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Status of two species of lac insects in the genus Kerria from China based on morphological, cellular, and molecular evidence

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Abstract

The taxonomic status of the Chinese lac insects *Kerria yunnanensis* (Ou and Hong) (Hemiptera: Kerridae) and K. ruralis (Wang, Yao, Teiu and Liang) were analyzed in this paper by comparing morphological, cellular, and molecular data. Cladistic analysis showed K. yunnanensis and K. ruralis to be distinct from other Kerria species such as K. lacca and K. chinensis. The karyotype of K. yunnanensis was 3A and the chromosome structure was K = 6m + 2sm + 10T, while in K. ruralis the karyotype was 3B and the chromosome structure was K = 8m + 10T. Kerria ruralis and K. yunnanensis had the closest relationship among species in the genus as they had the most similar karyotype homology. Based on the karyotype analysis, K. sindica and K. lacca formed a sister group with K. ruralis and K. yunnanensis. Kerria pusana and K. nepalensis were clustered as a sister branch, indicating the close relationship of these taxa. The karyotype of K. chinensis was however, different from the other six species and formed a separate branch. RAPD analysis also showed that K. yunnanensis and K. ruralis had distinct differences from other species of Kerria, although they did not form sister taxa. Molecular analysis based on the EF1 α gene using ML, MP, and Mr. Bayes' methods indicated that seven species of lac insects cluster in two major groups. In group 1, K. sindica and K. lacca formed a sister clade and were primitive members of the genus. In group 2, K. chinensis formed the earliest diverging branch followed by K. ruralis. Kerria yunnanensis was the next to diverge followed by the cluster containing K. pusana and K. nepalensis. Hybridization testing showed that crosses neither between K. yunnanensis and K. sindica, nor between K. yunnanensis and K. lacca could produce first generation larvae. This was indicative that K. yunnanensis had a distant genetic relationship from the other species. Morphological, cellular, molecular, and hybridization results confirmed the independent status of the Chinese endemic species K. yunnanensis and K. ruralis. Kerria ruralis was genetically closely related to K. yunnanensis, but relatively far from K. lacca. The main commercial species in China was K. yunnanensis, while in Thailand it was K. chinensis. The commercial species in Myanmar included K. nepalensis and K. pusana, the latter being most widely used in lac production.

Keywords: cladistic analysis, EFI α gene, karyotype, Kerria ruralis, Kerria yunnanensis

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Introduction

Lac insects are scale insects belonging to the family Kerriidae. These insects are considered to be of great economic value. Shellac that is produced by the lac insects of the genus Kerria is widely used in chemicals, electronics, military, food, and other industries (Chen 2005). The family Kerriidae contains nine genera and approximately 100 species (Ben-Dov 2002), while the genus Kerria, contains 19 species which have mainly been used in the lac industry. Kerria (Mahdihassan) chinensis (Hemiptera: Kerridae), K. ruralis (Wang, Yao, Teiu and Liang), K. yunnanensis (Ou and Hong), K. fici (Green), and K. greeni (Chamberlin) are reported from China, with K. yunnanensis and K. ruralis being endemic (Chamberlin 1923; Mahdihassan 1923; Varshney 1976, 1984; Wang 1982; Ou and Hong 1990; Chen 2005).

Shellac is mainly produced in China, India, Myanmar, Laos, and Thailand. The main commercial species of lac insect in India is K. lacca, while in Thailand it is K. chinensis (Chen 2005). There has long been controversy in the identification of commercial lac species in China. Lac insects in China were firstly named Laccifer lacca in the 1950s (Liu 1957, 1959). In 1982, the genus *Laccifer* was synonymized with Kerria and this has generally been accepted in accordance with taxonomic studies in India (Varshney 1976). The fresh collections in China were compared with K. lacca and several differences were found which resulted in the naming of the species K. yunnanensis (Ou and Hong 1990). Also, the commercial species in Thailand was regarded as K. yunnanensis (Ou and Hong 1990). However, lack of comparison with K. chinensis meant there were still certain aspects in need of clarification; and the Chinese commercial species was tentatively named as *K. chinensis* (Chen et al. 1992, 1998). Another endemic species in China was found and named *K. ruralis* (Wang et al. 1982), and this species showed a close relationship with *K. fici* and *K. lacca. Kerria fici* was first recorded in China (Chamberlin 1923), but has not been found there since 1930s (Chen et al. 2008). *Kerria greeni* was reported to occur in Yunnan and Fujian Provinces of China (Chen 2005).

