Description of a new species of the genus Neopseustis Meyrick, 1909 from China, with a new classification of the genus (Lepidoptera, Neopseustoidea, Neopseustidae)

Siyao Huang¹, Yongxiang Hou¹, Lijuan Zhu¹, Yongqiang Xu², Min Wang¹, Xiaoling Fan¹, Yang Long¹, Wa Da², Liusheng Chen³

¹ Department of Entomology, College of Plant Protection, South China Agricultural University, Guangzhou 510642, Guangdong, China ² Tibet Plateau Institute of Biology, Lhasa 850001, Xizang Autonomous Prefecture, China ³ Guangdong Academy of Forestry, Guangzhou 510520, Guangdong, China

Corresponding authors: Wa Da (tsea2@163.com), Liusheng Chen (lshchen2008@163.com)

Abstract
A new species of the genus Neopseustis Meyrick, 1909, Neopseustis chentangensis S.Y. Huang & Chen sp. nov., which was confirmed by both morphological and molecular methods, is described from Xizang, China. This is currently the westernmost species in Asia of the primitive lepidopteran family Neopseustidae. The new species is externally reminiscent of N. moxiensis Chen & Owada, 2009; however, it can be easily distinguished from the latter by comparison of the male genitalia and is further distinguished by the large genetic distance in DNA barcodes (COI). The adult and genitalia of the new and similar species have been illustrated. Utilizing our new data, a new classification of the genus is provided, with its members subdivided into four species groups: the meyricki-group, the moxiensis-group, the bicornuta-group, and the chentangensis-group, which are supported by both molecular and morphological evidence. A checklist of the genus and a key to the species groups are also provided.

Keywords
Classification, Himalaya, India, Neopseustina, new species, Sichuan, Xizang
Introduction

The family Neopseustidae is a small and archaic lepidopteran family known only by four genera and 14 species and with a peculiar disjunct distribution. Ten of these species are found in Southeast Asia, and the rest are found in South America (Davis 1975; Davis and Nielsen 1980; Davis and Nielsen 1985; Liao et al. 2021). Kristensen (1999) listed several probable autapomorphies for the family, mainly taken from the head, thorax, and abdomen, including the facial scales being restricted to paired lateral and usually swollen patches, the prominent apodemal plate invaginated from the upper base of the propectocoxal bridge, the male sternum VII with medial spinose process, etc. Faucheux et al. (2006) studied the antennal flagellum sensilla of several neopseustid species and stated that one sensillum type, called “multiporous large sensillum basiconicum” in their work, is unknown in other lepidopterans except Neopseustidae; thus, the presence of such a sensillum constitutes an autapomorphy of the family. Recent molecular studies have brought new knowledge concerning the phylogenetic position of the family. Mutanen et al. (2010) and Regier et al. (2013) recovered the clade Acanthopteroctetidae+Neopseustidae, but with weak support. Kristensen et al. (2015) found that with the discovery of Aenigmatineidae, the clade Acanthopteroctetidae+Aenigmatineidae+Neopseustidae (abbreviated as the AAeN clade in that work) was strongly supported, and Aenigmatineidae was found to be sister to Neopseustidae. The close relationship is supported by the sharing of a strong precoxal bridge between the prothoracic pleuron and sternum. Moreover, the AAeN clade was found to be sister to all the Heteroneura (Kristensen et al. 2015). Regier et al. (2015) also reported the grouping of Neopseustidae and Acanthopteroctetidae, and this clade was found to be supported by the presence of the smooth intercalary sclerotization and the alignment of antennal scale sockets in longitudinal rows in the antenna. Regier et al. (2015) also suggested that the former monotypic Neopseustoidea should include also the Acanthopteroctetidae and Aenigmatineidae, and that together they form the sister group to Heteroneura. Externally, Neopseustidae adults are small to medium-sized moths with long antennae and semitransparent, thin-scaled wings, and they resemble some families in the order Neuroptera. Little is known about the biology of Neopseustidae. Adults can be active during the day or night, flying above bushes or attracted to light traps (Liao et al. 2021; present study). As for the immature stages, Grehan (1991) suggested that the disjunct distribution of the plant family Lardizabalaceae fitted well with that of the family Neopseustidae, but no feeding had ever been recorded. Regier et al. (2015) reported an astonishing parasitoid immature stage on Limacodidae of Neopseustis meyricki Hering, 1925, but later the larvae of this Taiwanese species have been found to feed on Ampelopsis brevipedunculata var. hancei, family Vitaceae (DearLep 2021), suggesting that the former record was based on an error. For other Neopseustidae there is no information on immatures.

