# An "American" silkmoth endemic to Himalayas, part I: life history and natural distribution of Antheraea compta Rothschild, 1899 (Lepidoptera, Saturniidae) 

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#### Abstract

Females of the uncommon Antheraea compta Rothschild, 1899 were collected at the natural habitat in the Tibetan Sub-Himalayas, descendants of which were reared on Quercus yunnanensis (Fagaceae) successfully, with all the preimaginal instars recorded morphologically. Correlated characters revealed a close relationship between this taxon and New World Antheraea spp., suggesting more attention and protection towards this key species is necessary in the future.


## Key Words

chaetotaxy, ecoregion, evolution, Kachin, Meghalaya, morphology, protection, Quercus, SEM, silkworm, Telea, Tibet

## Introduction

Species of Antheraea Hübner, 1819 have long been known in the history of sericulture as excellent sources for non-mulberry silk (Peigler 1993). Research surrounding the genus has always been a focus in Lepidoptera, occupying most of the scientific works on the family Saturniidae around the world because of their spectacular appearances and considerable economic benefits. After a systematic review, Nässig (1991) proposed to classify these silkmoths into three subgenera, i.e., Antheraea, Antheraeopsis Wood-Mason, 1886 and Telea Hübner, 1819. Textiles called Chinese tussah (from Antheraea pernyi (GuérinMéneville, 1855)), Japanese tensan (from Antheraea yamamai (Guérin-Méneville, 1861)) and Indian tasar (from Antheraea paphia (Linnaeus, 1758)) make the subgenus Antheraea the center of attention, while the golden muga silk (from Antheraea assamensis (Helfer, 1837)) produced in Sub-Himalayas renders Antheraeopsis famous (Peigler 2020). The venerable polyphemus moth (Antheraea polyphemus (Cramer, 1775)) within Telea is a cultural icon in American entomology. From southern Siberia to the

Indonesian islands, from central Europe to the Japanese archipelago, the subgenus Antheraea occupies almost all the humid broad-leaved forests from temperate to tropical Eurasia. Antheraeopsis flies in Asia only south of the level of Qinling Mountains, but in any case, populations within these two subgenera are all entirely limited to the Palearc-tic-Indo-Malayan regions. The main habitats of Telea are in the Nearctic-Neotropical realms, but one putative member, namely Antheraea compta Rothschild, 1899 (Fig. 1), is endemic to the northeastern part of the Indian subcontinent (Peigler 1999), this disjunct distribution considered one of the core mysteries in saturniid evolution.

The original type series of $A$. compta was stated to be from "Khasia Hills, Assam, $8 \widehat{o}^{\hat{N}}$, no $q[$ sic $]$ ", regarded from the beginning to be an adelphotaxon allied to $A$. assamensis within Antheraeopsis (Rothschild and Jordan 1899). Such judgment may have been based largely on the superficial features, especially the similar eyespots and the shared natural habitat. Subsequently, Rothschild and Jordan (1901) provided a color photo of a male A. compta and supplemented "The $q$ is similar to the $\widehat{\lambda}$, but has a shorter and broader forewing [sic]". Watson (1912: pl. 2) first


Figure 1. Living adults of Antheraea compta, dorsal views. A. $\delta^{\lambda}$, calm condition; B. $\delta^{\lambda}$, frightened condition; $\mathbf{C} . \not \subset$, calm condition; D. $\rho$, frightened condition. Scale bar: 4 cm .
illustrated the female, captured at the type-locality. Packard (1914: 197) cited this taxon without additional comments, while Conte (1919: 14-15) and Seitz (1926-1928: 511) continued to associate $A$. compta with $A$. assamensis. The first entomologist who recognized the problem was Bouvier (1936: 167) who definitively stated that there is a fairly high morphological similarity between the male genitalia of Himalayan A. compta and Central American Antheraea godmani (Druce, 1892), especially their threelobed valvae, although he did not transfer the former into his checklist for Telea. After a comparison with A. polyphemus, Lemaire (1978: 207) further acknowledged the strong homology with $A$. compta about male genital structures, a taxonomic opinion recognized in more recent works (Nässig 1991; Nässig et al. 1996; d'Abrera 2012: 126-127; Naumann and Löffler 2015), but Meister (2011: 144) still regarded $A$. compta as a member of the gerontogeous subgenus Antheraea, while Paukstadt et al. (2000, 2003) suggested that the issue requires further clarification. Arora and Gupta (1979: 30) drew the venations of Indian Antheraea spp. and noticed the forewing R1 [radial vein 1] of A. compta ("Maghalaya, Cherra Punji, 1220$1524 \mathrm{~m} ., 1$ O$^{2}, 24 . v i i .1931$ [sic]") is relatively longer than that of $A$. assamensis, with a more medial arising point on the anterior margin of the discal cell, but those two authors did not discuss the New World taxa.

Except for Khasi Hills, specimens of $A$. compta have also been collected from "Assam, Jaintia Hills... 12 Juill. 1922 [sic]" and recorded by Bouvier and Riel (1931:53), subtropical forests of the two areas were formerly in Assam, but belong to Meghalaya today (Gupta 2000); furthermore, a male moth from the adjacent Hahim of Assam was reported by Peigler (1999). Bryk (1944) published a more distant record of specimens mentioned as $A$. compta discovered at "Kambaiti, $2000 \mathrm{~m}, 9 . \mathrm{VI}-17 . \mathrm{VI}[s i c] "$ [Kanpaikti Sub-Township, Kachin State] in northeastern Burma by the entomological expedition of René Malaise in 1934 (specimens viewed by

Richard S. Peigler in Stockholm in 2005, personal communication to Liu from Peigler on 31 Aug. 2022). The third core population of the species inhabits southeastern Tibet (Fig. 2), first noted as "Motuo [Metok] County... 2120 m, VII. 2013 [sic]" by Naumann and Löffler (2015) (a separate species? see below), and in 2019, living material from the same ecoregion was unsuccessfully reared on Quercus sp. (Fagaceae), died in $L_{3}$ without detailed description (Naumann and Nogueira 2021), but suggested the caterpillars "fit very well with typical habitus of Telea larvae [sic]".

The above mentioned moths were collected at upland mid-altitudes (ca. 1000-2500 m) in the height of summer, basically June-July (Bouvier and Riel 1931: 53: "11 Déc. 1903 [sic]" possibly invalid), so the species is considered to be a strictly univoltine Himalayan flyer. It must be admitted that specimens of $A$. compta are rare in collections worldwide, so obviously knowledge of its complete preimaginal stages is of great significance for our understanding of the evolutionary history of the genus Antheraea or even the family Saturniidae.

## Material and method

INSIZE 1108 -150C $0-150 \mathrm{~mm} / 0.01 \mathrm{~mm}( \pm 0.02 \mathrm{~mm})$ was used to measure general lengths. XINGYUN FA1204E $120 \mathrm{~g} / 0.0001 \mathrm{~g}( \pm 0.0002 \mathrm{~g})$ for recording weights. BENETECH GM1365 ( $\pm 2 \mathrm{RH} \%$; $\pm 0.3^{\circ} \mathrm{C}$ ) for recording air humidity and temperatures. BENETECH GM1020 ( $\leq 10000$ Lux $\pm 3 \% ; \geq 10000$ Lux $\pm 4 \%$ ) for recording illuminances. PHILIPS TL 6W (UV-A, peak: 365 nm ), was used for fluorescence photographs. All color figures were photographed by a NIKON D5500 with SIGMA $10-20 \mathrm{~mm} f / 4-5.6$ lens or LAOWA $60 \mathrm{~mm} f / 2.8-22$ lens. ZEISS GeminiSEM 360 was used for the SEM observations and photographs, based on sputter-coated samples of 2 ova, $6 \mathrm{~L}_{1}, 1 \mathrm{~L}_{5}, 1 \mathrm{~L}_{6}$ and 1 cocoon of $A$. compta. The


Figure 2. The natural habitat of Antheraea compta in southeastern Himalayas. Mêdog County, Tibet Autonomous Region, China, 2122 m. 28 Jun. 2021.
sources of botanical and other zoological materials will be given in the corresponding paragraphs below.

Terms involving continuity in time or space, the corresponding numbers use the subscript format, including: $\mathrm{L}_{1-6}=1^{\text {st }}-6^{\text {th }}$ larval instars; $\mathrm{T}_{1-3}=1^{\text {st }}-3^{\text {rd }}$ thoracic segments; $\mathrm{A}_{1-10}=1^{\text {st }}-10^{\mathrm{th}}$ abdominal segments.

Structures not having the above relationship between each other are cited: $\mathrm{S}=$ stemma [ocellus]; $\mathrm{BaS}=$ sensillum basiconicum; $\mathrm{ChS}=$ sensillum chaeticum; $\mathrm{TrS}=$ sensillum trichodeum; $\mathrm{PlS}=$ sensillum placodeum; $\mathrm{CaS}=$ sensillum campaniformium; DiS = sensillum digitiformium; $\mathrm{StS}=$ sensillum styloconicum. If necessary, ending with Arabic numerals (non-subscript) for coding (e.g., "S1", "ChS1"). Classification of the sensilla mostly consults some lepidopterous works, i.e., Dethier (1937, 1941), Schoonhoven and Dethier (1966), Grimes and Neunzig (1986a; 1986b) and Faucheux (1999: 247-280). Ryan (2002: 113-114) expressed that the term "chaetic" is applicable to thicker-walled bristles or spines, the word "trichoid" means slender and hair-like, and both can be used for chemo- or mechanosensilla (Zacharuk and Shields 1991; Keil 1999; Shields 2008), so they are sometimes hardly distinguishable externally. Schneider (1964) thought such definitions are strictly morphological, the former "distinguished by a specialized and flexible circular membrane at the base [sic]", and the latter "without any specialized basal cuticular ring serving as articulating membrane $[s i c]$ ". In addition, the word "sensillum (sensilla)" is used in figures only for some specified structures which are borne on antennae, maxillary palpi, maxillary mesal lobes and labial palpi. Here are labelled other primary setae and pores on labrum, mandibles and maxilla-hy-popharynx-labial complex into only "lateral (L)", "medial (M)", "dorsal (D)" and "ventral (V)" in figures, but does not rule out that they may have special sensory functions.