The purpose of this study is to clarify the taxonomic status of two species of Kerria in China. Fresh insects were collected from their host in the locations according the first record in publication. The two species were compared morphologically at the cellular level by studying their karyotypes, RAPD reactions were performed, they were sequenced with $EF1\alpha$ genes and hybridization to establish their identifications, and their relationship with other species in genus Kerria was analyzed.

Materials and Methods

Materials

The following samples used in this study are listed in Table 1. *Kerria yunnanensis* was identified by Ou Bingrong and Hong Guangji. *K. ruralis* was identified by Wang Ziqing. Both of them were collected in Yunnan, China. *K. chinensis* was collected in Thailand. *K. pusana* and *K. nepalensis* were collected in Burma. The three species were identified by authors after a discussion with Dr. Xie Yinping, who is an expert of scale insect in Shanxi University, China. *K. sindica* was identified by Mahdihassan S and collected from Pakistan. *K. lacca* was identified and presented by Indian Institute of Natural Resins and Gums.

Cladistic analysis based on morphology

Permanent slides of mature adult female lac insects were prepared by the method used by Chen et al. (2008). Twenty-two apomorphies which included brachia, anal tubercles, dorsal spine, anterior spiracles, posterior spiracles, and perivulvar pore clusters were selected as morphological characters. The average values for thirty individuals of each species were measured. Phylogenetic analyses performed under maximum parsimony (Nei and Kumar 2000) and analyses conducted using PAUP* 4.0b10 (Swofford 2001). To test for nodal reliabilities, heuristic bootstrap analyses (Felsenstein 1985; 1000 replicates) were applied with groups appearing in 50% or more of the trees in bootstrap analysis.

Karvotype analysis

Microphotographs of chromosomes different species were taken with a Nikon E800 optical system using the air-dried method (Chen et al. 2007) and karyotypic parameters were measured by IM50 software (Leica Ltd. 1992). Karyotype analysis was performed according to the standard method (Leven 1964; Stebbins 1971; Guo 1972). Phylogenetic relationship of lac insects were studied to built the dendrogram clustered using UPGMA by applying karyotype resemblance-near coefficients (λ) and the evolution distance with specific software (Li et al. 2005).

RAPD analysis

The technique of random amplified polymorphic DNA (RAPD) was used to study the relationships of 12 populations from 7 species of *Kerria*. The genetic distance and identity among species were generated by POPGENE32 (Yeh and Boyle 1997). The molecular dendrogram was constructed based

on Nei's genetic distance by MEGA3 using the UPGMA method (Kumar et al. 2004).

PCR and Sequencing

Total genomic DNA was isolated from whole insect body using a standard proteinase K, phenol/chloroform extraction technique (Marchant 1988; Tian 1999). EF1α! genes were amplified by polymerase chain reaction (PCR) using primer pair Forward (5'-ATGTGAGCAGTGTGGCAATCCAA-3') and Reverse (5'-GAACGT-GAACGTGGTATCAC -3') (Palumbi 1996).

DNA amplifications were carried out in the Bio-RAD MyCycler thermal cycler. Amplification cycles were as follows: 95"!# for 4 min as initial denaturation step; 35 cycles of 64"!# denaturation for 1 min, 72"!# annealing for 2 min, 72"!# extension for 7 min, and ended by cooling at 4"!#. The resulting sequences were assembled using Bioedit version 7.0.5.3 (Hall 1999) then deposited in GenBank (Accession numbers EU781492 - EU781498), and are listed in Table 1.

Alignments for the individual gene data matrices were generated using similarity calculated at the nucleotide level with ClustalW version 1.81 of Lasergene DNAstar software package (Thompson et al. 1997). Bayesian phylogenetic inference was used to estimate the tree topology by Mr. Bayes 3.1.2 (Huelsenbeck and Ronquist 2001). ML trees generated based on hierarchical likelihood ratio test (hLRT test) estimated by Modeltest 3.06. Statistical support for each node was evaluated by bootstrap analysis (Felsenstein 1985). Parsimony analyses were conducted in PAUP 4.0b software (beta 10th version, Swofford 2001). Tree visualization and drawing were carried out with TreeView version 1.5.2 (Page 1996).

Hybridization test

All male insects were manually removed from twigs harboring second instar larvae and the remaining females were covered by a synthetic net sleeve (80 mesh), which protected the insects from attack from parasitoids and predators. When the females in the sleeves developed into adults, they were copulated with males chosen from other species. The female insects not copulating were treated as controls.