To date, two genera and seven species have been recorded from mainland China, which are distributed in Henan, Sichuan, Guizhou, Hunan, Guangxi, and Guangdong provinces (Davis 1975; Chen et al. 2009; Liao et al. 2021). Xizang Autonomous
Prefecture, also known as Tibet, is a biological hotspot region located in southwestern China and is well known for its various biotopes. Due to the diverse vegetation types found at different altitudes, this area is home to many families of Lepidoptera, and new discoveries are frequently reported. Neopseustidae are currently unknown for the Xizang fauna. During a survey conducted in May 2021, the first author unexpectedly captured a strange looking individual of this intriguing family from Chentang Town, Xigaze City, located in southern Xizang. After careful examination, this individual has been proven to be an unknown species, which is described herein. This is currently the westernmost distribution record of the genus *Neopseustis* in Asia, and it is also the first record of the family in Xizang. Furthermore, we provide a new classification for the species in the genus *Neopseustis*, based both on molecular and morphological evidence.

**Materials and methods**

**Morphological study.**

Specimens examined were collected during daytime, using an insect net, or with a light trap at night and subsequently deposited in the collection of the South China Agricultural University (SCAU), Guangzhou. Photographs of the adult and the habitat of the new species were taken using a Sony DSC-RX100 v. 1.00 camera. The abdomens were removed and macerated in 10% NaOH for about 2 min at about 95 °C for dissection of the genitalia. The genitalia were removed from the abdomen and mounted in glycerin for photographing. Photographs of the genitalia of the new species were taken under a Keyence VHX-5000 digital microscope, and those of other taxa were taken under a Zeiss SteReo Discovery V.12 digital microscope. Photographs of adults and genitalia were processed using Adobe Photoshop CS5 software. The terminology for adults and genitalia follows Davis (1975) and Liao et al. (2021).

**Molecular analysis.**

Our molecular analysis comprised 19 samples, six of which are newly obtained COI sequences for DNA barcoding. Detailed information on these samples is provided in Table 1. Three COI sequences of three species of the genus *Apoplania* Davis, 1975, two sequences of the monobasic genus *Synemopora* Davis & Nielsen, 1980, and two sequences of one species of the genus *Neopseustis* were downloaded from BOLDSystem (www.boldsystems.org). Five sequences belonging to three species of the genus *Neopseustis* and a sequence of *Endoclita davidi* (Poujade, 1886), which was used as the outgroup in our phylogenetic analysis, were downloaded from NCBI (www.ncbi.nlm.nih.gov). The details of protocols for DNA extraction, amplification, and sequencing have been provided in previous publications (Fan et al. 2016; Tang et al. 2017; Huang et al. 2019). The sequences were aligned using Clustal W (Thompson et al. 1997) implemented in
MEGA v. 7.0 (Kumar et al. 2016) with default parameters, and genetic distances were calculated using Kimura-2-parameter models implied by the same software. Maximum likelihood analyses were performed using IQ-tree v. 2.1.3 (Minh et al. 2020) with the branch support values evaluated by 1000 ultrafast bootstrap (UFBS) replicates (Minh et al. 2013) on the web server (http://iqtree.cibiv.univie.ac.at/). We considered the branch support strong when the UFBS was 95 or higher. Genetic distances were calculated using the Kimura-2-parameter models implied by the same software. All sequences were submitted to GenBank under the submission numbers OK148463 to OK148468. The specimens with voucher numbers CT1, BX1, YJ1, YJ2, MX1, and SZ1 were deposited in SCAU.

**Taxonomy**

**Genus Neopseustis Meyrick, 1909**

*Neopseustis* Meyrick, 1909: 436.