Larval chaetotaxy, described below as far as possible, follows the universal terminology named in earlier Lepidoptera publications, primarily based on Heinrich
(1916), Gerasimov (1935) and Hinton (1946), with also the reviews by Chu (1956), Stehr (1987), Piao and Lee (1998) and Hasenfuss and Kristensen (2003). Additional works treating the standard setal coding of the family Saturniidae were consulted, of which Pease (1960), Heppner and Wang (1987) and Rougerie and Estradel (2008) are examples. The abbreviations of general areas are as follows: $\mathrm{O}=$ ocellar [stemmatal]; $\mathrm{SO}=$ subocellar [substemmatal]; $\mathrm{F}=$ frontal; $\mathrm{AF}=$ adfrontal; $\mathrm{C}=$ clypeal; $\mathrm{G}=$ genal [microgenal/midgenal]; $\mathrm{A}=$ anterior; $\mathrm{V}=$ ventral; SV = subventral; $\mathrm{L}=$ lateral; $\mathrm{M}=$ medial; $\mathrm{XD}=$ tactile dorsal (on the anterior margin of prothoracic shield and near the dorsal midline); $\mathrm{D}=$ dorsal; $\mathrm{SD}=$ subdorsal; $\mathrm{MD}=$ microdorsal [vertical]; $\mathrm{P}=$ parietal [posterodorsal]. Combined with these directional terms, Arabic numerals (non-subscript) and lowercase letters respectively code related primary setae and pores of the head (e.g., seta "SO1", pore "MDa"). The term "pore" for text below may appear to be depressed (pierce the cuticle, or not) or elevated. Further, the term "palpifer" is only used for the maxillae in this work, whereas "palpiger" is considered as a part of the labium, although some entomologists used the latter term within the maxillae (e.g., Peterson 1948: 119; Schoonhoven and Dethier 1966).

Nässig (1989) and Deml and Dettner (2002) classified saturniid scoli focusing on their basal morphology and functionality (secretions), but this article only divides related primary setae into the following four conditions expressed by Roman numerals on the setal map of $\mathrm{T}_{1}-\mathrm{A}_{10}$ (Fig. 3); each of them is considered as a "term" for modifying singular or plural nouns, hyphenated with a general area abbreviation (e.g., seta(e) "D-I", chalaza(e) "D-I", area(s) "D-I"), distinguished in detail with final Arabic numerals (non-subscript) if necessary (e.g., seta(e) "DI1", seta(e) "D-I2"):
$\mathrm{I}=$ For describing a single seta, or the structure/location in which it resides [uni-setal]. If used for the


Figure 3. $L_{1}$ of Antheraea compta, the setal map, lateral view, with only the primary chaetotaxy coded. The setae of cephalic regions and legs $\mathrm{T}_{1-3}$ are not shown, proleg $\mathrm{A}_{10}$ displays the mesal surface, the ventral midline constitutes the bottom margin of $\mathrm{A}_{1-9}$.
plural, for example, several setae within the designated area, then each seta is treated as an individual existing independently which could be objectively distinguished from others.
$\mathrm{II}=$ For describing a pair of setae $(\mathrm{n}=2)$, or the structure/location in which they reside collectively [bi-setal]. If used for the plural, for example, several setal pairs within the designated area, then each pair is treated as an individual existing independently which could be objectively distinguished from others.
III $=$ For describing a cluster of setae $(\mathrm{n}>2)$, or the structure/location in which they reside collectively [multi-setal]. If used for the plural, for example, several setal clusters within the designated area, then each cluster is treated as an individual existing independently which could be objectively distinguished from others.
IV = For describing uncertain condition on the designated structure/location [uni/bi/multi-setal].

## Results

## Morphology of preimaginal instars

Unless otherwise specified, numerical (metric) data was based on single normal individual in this chapter, and the fluorescence tests are only valid for naked eye vision.

## Ova (Figs 4, 8A)

Tri-axial ellipsoid, ca. 3.71 mm (length) $\times 3.32 \mathrm{~mm}$ (width) $\times 2.18 \mathrm{~mm}$ (height) (Fig. 4A-D). The micropylar zone is relatively flat, the external formation of rosette is combined by polygonal imprints of follicle cells (Fig. 4F, G). Other areas of the exochorion covered reticular crests [chorionic sculptures], walls generally heighted in ca. $18.7-30.4 \mu \mathrm{~m}$ and widened to ca. $27.1-34.8 \mu \mathrm{~m}$, the begirded regions shaped into irregular subcircles with comparatively smooth surfaces in most cases (Fig. 4E), but in the circumferential zone of the ovum (except the micropylar zone) (Fig. 4B, C), some of such fragments and their walls formed into special spongiform (Fig. 4H), decorated as darker color (Figs 4A, 8A). The aeropyles (widths $=c a$. 2.32-12.1 $\mu \mathrm{m}$ ) dispersedly located at intersections of the network except the micropylar part (Fig. 4E). Aeropyle crowns (widths $=$ ca. $16.6-37.9 \mu \mathrm{~m}$ ) are not fully developed (incomplete or missing) and only localized to the vertices of the spongy polygons (Fig. 4I). The hydrophilic ovum shell is colored mostly opaque white under dry conditions, but becomes translucent after being sprayed with water causing the internal embryo to be visible (Fig. 4A), while the aeropyles are the major pathways for water desorption possibly (Fig. 4E). The chorionic interior is multihole mainly superimposed by lichen-like fillers (Fig. 4J), such minute lobes are hung loosely in more external (Fig. 4K), and stacked densely as multiple tiers in more internal (Fig. 4L). Chorionic thicknesses (excluding re-


Figure 4. Ova of Antheraea compta. A. Fertilized ovum; a: dry condition; b: wet condition; B-D. Unfertilized ovum (the flat zones are critically sunken); B. Micropylar side of the circumferential zone; C. Non-micropylar side of the circumferential zone; D. Flat zone; E. Flat zone; a: dry aeropyle; b: wet aeropyle; F. Micropylar zone; G. Micropylar zone, the center; H. Circumferential zone, spongiform fragments and their walls; I. Circumferential zone, spongiform fragments and their walls; a: incomplete aeropyle crowns; b: spongiform structure; J. Chorionic cross section; a: exochorion; b: exo-fillers; c: endo-fillers; d: endochorion; K. Chorionic exo-fillers; L. Chorionic endo-fillers $2 \mu \mathrm{~m}$. Scale bars: $2 \mathrm{~mm}(\mathbf{A}) ; 500 \mu \mathrm{~m}(\mathbf{B}, \mathbf{C}) ; 1 \mathrm{~mm}(\mathbf{D}) ; 40 \mu \mathrm{~m}(\mathbf{E}, \mathbf{I}) ; 5 \mu \mathrm{~m}$ (G); $2 \mu \mathrm{~m}(\mathbf{K}, \mathbf{L}) ; 100 \mu \mathrm{~m}(\mathbf{F}, \mathbf{H}) ; 10 \mu \mathrm{~m}(\mathbf{J})$.
ticular crests and micropylar zone) are ca. 44.3-48.2 $\mu \mathrm{m}$. The glue secreted from adult accessory glands [colleterial glands] for affixing the ova is diluted into clear honey brown color, inconspicuously concentrated on the bottom external surface which contacts the vegetation; other areas seemed clean and without glue.

## Larvae

$\mathrm{L}_{1}$ (Figs 3, 5-7, 8A-C)
Head capsule is 1.78 mm width with shiny black appearance, whose anteclypeus is gray and longitudinally folded. Bearing 17 pairs of longer primary setae namely P1, $\mathrm{P} 2, \mathrm{~L} 1, \mathrm{AF} 1, \mathrm{AF} 2, \mathrm{~F} 1, \mathrm{C} 1, \mathrm{C} 2, \mathrm{~A} 1, \mathrm{~A} 2, \mathrm{~A} 3, \mathrm{O} 1, \mathrm{O} 2, \mathrm{O} 3$, $\mathrm{SO} 1, \mathrm{SO} 2$ and SO3, they are pointed and with a smooth surface and sometimes slightly helical-shaped (Fig. 5A, D, E), seta P1 is the longest (length $=1.16 \mathrm{~mm}$ ), seta F1 (length $=182.5 \mu \mathrm{~m}$ ) is the shortest. There are 4 other pairs of minute primary setae (lengths $=\mathrm{ca} .7 .4-8.1 \mu \mathrm{~m}$ ) erectly, i.e., MD1, MD2, MD3 and G1 which ended with rounded apexes (Fig. 5A, B, J). A total of 10 pairs of primary pores appeared on the head capsule, coded as $\mathrm{Pb}, \mathrm{La}, \mathrm{AFa}, \mathrm{Aa}$ ? [Pa?], $\mathrm{Fa}, \mathrm{Oa}, \mathrm{Ob}, \mathrm{SOa}, \mathrm{MDa}$, and Ga (Fig. 5A-E), among them, the pores Fa (width $=21.1 \mu \mathrm{~m}$ ), Ob ( width $=16.6 \mu \mathrm{~m}$ ) and SOa (width $=24.7 \mu \mathrm{~m}$ ) are slight raised (e.g., Fig. 5L), other ones are flat-bottomed or bowl pits with widths of ca. 9.4-12.5 $\mu \mathrm{m}$ (e.g., Fig. 5K, $\mathrm{M}-\mathrm{P}$ ), the deepest depression is pore Ga . A case was found that the sites of pores Pb are asymmetric, i.e., in the same specimen, they are respectively located on the inferolateral and superolateral sides to setae P2, but the former condition is normal. In addition, a few secondary setae and pores could be asymmetrically observed in individuals. S1 (width $=47.7 \mu \mathrm{~m}$ ), S2 (width $=46.3 \mu \mathrm{~m}$ ), S3 (width $=58.9 \mu \mathrm{~m}), \mathrm{S} 4($ width $=50.3 \mu \mathrm{~m}), \mathrm{S} 5($ width $=48.1)$ and S 6 (width $=46.7 \mu \mathrm{~m}$ ) have multiporous surface (e.g., Fig. 5I), their anisotropic wrinkles have observed on dry exuviae and alcohol-preserved specimens (Fig. 5A), but unclear if living larvae have the same feature. S1 and S6 are flatter, whereas S3 is the most elevated.

