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**Full Length Article**

**A graph theoretic model for prediction of reticulation events and phylogenetic networks for DNA sequences**



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Phylogenies are the commonly used tools for the prediction of ancestry of present day or- ganisms from the past decades. Several methods have been developed to construct phylogenetic trees that predict the history of species by direct linkage of edges. Very few studies have been developed for the phylogenetic networks (which is the generalization of trees). Presently, the methods used to determine phylogenetic networks are based on dis- tance measures or character measures of the sequences of species. It is a very challenging task for computational biologists to find the exact method that can predict the accurate networks of organisms. In this study, a phylogenetic network construction model based on basic graph theory concepts is reported. This model finds the distance matrix of every se- quence considered in the study. The two features (positioning and stack interactions) of every DNA sequence and their combined effect have been taken into account to calculate the dis- tances. Results suggested that reticulate events can be observed by using the distances obtained by the proposed method and no such event is predicted by using the distances calculated by the previous method. The important results obtained in the form of dis- tances are 1.637300, 2.000000, 0.932700, 2.331300, 2.829200 and the significance of these values is to represent the different reticulation events among the sequences for different fea- tures. Hence distances calculated by this model gives better insights to study the phylogenetic networks.

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# Introduction

The evolutionary and ancestry relationships have been inves- tigated among the number of species since several decades in

the past. The phylogenetic analysis of DNA or protein se- quences is performed by most of the studies to determine the evolutionary history of organisms [[1–3]](#_bookmark8). Phylogenies are the im- portant tools to predict the evolutionary history and relationships of different organisms. The estimation of divergence of species

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from a common ancestor can be predicted by various methods [[4–6]](#_bookmark9). This estimation permits one to construct a phylogenetic tree that represents the history of species according to their ancestry. The most commonly used methods for tree construc- tion are phenetic and cladistic [[7,8]](#_bookmark12). The former is based on distances and measures the pair-wise distances between or-

is applied over the data set of nine sequences and is vali- dated by comparison with the available phylogenetic network construction method. The basic concepts used in the present study are defined in the subsequent sections.

ganisms to construct the phylogenetic tree. The latter is based

on characters and constructs the tree by using the parsimo- nious methods [[8,9]](#_bookmark13). Lastly, the tree that optimizes the evolution of species most is considered [[1]](#_bookmark8).

However, most of the available methods are unreliable when the base composition of taxa or species varies among the se- quences. Various methods work on the computation of distances among the two or more sequences based on their nucleotide composition, binary composition or sequence similarity, etc. These methods generally use the Jukes Cantor (JC) distance, Kimura two parameter (K2P) distance, F84 distance, LogDet dis- tance, etc. to find the phylogeny among the taxas [[10–12]](#_bookmark14). Qi et al., 2011 reported a graph theory based method for similar- ity analysis. But no method based on graph theoretic concepts similar to those of Qi et al., 2011 is reported for identification of reticulation events and construction of phylogenetic

# Basic definitions

## *Directed graph*

A directed graph *G* consists of a set of vertices

*V*  *v*1, *v*2, *v*3,………… , a set of edges *E*  *e*1, *e*2, *e*3,  and a mapping that maps every edge onto some ordered pair of vertices *vi*, *vj*  [[16]](#_bookmark17).

## *Adjacency matrix*

Let *G* be a directed graph with *n* vertices and *E* edges [[16]](#_bookmark17). Then the adjacency matrix *M*  *xij*  of the diagraph *G* is an *n*  *n* matrix whose element

networks.

In the present study, the authors modify the method de- veloped by Qi et al., 2011 by incorporating two more features

*xij*

 1, if there is an edge directed from *i*th vertex to *j*th vertex.

 0, otherwise.

of a DNA sequence that are used to find the distance matrix of the sequence based on the graph theory and hence extend it to develop a model for identification of reticulation events and construction of phylogenetic network [[13]](#_bookmark15). Every nucleo- tide sequence is composed of four base pairs, namely, adenine (A), guanine (G), cytosine (C) and thymine (T). Hence the ad-

jacency matrix of the order 4 × 4 is obtained for both the features

of every sequence. After the computation of distance matri- ces of all the available sequences of the organisms, the Euclidean distance among all the matrices of the same order is calculated to determine the phylogeny [[10,14]](#_bookmark14).

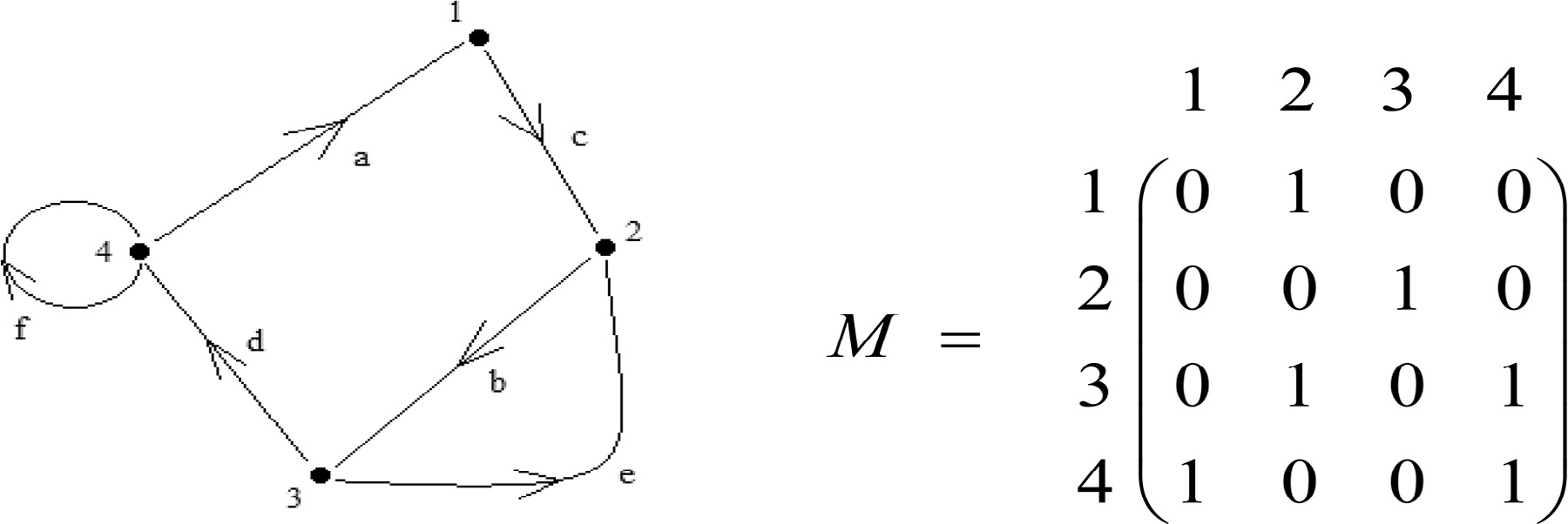
The phylogenetic tree is obtained by the average cluster- ing or neighbor-joining method [[5]](#_bookmark10) and then reticulation events are predicted among the taxas to construct the phylogenetic network and to predict the interactions between the differ- ent species. These interactions may be of the Lateral Gene Transfer (LGT), Homoplasy, Hybridization, Genetic Crossover, Recombination type, etc. [[14,15]](#_bookmark16). As an application, the method

A directed graph and its adjacency matrix is shown in [Fig. 1](#_bookmark1).