Results

Morphological Diagnosis Kerria yunnanensis (Ou and Hong) 1990: 15.

Adult female: 1.04-1.9 mm long, 0.69-1.38

mm wide, globe-like body, dark reddish brown. Anal tubercle heavily sclerotized with 0.08-0.38 mm long and 0.06-0.36 mm wide, nearly quadrate, apparently two-segmented and harboring 6-13 anal ring setae about 0.18-0.27 mm long. Branchial tube less than 0.09 mm high, brachial plate with crater about 0.10-0.15 mm long, 0.08-0.14 mm wide, and 0.03-0.05 mm² in the center. Dimples in crater are formed by the brachial pores, numbers vary from 8 to 15. Anterior spiracles are situated 0.02-0.11 mm to brachial plates with 0.18-0.31mm in length and 0.10-0.15mm in width, inside the keratinization trail is inconspicuous and less than 0.22 mm. Dorsal spines are found between the brachia and anal tubercle, and have two parts: a stout pedicel about 0.01-0.11 mm long and 0.04-0.05 mm wide and a conspicuous scletotized spine averaged 0.10-0.23 mm in length. Perivulvar pore clusters originate circularly near the anal

tubercle. Mouthparts have a labium about

0.31-1.40 mm long, 0.11-0.21 mm wide with inconspicuous segmentation and a pair of post

oral lobes 0.03-0.12 mm wide just behind the mouth.

Holotype: \bigcirc , paratypes, $8 \bigcirc \bigcirc$, 4 May 1987, Yunnan, P.R. China (Ou and Hong 1990).

Biological characteristics: Bi-voltine, summer (May-October) and winter crops (October-the next May). Life history of females and males are list in Table 1 and Table 2, respectively.

Host: Dalbergia obtusifolia (Baker) Prain.

Distribution: Subtropical areas of Pu'er and Lincang of Yunnan Province, P.R. China.

Kerria ruralis Wang 1982: 53.

Adult female: Length on slide 1.02-2.3 mm and 0.58-1.27 mm wide, globe-like body, two body color types, i.e. dark reddish brown or yellow. Anal tubercle heavily sclerotized, 0.06- 0.36 mm long and 0.21-0.37mm wide, apparently two-segmented and consist 1-13 anal ring setae about 0.17-0.31 mm long. Brachial tube less than 0.09 mm long, brachial plate with a crater about 0.08-0.13 mm long, 0.06-0.11 mm wide and 0.03-0.04 mm² in the center. Numbers of pores in crater vary from 4 to 11. Anterior spiracles are situated 0.02-0.11 mm to brachia 0.18-0.26 mm long and 0.10-0.15 mm wide, inside the inconspicuous keratinization trail is less than 0.12 mm. Dorsal spine heavily sclerotized 0.10-0.23 mm long; pedicel of dorsal spine 0.02-0.16 mm long, 0.04-0.13 mm wide. Perivulvar pore clusters circular, present near anal tubercle. Mouthparts with a labium about 0.21-0.82 mm long, 0.12-0.19 mm wide, inconspicuous segmentation and a pair of post oral lobes, each 0.06- 0.14 mm wide, behind the mouthparts.

Holotype: \bigcirc , paratypes, $7 \bigcirc \bigcirc$, 10 June 1969, Yunnan, P.R. China (Wang et al. 1982)

Biological characteristics: Bi-voltine, summer (March-July) and winter crops (August-the next March). Life history of female and male were list in Table 3 and Table 4, respectively. Two body color types, i.e. red and yellow. The ratio of red:yellow is about 12:1.

Host: Mallotus philippinensis (Lam.) Muell.

Distribution: Tropical and subtropical areas of Pu'er and Xishuangbanna of Yunnan Province, China.

Cladistic Analysis

Based on the morphological study of twentytwo apomorphies (Table 6), K. ruralis and K. yunnanensis should belong to different categories with obvious differences from other species such as K. lacca and K. yunnanensis (Chen et al. 2008). The phylogenetic relationships among seven species of lac insects using a cladistic approach showed that K. chinensis was the earliest diverging member in case of its special morphological characters. Kerria ruralis and K. yunnanensis were the newest species with similarity in morphology and form a sister group. Figure 1 shows the results. These specimens and Permanent slides were deposited in Research Institute of Resource Insect of China.