The antennal sclerites are mostly dark maroon linked with light gray antacoriae. The $2^{\text {nd }}$ antennal segment is multiporous and elongated peanut-shaped, whose lower part of the sclerotized wall is embossed with slight reticular crests forming dense polygons, its longish ChS1 (length $=372.2 \mu \mathrm{~m}$ ) and ChS2 (length $=90.7 \mu \mathrm{~m}$ ) dotted dispersedly with tiny wall-pores (more obvious on ChS 2 ) and standing lateroapically (Fig. 6A-C). Of the segment, BaS 1 is dorsoapical, BaS 2 and BaS 3 are ventroapical, CaS1 (width $=13.2 \mu \mathrm{~m}$ ) is lateroproximal (Fig. 6D), CaS2 and CaS3 (widths $=$ ca. $4.5-4.9 \mu \mathrm{~m}$ ) are lateroapical. Relatively, the $1^{\text {st }}$ and $3^{\text {rd }}$ segments of antenna have fewer pores on the walls, the latter is implanted distal BaS4, BaS5, BaS6 and StS (Fig. 6B, C). Antennal BaS1, BaS 2 and BaS 4 showing fingerprint-like surfaces (lined pores) and rounded tips with similar lengths of ca. 20.2$25.4 \mu \mathrm{~m}$, alike epicuticle also observed on BaS3, BaS6 and BaS5, but the former two owning varying degrees of
forky apexes (BaS3 usually has 2-4 forks and asymmetrically presents in a specimen, BaS6 is always bifurcated) with respective maximum lengths of $15.6 \mu \mathrm{~m}$ and $8.7 \mu \mathrm{~m}$, BaS 5 is an elevated pimple in $5.8 \mu \mathrm{~m}$ height. The external surface of uniporous StS (length $=18.0 \mu \mathrm{~m}$ ) is smoother, exhibiting sharp or uni/bis-papilliform terminals.

The labral anterior surface is sepia, shiny, arrayed primary setae L1, L2, L3, M1, M2 and M3 in pairs with plane ends, seta M2 is the longest with $204.4 \mu \mathrm{~m}$ length and other ones are ca. $84.7-178.5 \mu \mathrm{~m}$. The primary pores $\mathrm{Ma}, \mathrm{Mb}$ and Mc are ca. $9.5-16.11 \mu \mathrm{~m}$ widths, the sizes one by one incrementally (Fig. 5F). The labral posterior surface [epipharynx] is off-white principally but the mid-notch [groove] is sepia. Except the smooth margin, other area covered by white and short denticulations densely, exhibiting pointed and flatted setae M4, M5 and L4 (lengths $=82.9-93.4 \mu \mathrm{~m}$ ), with a digitiform seta L5 in size ca. $56.8 \mu \mathrm{~m}$ (length) $\times 24.3 \mu \mathrm{~m}$ (width), they are aporous. Symmetrically paired pores Md, Me and Mf are obvious, the latter is the biggest up to $17.5 \mu \mathrm{~m}$ width, while other two are reduced to ca. 12.3-14.0 $\mu \mathrm{m}$ (Fig. 5G). Setae L1-5 and M1-5 are clear brown, pores Ma-f are campaniform.

Each mandible has 6 teeth ( 4 outer and 2 inner), primarily bears a pair of neighboring setae L1 (length $=295.9 \mu \mathrm{~m}$ ) and L2 (length $=134.2 \mu \mathrm{~m}$ ), with a pimple-like pore La (width $=27.8 \mu \mathrm{~m}$ ) which is surrounded by dense subcircular pits. The posterior condyle [postartis] is a prominent smooth knob (Fig. 5F, H).

The weakly sclerotized cardo is nearly a semicircle, the stipes forms a " $F$ " shaped sclerite whose longer margin is lying medially and the arm points to the lateral side, its anterior and posterior edges respectively array the primary setae L1 (length $=202.2 \mu \mathrm{~m}$ ) and L2 (length $=213.3 \mu \mathrm{~m}$ ). Setae M1 (length $=218.3 \mu \mathrm{~m}$ ) and M2 (length $=90.2 \mu \mathrm{~m}$ ) could be invariably found on the apical margin of the incomplete annular sclerites, separately belongs to the palpifer and the $1^{\text {st }}$ maxillary palpal segment; the latter has a striate-wrinkled top and connecting with the maxillary mesal lobe [galea], their medial junction zone (facing to the hypopharynx) is membranous and placing many toothed setae (Figs 6E, 7A-C). The sclerotized walls encircling maxillary palpus and mesal lobe are distally multiporous, the apical membranes of the latter and the $2^{\text {nd }}-3^{\text {rd }}$ segments of the former are knot-wrinkled (Fig. 7B, G, I). Maxillary palpal CaS1-4 are lateral, thereinto CaS1 and CaS2 are pits while CaS3 (height $=1.8 \mu \mathrm{~m}$ ) is elevated, but both forms are applicable to CaS4 (Fig. 7A, C, D, H). Each maxillary palpus ended with BaS1-8 (lengths $=$ ca. $5.2-5.6 \mu \mathrm{~m}$ ) which have rounded tips and many campaniform olfactory pores (Fig. 7G). Dorsal sensilla of the $3^{\text {rd }}$ maxillary palpal segment are rough, PlS1, PlS2 and PlS3 in widths of ca. $5.4-7.2 \mu \mathrm{~m}$ always surround the lateral to proximal sides of DiS (length $=11.8 \mu \mathrm{~m}$; widths $=6.6 \mu \mathrm{~m}$ ), distal morphology of the latter is alterable with $2-4$ slight apical forks in different individuals and sometimes showing a sagittal concave midline (e.g., Fig. 7E, F). Each max-


Figure 5. $L_{1}$ of Antheraea compta. A. Ventrolateral view; a: head capsule; b: antenna; c: cervacoria; B. Head capsule, superolateral view; C. Head capsule, lateral view; D. Head capsule, superofrontal view; E. Head capsule, frontal view; a: frons; b: clypeus; a + b: frontoclypeus; c: anteclypeus; d: other area; F. Mouthpart (partly), frontal view; a: anteclypeus; b: labrum; c: mandible; G. Labrum, posterior view [epipharynx]; a: the smooth marginal area; b: the hirsute area; H. Mandible, the inner surface; a: inner teeth; b: posterior condyle; c: adductor; d: abductor; I. The external surface of S3, vertical view; J-P. Primary seta and pores of head capsule; J. Seta MD3, lateral view; K. Pore Pb, vertical view; L. Pore SOa, vertical view; M. Pore Aa? [Pa?], lateroapical view; N. Pore MDa, lateroapical view; O. Pore La. lateroapical view; P. Pore Ga, lateroapical view. Scale bars: $200 \mu \mathrm{~m}(\mathbf{A}, \mathbf{D}, \mathbf{E}, \mathbf{H}) ; 50 \mu \mathrm{~m}(\mathbf{B})$; $150 \mu \mathrm{~m}(\mathbf{C}, \mathbf{F}) ; 80 \mu \mathrm{~m}(\mathbf{G}) ; 2 \mu \mathrm{~m}(\mathbf{I}) ; 6 \mu \mathrm{~m}(\mathbf{J}) ; 5 \mu \mathrm{~m}(\mathbf{K}, \mathbf{N}, \mathbf{P}) ; 15 \mu \mathrm{~m}(\mathbf{L}) ; 7 \mu \mathrm{~m}(\mathbf{M}, \mathbf{O})$.


Figure 6. $L_{1}$ of Antheraea compta. A-D. Antennae. A. a: antacoriae; b-d: the $1^{\text {st }}-3{ }^{\text {rd }}$ segments; B. a, b: the $2^{\text {nd }-} 3^{\text {rd }}$ segments; C. a, b: the $2^{\text {nd }}-3^{\text {rd }}$ segments; D. a: the $2^{\text {nd }}$ segment; E. Ventral view; a: head capsule; b: antenna; c: mandibles; d: stipites; e: palpifer; f-h: the $1^{\text {st }}-3^{\text {rd }}$ maxillary palpal segments; i: maxillary mesal lobe; j : postmentum; k: mentum; l: prementum (posterior); m: spinneret; n: prementum (anterior); o: hypopharynx; F. Ventrolateral view; a: mentum; b: prementum (posterior); G. Anterolateral view; a: prementum (anterior); b: the lateral lobe of spinneret; c: labial palpus; H. Lateral view; $a$ : prementum (anterior); b: the medial lobes of spinneret; c: the lateral lobe of spinneret; I. Anteroventral view; a: prementum (anterior); b: hypopharynx; J. The lateral margin of hypopharynx (faces to the maxilla), lateral view; K. Legs $T_{1}$ (seta L2 is obscured), anterior view; a: coxae; b: femur; c: tibia; d: tarsus; e: pretarsus; f: secondary [subprimary?] seta; L. Pretarsus of leg $\mathrm{T}_{1} ; \mathbf{M}$. Crochets of proleg $\mathrm{A}_{4}$; ventral view;. Scale bars: $100 \mu \mathrm{~m}(\mathbf{A}, \mathbf{E}) ; 20 \mu \mathrm{~m}(\mathbf{B}, \mathbf{D}, \mathbf{F}-\mathbf{I}, \mathbf{M}) ; 10 \mu \mathrm{~m}(\mathbf{C}, \mathbf{J}) ; 200 \mu \mathrm{~m}(\mathbf{K}) ; 60 \mu \mathrm{~m}(\mathbf{L})$.