## *Stack interactions*

The role of hydrogen bonding and stack interactions is con- sidered as an important role in biological structures [[17]](#_bookmark18). So, the DNA structure is also maintained by intra-strand base stack- ing interactions. Henceforth, the stacking interaction and length of sequence play vital roles in the stability of DNA sequence. The average stacking interactions of GC base pairs are two or three times stronger than AT base pairs [[18,19]](#_bookmark19).

In comparison to other interactions (hydrogen bonds and hydrophobic interactions) involved in holding double stranded DNA sequence together, the energies of the stacking interac- tions in DNA sequence are significantly large [[18,19]](#_bookmark19). The probability of these interactions is considered in the study to calculate the distances among sequences.



**Fig. 1 – Directed graph and its corresponding adjacency matrix.**

## *Euclidean distance matrix*

A matrix *M* of order *n*  *n* is called a Euclidean distance matrix if it satisfies the following conditions [[20]](#_bookmark20):

sequence is constructed by adding to it the content of prob- ability of interactions among nucleotides [[14]](#_bookmark16). The rule used to construct the DWG for the feature of stack interaction is defined as:

* + 1. *M* is a symmetric matrix: *dij*  *dji*

1, 2, , *n* .

* + 1. Diagonal elements are all zero: *dii*  0 1, 2, , *n* .

 *i*, *j* 

 *i*, *j* 

Stack Interaction

 Probability of interaction among nucleotides Position among them

* + 1. There exist *n* points: *p*1, *p*2,………… . . , *pn* on

*m*-dimensional space such that

The three probabilities for the interactions among the nucle- otides are considered which are given below:

*d*  *p*  *p* 2 : 1  *i*, *j*  *n* .

*ij i j*

## *Phylogenetic tree*

Any connected graph with a unique path between any two dis- tinct vertices *u* and *v* and no cycles is called a tree and is denoted as *T*. A tree *T* has exactly *V*  1 edges. The degree of a vertex *v*, deg *v* , is defined to be the number of edges at- tached to *v*. In a tree, any vertex *v* with deg *v*  1 is called a leaf [[21,22]](#_bookmark21).

## *Phylogenetic network*

Any phylogenetic tree with edge lengths and cycles is called a network. It is represented as a triplet *V*, *E*, *l*, where *l* is a

function of edge lengths assigning real nonnegative numbers to the edges. The creation of new sequence *x* from parent nodes *Z* and *z* at a recombination node is called a “recombination

or reticulation event” [[23,24]](#_bookmark22).

# Methodology

Probability of conversion from purines (A, G) to pyrimi- dines (C, T) or vice-versa which is called transversions is 1/2.

Probability of conversion from purines to purines or py- rimidines to pyrimidines which is called transitions is 1/3. Probability of conversion into the same nucleotide is 1/6.

These probabilities are assumed according to the biologi- cal effects of these conversions on the evolution of species and the total probability is considered as 1.

Let us take a sequence of length five as an illustration to construct the directed weighted graph for both the features ana- lyzed in the present study.

Sequence  CGATA

Theorem 3.1. There exists one–one mapping between the DNA sequence and its corresponding directed weighted graph with respect to the positioning of nucleotides [[13]](#_bookmark15).

Proof. Let the DNA sequence for which a graph is to be con- structed with respect to the position of nucleotides be CGATA. Then the graph represented by this sequence with the mapping from vertices (nucleotides) to edges (pair of nucleotides) is

The DNA sequences are composed of the four bases adenine, guanine, cytosine and thymine, i.e. A, G, C, T. Let us assume

defined by 1

*j*  *i*

. The graph obtained by using this mapping

that any sequence of length *n* having the four characters can be represented as:

is shown in [Fig. 2](#_bookmark2).

*S*  *s*1, *s*2, *s*3, , *sn*

such that

*si* *A*, *C*, *G*, *T*.

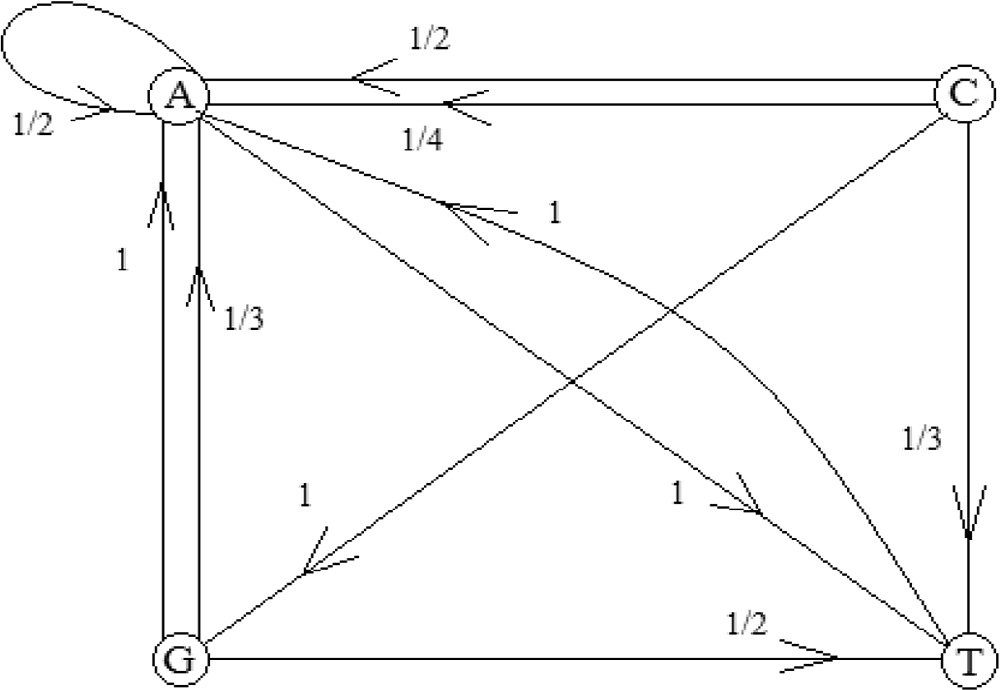
To describe the process of getting the distance matrix of a sequence by the concept of adjacency of graph theory, authors will construct the directed weighted graph (DWG) for *s*1, *s*2, *s*3, , *sn* which is denoted by

*Gw*  *V* *Gw* , *E**Gw*  . The vertex set *V* *Gw*   *A*, *C*, *G*, *T*and the

edge set *E**Gw*  comprise pairs of two vertices. For each pair of nucleotides *Si* and *Sj* in *S* with *i*  *j*, put an arc from *Si* to *Sj* with the weight as 1 *j*  *i* so that 1 *j*  *i* is a decreasing func-

tion of  *j*  *i* which reflects the fact that the two nucleotides with smaller distance would have stronger interactive rela- tionship than those with bigger distance.