Karyotype analysis

The chromosomes of seven species including K. yunnanensis, K. ruralis, K. lacca, K. chinensis, K. sindica, K. pusana, and K. nepalensis were composed of metacentric (or sub-metacentric) and telocentric chromosomes, showing the consistency of the genus while there were four kinds of chromosomes structure shape: K = 10m + 8T, K = 8m (6m + 2sm) + 10T, K = 6m (4m + 10m)

2sm) + 12T, K = 4m + 14T and the data are shown in Table 7 (Chen et al. 2007). The chromosomes of K. yunnanensis are made of six metacentric (or sub-metacentric) and ten telocentric chromosomes. And K. ruralis were formed with eight metacentric and ten telocentric chromosomes. Both differ from other species and have a certain degree of uniqueness. Differences of interspecific relationship were reflected in the centromere position of chromosomes. The cluster analysis of karyotype resemblance-near method coefficient indicated K. ruralis and K. yunnanensis had the highest identity in karyotype (0.9688) and nearest distance in evolution (0.0317), which showed they were the latest species of the seven grouped in the dendrogram (Table 8, Figure 2).

RAPD analysis

The results of RAPD analysis showed the genetic distance inter-species were 0.1854-0.7917, in which the average genetic distance among species was 0.4430 (Table 9) (Chen et al. 2006). The value of genetic distance between K. yunnanensis and K. chinensis, and *K*. lacca 0.6297 0.5789. were and respectively; obviously for different categories. The results of UPGMA showed that the seven species could be divided into two natural groups (Figure 3). In group 1, K. sindica and K. lacca had the closest relationship and were sister taxa. In group 2,

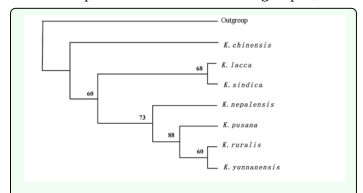


Figure 1. Phylogenetic hypotheses among seven species of lac insects based on the morphological characters by the MP method (The numbers above branches are bootstrap values, Chen et al., 2008). The aphid Stomaphis japonica (Hemiptera: Aphididae) is the outgroup. High quality figures are available online.

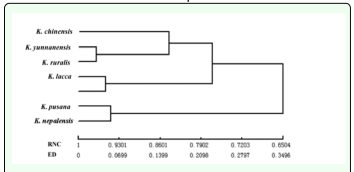


Figure 2. Dendrogram of seven species of lac insect based on the karyotype resemblance -near coefficients and evolutionary distance. RNC = Resemblance -near coefficients. ED = Evolutionary distance. High quality figures are available online.

K. nepalensis was the nearest taxon to the K. ruralis and K. pusana sister group followed by K. yunnanensis. K. chinensis was the earliest diverging member of this group and is placed as the base branch in the group.

Phylogeny of EF1a gene

The phylogenetic trees are based on EF1α gene data of seven lac species, the topologies made by using of MP, ML, and Bayesian methods were similar with different bootstrap values varying from moderate to high for different branches (Figure 4). In the MP tree, all the branches received high bootstrap values (>70%) except the weakly supported branch of K. yunnanensis (57%), in which MP tree length = 377, CI = 0.976, RI = 0.813, and RC = 0.793.Using the maximum likelihood (ML) to build the phylogenetic tree, the GTR + G model was selected as the best model for phylogenetic analysis in accordance with the hLRT test (-InL = 2913.3172). The support value of all branches was in excess of 70%. Bayesian analysis results were consistent with the systematic relationships of MP and ML trees. Except the branch of K. chinensis, which was 92%, other branches all received high posterior probability (PP > 95%).

The seven lac species clustered into two distinct clades (group 1 and 2 in Figure 4), one containing *K. sindica* and *K. lacca* (Bayes, 99%; MP, 100%; ML, 100%) and the

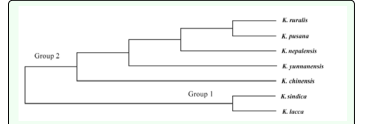


Figure 3. Dendrogram of lac insect among seven species based on genetic distances using the method of UPGMA. High quality figures are available online.