Figure 7. L of Antheraea compta. A. Dorsolateral view; a-c: the $1^{\text {st }}-3^{\text {rd }}$ maxillary palpal segments; d: maxillary mesal lobe; B. Medial view; a: mandible; $b$ : palpifer; c : the hirsute membranous zone between the $1^{\text {st }}$ maxillary palpal segment and mesal lobe; d , e: the $2^{\text {nd }}-3^{\text {rd }}$ maxillary palpal segments; f: maxillary mesal lobe; g: hypopharynx; C. Lateral view; a: palpifer; b-d: the $1^{\text {st }}-3^{\text {rd }}$ maxillary palpal segments; D. Part of the apex of the lateral area of the $2^{\text {nd }}$ maxillary palpal segment, apical view; E. Apical view; a, b: the $2^{\text {nd }}-3^{\text {rd }}$ maxillary palpal segments; F. Part of the dorsal area of the $3^{\text {rd }}$ maxillary palpal segment; G. The apex of the $3^{\text {rd }}$ maxillary palpal segment, apical view; H. Ventromesal view; a, b: the $2^{\text {nd }}-3^{\text {rd }}$ maxillary palpal segments; c: maxillary mesal lobe; d: hypopharynx; I. Part of the apex of the maxillary mesal lobe, medioapical view. Scale bars: $40 \mu \mathrm{~m}(\mathbf{A}-\mathbf{C}) ; 4 \mu \mathrm{~m}(\mathbf{D}, \mathbf{G}) ; 10 \mu \mathrm{~m}(\mathbf{E}) ; 5 \mu \mathrm{~m}(\mathbf{F}, \mathbf{I}) ; 20 \mu \mathrm{~m}(\mathbf{H})$.
illary mesal lobe arranging flat-topped $\mathrm{ChS} 1, \mathrm{ChS} 2$ and ChS 3 from the lateral to medial, they are multiporous but such perforations of the former are more obvious and on the latter two are thinner. The lengths of ChS 1 are unstable and the measurements based on a single specimen are $54.7 \mu \mathrm{~m}$ and $84.9 \mu \mathrm{~m}, \mathrm{ChS} 2$ and ChS 3 are special chisel shaped and ca. 37.8-41.2 $\mu \mathrm{m}$ lengths. The elevated basal columns of StS1 and StS2 are relatively smooth (less pores) with lengths of ca. 23.6-27.8 $\mu \mathrm{m}$, their terminal dome topped cones are morphologically look like
the maxillary palpal BaS1-8, but lengthened to ca. 8.3$9.0 \mu \mathrm{~m}$ (Fig. 7I) with finer pores. For each maxillary mesal lobe, BaS 1 is terminally uniporous and tapered (length $=14.2 \mu \mathrm{~m}$ ), while BaS2 is rough and reduced to a small wart (height $=3.4 \mu \mathrm{~m}$ ), and, an indeterminate shaped CaS (in the same larva, respectively crater-like and spine-like) was fixedly observed on the ventroapical margin and located close to StS1 (Fig. 7B, H). On the ventral area is a very large subcircle forming a slight protrusion with weak wave-like margins marked as CaS? (Fig. 7H), it is
aporous and seems to only be observed at specific angles with certain light conditions, and its function is unclear.

The postmentum [submentum] is smooth and membranous largely, the proximolateral sclerite [submentales] is adjacent with the maxillary cardo, primary seta V1 (length $=104.1 \mu \mathrm{~m}$ ) is close to the border with the mentum, the latter is a semicircular membrane covered by minute noncellular processes (Fig. 6E, F). The prementum [stipulae] is lyre-shaped with most areas membranous, but a crescent sclerite is inlaid proximally of the posterior surface which abutted with the mentum, primary seta V2 (length $=14.6 \mu \mathrm{~m}$ ) is smooth and rounded. Campaniform pores $\mathrm{Va}, \mathrm{Vb}$ and Vc are ca. $4.9-5.4 \mu \mathrm{~m}$ widths, always present on the posterolateral area of the prementum (Fig. 6F). The cleft spinneret is fleshy and semitransparent, whose linguiform medial lobes [glossae] are hirsute and the pliciform lateral lobes [paraglossae] are fine-grained surfacely (Fig. 6G, H). The distance between the two labial palpi is $238.8 \mu \mathrm{~m}$, each of them dotted with many micropores but more remarkable on the lateral $\operatorname{TrS}$ (length $=47.2 \mu \mathrm{~m}$ ) and medial StS, the basal and setal parts of latter are respectively $16.5 \mu \mathrm{~m}$ and $68.0 \mu \mathrm{~m}$ lengths (Fig. 6G). The distal area of the hypopharynx is smooth but the posterior lingua is covered by numerous spines (Fig. 6I, J).

The setal map and related statistics are provided here (Fig. 3; Table 1). Grounded with bright yellow on $\mathrm{T}_{1}-\mathrm{A}_{10}$ generally but the cervacoria and venter are lower saturation. The dorsal area of $\mathrm{T}_{1}$ dotted with three black spots, one shown betweenthe pair of scoli XD-III and other two exhibited on the dorsolateral respectively. Each of $\mathrm{T}_{2}-\mathrm{A}_{8}$ decorated two black dots along the dorsal midline, these segments have also lateral black stripes between the levels D and L. Level SV of $\mathrm{A}_{9}$ lying a big black dapple on each side (divided along the dorsal and ventral midlines), while the anal shield, and the lateral surface of each proleg $\mathrm{A}_{10}$ appearing as a large black patch shaping nearly semicircular, the latter structure distributed 13-18 bristles on the posterior to ventral margin. All thoracic and abdominal setae are translucent, to be the longest on scoli L-III of $\mathrm{T}_{1}$. Scoli XD-III, D-III, and SD-III bear stout brown to black setae and the latter two have significantly elevated bases; other chalazae/scoli are developed with white setae. A pair of scoli D-III are medially fused on $\mathrm{A}_{8}$, whose fusional characteristic is similar to the case of scoli XD-III and SDIII on the anterior margin of prothoracic shield on each side. Spiracle $T_{1}$ is the largest whereas spiracles $\mathrm{A}_{1-8}$ are relatively small, all brownish colors. The paired sclerites of coxae are unclosed annular shaped and medially fused on $T_{1}$. Coxal setae L1 and L2 are respectively located on the antero/postero-terminal areas of the sclerite, other primary setae coded as M1, M2 and M3. The femur apically bears setae M4 and M5; each tibia has 6 setae labeled M6, M7, M8, M9, L3 and L4. On the top of the tarsus, setae M10, M11 and M12 are widened with slight longitudinal crests, seta L5 is a weak hair (Fig. 6K). A thick basal lobe is displayed on the pretarsus (Fig. 6L), sometimes there are secondary [subprimary?] setae occurring on legs $\mathrm{T}_{1-3}$

Table 1. $\mathrm{L}_{1}$ of Antheraea compta, a quantity statistic of chalazae/ scoli of the single side (divided along the dorsal and ventral midlines), and the numbers of primary setae borne on each them (in brackets). The "?" means inconclusive.

| Structures Segments | $\mathbf{T}_{1}$ | $\mathbf{T}_{2-3}$ | $\mathbf{A}_{\mathbf{1}}$ | $\mathbf{A}_{\mathbf{2}}$ | $\mathbf{A}_{3-6}$ | $\mathbf{A}_{\mathbf{7}}$ | $\mathbf{A}_{\mathbf{8}}$ | $\mathbf{A}_{\mathbf{9}}$ | $\mathbf{A}_{10}$ |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Scoli D-III |  | $1(8)$ | $1(5)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(9 ?)$ |
| Chalazae D-I | $2(1)$ |  | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ |  |
| Scolus XD-III | $1(3-4)$ |  |  |  |  |  |  |  |  |
| Scoli SD-III | $1(4)$ | $1(6)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(4)$ | $1(6)$ |  |
| Scoli L-III | $1(8-10)$ | $1(6)$ | $1(6)$ | $1(6)$ | $1(6)$ | $1(6)$ | $1(6)$ |  |  |
| Chalazae L-I |  |  | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ |  |  |
| Scoli SV-III | $1(4-5)$ | $1(3)$ |  |  |  |  |  |  |  |
| Chalazae SV-I |  |  | $1(1)$ | $1(1)$ | $2-3(1)$ | $2(1)$ | $2(1)$ | $2(1)$ |  |
| Chalazae V-I |  |  | $2(1)$ | $2(1)$ | $1(1)$ | $1(1)$ | $1(1)$ | $1(1)$ | $4 ?(1)$ |

(e.g., Fig. 6 K ). The plantae of prolegs are rosybrown in lateral view, each of them arranged dark maroon crochets in uniserial heteroideous mesoseries, the numbers are 23-$27\left(\mathrm{~A}_{3-6}\right)$ and 26-29 $\left(\mathrm{A}_{10}\right)$, such hooks have solid insides and longitudinally plicated epicuticle (Fig. 6M).

## $\mathrm{L}_{2}$ (Fig. 8D-F)

The primary chaetotaxy and related fusions are unchanged to $\mathrm{L}_{1}$. The head capsule ( width $=2.61 \mathrm{~mm}$ ) turned into sepia color with darker ocellar area. The antennal sclerotized parts are sienna connecting with golden antacoriae, the labrum, anteclypeus and frontoclypeus are bronzed. The cervacoria is lemon, the integumentary ground color of thorax-abdomen is largely pure or yellowish green but $\mathrm{T}_{1}$ is always paler, the prothoracic shield (with scoli XDIII and SD-III) is gradually changing to light yellow with aquamarine scoli L-III. Chalazae SV-I of $\mathrm{A}_{1-2}$ and scoli SV-III and are blue to yellow-green, other scoli L-III, SDIII, and all scoli D-III owning ultramarine to blueviolet bases in smooth, but the proximal halves of the latter two borne on $\mathrm{T}_{2}-\mathrm{A}_{8}$ have shiny silvery appearance, the maximum size always showing on scoli SD-III of $\mathrm{A}_{2-7}$. Some setae are elongated to form clubbed tips which displayed on scoli XD-III of $\mathrm{T}_{1}$, scoli D-III of $\mathrm{T}_{2}$ (sometimes $\mathrm{T}_{2-3}$ ) and $\mathrm{A}_{2-9}$, as well as on all scoli SD-III and L-III, such special bristles lengthened on scoli D-III of $\mathrm{T}_{2}$ to be longer than any others. Since this larval instar, secondary hairs beginning to obviously emerge from the cephalic regions and the venter of $\mathrm{T}_{1}-\mathrm{A}_{10}$, accompanied by the harder observing on chalazae D-I, L-I, V-I and partial SV-I. Spiracles are surrounded by amber integumentary edges. The sclerites of legs $\mathrm{T}_{1-3}$ are sepia, the distolateral surfaces of proleg bases of $\mathrm{A}_{3-6}$ developed many black dots with secondary setae, while prolegs $\mathrm{A}_{10}$ also stretched multidirectionally their marginal bristles. The lateral plate of each anal proleg is dyed with posterior ochre, central black and anterior yellow (sometimes displaying only the former two colors), the latter two colors usually corresponding multi-pitted surfaces, the feature similarly arises in the yellowish triangular area of the anal shield. The plantar colors of prolegs $\mathrm{A}_{3-6}$ and $\mathrm{A}_{10}$ are the same as $\mathrm{L}_{1}$, but the crochets rowed into uniserial homoideous mesoseries with the quantities of 26-30.