In this work, the two features of the nucleotide sequences have been incorporated in the model; one is based on the po- sition of the nucleotides and the other is based on the stack interactions among the nucleotides of the sequences. In case of second feature, the directed weighted graph of the



**Fig. 2 – Directed weighted graph with respect to the position of the nucleotides in the sequence CGATA** [**[5]**](#_bookmark10)**.**

From the graph, one can find the first and the last nucle- otides of the sequence by using the lowest weight of the arc in the graph and the direction of the arc signifies which nucleo- tide comes first in the sequence. Also, the loop in the graph signifies how many times the same nucleotide comes in the sequence. Further, the other positions of the nucleotides can be determined by using the weights of the particular arcs in the graph. Hence, it is easy to find the sequence from the di- rected weighted graph. Thus there exists one–one correspondence between DNA sequence and directed weighted graph.

## *The resultant graph*

matrix, the higher will be the interaction between them and vice-versa.

*A C G T*

*A*

*a*11 *a*12 *a*13 *a*14 *a*21 *a*22 *a*23 *a*24 *a*31 *a*32 *a*33 *a*34 *a*41 *a*42 *a*43 *a*44

*M*  *C*

*G T*

The matrices obtained for the parameter of position of nucle- otides of the sequence CGATA and the parameter of stack interactions among them are as below:

Since there are more than one arc between the same nucle- otides in the same direction, to make this graph simple, authors

 0.5 0 0 1.0 

 0.75 0 1.0 0.3334

 0.0833 0 0 0.5 

 0.375 0 0.5 0.1111

*MP*  1.3334 0 0 0.5  *MSI*   0.4444 0 0 0.25 

added all the weights of the arcs occurring in the same direc- tion. The mathematical equation for the addition of all these weights is as [[13]](#_bookmark15):

 

 1.0 0 0 0 

 

 0.5 0 0 0 

*WR* *u*, *v* 

 *Ww* *u*, *v*

(1)

So, by using these matrices obtained for every sequence con- sidered in the study, one can find the distances among the two sequences by using the appropriate distance metrics.

So, based on this rule, the resultant graph is shown in [Fig. 3](#_bookmark3). It is also observed in this study that there is no one–one correspondence between the given sequence and its weighted graph obtained for the parameter of stack interactions among the nucleotides of the sequence. But the process of construc- tion of graph remains same and here the probabilities of conversion of nucleotides (transversion to transversion or tran- sition to transition or vice versa) are included with the position

of the nucleotides in the sequence.

Now from the resultant graph of DNA sequence obtained from the above discussion, one can obtain the 4 × 4 adja- cency matrix of the sequence which is represented as below. This matrix will also show the interactions among the differ- ent nucleotides of the sequence. The higher the value of the particular entry corresponding to the pair of nucleotides in the

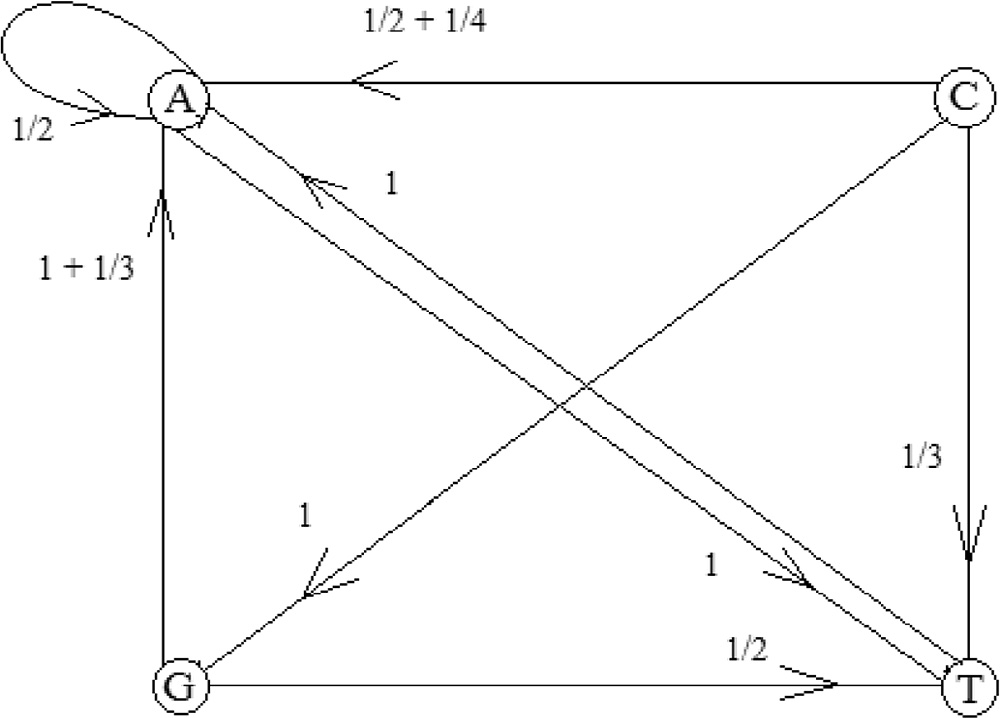
## *Euclidean matrix distance metric to calculate* dissimilarity

In the above section, a distance matrix for a DNA sequence is derived based on two parameters, i.e. positioning and stack in- teractions, by using the graph-theoretic method. So, the dissimilarity between two DNA sequences becomes the same as between the matrices formed by them. The smaller the dis- tance between two sequences, the less dissimilar they are and vice-versa. Let us consider two DNA sequences A and B and the two matrices corresponding to them are as MA and MB for different parameters.