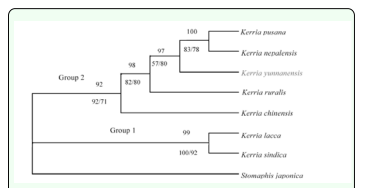


Figure 4. Majority-rule consensus tree resulting from Bayesian analysis of EFI α gene (model =GTR+G) from seven species of lac insects and the aphid *Stomaphis japonica* (Hemiptera: Aphididae) as outgroup. Branches represented are based on the maximum likelihood topology. Numbers above internodes indicate Bayesian posterior probabilities; numbers below internodes indicate non-parametric bootstrap proportions for the parsimony analysis (left) and likelihood analysis (right). Thickened branches indicate Bayesian posterior probabilities \geq 95%. High quality figures are available online.

other containing the remain five species (Bayes, 92%; MP, 92%; ML, 71%). In group K. sindica and K. lacca had a close relationship and were sister taxon. In group 2, K. nepalensis and K. pusana formed a sister branch, showing a close relationship, and indicated that they were the most recently evolved species of the seven. K. chinensis was the earliest diverging member of group 2, and has a distant relationship with the others. K. yunnanensis and K. ruralis were earlier evolved species, but they diverge later than K. chinensis. K. pusana and K. nepalensis were the most recently evolved species of the taxon, forming a close sister group with high support value (Bayes, 100%; MP, 83%; ML, 78%).

Hybridization test

Hybridization tests showed that the mating between K. lacca and K. sindica produced first filial generation, indicating the close genetic relationship of the two species. crossbreeding However, between Κ. yunnanensis and K. sindica, or that of K. yunnanensis and K. lacca, did not produce any first filial generation, which indicated K. yunnanensis had distant relationship to K. sindica and K. lacca (Table 10) (Chen et al. 1992). Five kinds of hybridization could not produce normal offspring (Table 10), which indicated that three populations belong to separate species with the existence of reproductive isolation among them.

Discussion

In this study, K. ruralis and K. yunnanensis should belong to different categories, with obvious differences from other five species on the morphological characters. The results accord were in with the study of morphological characteristics bv using scanning electron microscopy, which confirmed lac commercial species in China was a new species with clear differences from K. lacca and nominated as K. yunnanensis (Ou and Hong 1990, 1991). And the two species also differ from other species with the unique chromosomes structure as K = 8m (6m + 2sm) + 10T (Table 7). In RAPD analysis, the genetic distance between K. ruralis and K. chinensis, and K. lacca were 0.5604 and 0.5110, respectively (Table 9), both higher than the average value of inter-species, which proved the three individuals should be classified into different species. This was in contrast to the idea that lac production species in China were the same as K. lacca in India (Liu 1957, 1959). On the phylogenetic trees based on EF1 α !gene data, the branches of K. yunnanensis and K. ruralis were highly supported with MP, ML, and Bayes analysis in diverging with other species (Figure 4). Results of hybridization also confirmed that the species used for lac production in China was not *K. lacca*, but *K. yunnanensis* (Chen et al 1992), which is the commercial species in China that is very different from *K. chinensis*, the commercial species of Thailand.

Morphology, cytology, and molecular biology evidence consistently indicated Κ. yunnanensis and K. ruralis, the two Chinese endemic species, had significant differences with the other five species examined in this study. The relationships of the seven species were basically consistent, with a few phylogenetic positions being incomplete. However, with no matter which methods, K. lacca and K. sindica, and K. pusana and K. nepalensis, always clustered together and formed two sister groups indicating a close genetic relationship between them. With the exception of RAPD, K. nepalensis and K. pusana always clustered together, indicating a close genetic relationship. K. chinensis was rather special and always stood alone in a separate branch.

The phylogenetic analysis also found that species distributed in similar ecological environments usually clustered indicating a near relationship. K. lacca and K. sindica were both distributed in tropical regions, in which K. lacca was originally found (India, Pakistan, Nepal, Bangladesh, and Sri Lanka). K. sindica was originally located in Sindh Province of Pakistan, mainly located in the lower reaches of the India River in Hyderabad and Karachi, with an annual temperature above 24"# and an annual rainfall of 250-400 mm, which both indicate a tropical monsoon climate (Table 11). The phylogenetic tree (Figure 4) showed K. sindica and K. lacca had the closest relationship and were sister taxa with high support values (Bayes, 99%; MP, 100%; ML, 92%), in the most basal branch of seven species.

Kerria chinensis was found in the north and northeastern of Thailand, where the average annual temperature was 22-28° C and the annual precipitation is 1300-1900 mm (Table 11). The environment was different from that of other species, and this taxon stood alone as a separate branch (Figure 4). K. yunnanensis and K. ruralis diverged later than K. chinensis in this group. While K. pusana and K. nepalensis were the most recent of the genus, forming a close sister group with high support values (Bayes, 100%; MP, 83%; ML, 78%). These four more advanced species were mainly distributed in edge of south subtropical and the north tropical regions.

