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Phylogenetic Relationship of Some Species of Genus Junonia Butterfly Using RAPD

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Author's contribution

The sole author designed, analysed, interpreted and prepared the manuscript.

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ABSTRACT

The present investigation was carried out to find out similarities and diversities between four species of the genus *Junonia* belonging to Nymphalidae, the largest family of butterflies. The genomic DNA eluded from the species, *J. lemonias, J. alamana, J. hierta and J. orithiya* were screened by nine universal primers OPA-1, OPA-2, OPA-3, OPA-4, OPA-5, OPP-9, OPP-18, OPN-17, OPN-16 through RAPD - PCR. Primer OPN-17 did not produce any bands, whereas, the discrete banding pattern was observed in the remaining 8 primers. Primer OPP-18 distinctly highlights only species-specific bands ranging below 200 base pairs. Primer OPA-2 that produced significant banding patterns was taken into consideration for further analysis. The dendrogram was constructed by using binary data interpreted from the RAPD gel image. Two clades were obtained, where in one clade taxa *J. hierta* and *J. lemonias* cluster as sister taxa, and in the other clade taxa *J. alamana* and *J. orithiya* cluster as sister taxa. The investigation was supported by the Jaccard similarity and distance indices matrix reveals that there is closeness within two species in their genetic makeup, whereas some genetic characters are expressed as species-specific. Henceforth, these patterns produced by respective primers can be considered as diagnostic bands and may

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contribute to molecular markers for *Junonia* species identification. Thus, in related evolutionary studies whenever be under investigation these markers will play a pivotal role in concluding the direction of evolution.

Keywords: Dendrogram; DNA; Junonia; molecular marker; PCA; polymorphism; RAPD-PCR.

1. INTRODUCTION

Butterflies the most magnetic and captivating colourful creatures belong to the largest orderlepidoptera under class- Insecta. Nymphalidae also known as brush-footed or four-footed butterflies is the largest family of butterfly. Out of 18.000 species of butterflies found worldwide. 6000 species belong to nymphalidae and 521 nymphalidae inhabit India [1]. Butterflies are key taxa for monitoring biodiversity as their population is susceptible to alternation in surrounding habitat specifically leading to the loss of species with restricted geographical distribution [2,3]. Several studies revealed that habitat specificity is directly linked to the availability of host plants for larvae and adults [4]. Thus to keep a check on the richness or destruction caused to the environment there arises an urgency to conserve butterflies [5]. Molecular-based work started by William [6] is since then continued and applied in various fields like forensic investigation [7]; linkage map construction to state phylogenetic relations and morphological evolution among Nymphalidae subfamily. Phylogenetic studies of taxa that exhibit adaptive phenotypic variation provide valuable insiahts into the evolutionarv mechanisms driving the origins of biodiversity [8]. Butterfly species richness studied by Principal component analysis (PCA), is a powerful tool for analysing data that displays similarity between observation and variables as points in spot maps. PCA extracts most important information from the data table and thus compresses the provided data set [9,10]. The main aim of the investigation is to study species-specific genetic characters and closeness within two species in their genetic makeup supported by PCA.

2. MATERIALS AND METHODS

2.1 Collection and Identification of Butterflies

Nymphalidae butterflies were collected from different region of Amravati by sweep netting and hand-picking methods by observing their preliminary morphological data, like the presence of reduced first pair of legs highlighting the specific character of family Nymphalidae and later their taxa were identified by observing key characters as mentioned in Books of Indian Butterflies, like location and colour of eye spots on the wings, demarcation of costa (costal margin) and termen (outer margin) of the wings and shape of the apex [1,5,11].

2.2 Preservation of Specimen

The butterfly legs and thorax tissues were preserved in 70% alcohol [12] before post-mortal changes.

2.3 Extraction of DNA

Genomic DNA was extracted from freshly collected leg tissues of butterflies by using Genetix DNAsure Tissue Mini Kit and stored at 4° C for further use. Later it was quantified by using UV spectrophotometry and gel electrophoresis.

2.4 Amplification of Genomic DNA through the RAPD-PCR Method

Genomic DNA was amplified in thermocycler gradient by using RAPD marker with primer ranged from OPA 1 to OPA 5, OPN 16, OPP 9 & OPP 18. PCR proves to be boon for molecular biologist as it is an indispensable technique in the biological and medical field and allows automated DNA sequencing, thus readily producing markers for further investigation [2]. Primers used with different percentage of base content are highlighted in Table 1.