**Type species.** *Neopseustis calliglauca* Meyrick, 1909, by monotypy. [Type locality: Khasi Hills, Assam, India].

**Table 1.** Voucher information and GenBank accession numbers for COI sequences of the Neopseustidae specimens and outgroup in this study. Newly obtained sequences are indicated by an asterisk (*).

| Taxon                          | Locality       | Date   | Voucher Number | Accession Number |
|-------------------------------|----------------|--------|----------------|------------------|
| *Neopseustis chentangensis* S.Y. Huang & Chen sp. nov. | Xizang, China | V.2021 | CT1            | OK148463*        |
| *Neopseustis rectangnatha* Liao, Chen & Huang, 2021 | Hunan, China  | VIII.2020 | HAUHL039474 | MW804623         |
| *Neopseustis rectangnatha* Liao, Chen & Huang, 2021 | Hunan, China  | VIII.2020 | HAUHL039473 | MW804622         |
| *Neopseustis rectangnatha* Liao, Chen & Huang, 2021 | Hunan, China  | VI.2020  | HAUHL040282 | MW804609         |
| *Neopseustis archiphenax* Meyrick, 1928 | Henan, China  | VII.2002 | LNAUT030–14   | N/A              |
| *Neopseustis archiphenax* Meyrick, 1928 | Henan, China  | VII.2002 | LNAUT031–14   | N/A              |
| *Neopseustis sinenii* Davis, 1975 | Sichuan, China | VII.2009 | BX1           | OK148464*        |
| *Neopseustis sinenii* Davis, 1975 | Sichuan, China | VII.2009 | YJ1           | OK148465*        |
| *Neopseustis meyricki* Hering, 1925 | Taiwan, China | N/A     | LS-06-0068   | GU828566         |
| *Neopseustis mociensis* Chen & Owada, 2009 | Sichuan, China | VIII.2004 | MX1           | OK148466*        |
| *Neopseustis fanjingshana* Yang, 1988 | Hunan, China  | VIII.2019 | HAUHL041880  | MW804624         |
| *Neopseustis fanjingshana* Yang, 1988 | Hunan, China  | VIII.2008 | SZ1           | OK148467*        |
| *Neopseustis bicornuta* Davis, 1975 | Sichuan, China | VII.2009 | YJ2           | OK148468*        |
| *Apoplania valdiviana* Davis & Nielsen, 1985 | Cautín, Chile | XII.1982 | LNAUT029–14  | N/A              |
| *Apoplania penai* Davis & Nielsen, 1980 | N/A            |        |               |                  |
| *Apoplania chilensis* Davis, 1975 | Curico Las Tablas, Chile | II.1985 | LNAUT019–14  | N/A              |
| *Synemopa andaeae* Davis & Nielsen, 1980 | Sagrario Puerto, Argentina | II.1979 | LNAUT041–14  | N/A              |
| *Synemopa andaeae* Davis & Nielsen, 1980 | Aguas Calientes, Argentina | II.1979 | LNAUT042–14  | N/A              |
| *Endoclita davidi* (Poujade, 1886) | Hunan, China  | XI.2015 | HN20170409020 | KY928030         |
Neopseustis chentangensis S.Y. Huang & Chen sp. nov.

http://zoobank.org/9E16636E-F0EE-4738-9DD9-4A6259EB96B6

Figures 1, 3–10

Type material. Holotype: male, altitude 2600 m, 23.V.2021, Chentang Town, Dingjie County, Xigaze City, Xizang Autonomous Prefecture, P.R. China, leg. Siyao Huang, voucher number and dissection number CT1 (SCAU).

Diagnosis. Externally, *N. chentangensis* resembles *N. moxiensis* Chen & Owada, 2009 (Fig. 2, 11–12) from Moxi, western Sichuan, share a fuscous ground colour on both wings. However, the new species can be immediately distinguished from *N. moxiensis* by the combination of the following characters: smaller size (length of forewing 8.7 mm vs 9 mm in holotype of *N. moxiensis*), narrower forewing (slightly broader in *N. moxiensis*), patches along forewing costa slenderer and darker (patches along forewing costa thicker and lighter in *N. moxiensis*), narrower hindwing and light fuscous ground colour (broader hindwing and light yellowish brown ground colour in *N. moxiensis*), and more uniform fringe in both wings (cilia clearly chequered, especially in hindwing in *N. moxiensis*). In the male genitalia, *N. chentangensis* can be easily distinguished from *N. moxiensis* by the shape of the latero-posterior process of anellus, which is long, robust, and L-shaped; the distal end is deeply bifurcated and forms two sharp processes bending anteriorly (in *N. moxiensis*, the latero-posterior process of anellus is not L-shaped and bent anteriorly at the tip.). The tegumenal lobe is significantly slenderer after it is flattened (in *N. moxiensis* the tegumenal lobe is much broader when it is flattened), the valvae lack the uncinate process apically, and long and thick processes ventrally (both processes present in *N. moxiensis*). The anterior arms of the vinculum are more slender (these arms are broader and shorter in *N. moxiensis*). From the other congeners, *N. chentangensis* can be simply distinguished by the shape of its latero-posterior process of anellus mentioned above.

