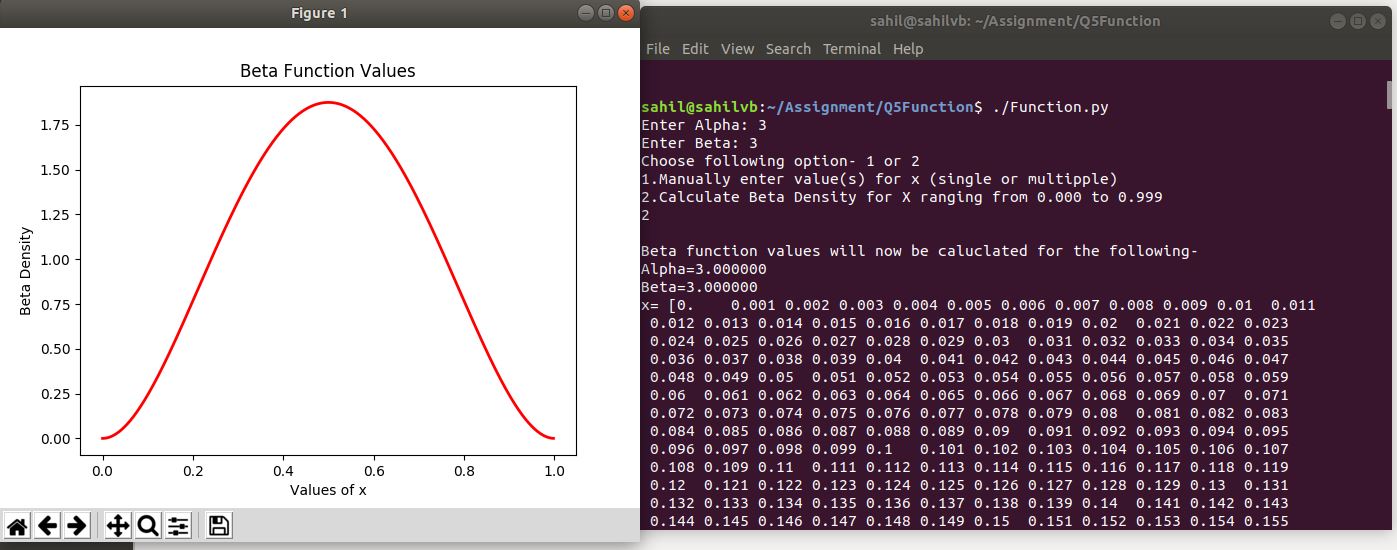
****

**Q5. Beta Function**

**A. Using input ‘Alpha’, ‘Beta’, ‘x’ values (total x values can be greater than or equal to 1)**

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**B. Plotting for input ‘Alpha’, ‘Beta’ and ‘x’= Array from 0.000 to 0.999**

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