PCR conditions were as follows [Table 2] and the amplicons gained were stored at 4°C.

The PCR product was run at 100 volts on 2% agarose gel for 3- 4 hours. 10 µl amplicon along with 1µl of DNA loading dye were separated on a gel plate stained with Ethidium Bromide and loaded with 1kb DNA marker. Bands on the gels were viewed under Imaging System Unit and photographed for further analysis.

Primers	Primer Sequence (5'to 3')	% Of GC content	Molecular weight (bp)
OPA 1	CAGGCCCTTC	70%	2964
OPA 2	TGCCGAGCTG	70%	3044
OPA 3	AGTCAGCCAC	60%	2997
OPA 4	AATCGGGCTG	60%	3068
OPA 5	AGGGGTCTTG	60%	3099
OPP 9	GTGGTCCGCA	70%	-
OPN 16	AAGCGACCTG	60%	-
OPP 18	GGCTTGGCCT	70%	-

Table 1. Primers used for RAPD-PCR profiling of Nymphalidae butterflies

Table 2. PCR conditions for amplification of Nymphalids DNA

Stage	Temperature	Duration
Pre-denaturation	94 ⁰ C	5 min
Denaturation	94 [°] C	1 min
Annealing	37.6 ⁰ C	1 min
Extension	72 ⁰ C	1 min
Final extension	72 ⁰ C	5 min
Total Cycles	35	

2.5 Bioinformatics Aspect

Bioinformatics is an interdisciplinary research area that is the interface between the biological and computational sciences. It is the application of computer technology to the management and analysis of biological data, as a result computers are being used to gather, store, analyse and merge biological data [13].

2.6 Application of MEGA-5 for Analysis of Brushfooted Butterflies RAPD-PCR Data

A computer program package called MEGA-5 is developed for estimating the evolutionary distance, reconstructing phylogenetic trees and computing basic molecular data. For the estimation of characters-based distance, some methods like Jukes and Cantor [14] distance, Tamura [15] distance, p-distance, UPGMA and Neighbor joining methods require matrix of pair wise distance. MEGA is an integrated tool for conducting automatic and manual sequence alignment, inferring phylogenetic trees, mining web-based databases, estimating rates of inferring molecular evolution. ancestral sequences, and testing evolutionary hypotheses. MEGA is a multi-threaded Windows application. It runs on all releases of the Microsoft Windows operating system [16].

2.7 Phylogenetic Analysis

Phylogenetic trees are the most convenient way of visually presenting evolutionary relationships among a group of organisms. It can be drawn in various ways. The relationships established by phylogenetic trees describe а species evolutionary its historical history and relationships among lineages or organisms or their parts, such as their genes. In the construction of phylogenetic trees, the principle of minimum evolution or maximum parsimony is often used [16]. Different tree-building methods exist for the study of phylogenetic relatedness between DNA samples [18].

2.8 Principal Component Analysis (PCA)

It is a multivariate technique that analyse data table that describes inter-correlated quantitative dependent variables. It is mode of identifying pattern in data, and expressing the data to highlight similarities and differences [10].

3. RESULTS AND DISCUSSION

Preceding the amplification process of fragments of DNA, a data was scored in form of binary code, '1' for the presence and '0' for the absence of band for further RAPD analysis. Bands represent the genotypic characters. Nine primers were screened of that OPN 17 did not produce any bands. The remaining 8 primers showed the different frequencies of combinations of monomorphic and polymorphic bands. Whereas Primer OPP-18 distinctly highlights only speciesspecific bands ranging below 200 base pairs. Primer OPA 2 producing a significant banding pattern of the four Junonia species was taken into consideration for further investigation.