Description. Adult: length of forewing 8.7 mm. Antennae brownish dorsally. Head, thorax, and abdomen uniformly brownish. Forewing nearly oval, apex slight-
Figures 3–12. Male genitalia of *Neopseustis* spp. 3–10 *Neopseustis chentangensis* sp. nov., holotype, dissection number CT1 3 genitalia capsule in natural shape with anellus-juxta-parameres removed, dorsal view 4 same, in ventral view 5 genitalia capsule flattened with anellus-juxta-parameres removed 6 genitalia capsule in natural shape with anellus-juxta-parameres removed, in lateral view 7 anellus-juxta-parameres in natural shape, in dorsal view 8 same, in ventral view 9 same, in lateral view 10 anellus-juxta-parameres flattened, in ventral view 11, 12 *Neopseustis moxiensis*, holotype, dissection number MX1 11 anellus-juxta-parameres flattened, in ventral view 12 genitalia capsule flattened with anellus-juxta-parameres removed. J = Juxta; LPA = lateroposterior process of anellus; PE = parameres; TB = transverse bar. Scale bar: 1 mm (Figures 3–10).

Forewing ground color pale yellowish fuscous, with four fuscous patches along costa to apex. Several irregular black or brownish transverse lines present in the median and submarginal zones. A row of brownish spots extending from apex to anal angle along termen. Fringe fuscous from apex to anal angle, slightly checkered with
creamy white in dorsum. Hindwing oval, ground color uniformly light fuscous. Hindwing apex with light yellowish spot at the marginal zone. Fringe generally fuscous from apex to anal angle and slightly checkered with creamy white around anal angle.

**Male genitalia:** Uncus fused with tegumen, bifurcate basally and forming two short and distally rounded lobes. Gnathos strongly sclerotized thoroughly, consisting of a medially curved, short, and robust distal process and a large and thick base. Socii rounded, densely setose. Tegumen lobe slightly curved outwards beyond the base and gradually narrowing towards its tip. Valvae totally fused with vinculum, broad and nearly trapezoid in natural shape. Vinculum broad posteriorly, abruptly narrowing anteriorly and forming long and slender arms. Lateroposterior process of anellus generally L-shaped, thick, and robust, with the tip deeply bifurcate and forming two sharp processes bending anteriorly. Two denticles present at the upper margin of dorsal process. Paired processes of anellus absent. Transverse bar in lateral view obtuse-triangular and slightly bending upwards near tip, while in dorsal and ventral views generally triangular with the lower angles shallowly bifurcate. Juxta in lateral view slightly curved outwards and nearly broad Y-shaped in dorsal and ventral views. Parameres short and setose-like, weakly sclerotized, situated between the two lateroposterior processes of anellus.

**Female.** Unknown at present.

**Bionomics.** The holotype of *N. chentangensis* was spotted weakly flying above bushes during the daytime at an altitude about 2600 m. The collecting site (Fig. 13) is located at the edge of a forest along a road in a valley.

**Distribution.** Currently only known from the type locality, Chentang Town (Fig. 14).

**Etymology.** The specific epithet chentangensis is derived from the type locality, Chentang Town.

**Molecular analysis.** The Kimura-2-parameter distance of the genus *Neopseustis*, based on COI barcoding, is given in Table S1. The maximum interspecific divergence occurred between *N. chentangensis* and *N. moxiensis*, which was 11.7%, and the minimum interspecific divergence occurred between *N. fanjingshana* and *N. bicornuta*, which was 1.5%. According to the table, *N. chentangensis* is genetically distinct from its congeners, with the genetic divergence varying from 7.6 to 11.5%. Based on the ML tree (Fig. 15) constructed using the COI barcoding region, the genus *Neopseustis* was monophyletic (UFBS = 98), and subsequently diverged into four clades, with three of them receiving strong support (UFBS > 95). *Neopseustis chentangensis* was found to be sister to all the remaining taxa in the current study.