The distance measurement for the two sequences is based on the Euclidean distance matrices among the sequences and

is denoted by *d**A*, *B* [[20]](#_bookmark20). The distance *d**A*, *B* is defined

as:

*d**A*, *B* 

 *ij*

4 4

*M*  *M* 

*A*

2

*ij*

*B*

*i*1 *j*1

(2)

where *MA*  *MB* represents the difference between the same

*ij ij*

**Fig. 3 – Resultant directed weighted graph with respect to the position of the nucleotides in the sequence CGATA.**

entries of the matrices A and B.

In the present work, the two features are considered to cal- culate the distances among the sequences. As a result of which the total dissimilarity *dT* *A*, *B* among the sequences A and B is defined as [[25]](#_bookmark23):

*dT* *A*, *B*  *dP* *A*, *B*  *dSI* *A*, *B* (3)

where *dP* *A*, *B* is the dissimilarity due to position and *dSI* *A*, *B* is the dissimilarity due to stack interactions among sequences. The comparison between the DNA sequences for their dissimilarity is measured by the distances among them. The distances based on all three features are considered for the detection of the phylogenetic network among the given

sequences.

Based on the above graph-theoretic method, the 4 × 4 ma- trices for all the sequences of [Table 1](#_bookmark4) are computed manually. Then these matrices were used to calculate the distances among all the sequences by using the above metric. The distance tables obtained for the two parameters of the sequences and the com- bined result of these parameters are represented in [Tables 2–4](#_bookmark4). After getting these distances, authors were able to calcu- late the relationships among the DNA sequences based on the

|  |
| --- |
| **Table 1 – Input data sequences with their labels.** |
| Label Sequence |
| I ACAAG  II ACCAG  III GACAA  IV AGCAA  V AGACA  VI AGATA  VII CGATA  VIII AGGTA  IX CGGTA |

above technique.

# Results and discussion

1. **Implication of proposed method**

## *Data analysis*

The data to be analyzed for this study are composed of a set of nine sequences which is provided in the [Table 1](#_bookmark4).

The dissimilarity matrices obtained by the proposed method for different features are used to analyze the sequences for their relationships. Various methods can be used to find the dis- tances but these are based on multi-alignment of sequences where the chances of losing information about sequences in- crease. The proposed study finds the distance matrix of every

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 2 – Dissimilarity distances among nine sequences based on their positions of nucleotides.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.0000 | 2.0000 | 2.4353 | 1.6373 | 1.8448 | 3.0116 | 3.3146 | 3.3146 | 3.9247 |
| II | 2.0000 | 0.0000 | 2.9179 | 2.4523 | 2.2142 | 3.4137 | 3.0935 | 3.3645 | 3.4621 |
| III | 2.4353 | 2.9179 | 0.0000 | 1.6583 | 1.3280 | 2.8161 | 2.6484 | 3.6305 | 3.6535 |
| IV | 1.6373 | 2.4523 | 1.6583 | 0.0000 | 1.2473 | 2.6247 | 2.7563 | 3.1447 | 3.5959 |
| V | 1.8448 | 2.2142 | 1.3280 | 1.2473 | 0.0000 | 2.4608 | 2.6484 | 2.9815 | 3.4662 |
| VI | 3.0116 | 3.4137 | 2.8161 | 2.6247 | 2.4608 | 0.0000 | 1.8296 | 2.1213 | 3.0024 |
| VII | 3.3146 | 3.0935 | 2.6484 | 2.7563 | 2.6484 | 1.8296 | 0.0000 | 2.5847 | 2.0000 |
| VIII | 3.3146 | 3.3645 | 3.6305 | 3.1447 | 2.9815 | 2.1213 | 2.5847 | 0.0000 | 2.2017 |
| IX | 3.9247 | 3.4621 | 3.6535 | 3.5959 | 3.4662 | 3.0024 | 2.0000 | 2.2017 | 0.0000 |

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 3 – Dissimilarity distances among nine sequences based on the stack interactions of nucleotides.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.0000 | 0.4930 | 0.8421 | 0.6940 | 0.6709 | 1.4539 | 1.2898 | 1.3646 | 1.5311 |
| II | 0.4930 | 0.0000 | 0.9171 | 0.9104 | 0.7489 | 1.5994 | 1.2979 | 1.5096 | 1.4827 |
| III | 0.8421 | 0.9171 | 0.0000 | 0.5481 | 0.4668 | 1.3265 | 1.1551 | 1.4578 | 1.5080 |
| IV | 0.6940 | 0.9104 | 0.5481 | 0.0000 | 0.5740 | 1.3365 | 1.2039 | 1.3492 | 1.5162 |
| V | 0.6709 | 0.7489 | 0.4668 | 0.5740 | 0.0000 | 1.2304 | 1.2142 | 1.3071 | 1.4984 |
| VI | 1.4539 | 1.5994 | 1.3265 | 1.3365 | 1.2304 | 0.0000 | 0.7466 | 0.7818 | 1.2227 |
| VII | 1.2898 | 1.2979 | 1.1551 | 1.2039 | 1.2142 | 0.7466 | 0.0000 | 1.0350 | 0.8292 |
| VIII | 1.3646 | 1.5096 | 1.4578 | 1.3492 | 1.3071 | 0.7818 | 1.0350 | 0.0000 | 0.9327 |
| IX | 1.5311 | 1.4827 | 1.5080 | 1.5162 | 1.4984 | 1.2227 | 0.8292 | 0.9327 | 0.0000 |

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 4 – Total dissimilarity distances among sequences based on position and stack interaction of their nucleotides.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.0000 | 2.4930 | 3.2774 | 2.3313 | 2.5157 | 4.4655 | 4.6044 | 4.6792 | 5.4558 |
| II | 2.4930 | 0.0000 | 3.8350 | 3.3627 | 2.9631 | 5.0131 | 4.3914 | 4.8741 | 4.9448 |
| III | 3.2774 | 3.8350 | 0.0000 | 2.2064 | 1.7948 | 4.1426 | 3.8035 | 5.0883 | 5.1615 |
| IV | 2.3313 | 3.3627 | 2.2064 | 0.0000 | 1.8213 | 3.9612 | 3.9602 | 4.4939 | 5.1121 |
| V | 2.5157 | 2.9631 | 1.7948 | 1.8213 | 0.0000 | 3.6912 | 3.8626 | 4.2886 | 4.9646 |
| VI | 4.4655 | 5.0131 | 4.1426 | 3.9612 | 3.6912 | 0.0000 | 2.5762 | 2.9031 | 4.2251 |
| VII | 4.6044 | 4.3914 | 3.8035 | 3.9602 | 3.8626 | 2.5762 | 0.0000 | 3.6197 | 2.8292 |
| VIII | 4.6792 | 4.8741 | 5.0883 | 4.4939 | 4.2886 | 2.9031 | 3.6197 | 0.0000 | 3.1344 |
| IX | 5.4558 | 4.9448 | 5.1615 | 5.1121 | 4.9646 | 4.2251 | 2.8292 | 3.1344 | 0.0000 |