Among the four species of group 2 in Figure 4, K. yunnanensis was found in semi-arid and semi-humid area of Southern subtropical zone, with 600-1500 m elevation and 18-20 "# average annual temperature. K. ruralis occured in Xishuangbanna, Yunnan Province in the type humid subtropical climate, close to tropical north border, where the average annual temperature was 19-21"! # and the annual precipitation was 1200-1700 mm. K. pusana and K. nepalensis were distributed in different altitudes in Myanmar. K. pusana distributed in 800-1400 m altitude area of Taunggyi, Lashio, Meimiao, which belonged to subtropical climate with average annual temperature of 19-20"!# and 1200-1500 mm average annual rainfall. While K. nepalensis was located at low elevations in south Mandalay, about 200 m above sea level, in a tropical monsoon climate with annual average temperature of 23-29"!#!and an average of 800-1000 mm annual precipitation.

Based on the above results we concluded that the climate was possibly the main factor that deduced the inter-species diversity of genus Kerria in different environment. The unique taxonomic statuses of two species in China were proved by comparison in morphological, cellular, and molecular levels with other species in genus Kerria. Evidence from this study showed the commercial species in China was K. yunnanensis and this species was very different from K. chinensis, the commercial species of Thailand. K. ruralis and K. lacca, the main commercial species in India, differed in morphological characters, ecological location, chromosome structure, and genetic material. The two species of lac production in Myanmar were K. pusana and K. nepalensis, of which, K. pusana is most widely used in lac industry. The main commercial species of lac insect in various countries, with differences in climate and environment, belong to different species.

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Species	Host	Sample size	Location	GenBank accession No. of EF1α gene
K.yunnanensis Ou et Hong	Dalbergia assamica Benth	50♀	China	EU781498
K.lacca Kerr.	Zizyphus mauritiana Lam.	50♀	India	EU781493
K.chinensis Mahdihassan	Litchi chinensis Sonn.	50♀	Thailand	EU781492
K.nepalensis Varshney	Ficus.spp.	50♀	Burma	EU781494
K.pusana Misra	Samanea saman (Jacq.) Merr.	50♀	Burma	EU781495
K.ruralis Wang	Pueraria tonkinensis Gagn.	50♀	China	EU781496
<i>K.sindica</i> Mahdihassan	Ficus racemose Linn.	50♀	Pakistan	EU781497

able 2. Life history	of Kerria yunanensis (Female)			
	1st larval stage	2nd larval stage	3rd larval stage	Adult
Summer Crop	20d	15d	15d	100d
Winter Crop	50d	45d	30d	90d

Table 3. Life histor	y of Kerria yunanensis (Mal	e)		
	1st larval stage	2nd larval stage	Pupae (Propupa)	Adult
Summer Crop	20d	18d	12d	8d
Winter Crop	50d	60d	20d	15d

	1st larval stage	2nd larval stage	3rd larval stage	Adult
Summer Crop	40d	20d	15d	90d
Winter Crop	20d	15d	10d	170d

Table 5. Life histo	ry of Kerria ruralis (Male)			
	1st larval stage	2nd larval stage	Pupae (Propupa)	Adult
Summer Crop	40d	30d	15d	25d
Winter Crop	20d	20d	15d	15d