**Discussion**

Although Davis (1975) and Liao et al. (2021) considered that the genus *Neopseustis* should be subdivided into two groups based on the morphology of male genitalia and molecular phylogenetic analysis, we consider that this genus may actually comprise of at least four groups, after utilizing more data from previously unsampled taxa. The first group, as already recognized by Davis (1975) and Liao et al. (2021), consists of
Figures 13, 14. 13 Collecting site of *Neopseustis chentangensis* in Chentang Town, Xizang 14 Distribution map of some *Neopseustis* spp. in Asia.
Figure 15. Phylogenetic tree of Neopseustidae based on an analysis of the COI barcoding region, using the maximum likelihood method. The genus Neopseustis is divided into four groups with their corresponding anellus-juxta-parameres illustrated on the left side.
N. rectagnatha Liao, Chen & G.H. Huang, 2021; N. meyricki Hering, 1925; N. archiphænax Meyrick, 1928; and N. sinensis Davis, 1975, and is called the meyricki-group. The second group consists only of N. moxiensis Chen & Owada, 2009, and is called the moxiensis-group. The third group, consisting of N. fanjingshana Yang, 1988 and N. bicornuta Davis, 1975, is called the bicornuta-group and probably also includes the unsampled type species, N. calliglauca Meyrick, 1909, based on the morphology of its anellus-juxta-parameres. The fourth group consists of only N. chentangensis S.Y. Huang & Chen and is called the chentangensis-group. Among these four groups, except for the meyricki-group which is unique in having well-developed parameres and a narrow, short, and forked lateroposterior process on the anellus, the moxiensis, bicornuta, and chentangensis groups all share ill-developed parameres, but they can be distinguished from each other by the combination of features in the male genitalia. The moxiensis-group is characterized by the latero-posterior process of the anellus covered by dense spinules from middle to distal end, in addition to the valvae which have an uncinate process apically and a long and thick process ventrally. The bicornuta-group is characterized by the latero-posterior process of the anellus smooth from middle to distal end and the absence of ventral process in the valvae. The chentangensis-group is characterized by latero-posterior process of anellus long, L-shaped with apex deeply bifurcating and bending anteriorly, and gnathos with a large and thick base.

It is rather intriguing that although N. chentangensis is similar externally only to N. moxiensis in Neopseustis, among the whole genus, they have the greatest genetic divergence. Their male genitalia structures are also considerably different from each other, suggesting that the relationship between them is distant. We believe that their external similarity may probably due to their parallel evolution under similar environments. Unlike their relatively whitish congeners inhabiting the mid- and lower-elevation mountainous areas in India, mainland China, and Taiwan, these two species all inhabit high mountainous areas above 2500 m, and the similar cool climate in high elevation areas in western Sichuan and southern Xizang. This probably may have led to the evolution of their dark wing coloration which can help them absorb heat faster. This assumption is also supported by the studies of Wu et al. (2019), Trullas et al. (2007), and Pereboom and Biesmeijer (2003), who produced similar conclusions.

The former westernmost record of the genus Neopseustis is the type species N. calliglauca, which is found in the Khasi Hills in India. The current record of this new species is situated about 520 km northwest of Khasi Hill, and thus is currently the westernmost record of the genus. The discovery of N. chentangensis in Chentang, on the southern slope of the Himalayas in Xizang, suggests that the investigation of the microlepidopteran fauna is still inadequate in remote areas along the Himalaya. The collection site of the new species is very close to the border of China and Nepal, and Neopseustidae are unknown in Nepal. It can be expected that this species or other new species will someday be discovered in Nepal or Bhutan. Moreover, Neopseustidae are also expected in the southeastern part of Xizang, where no species are currently found. It is possible that the absence of this family there is only due to a lack of surveys, as poor transportation conditions in past decades makes this paradise of moths difficult to access.
Checklist of the genus *Neopseustis* Meyrick, 1909

**meyricki-group**

*N. archiphenax* Meyrick, 1928  
Distribution. Myanmar, China (Sichuan, Henan)

*N. meyricki* Hering, 1925  
Distribution. China (Taiwan)

*N. rectagnatha* Liao, Chen & G.H. Huang, 2021  
Distribution. China (Hunan, Guangxi, Guangdong)

*N. sinensis* Davis, 1975  
Distribution. China (Hunan, Sichuan)