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 5 – Phylogenetic network distance matrix for the data of positions of nucleotides in the sequences.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.000000 | 1.999973 | 2.390675 | **1.637300** | 1.976846 | 2.958965 | 2.966851 | 3.416662 | 3.697089 |
| II | 1.999973 | 0.000000 | 2.595716 | 2.280868 | 2.181887 | 3.164006 | 3.171893 | 3.273086 | 3.553514 |
| III | 2.390675 | 2.595716 | 0.000000 | 1.659678 | 1.328004 | 2.977556 | 2.985442 | 3.435252 | 3.715680 |
| IV | **1.637300** | 2.280868 | 1.659678 | 0.000000 | 1.245849 | 2.662707 | 2.670593 | 3.120404 | 3.400831 |
| V | 1.976846 | 2.181887 | 1.328004 | 1.245849 | 0.000000 | 2.563727 | 2.571613 | 3.021423 | 3.301851 |
| VI | 2.958965 | 3.164006 | 2.977556 | 2.662707 | 2.563727 | 0.000000 | 1.924397 | 2.374207 | 2.654635 |
| VII | 2.966851 | 3.171893 | 2.985442 | 2.670593 | 2.571613 | 1.924397 | 0.000000 | 2.152155 | **2.000000** |
| VIII | 3.416662 | 3.273086 | 3.435252 | 3.120404 | 3.021423 | 2.374207 | 2.152155 | 0.000000 | 2.201702 |
| IX | 3.697089 | 3.553514 | 3.715680 | 3.400831 | 3.301851 | 2.654635 | **2.000000** | 2.201702 | 0.000000 |

sequence by which the complete information about the se- quences can be availed. Then, the phylogenetic tree is calculated for all the distance measures by using the neighbor-joining method of T-Rex software [[6]](#_bookmark11). After finding the trees, the tree distance forms the basis for the construction of the network. The phylogenetic networks are calculated to predict the rela- tionships among different sequences of organisms. [Table 5](#_bookmark5) shows the phylogenetic network distances for the input set of sequences given in [Table 1](#_bookmark4), for the positioning feature of the nucleotides in the sequences.

The significant results of [Table 5](#_bookmark5) are made bold and have values 1.637300 and 2.000000. These values represent branch lengths among the pairs of sequences corresponding to them and there is also the possibility of reticulation events among these sequences. The sequences corresponding to 1.637300 are sequence I and sequence IV. The sequences corresponding to 2.000000 are sequence VII and sequence IX. Phylogenetic network related to this dissimilarity data of [Table 5](#_bookmark5) is also shown in [Fig. 4](#_bookmark5) in which red dotted branches show the reticu- lated events among sequences.

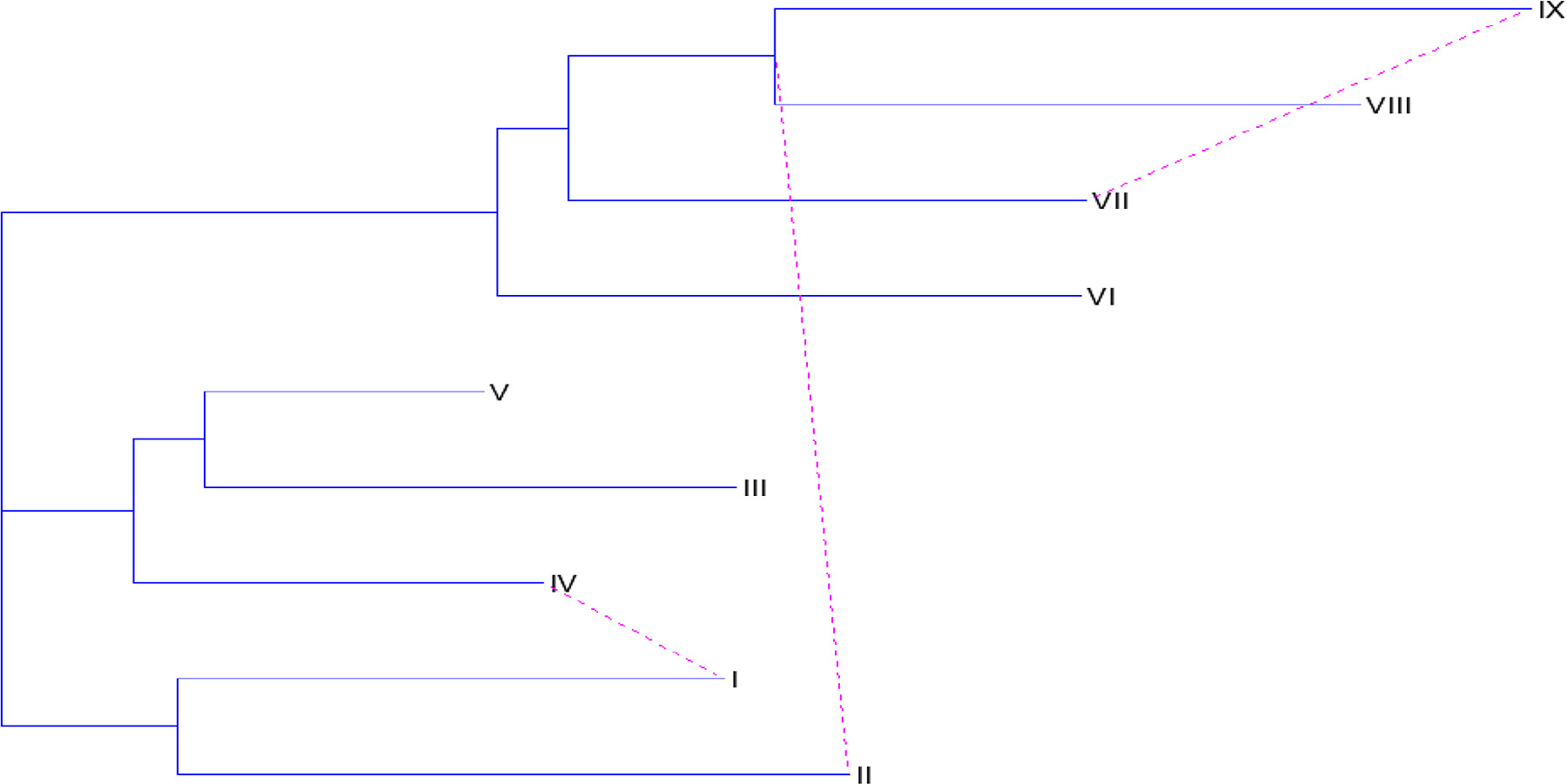
In a similar fashion, the dissimilarity network distances for the stack interactions feature of nucleotides in the sequences is given in [Table 6](#_bookmark6). The significant value for the feature of stack interactions is 0.932700 (in bold) among the sequences VII and

