## Abstract and Introduction summary.

## Abstract:

- Nowadays bioinformatics is considered as an active research field that acts as an interface between the computer science and biology.
- We have two sequences, A subsequence is a sequence that appears in the same relative order, but not necessarily contiguous.
- The longest common subsequence (LCS) is one of the essential issue to be solved viably in computational science.
- In this paper, we will discuss new calculation for LCS of two DNA sequences, will find all common subsequences in the Dna Sequences and determine their location and length then will determine the length and location of longest subsequence present between all sequences and the time complexity will be computed too.
- To achieve this, we store DNA sequences in an array then start to compare them using matching algorithm.
- nowadays DNA sequence comparison is an important and upcoming area in computational molecular biology Intro:
- Bioinformatics helps us to understand and organize the information associated with biological molecules and to build up a guide in the examination of organic information.
- Cell is the basic unit of all living tissue, each Cell performs several operations like molecule transportation, energy conversion and reproduction.
- Each cell contains the genetic information as a form of (DNA) sequence. DNA contains the instructions for making proteins(genes).
- Human genome contains approximately 3 billion base pairs and about 20,000 genes.
- DNA sequence analysis and gene analysis are used for finding forensics related problems and discovering changes in human DNA sequence or genes and biotechnology related problems .
- To discover changes in DNA sequence we compare the affected DNA sequence with the normal one, and these kind of comparisons would take long time for processing because human genome contains three million nucleotide bases.
- biological data are big and very complex, and we need efficient ways to analyze it
- Computational methods can be useful to reduce the time and space complexity.
- Subsequences are used to determine how similar the two strands of DNA are , so it will help in discovering diseases.
- Effective subsequence match still remains challenging.

## **Assignment 2:** Related works

- Tripathi and Pandey (2016): using two similarity algorithms Rabin-Karp and (MCS) maximum common sub stream to find LCS. They found that Rabin-Karp approach is better than MCS calculation.
- Rubi and Arockiam (2012): proposed Positional LCS for decreasing the time complexity.
- Alsmadi and Nuser (2012): evaluated two algorithms longest common subsequences (LCSS) and longest common substring (LCS) for comparing DNA sequences. Performance evaluation has been done by using different kind of code.
- Wang et al. (2011): used the dominant point approach (dominant points are computed using divide and conquer technique) for finding LCS of any number of strings.
- **Beal et al. (2015):** using suffix tree method for recognizing Common substrings in two sequences.
  - then by using the direct acyclic graph(DAG) longest path of the string is determined.
- Lavanya and Murugan (2013): first approach for finding Multidimensional Shortest Common Subsequences (MSCS) and Multidimensional Longest Common Subsequences (MLCS) is using support vector ,second one using Positional Weight Matrix .for testing they used DNA sequences of length (100 to 25,000).
- **Peng and Wang (2017):** using graph-based model(levelled-DAG) for finding MLCS .this way takes out the hubs that cannot add to the development of MLCS. last graph have a single hub which incorporates all the required MLCS. Tests tells that levelled-DAG needs less amount of space and time.
- Chen et al. (2017): patients reports may contain mistakes but using LCS based on semantic similarity will help patients to get accurate reports. Two dimensional matrices are used to store the LCS and recursive relation is used for semantic similarity process.