**moxiensis-group**

*N. moxiensis* Chen & Owada, 2009  
Distribution. China (Sichuan)

**bicornuta-group**

*N. bicornuta* Davis, 1975  
Distribution. China (Sichuan)

*N. calliglauca* Meyrick, 1909  
Distribution. India (Khasi Hills, Meghalaya)

*N. fanjingshana* Yang, 1988  
Distribution. China (Guizhou, Hunan)

**chentangensis-group**

*N. chentangensis* S.Y. Huang & Chen, sp. nov.  
Distribution. China (Xizang)

Key to the species-groups of the genus *Neopseustis* based on male genitalia structures

1. Parameres well developed and narrow; latero-posterior process of anellus short, and forked ................................. *Neopseustis meyricki-group*
   - Parameres poorly developed ........................................ 2

2. Latero-posterior process of anellus long, L-shaped with apex deeply bifurcate, bending anteriorly ............................. *Neopseustis chentangensis-group*
   - Latero-posterior process of anellus apex not bifurcate and pointed posteriorly .........................................................3

3. Latero-posterior process of the anellus covered by dense spinules from middle to distal end; valvae with an uncinate process apically and a long and thick process ventrally........................................... *Neopseustis moxiensis-group*
   - Latero-posterior process of the anellus smooth from middle to distal end; valvae without a ventral process ..................... *Neopseustis bicornuta-group*
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References

Chen LS, Owada M, Wang M, Long Y (2009) The genus Neopseustis (Lepidoptera: Neopseustidae) from China, with description of one new species. Zootaxa 2089(1): 10–18. https://doi.org/10.11646/zootaxa.2089.1.2

Davis DR (1975) Systematics and zoogeography of the family Neopseustidae with the proposal of a new superfamily (Lepidoptera: Neopseustoidea). Smithsonian Contributions to Zoology 210: 1–45. https://doi.org/10.5479/si.00810282.210

Davis DR, Nielsen ES (1980) Description of a new genus and the two new species of Neopseustidae from South America, with discussion of phylogeny and biological observations (Lepidoptera: Neopseustoidea). Steenstrupia 6(16): 253–289.

Davis DR, Nielsen ES (1985) The South-American neopseustid genus Apoplania Davis: a new species, distribution records and notes on adult behavior (Lepidoptera: Neopseustidae). Entomologica Scandinavica 15: 497–509. https://doi.org/10.1163/187631284X00325

DearLep (2021) http://dearlep.tw/species.html?namecode=345921 [Accessed on: 2021–9–12]

Fan XL, Chiba H, Huang ZF, Fei W, Wáng M, Sáfánn S (2016) Clarification of the phylogenetic framework of the tribe Baorini (Lepidoptera: Hesperiidae: Hesperiinae) inferred from multiple gene sequences. PLoS ONE 11: e156861. https://doi.org/10.1371/journal.pone.0156861

Faucheux MJ, Kristensen NP, Yen SH (2006) The antennae of neopseustid moths: morphology and phylogenetic implications, with special reference to the sensilla (Insecta, Lepidoptera, Neopseustidae). Zoologischer Anzeiger 245: 131–142. https://doi:10.1016/j.jcz.2006.05.004

Grehan JR (1991) A panbiogeographic perspective for pre-cretaceous angiosperm-Lepidoptera coevolution. Australian Systematic Botany 4(1): 91–110. https://doi.org/10.1071/SB9910091

Hering M (1925) Ueber die Gattung Neopseustis Meyrick (Lep.). Mitteilungen aus dem Zoologischen Museum, Berlin 12(1): 141–147.

Huang ZF, Chiba H, Jin J, Kizhakke AG, Wáng M, Kunte K, Fan XL (2019) A multilocus phylogenetic framework of the tribe Aeromachini (Lepidoptera: Hesperiidae: Hesperiinae), with implications for taxonomy and biogeography. Systematic Entomology 44: 163–178. https://doi.org/10.1111/syen.12322

Kristensen NP (1999) The homoneurous Glossata. In: Kristensen NP (Ed.) Lepidoptera, Moths and Butterflies. Vol. 1. Evolution, Systematics, and Biogeography. Walter de Gruyter, Berlin/New York, 51–63. https://doi.org/10.1515/9783110804744.51