Figure 8. $\mathrm{L}_{1-6}$ of Antheraea compta. A. $\mathrm{L}_{1}$, newly hatched individual feeding on the eggshell, lateral view; $\mathbf{B} . \mathrm{L}_{1}$, drinking water, dorsal view; C. $\mathrm{L}_{1}$, feeding on the edge of leaf (Quercus yunnanensis), lateral view; D. $\mathcal{+}, \mathrm{L}_{2}$, newly molted, lateral view; $\mathbf{E} . \mathcal{Q}, \mathrm{L}_{2}$,

 dorsal view; K. $\widehat{O}, \mathrm{~L}_{4}$, lateral view; L. $\widehat{O}$, the late period of the $4^{\text {th }}$ pre-molting, the dorsal area of $\mathrm{T}_{2}$, frontal view; M. $\widehat{O}, \mathrm{~L}_{5}$, dorsal view; N. |  |
| :--- | $\mathrm{L}_{5}$, lateral view; O, P. $\widehat{\delta}, \mathrm{L}_{5}$, epicuticle of dry exuviae, the medial surface of scolus SD-III of $\mathrm{A}_{1}$; O. a: spiny seta; b: shiny area; c: green integument; $\mathbf{P}$. a: shiny area; b: green integument; $\mathbf{Q}, \mathbf{R} . \widehat{\delta}, \mathrm{L}_{6}$, epicuticle of dry exuviae (sampling from cocoon), the clubbed setal apex of scolus D-III of $\mathrm{T}_{3} ; \mathbf{Q}$. Centeral area; R. Marginal area. Scale bars: $3 \mathrm{~mm}(\mathbf{A}-\mathbf{C}, \mathbf{I}, \mathbf{L}) ; 5 \mathrm{~mm}(\mathbf{D}-\mathbf{F})$; $1 \mathrm{~cm}(\mathbf{G}, \mathbf{H}, \mathbf{J}, \mathbf{K}) ; 2 \mathrm{~cm}(\mathbf{M}, \mathbf{N}) ; 200 \mu \mathrm{~m}(\mathbf{O}) ; 30 \mu \mathrm{~m}(\mathbf{P}) ; 4 \mu \mathrm{~m}(\mathbf{Q}) ; 20 \mu \mathrm{~m}(\mathbf{R})$.


 prolegs, lateral view; F. $\widehat{o}^{\lambda}$, lateral margin of anal shield, dorsolateral view; $\mathbf{G}$. $\widehat{\delta}^{\lambda}$, head and $\mathrm{T}_{1-2}$, anterolateral view; $\mathbf{H} . \delta^{\lambda}$, spiracle $\mathrm{T}_{1}$, vertical view; I. $\delta^{\hat{1}}$, legs $\mathrm{T}_{1-3}$, ventral view; J. $\widehat{\delta}^{\lambda}$, proleg $\mathrm{A}_{10}$, ventral view; K. $\boldsymbol{\delta}^{\hat{1}}$, ventral view; a-c: $\mathrm{A}_{8-10}$; d: sexual gland; L..+ , ventral view; a-c: $\mathrm{A}_{8-10}$; d: sexual gland; M-T. Latero-medioapical views; M. + , scolus D-III of $\mathrm{T}_{2} ; \mathbf{N} .+$, scolus D-III of $\mathrm{A}_{6} ; \mathbf{O} .+$,
 of $\mathrm{T}_{3} ; \mathbf{T} .{ }^{\lambda}$, chalaza SV-I of A. Scale bars: $2 \mathrm{~cm}(\mathbf{A}, \mathbf{B}) ; 1 \mathrm{~cm}(\mathbf{C}, \mathbf{D}) ; 5 \mathrm{~mm}(\mathbf{E}, \mathbf{G}, \mathbf{I}) ; 2 \mathrm{~mm}(\mathbf{F}, \mathbf{M}-\mathbf{T}) ; 1.5 \mathrm{~mm}(\mathbf{H}) ; 2.5 \mathrm{~mm}(\mathbf{J}-\mathbf{L})$.

## $\mathrm{L}_{3}$ (Fig. 8G, H)

General morphology close to $L_{2}$, but the head capsule is paler and widened to 3.73 mm . The flash basal parts have enlarged the volume further on scoli SD-III of $\mathrm{A}_{2-7}$, especially. Supplemented long setae with club-like apexes on scoli D-III of $\mathrm{T}_{3}-\mathrm{A}_{1}$, as well as on chalazae SV-I of $\mathrm{A}_{1-2}$ and scoli SV-III. Dense coryneform bristles initiating to present on the posterior zones of the middorsal $\mathrm{A}_{2-6}$, colored as yellow vividly; they are minute and cross a transverse row combined by longer and pointed setae in the same color but borne on the anterior zones of $\mathrm{A}_{3-7}$. Tricolored lateral plates of prolegs $\mathrm{A}_{10}$ observed in most of $L_{2}$ now shared in all individuals. More secondary hairs are exhibited principally on ventral integument. Some of them are based on dollar spots, such dots originating since $L_{2}$ are now widely distributed to level $L$, gradually colored from black to light brown. The marginal setae of $\mathrm{A}_{10}$ are pronouncedly elongated and orientated to the posterior side. The numbers of $50-55$ crochets arranged as biordinal mesoseries on each planta.

## $\mathrm{L}_{4}$ (Fig. 8I-L)

The macroscopic pattern is essentially same as $L_{3}$, but the 5.47 mm width head capsule begins to appear more yellowish in proportions, mainly enriched on its ventrolateral and frontal zones, but the dark sepia strip ornamented on ocellar areas are still visible. The color of cervacoria is more vivid, slanted, parallel dark olive-green streaks arising between the levels SD and L of $\mathrm{A}_{1-8}$. For $\mathrm{T}_{2}-\mathrm{A}_{10}$, the venter is clearly darker than dorsa, the dense dark spots are developing to the level SD with some secondary hairs, and the coxal apexes of legs $\mathrm{T}_{2-3}$ are colored lightly like $\mathrm{T}_{1}$. The clubbed setal apexes already occur on the longer bristles on prolegs, starting from this instar, each lateral plate of anal prolegs shows a very minute black dot on the yellow tint area with relatively fixed position, and each lateral margin of the anal shield formed into vitreum, enveloping an internal dark brown band. Plantar crochets are still biordinal mesoseries but increased to numbers of 54-60. Take $\mathrm{L}_{4}$ as an example, for a fresh larva or during the pre-molting, the oblate setal tip could be seen white translucent, whereas the silver-reflective zones of scoli are transparent just after ecdysis but fully colored after approximately about 18-22 hours (Fig. 8I, L).

## $\mathrm{L}_{5}$ (Fig. 8M-P)

Head capsule (width $=7.33 \mathrm{~mm}$ ) is paler distinctly, but the distribution patterns of colors are basically the same as $\mathrm{L}_{4}$. The cephalic setae are more strongly developed, the dotted integument reaching the level D and all the secondary hairs turned into brownish to black with longer sizes. Parallel oblique stripes lying around spiracles $\mathrm{A}_{1-8}$ are more evident, each spiracle is outlined by pale aquamarine margin. Scoli XD-III and SD-III on each side of prothoracic shield are highly fuse to be a unified base outwardly. The band inlaid inside the vitric margin of anal
shield is now darker. The shining silvery bases of scoli D-III of $\mathrm{T}_{2}-\mathrm{A}_{9}$, and SD-III of $\mathrm{T}_{2}-\mathrm{A}_{8}$ are already spectacular (largest on scoli SD-III of $\mathrm{A}_{2-7}$ still), their epicuticles are extraordinary smooth in stark contrast to the green integument, the latter is densely covered noncellular processes (Fig. 80, P). Proleg plantae are surfaced with lateral dark gray and medial white with 61-64 crochets in biordinal mesoseries. Each of the lateral plates of prolegs $\mathrm{A}_{10}$ is constituted by the wrinkled bright yellow-green area, the smooth dark ochre posterior margin, and a narrow black stripe between the former two.

## $\mathrm{L}_{6}$ (Figs $8 \mathrm{Q}, \mathrm{R}, 9,10$ )

The final larval instar with giant body, head capsule is 10.32 mm width and the overall color pattern is near $\mathrm{L}_{5}$ but more pallid (Fig. 9C, G). Combined with the high-saturation goldenrod cervacoria and the pale lemon coxae, the front margin of $\mathrm{T}_{1}$ forms a bright yellow annular zone, which showed intensely fluorescent enrichment under the excitation by UV 365 nm (Figs 9G, 10B). Fluorescence was not detected in the flash bases with the clubbed setae (Fig. 10C), but weakly appeared on the yellow-green regions of the central anal shield and the lateral plates of prolegs $\mathrm{A}_{10}$ (Fig. 10D), the wrinkles of the latter are more obvious than those during the prior stage (Fig. 9E). Both scoli XD-III and SD-III have different numbers of primary setae during $\mathrm{L}_{2-5}$ which appeared as short spines, but all of these are elongated into hair-like now (Fig. 9C, G). Similar to $\mathrm{L}_{2-5}$, for all scoli, only partial setae on $\mathrm{T}_{1}-\mathrm{A}_{9}$ have clubbed tips (Fig. 9M-T), statistic on the single larva displayed that most setal quantities of the corresponding sites are same as $L_{1}$, but that on 12 scoli are decreased and 2 scoli are increased (Table 2). The rod-like part of the clubbed seta is smooth helical, the flaky end is paved by shallow pits densely but strongly reflection (Fig. 8Q, R). There is a significant difference in general integumentary color, i.e., 2 it $q$ are more yellowish than $3 \widehat{\sigma}^{\top} \widehat{\sigma}^{\lambda}$, and the shiny scoli of females are tending to be purple-red but that of males look more purple-blue (Figs 9M-T, 10A). Observed the male sexual gland ["Herold's gland"] to be a slight mark localizing on ventral midline near to $\mathrm{A}_{8} / \mathrm{A}_{9}$ boundary (Fig. 9K), the female sexual gland ["Ishiwata's gland"] arising as two pairs of dots on ventral $\mathrm{A}_{8}$ and $\mathrm{A}_{9}$ (Fig. 9L). Distinguished from $\mathrm{L}_{5}$, the dark spots are added to the middorsal area except the prothoracic shield, and the secondary hairs are increased on the whole body at present, but except the sizes, the glassy margins of anal shield as well as the spiracles are morphologically almost unchanged (Fig. 9F, H). Vesicular processes were observed on the distomedial areas of each tibia and tarsus, and all the medial setae (both primary and secondary) of legs $\mathrm{T}_{1-3}$ form white thick spines (Fig. 9I), unfortunately, it lacked a leg observation for $\mathrm{L}_{2-5}$. Plantae of $\mathrm{A}_{3-6}$ and $\mathrm{A}_{10}$ bear 59-65 crochets within biordinal mesoseries (e.g., Fig. 9J). The liquid feces after feeding ended (all larval instars fed from Q. yunnanensis) is bronzed mucus (Fig. 10E), after that, $1 \delta^{1}$ was recorded the weight as 14.77 g .