No.	Characters	K.yunnanensi	K.lacca	K.pusana	K.nepalensis	K.sindica	K.chinensis	K.ruralis
1	Length of anal tubercle	0.22±0.097	0.48±0.131	0.10±0.060	0.62±0.245	0.31±0.055	0.71±0.276	0.16±0.0
2	Width of anal tubercle	0.22±0.103	0.43±0.060	0.17±0.059	0.41±0.116	0.40±0.029	0.45±0.123	0.30±0.04
3	Length/ Width ratio of anal tubercle	1.00±1.273	1.12±0.227	0.59±0.381	1.51±0.428	0.78±0.128	1.58±0.570	0.53±0.2
4	Numbers of anal ring setae	8.80±1.710	3.90±3.356	8.03±1.903	9.66±2.663	5.00±2.853	6.40±2.908	8.90±2.2
5	Length of anal ring setae	0.22±0.025	0.16±0.059	0.17±0.036	0.22±0.048	0.16±0.038	0.22±0.033	0.22±0.0
6	Body length	1.41±0.243	2.51±0.595	1.00±0.213	2.70±0.481	2.62±0.454	6.34±0.664	1.36±0.3
7	Body width	1.04±0.163	2.51±3.973	0.80±0.166	1.69±0.293	1.83±0.373	2.25±0.446	0.99±0.1
8	Body length/ Body width ratio	1.35±0.335	1.00±0.249	1.25±0.167	1.60±0.305	1.43±0.302	2.82±0.616	1.37±0.3
9	Length of dorsal spine	0.18±0.032	0.17±0.054	0.13±0.060	0.18±0.060	0.12±0.033	0.19±0.022	0.14±0.0
10	Length of dorsal spine/ Length of branchial crater ratio	1.50±0.301	1.06±0.342	1.63±0.753	1.80±0.474	0.92±0.291	1.58±0.342	1.40±0.2
11	Width of dorsal spine pedicel	0.07±0.022	0.13±0.037	0.04±0.020	0.10±0.021	0.13±0.021	0.12±0.015	0.06±0.0
12	Length of dorsal spine pedicel	0.04±0.020	0.18±0.060	0.04±0.025	0.11±0.027	0.16±0.039	0.24±0.112	0.05±0.0
13	Length of dorsal spine/ Length of dorsal spine pedicel ratio	4.50±4.559	0.94±0.551	3.25±2.703	1.64±0.764	0.75±0.231	0.79±0.275	2.80±1.1
14	Length of keratinization trail of anterior spiracle	0.08±0.076	0.12±0.043	0.01±0.019	0.18±0.060	0.14±0.031	1.26±0.452	0.00±0.0
15	Distance between anterior spiracle and branchia tube	0.06±0.030	0.37±0.133	0.07±0.024	0.25±0.067	0.15±0.068	0.36±0.086	0.08±0.0
16	Width of anterior spiracle	0.22±0.029	0.19±0.030	0.16±0.056	0.18±0.043	0.16±0.016	0.16±0.037	0.22±0.0
17	Length of anterior spiracle	0.12±0.012	0.11±0.017	0.09±0.035	0.12±0.024	0.11±0.011	0.11±0.030	0.14±0.0
18	Height of branchia tube	0.03±0.033	0.31±0.220	0.00±0.000	0.29±0.081	0.18±0.061	0.60±0.106	0.01±0.0
19	Numbers of branchial pores	11.07±1.780	9.33±1.446	6.23±1.104	0.00±0.000	5.93±1.701	7.10±2.264	6.10±1.6
20	Length of branchial crater	0.12±0.014	0.16±0.015	0.08±0.023	0.10±0.024	0.19±0.016	0.12±0.020	0.10±0.0
21	Width of branchial crater	0.10±0.014	0.14±0.015	0.07±0.020	0.07±0.024	0.13±0.013	0.09±0.020	0.08±0.0
22	Area of branchial crater	0.04±0.006	0.07±0.015	0.02±0.010	0.04±0.014	0.05±0.005	0.05±0.020	0.03±0.0
23	Numbers of marginal pore clusters	24.60±5.654	16.03±3.943	14.57±11.944	19.24±6.162	15.17±3.770	16.07±4.806	23.73±6.
24	Length of mouth parts	0.64±0.274	0.37±0.149	0.70±0.242	0.46±0.184	0.51±0.272	0.60±0.209	0.39±0.2
25	Width of tentorium	0.17±0.018	0.15±0.017	0.13±0.034	0.15±0.026	0.16±0.014	0.16±0.020	0.16±0.0
26	Width of post oral lobes	0.09±0.021	0.08±0.016	0.08±0.028	0.08±0.017	0.09±0.014	0.09±0.017	0.09±0.0

Table 7. The comparison of	Raryotypic analysis among seven species of lac insect
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S	pecies	Centromere location type	Constitution of relative length	Karyotype type
K	. lacca	6m+12T	2L + 6M2 + 6M1 + 4S	3B
K	. sindica	4m+2sm+12T	2L + 8M2 + 4M1 + 4S	3B
K	. ruralis	8m+10T	4L + 4M2 + 4M1 + 6S	3B
K	. yunnanensis	6m+2sm+10T	10M2 + 4M1 + 4S	3A
K	. pusana	4m+14T	2L + 6M2 + 8M1 + 2S	3A
K	. nepalensis	4m+14T	4L + 4M2 + 4M1 + 6S	3B
K	. chinensis	10m+8T	2L + 8M2 + 4M1 + 4S	2B