Kristensen NP, Hilton DJ, Kallies A, Milla L, Rota J, Walberg N, Wilcox SA, Glatz RV, Young DA, Cocking G, Edwards T, Gibbs GW, Halsey M (2015) A new extant moth family from Kangaroo Island and its significance for understanding early Lepidoptera evolution (Insecta). Systematic Entomology 40: 5–16. https://doi.org/10.1111/syen.12115
Huang et al. *Neopseustis chentangensis* sp. nov. from Xizang, China

Kumar S, Stecher G, Taamura K (2016) MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Molecular Biology and Evolution 33 (7): 1870–1874. https://doi.org/10.1093/molbev/msw054

Liao CQ, Chen LS, Huang GH (2021) Notes on the genus *Neopseustis* Meyrick with description of a new species from China (Lepidoptera: Neopseustidae). Zootaxa 4970(2): 340–352. https://doi.org/10.11646/zootaxa.4970.2.7

Meyrick E (1909) Descriptions of Indian Microlepidoptera. Journal of the Bombay Natural History Society 19(2): 410–437.

Minh BQ, Nguyen MAT, von Haeseler A (2013) Ultrafast approximation for phylogenetic bootstrap. Molecular Biology and Evolution 305: 1188–1195. https://doi.org/10.1093/molbev/msr024

Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, von Haeseler A, Lanfear R (2020) IQ-TREE 2: new models and efficient methods for phylogenetic inference in the genomic era. Molecular Biology and Evolution 37: 1530–1534. https://doi.org/10.1093/molbev/msaa015

Mutanen M, Wahlberg K, Kaila L (2010) Comprehensive gene and taxon coverage elucidates radiation patterns in moths and butterflies. Proceedings of the Royal Society B 277: 2839–2849. https://doi.org/10.1098/rspb.2010.0392

Pereboom J, Biesmeijer J (2003) Thermal constraints for stingless bee foragers: the importance of body size and coloration. Oecologia 137: 42–50. https://doi.org/10.1007/s00442-003-1324-2

Poujade MG-A (1886) [No title]. Annales de la Société entomologique de France, 6e Série 6: cl–cli.

Regier JC, Mitter C, Zwick A, Bazinet AL, Cummings MP, Kawahara AY, Sohn JC, Zwickl DJ, Cho S, Davis DR, Baixeras J, Brown J, Parr C, Weller S, Lees DC, Mitter KT (2013) A large-scale, higher-level, molecular phylogenetic study of the insect order Lepidoptera (moths and butterflies). PLoS ONE 8: e58568. https://doi.org/10.1371/journal.pone.0058568

Regier JC, Mitter C, Kristensen NP, Davis DR, van Nieukerken EJ, Rota J, Simonsen TJ, Mitter KT, Kawahara AY, Yen SH, Cummings MP, Zwick A (2015) A molecular phylogeny for the oldest (nonditrysian) lineages of extant Lepidoptera, with implications for classification, comparative morphology and life-history evolution. Systematic Entomology 40: 671–704. https://doi.org/10.1111/syen.12129

Tang J, Huang ZF, Chiba H, Han YK, Wang M, Fan XL (2017) Systematics of the genus *Zinaida* Evans, 1937 (Hesperiidae: Hesperiinae: Baorini). PLoS ONE 12(11): e0188883. https://doi.org/10.1371/journal.pone.0188883

Thompson JD, Gibson TJ, Plewniak F, Jeanmougin F, Higgins DG (1997) The CLUSTAL_X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Research 25: 4876–4882. https://doi.org/10.1093/nar/25.24.4876

Truallas SC, van Wyk JH, Spotila JR (2007) Thermal melanism in ectotherms. Journal of Thermal Biology 32: 235–245. https://doi.org/10.1016/j.jtherbio.2007.01.013

Wu S, Chang CM, Mai GS, Rubenstein DR, Yang CM, Huang YT, Lin HH, Shih LC, Chen SW, Shen SF (2019) Artificial intelligence reveals environmental constraints on colour diversity in insects. Nature Communication 10: e4554. https://doi.org/10.1038/s41467-019-12500-2

Yang CK (1988) Notes on family Neopseustidae and a new species from Fanjingshan (Lepidoptera: Homoptera). Guizhou Science S1–018: 148–153.
Supplementary material I

Table S1. The Kimura-2-parameter distance on COI sequences between different taxon of the genus *Neopseustis* sampled for the current study
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Data type: molecular data
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