 C. $\mathrm{A}_{7}$, lateral view; D. $\mathrm{A}_{10}$, lateral view; E. $\widehat{\Omega}$, the last defecation [liquid defecation] from $\mathrm{A}_{10}$, lateral view; $\mathbf{F}$. $\widehat{0}$, spinning the peduncle of cocoon; $\mathbf{G} .{ }^{\lambda}$, spinning the external part [floss] of cocoon. Scale bars: $2 \mathrm{~cm}(\mathbf{A}) ; 1 \mathrm{~cm}(\mathbf{B}, \mathbf{F}, \mathbf{G}) ; 5 \mathrm{~mm}(\mathbf{C}-\mathbf{E})$.

Table 2. $\mathrm{L}_{6}$ of Antheraea compta, a statistic based on primary chalazae/scoli which have clubbed setae, with the format "short spines + elongated hairs (how many clubbed ones)". The " $<$ " is lower, the " $=$ " is same, and the " $>$ " is higher than the normal value of the corresponding locations during $L_{1}$ (see Table 1). Merged cells means the scoli are medially fused.

| Segments\Areas | SV-I/III | L-III | SD-III | XD/D-III | XD/D-III | SD-III | L-III | SV-I/III |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| T ${ }_{1}$ | $0+4(1)=$ | $1+7(5)=$ | $0+4(1)=$ | $0+4(2)=$ | $0+3$ (2) $=$ | $0+4(2)=$ | $2+7(4)=$ | $0+4(0)=$ |
| $\mathrm{T}_{2}$ | $0+1(1)<$ | $4+2(2)=$ | $3+3(3)=$ | $5+3(3)=$ | $5+3(3)=$ | $3+2(2)<$ | $4+2(2)=$ | $2+1(1)=$ |
| T3 | $2+1(1)=$ | $4+2(2)=$ | $3+3(3)=$ | $5+3(3)=$ | $2+4(4){ }^{<}$ | $3+2(2)<$ | $4+2(2)=$ | $2+1(1)=$ |
| $\mathrm{A}_{1}$ | $0+1(1)=$ | $4+2(2)=$ | $3+1(1)=$ | $3+2(2)=$ | $1+1(1){ }^{<}$ | $4+1(1){ }^{>}$ | $4+1(1)<$ | $0+1(1)=$ |
| $\mathrm{A}_{2}$ | $0+1(1)=$ | $3+1(1)<$ | $3+1(1)=$ | $2+2(2)=$ | $3+1(1)=$ | $2+2(2)=$ | $4+2(2)=$ | $0+1(1)=$ |
| $\mathrm{A}_{3}$ |  | $3+3(3)=$ | $3+1(1)=$ | $3+1(1)=$ | $3+1(1)=$ | $3+1(1)=$ | $4+2(2)=$ |  |
| $\mathrm{A}_{4}$ |  | $3+2(2)<$ | $3+1(1)=$ | $3+1(1)=$ | $3+2(2){ }^{\text {> }}$ | $3+1(1)=$ | $4+2(2)=$ |  |
| $\mathrm{A}_{5}$ |  | $3+2(2)<$ | $3+1(1)=$ | $3+1(1)=$ | $3+1(1)=$ | $3+1(1)=$ | $4+2(2)=$ |  |
| $\mathrm{A}_{6}$ |  | $4+2(2)=$ | $3+1(1)=$ | $3+1(1)=$ | $3+0(0)<$ | $3+1(1)=$ | $4+2(2)=$ |  |
| $\mathrm{A}_{7}$ |  | $4+2(2)=$ | $3+1(1)=$ | $2+2(2)=$ | $3+1(1)=$ | $3+1(1)=$ | $4+2(2)=$ |  |
| $\mathrm{A}_{8}$ |  | $3+2(2)<$ | $3+1(1)=$ | $2+2(2)=$ | $2+2(2)=$ | $3+1(1)=$ | $3+2(2)<$ |  |
| $\mathrm{A}_{9}$ |  |  | $3+3(3)=$ | $2+2(2)=$ | $2+2(2)=$ | $4+2(3)=$ |  |  |

## Pupae (Figs 11, 12C)

The epicuticle is dark maroon to black except the translucent epicranial plate [frontoclypeus]. Females are larger (lengths: 3 đ $\widehat{ }=40.58 \mathrm{~mm}, 39.64 \mathrm{~mm}, 38.68 \mathrm{~mm}$; $2 q Q=42.63 \mathrm{~mm}, 44.48 \mathrm{~mm}$ ) and heavier (weights of during overwintering diapause, measured on 06 Feb. 2023: 3 ふた $=8.16 \mathrm{~g}, 7.46 \mathrm{~g}, 7.54 \mathrm{~g} ; 2$ 우 $=9.29 \mathrm{~g}$, 8.51 g ) than males. The antennal margins in females are more flattened than males, the pair of antennae of the latter are not medially touching along the ventral midline, the maxillae and legs $T_{1-2}$ are visible in both sexes (Fig. 11C, E). Spiracles $T_{1}$ is gapped on the boundary of $\mathrm{T}_{1} / \mathrm{T}_{2}$ (Fig. 11F), spiracle $\mathrm{A}_{2-7}$ are functional and large, but that of $\mathrm{A}_{8}$ is closed as a remnant. A pair of forewing tubercles [adult spurs] is rendered on the dorsolateral areas of $\mathrm{T}_{2}$ specifically (Fig. 11F). The biserial larval scoli D-III had been reduced to smooth vestiges on pupal
stage, in only one case that such bulges on $\mathrm{T}_{3}-\mathrm{A}_{4}$ are obvious (Fig. 11 H ). The unsclerotized joints are junction zones between $\mathrm{A}_{4} / \mathrm{A}_{5}, \mathrm{~A}_{5} / \mathrm{A}_{6}$, and $\mathrm{A}_{6} / \mathrm{A}_{7}$, the tip of $\mathrm{A}_{10}$ is rounded without cremaster (Fig. 11G). Genital pores are navel-shaped scars apertured midventrally for males $\left(\mathrm{A}_{9}\right)$ and females $\left(\mathrm{A}_{8}\right.$ [ostium bursae] and $\mathrm{A}_{9}$ [ostium oviductus]) (Fig. 11B, D).

## Cocoons (Figs 10F, G, 12)

Large (max-lengths without peduncles: $3 \widehat{\delta} \hat{=}=$ $63.49 \mathrm{~mm}, 52.14 \mathrm{~mm}, 56.44 \mathrm{~mm} ; 2$ Q $Q=50.37 \mathrm{~mm}$, 60.41 mm ), single layered (thicknesses $=\mathrm{ca} 0.37-$. 1.16 mm ) and fully closed (no pre-formed exit), the external part was woven by floss for attaching to leaves (Fig. 10G), the general shape is irregular polyhedron but the silky pupal cell is prolate spheroid. White colored in dry environment (weak fluorescence under UV


Figure 11. Pupae of Antheraea compta. A. a: $\widehat{\delta}^{\lambda}$, lateral view; b: $\widehat{\delta}^{\lambda}$, ventral view; c: $\widehat{\jmath}^{\wedge}$, dorsal view. d: $\varphi$, lateral view; e: $\ell$, ventral view; f: $\rho$, dorsal view; B. $\delta^{\star}$, ventral view; a-d: A $\mathrm{A}_{7-10}$; e: genital pore; C. $\delta^{\lambda}$, head and $\mathrm{T}_{1-2}$, ventrolateral view; D..+ , ventral view; a-d: $\mathrm{A}_{7-10}$; e, f: genital pores; E. + , head and $\mathrm{T}_{1-2}$, ventrolateral view; $\mathbf{F} . \widehat{\delta}^{\lambda}, \mathrm{T}_{1-2}$, dorsolateral view; a: spiracle $\mathrm{T}_{1} ;$ b: forewing tubercle; $\mathbf{G} . \widehat{\delta}^{\hat{}}$, posterolateral view; a, b: $\mathrm{A}_{9-10} ; \mathbf{H} . \widehat{\delta}^{\lambda}$, dorsolateral view; a-e: $\mathrm{T}_{3}-\mathrm{A}_{4}$. Scale bars: $1 \mathrm{~cm}(\mathbf{A}) ; 2 \mathrm{~mm}(\mathbf{B}, \mathbf{D}) ; 5 \mathrm{~mm}(\mathbf{C}, \mathbf{E}) ; 3 \mathrm{~mm}(\mathbf{F}-\mathbf{H})$.