Table 8. Karyotype resemblance-near coefficients (below diagonal) and evolutionary distance (above diagonal) in seven species of lac insects

	K. chinensis	K. yunnanensis	K. lacca	K. ruralis	K. pusana	K. nepalensis	K. sindica
K. chinensis	-	0.1772	0.3565	0.1584	0.6428	0.618	0.3951
K. yunnanensis	0.8376	-	0.1741	0.0317	0.431	0.4125	0.2107
K. lacca	0.7001	0.8402	=	0.1649	0.3291	0.3234	0.0478
K. ruralis	0.8535	0.9688	0.848	-	0.4545	0.4335	0.2011
K. pusana	0.5258	0.6498	0.7196	0.6348	-	0.0576	0.2711
K. nepalensis	0.539	0.662	0.7237	0.6482	0.9441	-:	0.2751
K. sindica	0.6736	0.81	0.9533	0.8178	0.7625	0.7595	-

Table 9. Nei's genetic distance (below diagonal) and genetic identity (above diagonal) among seven species of lac insect generated by RAPD

	K.	K. lacca	K. pusana	K.	K. ruralis	K. sindica	K. chinensis
	yunnanensi	1	100000000000000000000000000000000000000	nepalensis	The state of the state of	100000000000000000000000000000000000000	and a second second
K. yunnanensis	****	0.5328	0.7739	0.777	0.796	0.624	0.5605
K. lacca	0.6297	****	0.5718	0.6249	0.571	0.6651	0.5875
K. pusana	0.2563	0.5589	***	0.7766	0.8308	0.6021	0.5186
K. nepalensis	0.2523	0.4702	0.2529	****	0.7732	0.657	0.63
K. ruralis	0.2282	0.5604	0.1854	0.2572	****	0.6243	0.5999
K. sindica	0.4716	0.4079	0.5037	0.4201	0.4711	****	0.5292
K. chinensis	0.5789	0.5319	0.6567	0.4621	0.511	0.6365	***

Test	Growth status	Lac secretion status	F1	F2
ÇK. lacca ×∂K. sindica	Normal growth with complete life cycle	Producing contiguous encrustation, capacity to produce less than natural individualS	producing first filial generation but dead in adult stage	None
$\cQentline K$. sindica $ imes \cap{3}$ K. lacca	Normal growth with complete life cycle	Producing contiguous encrustation, capacity to produce less than natural individual	producing first filial generation but dead in adult stage	None
♀K. yunnanensis ×♂K. lacca	Mortality in adult stage with incomplete life cycle	Secreting seedlac	None	None
$ abla K$. yunnanensis $ imes \partial K$. sindica	Mortality in adult stage with incomplete life cycle	Secreting thin layer of shellac	None	None
♀K. lacca × &K.yunnanensis	Mortality in adult stage with incomplete life cycle	Secreting seedlac	None	None
Parthenogenesis (Control)				
♀ K. lacca	Mortality in adult stage with incomplete life cycle	Secreting thin layer of shellac LAC	None	None
♀ K. sindica	Mortality in adult stage with incomplete life cycle	Secreting thin layer of shellac LAC	None	None
♀ K.yunnanensis	Mortality in adult stage with incomplete life cycle	Secreting thin layer of shellac LAC	None	None

Table 11. Environment comparison of seven species of Lac insects

1/	Climate	Distribution
K.yunnanensis	Subtropical climate. Average annual temperature 18.0-20.0 °C, annual precipitation 1200 - 1700mm	China
K. ruralis	Subtropical climate. Average annual temperature 19.0-21.0 °C, annual precipitation 1200 - 1700mm	China
K. lacca	Tropical monsoon climate. Average annual temperature 23.5-26.5 °C, annual precipitation 500-800mm	India, Pakistan , Nepal
K. sindica	Tropical monsoon climate. Average annual temperature above 24.0 °C, annual precipitation 250-400mm	Pakistan, Bangladesh
K. pusana	Subtropical climate. Average annual temperature 19.0-20.0°C, annual precipitation 1200 - 1500mm	India, Burma
K. nepalensis	Tropical monsoon climate. Average annual temperature 23.0-29.0°C, annual precipitation 800-1000mm	India,Burma, Nepal
K. chinensis	Tropical monsoon climate. Average annual temperature 22.0-28.0°C, annual precipitation 1300 - 1900mm	Thailand