IX. Corresponding to these data, the visualization of the network relationships is represented in [Fig. 5](#_bookmark6).

[Table 7](#_bookmark6) shows the reticulated network distances for the com- bined result of both the features of the nucleotides in the sequences. The values 2.331300 and 2.829200 (in bold) of [Table 7](#_bookmark6) represent the occurrence of reticulation events among the se- quences corresponding to them. In response to these distances, the representation of phylogenetic network is shown in [Fig. 6](#_bookmark7). Finally, the phylogenetic network distances obtained by the

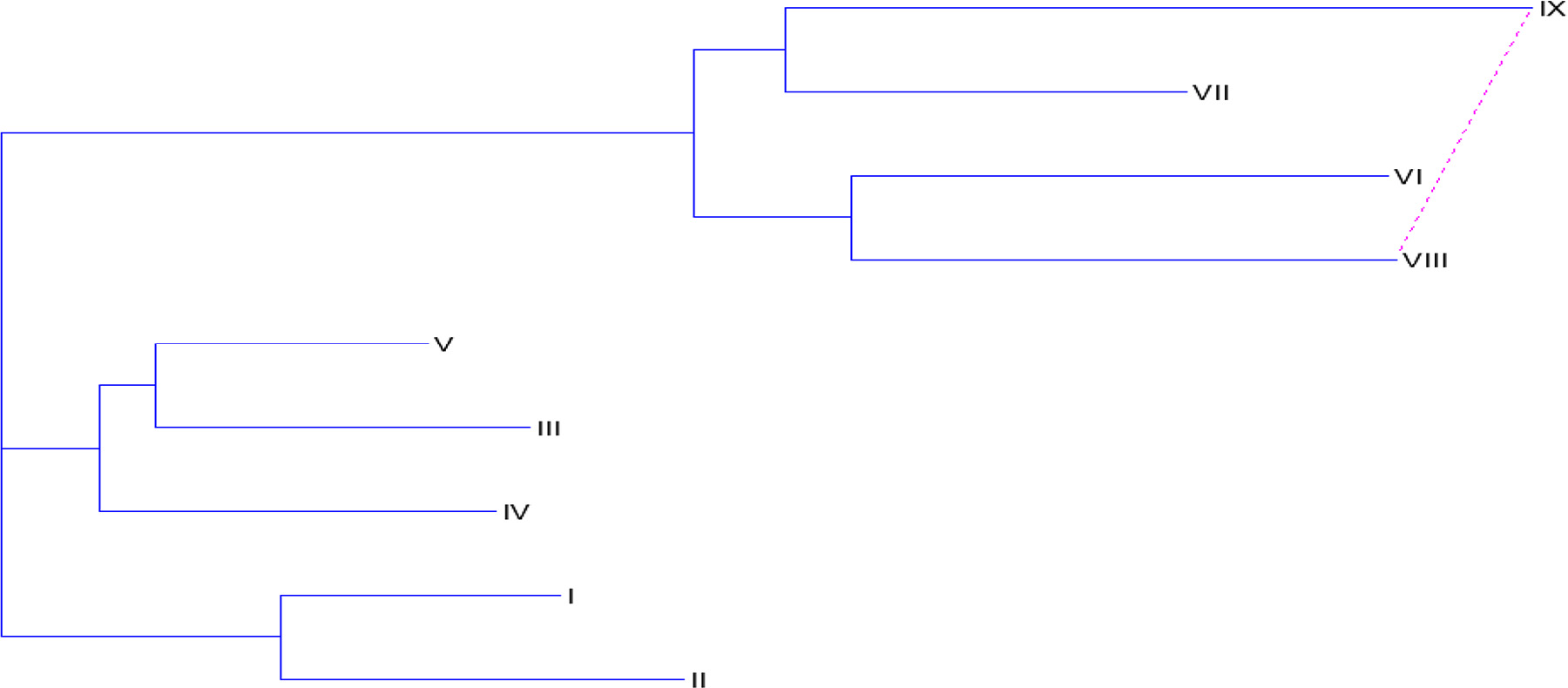
T-Rex package for the input data sequences are shown by [Table 8](#_bookmark6) and [Fig. 7](#_bookmark7) depicts the network for these sequences which is the same as Phylogenetic tree for the input sequences.

Now, it is observed that [Fig. 4](#_bookmark5) represents three main clus- ters and the three reticulation events while [Fig. 7](#_bookmark7) shows no reticulation event with two main clusters and one species as an out-group. This information tells us that positioning feature gives us the close picture of the evolutionary history of the or- ganisms. The events marked by dotted lines in [Fig. 4](#_bookmark5) show that there exist some important relationships among the non- sister species. This information could be useful for prediction of pathogens and antigens based on evolutionary relationships. In the same way, [Fig. 5](#_bookmark6) depicts some results that are dif- ferent from [Fig. 7](#_bookmark7) in that one reticulation event and three main clusters occur due to the parameter of stack interactions among