365 nm , barely visible) but metamorphosing into brown tones in wet (fluorescence disappeared). $2 \delta^{\lambda}$ and 1 q spun on erect vegetations with wide and loose peduncles of inferior structure (e.g., Figs 10F, 12A-C), 1 ठ and $1 q$ spun cocoons within fallen leaves and branches on the ground without peduncle (e.g., Fig. 12D). Each filament is a double-strand flat belt, wrinkles formed by longitudinal stretching could be observed on epi-sericin (Fig. 12F), the part of fibroin is combined by dense elongated fibers longitudinally to be almost non-porous within (less voids) (Fig. 12E). Microscopically surveyed three pieces (ca. $6.5-8.4 \mathrm{~mm}^{2}$ ) of silk-layer sliced from random sites of a cocoon, showing the external surface is rougher, the filaments are ca. 83.8-94.4 $\mu \mathrm{m}$ wide and ca. 4.7-9.3 $\mu \mathrm{m}$ thick (Fig. 12H). The internal surface [pupal cell] is smooth and flat relatively, widths of the filaments are ca. $56.4-92.8 \mu \mathrm{~m}$ with the thicknesses of ca. 2.3-3.9 $\mu \mathrm{m}$ (Fig. 12I). Many crystals (mainly calcium oxalate) granulated from the prepupal discharge liquor are scattered or concentrated in the silk-layer (Fig. 12G), due to gravity, the precipitation is more obvious in the outer surface of the cocoon that lies closer to the ground.

## Rearing report

During my first expedition on Tibetan Nyingchi, $1 q$ of A. compta was collected on 28 Jun. 2021 from Mêdog 2122 m (Fig. 2), her 60 ova were hatched into 37 larvae in the experimental site in Kunming City of Yunnan Province, 1940 m , during mid Jul. to early Aug., these fresh $\mathrm{L}_{1}$ rejected any Lauraceae, i.e., Cinnamomum camphora, Cinnamomum burmannii, Machilus spp., Lindera sp., Neolitsea sp., Litsea pungens and Litsea populifolia, moreover, Magnolia delavayi (Magnoliaceae), Prunus cerasoides (Rosaceae), Salix babylonica (Salicaceae) and Liquidambar formosana (Altingiaceae) were likewise not accepted. Finally, three individuals nibbled the leaves of Quercus variabilis and Q. yunnanensis, but only one grew into the $3^{\text {rd }}$ pre-molting with the latter, and because it was reared outdoors without a cage, this larva was preyed upon by a Myophonus caeruleus (Aves: Passeriformes, Muscicapidae).

After the first failed attempt, I recaptured $3 q$ adults with some males (Fig. 1) from the same natural habitat; during the rainy nights of 25-26 Jun. 2022, many large Quercus lamellosa were noticed at the site. Successively, 3 (coded as Gi, laid on 26 Jun.), 102 (coded as Gii, laid


Figure 12. Cocoons of Antheraea compta. A-C. $\widehat{0}$, spun on the erect vegetation, lateral view; A. Original condition; B. Removed some leaves and branches; C. Opened; D. $\mathcal{Y}$, spun on the bottom of the net cage under a fallen leaf, lateral view [the underside]; E. $\widehat{\sigma}$, filaments of the external part, cross sectional view; a: transverse [horizontal] plane; b: diagonal plane; c: sagittal plane; F. $\widehat{0}$, filament of the external part, vertical view; G. $\widehat{\delta}$, the crystals precipitated from the prepupal discharge liquor, vertical view; H. $\widehat{\delta}$, filaments of the external surface, vertical view; I. $\begin{gathered} \\ \text {, filaments of the internal surface, vertical view. Scale bars: } 3 \mathrm{~cm}(\mathbf{A}-\mathbf{D}) ; 10 \mu \mathrm{~m}, ~\end{gathered}$ (E); $20 \mu \mathrm{~m}(\mathbf{F}) ; 2 \mu \mathrm{~m}(\mathbf{G}) ; 100 \mu \mathrm{~m}(\mathbf{H}, \mathbf{I})$.
on 25 Jun.-01 Jul.) and 16 (coded as Giii, laid on 26-30 Jun.) eggs severally into paper envelopes. Each of Gi and Giii was oviposited naturally into a single tier without longitudinal accumulation, Gii was the same condition in the early period but later heaped up to lump because of larger quantity and lack of available space for adult abdomen. The material was brought back to the same site in Kunming indoor and then grouped on three mesh pieces, a cotton pad was placed below with interval ca. 1 cm , water sprayed the ova every day, the circumambient air was ca. $17-21^{\circ} \mathrm{C}$ with ca. $80-95 \% \mathrm{RH}$. This method helped prevent water from accumulating around the eggs; excess water drained through the mesh openings to the cotton pad below, while the space between helped ventilate the bottom of the eggs to prevent mold.

At Beijing Time 06:00 to 16:00 every day, 3 larvae of Gi hatched on 13 Jul. (ovum period 17 days), 81 larvae of Gii hatched during 12-19 Jul. (ordered as: $10,27,12$, $5,15,6,5,1$; ovum period 17-18 days) and 14 larvae of Giii hatched during 13-18 and 20 Jul. (ordered as: 3, 4, 3, $1,1,1,1$; ovum period $17-18$ or 20 days, but the last individual was unhealthy and died on the day of hatching). Newly hatched $L_{1}$ usually fed on eggshells preferentially. Checking unhatched material on 31 Jul., 17 ova of Gii and 1 ovum of Giii were devoid of embryonic development and shriveled; mature embryos had died in 4 ova of Gii and 1 ovum of Giii.

Fresh $L_{1}$ were moved to a semi-open balcony for rearing, without cage protection. The plant stems were inserted into the bottle filled with water, and the bottleneck blocked with paper towels to prevent larval drowning. Some newly hatched individuals did not bite the leaves of Quercus schottkyana and Magnolia denudata within two days, so all larvae were given $Q$. yunnanensis, which was readily accepted. These caterpillars had weak positive phototaxis, usually did not climb to the apex of host plant and did not exhibit gregarious behavior, but there were often $2-5$ scattered $L_{1}$ on a single leaf. They rested on the undersides of the leaves and moved to the edges only when feeding, if a branch had fresh terminal buds and thick mature leaves, newly hatched larvae did not distinguish between them, as they are perfectly capable of gnawing on harder leaves, which is apparently related to their larger head with powerful mandibular adductors. Although they did not have strict requirements for air humidity during the larval stage (kept in ca. $64-98 \% \mathrm{RH}$ ), the temperature should not exceed $28^{\circ} \mathrm{C}$, otherwise the larvae will die in large numbers. When the environment reached $26^{\circ} \mathrm{C}$, they usually unclasped the legs $\mathrm{T}_{1-3}$ and prolegs $\mathrm{A}_{3-5}$ from host plant and suspended in air to enhance ventilation for heat dissipation. This may cause many larvae to fall to the ground. The whole larval stage accompanied a strong need for drinking water, and the leaves must be sprayed with water several times each day to ensure normal development of caterpillars, except on rainy days.
A. compta is a rather difficult species to rear in captivity with a very high mortality in $L_{1}$, in this study, only four individuals of Giii and five of Gii survived to the $1^{\text {st }}$ pre-molting. However, two larvae within the latter group
died during pharate and two died in early $\mathrm{L}_{2}$. In the five surviving $\mathrm{L}_{2}$, the individual "Giii1" was caged in outdoor and other four still fed semi-openly, until they cocooned respectively. Although the weather and nutritional conditions are identical, ontogenic time-consumptions are quite different in the same gender (Table 3).

Before entering each pre-ecdysis, larvae always moved away from their last feeding site. In late $\mathrm{L}_{5}$ they cut the petiole to make the leaf fall to the ground or break it off completely to eliminate signs of feeding, which apparently helps to avoid some natural enemies which could find the caterpillar location by defoliation. Some individuals would sometimes entangle silk in selected locations to anchor themselves before shifting into apolysis state; such behavior was not consistent, but each of them always preferentially fed on their exuviae except the head capsule.

During $L_{2-3}$, larvae usually rested on the leaf midrib after a period of feeding of the lamina and secondary veins. For $\mathrm{L}_{4}$ to medium $\mathrm{L}_{5}$ they rested intermittently near the petiole and gradually nibbled along the midrib. Since late $\mathrm{L}_{5}$, the weight only allowed the activities on branches, before feeding a new leaf, the larva often first bit a large pit on the petiole and then ate along the basal area, if necessary, gnawed more gaps on midrib for further folding the leaf apex closer to itself. For mature larvae, feeding behavior mostly occurred at night, probably to avoid detection by birds. A continuous observation on single $\mathrm{L}_{6}$ was recorded for reference (Suppl. material 1).

All larval instars were sensitive to light, i.e., in resting states, they always faced toward the side where the illuminance is higher (Suppl. material 2), and this behavior would draw predators' attention to the bright yellow ring on the front of $\mathrm{T}_{1}$, whose fluorescent was sufficient to indicate it plays a role in defensive warning. These caterpillars were extremely alert, and every slight shake would interrupt the feeding behavior. Under this state, larvae usually detached their legs $\mathrm{T}_{1-3}$ and prolegs $\mathrm{A}_{3}$ (sometimes $\mathrm{A}_{3-4}$ ) from the host plant, which may be an effective mimicry to make the lateral views more like leaves, sometimes clicking the mouthparts as a popping noise. The most alarmed individuals would swing from side to side, even triggering biting, whereas the special clubbed setae may help to enhance the spatial scope of tactile cue

Table 3. Developmental data of different individuals of Antheraea compta, fed from Quercus yunnanensis.

| Stages\Codes | Giii1, ${ }^{\text {® }}$ | Giii2, ${ }^{\text {T}}$ | Giii3, ${ }^{\text {T}}$ | Giii4, + | Gii1, , |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Oviposited | 30 Jun. | 27 Jun. | 28 Jun. | 28 Jun. | 25 Jun. |
| Hatched into $\mathrm{L}_{1}$ | 17 Jul . | 14 Jul. | 15 Jul. | 15 Jul . | 12 Jul . |
| $1{ }^{\text {st }}$ pre-ecdysis | 27 Jul. | 25 Jul. | 26 Jul. | 26 Jul. | 23 Jul. |
| Molted into $\mathrm{L}_{2}$ | 29 Jul. | 27 Jul. | 28 Jul. | 28 Jul. | 25 Jul. |
| $2^{\text {nd }}$ pre-ecdysis | 05 Aug. | 07 Aug. | 08 Aug. | 08 Aug. | 06 Aug. |
| Molted into $\mathrm{L}_{3}$ | 07 Aug. | 09 Aug. | 10 Aug. | 10 Aug. | 08 Aug. |
| $3{ }^{\text {rd }}$ pre-ecdysis | 14 Aug. | 16 Aug. | 18 Aug. | 18 Aug. | 17 Aug. |
| Molted into $\mathrm{L}_{4}$ | 16 Aug. | 18 Aug. | 20 Aug. | 20 Aug. | 19 Aug. |
| $4^{\text {th }}$ pre-ecdysis | 23 Aug. | 26 Aug. | 28 Aug. | 30 Aug. | 28 Aug. |
| Molted into $\mathrm{L}_{5}$ | 26 Aug. | 29 Aug. | 31 Aug. | 01 Sep. | 31 Aug. |
| $5{ }^{\text {th }}$ pre-ecdysis | 05 Sep . | 10 Sep. | 14 Sep. | 15 Sep. | 17 Sep. |
| Molted into $\mathrm{L}_{6}$ | 09 Sep. | 15 Sep. | 18 Sep. | 19 Sep. | 21 Sep. |
| Feeding ended | 03 Oct. | 09 Oct. | 12 Oct. | 16 Oct. | 23 Oct. |
| Spun cocoon | 04 Oct. | 11 Oct. | 13 Oct. | 17 Oct. | 24 Oct. |

(Suppl. material 3). Even so, no secretions exude from the scoli under stimulated conditions, and no urticating reaction was detected on human skin after touch.