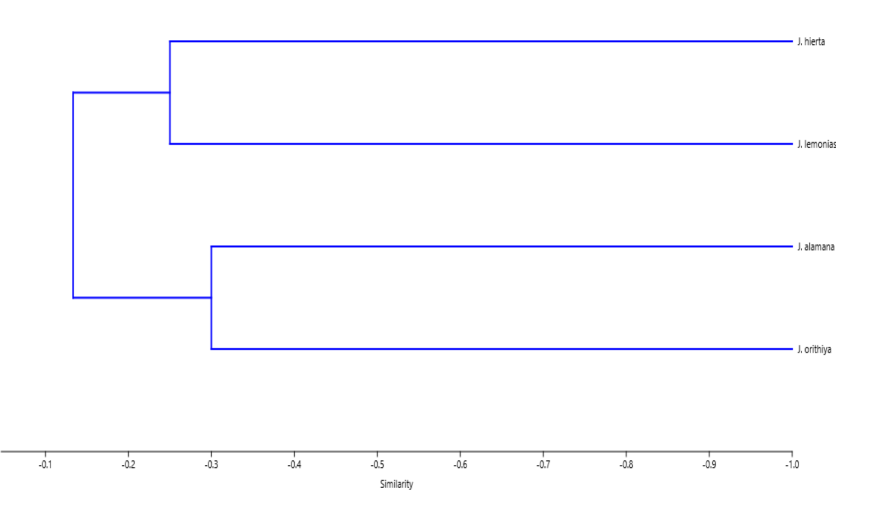


Fig. 1. Jaccard Similarity coefficient of four species of Junonia

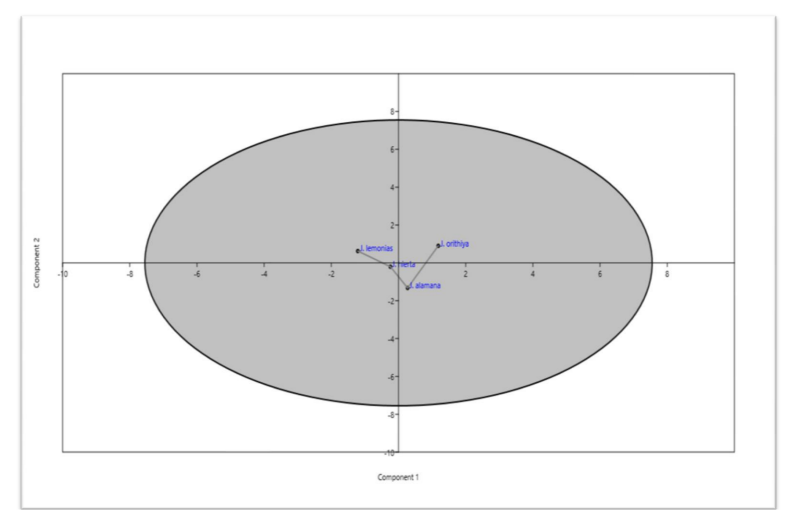


Fig. 2. Principal component analysis (PCA) of four species of Junonia

Species	J. lemonias	J. alamana	J. hierta	J. orithiya
J. lemonias	1			
J. alamana	0.1	1		
J. hierta	0.25	0.16666667	1	
J. orithiya	0.066666667	0.3	0.2	1

Table 3. Distance matrix based on RAPD-PCR of four species of Junonia

The similarity coefficient reveals maximum genetic similarity (0.3) between *J. alamana* and *J. orithiya* and minimum genetic similarity (0.066666667) between *J. lemonias and J. orithiya* [Table 3]. This data is strongly supported by Principal Component Analysis (PCA) dot map [Fig. 2], where *J. orithiya is* completely diverging out and is distantly located from *J. lemonias*.

The species of Junonia were clustered by using Jaccard similarity indices and dendrogram [Fig. 1] created based on binary molecular data shows two clades, where in one clade taxa J. hierta and J. lemonias cluster as sister taxa, the second clade exhibits the association between taxa J. alamana and J. orithiya. Similar clustering of taxa J. alamana and J. orithiya was explained by Vanlalruati et al. in 2011 highlighting close relatedness between them. The Principal Component Analysis (PCA) also showed the similar relationship between species based on their placement in principal component plot. Henceforth, all the above analysis reveals that there is closeness within two species in their makeup, genetic whereas some genetic characteristics are expressed as speciesspecific. These patterns produced by respective primers can be considered as diagnostic bands and may contribute to molecular markers for Junonia species identification. Thus, related evolutionarv studies whenever under investigation these markers will play a pivotal role in concluding the direction of evolution.

4. CONCLUSION

All the primers produced a large number of bands with different intensities and banding patterns suggested that the amplified fragments were repeated in the genome in varying degrees. Results suggested that the RAPD-PCR technique could provide a powerful tool to improve species identification and to better understand genetic variability and the point where divergence is occurring within the genus. The presence of species-specific bands among some species suggested the interspecific genetic relatedness between them.

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COMPETING INTERESTS

Author has declared that no competing interests exist.

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