**Fig. 4 – Phylogenetic network for the data of positions of nucleotides in the sequences.**

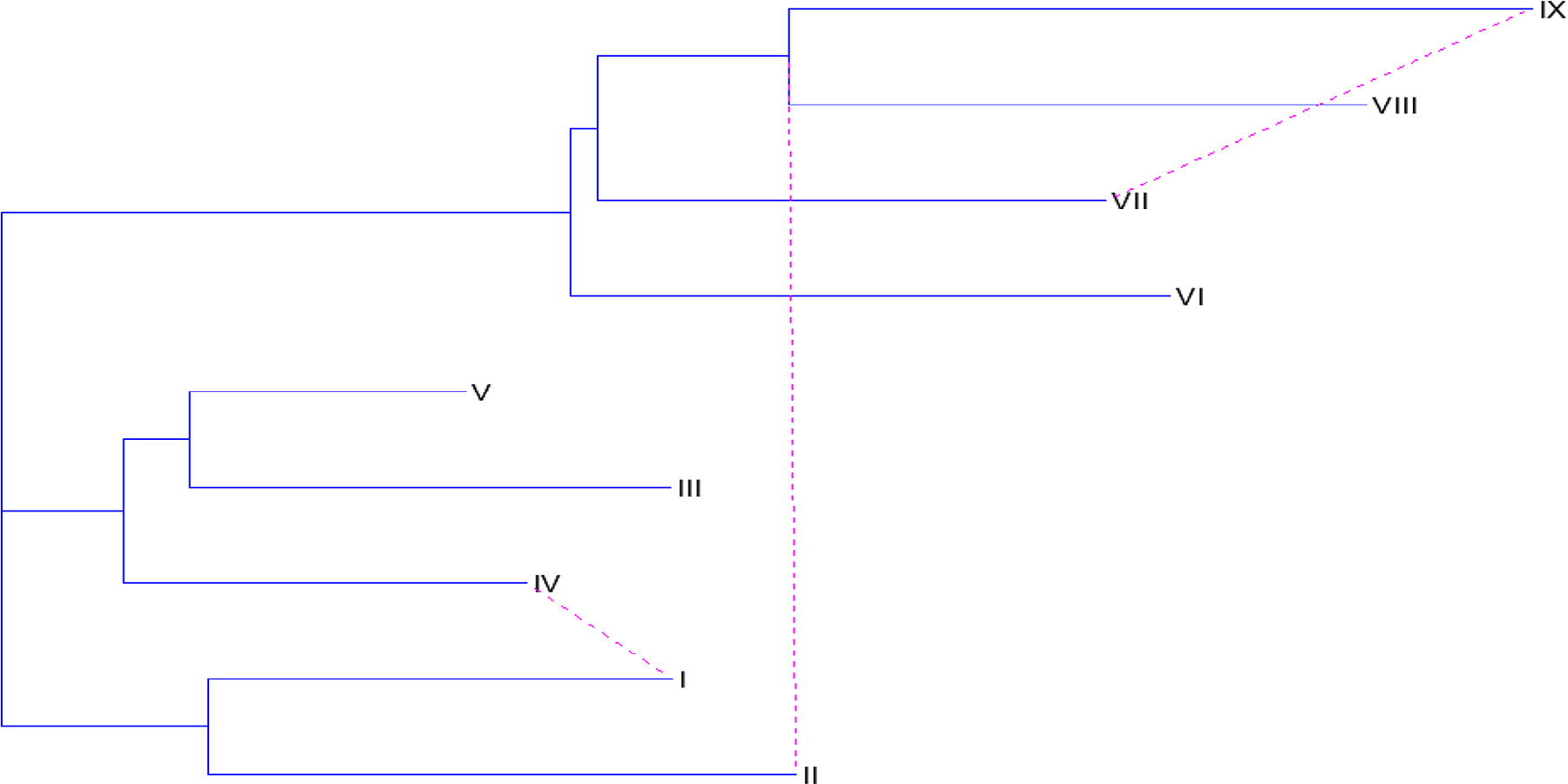
|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 6 – Phylogenetic network distance matrix for the data of stack interactions of nucleotides in the sequences.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.000000 | 0.492997 | 0.785467 | 0.760973 | 0.712495 | 1.405895 | 1.260032 | 1.411610 | 1.509931 |
| II | 0.492997 | 0.000000 | 0.873981 | 0.849487 | 0.801009 | 1.494409 | 1.348546 | 1.500124 | 1.598445 |
| III | 0.785467 | 0.873981 | 0.000000 | 0.597528 | 0.466801 | 1.383459 | 1.237596 | 1.389174 | 1.487495 |
| IV | 0.760973 | 0.849487 | 0.597528 | 0.000000 | 0.524556 | 1.358965 | 1.213102 | 1.364680 | 1.463001 |
| V | 0.712495 | 0.801009 | 0.466801 | 0.524556 | 0.000000 | 1.310488 | 1.164625 | 1.316202 | 1.414524 |
| VI | 1.405895 | 1.494409 | 1.383459 | 1.358965 | 1.310488 | 0.000000 | 0.856443 | 0.781800 | 1.106342 |
| VII | 1.260032 | 1.348546 | 1.237596 | 1.213102 | 1.164625 | 0.856443 | 0.000000 | 0.862157 | 0.829205 |
| VIII | 1.411610 | 1.500124 | 1.389174 | 1.364680 | 1.316202 | 0.781800 | 0.862157 | 0.000000 | **0.932700** |
| IX | 1.509931 | 1.598445 | 1.487495 | 1.463001 | 1.414524 | 1.106342 | 0.829205 | **0.932700** | 0.000000 |



**Fig. 5 – Phylogenetic network for the data of stack interactions of nucleotides in the sequences.**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 7 – Phylogenetic network distance matrix for the combined data of positions and stack interactions of nucleotides in the sequences.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.000000 | 2.492960 | 3.176137 | **2.331300** | 2.689338 | 4.363562 | 4.210355 | 4.830406 | 5.222720 |
| II | 2.492960 | 0.000000 | 3.469693 | 3.130351 | 2.982893 | 4.657118 | 4.503911 | 4.713293 | 5.105607 |
| III | 3.176137 | 3.469693 | 0.000000 | 2.257206 | 1.794805 | 4.359715 | 4.206508 | 4.826559 | 5.218873 |
| IV | **2.331300** | 3.130351 | 2.257206 | 0.000000 | 1.770407 | 4.020373 | 3.867166 | 4.487217 | 4.879530 |
| V | 2.689338 | 2.982893 | 1.794805 | 1.770407 | 0.000000 | 3.872916 | 3.719709 | 4.339759 | 4.732073 |
| VI | 4.363562 | 4.657118 | 4.359715 | 4.020373 | 3.872916 | 0.000000 | 2.690632 | 3.310683 | 3.702996 |
| VII | 4.210355 | 4.503911 | 4.206508 | 3.867166 | 3.719709 | 2.690632 | 0.000000 | 3.028319 | **2.829200** |
| VIII | 4.830406 | 4.713293 | 4.826559 | 4.487217 | 4.339759 | 3.310683 | 3.028319 | 0.000000 | 3.134398 |
| IX | 5.222720 | 5.105607 | 5.218873 | 4.879530 | 4.732073 | 3.702996 | **2.829200** | 3.134398 | 0.000000 |

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Table 8 – Phylogenetic network distance matrix for the original input sequences of nucleotides.** | | | | | | | | | |
| Sequence labels | I | II | III | IV | V | VI | VII | VIII | IX |
| I | 0.000000 | 0.450416 | 1.229796 | 1.131206 | 1.257645 | 1.108086 | 1.112828 | 1.178000 | 1.135730 |
| II | 0.450416 | 0.000000 | 0.779400 | 0.680810 | 0.807249 | 0.657690 | 0.662432 | 0.727604 | 0.685335 |
| III | 1.229796 | 0.779400 | 0.000000 | 0.785452 | 0.911891 | 0.762332 | 0.767074 | 0.832246 | 0.789977 |
| IV | 1.131206 | 0.680810 | 0.785452 | 0.000000 | 0.748414 | 0.598855 | 0.603597 | 0.668769 | 0.626500 |
| V | 1.257645 | 0.807249 | 0.911891 | 0.748414 | 0.000000 | 0.255186 | 0.487507 | 0.631705 | 0.589435 |
| VI | 1.108086 | 0.657690 | 0.762332 | 0.598855 | 0.255186 | 0.000000 | 0.337948 | 0.482146 | 0.439877 |
| VII | 1.112828 | 0.662432 | 0.767074 | 0.603597 | 0.487507 | 0.337948 | 0.000000 | 0.486888 | 0.444618 |
| VIII | 1.178000 | 0.727604 | 0.832246 | 0.668769 | 0.631705 | 0.482146 | 0.486888 | 0.000000 | 0.239156 |
| IX | 1.135730 | 0.685335 | 0.789977 | 0.626500 | 0.589435 | 0.439877 | 0.444618 | 0.239156 | 0.000000 |



**Fig. 6 – Phylogenetic network for the data of positions and stack interactions of nucleotides in the sequences.**

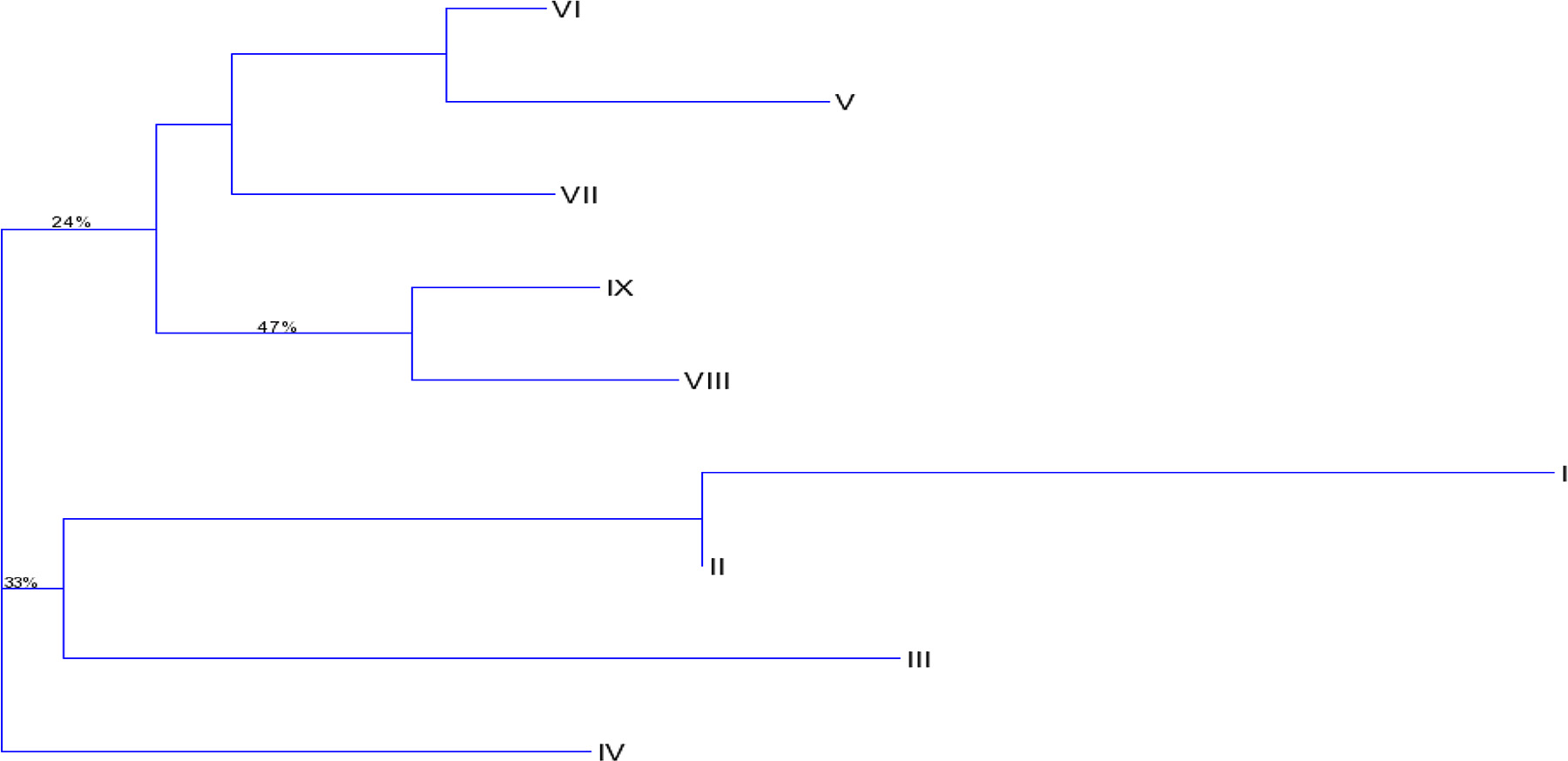
the nucleotides. Irrespective of the same number of clusters, the species change their place of interactions with the other species in the main cluster of four species.

Likewise, the combined result of both the features (posi- tions and stack interactions of nucleotides) highlighted by [Fig. 6](#_bookmark7) shows three reticulation events among the clusters while [Fig. 7](#_bookmark7) has no such type of events.

At last, authors can comment that the loss of reticulation events in [Fig. 7](#_bookmark7) when data sequences are inputted is due to the alignment of sequences and then converting them into dis- tances. But the proposed graph theory method converts each sequence into the form of matrix distances without aligning them. Hence, the possibility of losing information is decreased and the clear picture for the evolution of species can be predicted.

# Conclusion

The proposed method gives better results and better insights about the history of organisms. It also provides the indi- vidual information of every sequence in the form of distances and thus saves the time of multi-alignment of the sequences. The results obtained by this study were compared with the existing method of T-Rex Package [[6]](#_bookmark11) which shows the loss of reticulation events in network. The results obtained are dif- ferent for different features of nucleotide sequences. The reticulation events predicted for positioning, stack interac- tions and combined effect of both of them are three, one and three which are shown in [Figs. 4–6](#_bookmark5). Henceforth, the proposed



**Fig. 7 – Phylogenetic network for the input sequences of nucleotides.**

model performed well in determining the reticulation events and gives us better results as well.

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