Larvae never left the host plant voluntarily if the leaves were plentiful. However, they quickly climbed or fell to the ground after feeding ended [liquid defecation] (happened at night usually), whereupon they were caged with upright living vegetation and dead leaves at the bottom for observation. Almost all individuals moved restlessly for about 24 hours before choosing a spinning location. The species undoubtedly constructs cocoons away from the host plant in the wild, either in low shrubs under the oak canopies or in the leaf litter on the ground.

The flight time in the natural habitat and captivity and reproductive behavior of adults will be recorded and published after the fresh adults emerge from existing pupae.

## Discussion and conclusion

This article provides the first report of the complete life history of the enigmatic $A$. compta, whose lateral stripes of mature larvae are a feature close to that of North American Antheraea oculea (Neumoegen, 1883) (Tuskes et al. 1996: pl. 6; Lampe 2010: 121) and A. polyphemus (Bouseman and Sternburg 2002: 51; Hall 2021), the scale and distribution of the shiny silver spots closely resemble those in the Mexican Antheraea montezuma (Sallé, 1856) and Antheraea meridiana Naumann \& Nogueira, 2021 (Paukstadt and Paukstadt 2020; Naumann and Nogueira 2021), while the tricolored lateral plate of anal proleg of Neotropical A. godmani (Nässig et al. 1996; Goossens et al. 2015) is similarly shared in $A$. compta. These observations preliminarily suggest that Himalayan A. compta should be assigned to the subgenus Telea.

Due to $\mathrm{L}_{1}$ rejecting $Q$. schottkyana in captivity, it is inferred that $Q$. lamellosa is observed in the Tibetan habitat, and more cycle-cup oaks may not be the correct host plants to $A$. compta, namely the section Cyclobalanopsis. Additionally, in the eastern Himalayas, native specimens Quercus lodicosa, Quercus lanata, Quercus engleriana, Quercus semecarpifolia, Quercus aquifolioides, Quercus gilliana, Quercus rehderiana and Quercus senescens within the section Ilex, as well as Quercus acutissima of the section Cerris were also identified in recent works (Zhou and Sun 1996; Denk and Grimm 2009; Yang and Zhou 2015: 130; Lahiri et al. 2017), the three sections have been confirmed to form a monophyletic subgenus Cerris (Denk et al. 2017; Hipp et al. 2019). Because of the observed rejection of Q. variabilis (a single larva reluctantly ate a small gap, but did not continue), its sister taxon Q. acutissima would probably also be rejected. There is currently no experimental result on infrasubgeneric Ilex so it is uncertain whether $A$. compta can partially accept some species of the subgenus, but probably, Q. lodicosa and Q. lanata would be accepted as foods by such caterpillars; the former tree is common in southeastern Tibet and the latter is dominant from Mêdog to Tengchong County, Yunnan (bordered with Myanmar

Kanpaikti), all having humid mid-altitudinal distributions (Zhou et al. 1995). In any case, the above species are clearly distinguished from $Q$. yunnanensis, a deciduous oak within the subgenus Quercus. Another roburoid white oak Quercus griffithii is widely known in northeastern India (Negi and Naithani 1995: 50-53; Singh et al. 2009; Singh et al. 2015), likewise naturally occurring in mountains from northern Yungui Plateau to sections of Indochina (Huang et al. 1999; Menitsky 2005: 47-49), with a high probability to be a primary host to $A$. compta at the corresponding biotopes.
A. assamensis is a well-known Lauraceae feeder in the wild (Seidel and Peigler 2018; Devi et al. 2021) but accepted beech (Lampe 2010: 358) and oak (Crotch 1956: 55) in captivity. Interestingly, the current known hostplants of A. godmani are limited to oaks (Meister 2011: 148), while A. compta is also oligophagous and feeding on certain Quercus only. This means that the empty cocoons collected from Assam soalu trees Litsea monopetala and then sent to Peigler (1999) were actually spun by muga silkworms. Therefore, Jolly's (1981) conclusion that "A. compta is almost identical with A. assamensis [sic]" is completely incorrect, while native Indian material for study of chromosomes (Gupta and Narang 1981) and silk (Luikham et al. 2017) using the former name but based on Lauraceae host plants are surely misidentifications to $A$. assamensis or its sister species Antheraea castanea Jordan, 1910. In fact, no Indian author has ever provided an adult image to prove the true identity of so-called " $A$. compta", except Arora and Gupta (1979) who described a pinned male from the British India period. In recent years, there have been a few indigenous records from the country but limited to only checklists without reliable morphological description and often confusingly called "wild muga" or "oak tasar" (e.g., Bhatia et al. 2010; Devi et al. 2011; Kumar et al. 2016; Gogoi and Goswami 2016; CSB 2018; Marepally 2018: 4; Boro and Borah 2020; Kumar et al. 2020; Keisa et al. 2022). This confusion suggests that the current population of the species may be nearly extirpated because virtually no local researchers have actually collected A. compta. Coincidentally, in works in which illustrated moth photographs corresponding to the scientific names, no specimen of $A$. compta was recaptured in recent surveys of northeastern India (Gogoi et al. 2014; Kakati and Chutia 2009; Kalita and Dutta 2014; Lalhmingliani 2015; Sondhi et al. 2021a) or even its type-locality Meghalaya (Shangpliang and Hajong 2015a, 2015b), whereas also does not turn up in the long-term monitoring of the Himalayas and Indian subcontinent (Shubhalaxmi 2018; Chandra et al. 2019; Sondhi et al. 2021b). To date, no Chinese publications included this species, and no records from Bhutan, Nepal and Uttarakhand are available, whereas Schüssler (1933: 175) added Sikkim to the distribution without supporting data.

Actually, the moths flying in southeastern Tibet were separated taxonomically from Meghalaya $A$. compta, the former was named Antheraea discata Naumann \& Löffler, 2015, whose type locality is the same as the origin of my material. Unfortunately, the authors of $A$. discata
overlooked the study of specimens from mountainous Burma, and although they discovered some morphological and COI differences between Mêdog and Khasi Hills, the Myanmar samples are still crucial in resolving the phylogenetic relationships in this case. Unlike the natural barrier of the Assam Valley, the mountains of SagaingKachin are connected to the Shillong Plateau and Himalayas, making it a logical corridor for the ancestral colonial activity of these moths. This suggests that the "species boundary" of $A$. discata is possibly fuzzy (transitional or gradual) or even not able to be located. Therefore, this paper rejects to use the name subjectively and treats it as "data-deficient" temporarily, however, it does not imply any effective taxonomic (synonymous) treatment for this taxon here. There is still a possibility in the future that the three ecoregional populations will be accepted as different species or subspecies to form a complex; the Sub-Himalayan genus Sinobirma Bryk, 1944 is a known case (Rougerie et al. 2012).

At present, the sources of published specimens of A. compta are localized within the following three restricted ecoregions of Indo-Malayan realm, the zoning based on Dinerstein et al. (2017):

- Meghalaya subtropical forests (Rothschild and Jordan 1899; Rothschild and Jordan 1901; Bouvier and Riel 1931: 53; Arora and Gupta 1979; Peigler 1999; Naumann and Löffler 2015).
- Northern triangle subtropical forests (Bryk 1944).
- Eastern Himalayan broadleaf forests (Naumann and Löffler 2015; Naumann and Nogueira 2021; this article).

Diverse evidence stresses the importance and necessity for conservation of wild $A$. compta in the current situation. As one of the central clues regarding the evolutionary history of the economically important genus Antheraea, its ecological and taxonomic significance is critical.

Larval feeding preference and morphological characters shared by the montane A. godmani, A. montezuma and $A$. meridiana with the Himalayan $A$. compta seemingly suggest that those Neotropical species are closer to the New World ancestor than are the Nearctic species (A. polyphemus, A. oculea). Three centuries of intense study of the genus Antheraea make more rigorous comparisons possible, and detailed phylogenetic studies on preimaginal morphologies and complete mitochondrial genomes of $A$. compta with related species are in preparation, to further confirm its evolutionary status.

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## Supplementary material 1

## Feeding schedule of Antheraea compta $\mathrm{L}_{6}$

Author: Zhengyang Liu
Data type: table (.xlsx file)
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Link: https://doi.org/10.3897/dez.70.102952.suppl1

## Supplementary material 2

## Light sensitivity (the darker side = ca. 5441185 Lux, the brighter side = ca. 9300-13600 Lux) of Antheraea compta $\mathrm{L}_{6}($ speed $\times 12)$

Author: Zhengyang Liu
Data type: video (.mov file)
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Link: https://doi.org/10.3897/dez.70.102952.suppl2

## Supplementary material 3

## Defensive behavior of Antheraea compta $\mathrm{L}_{6}$

## Author: Zhengyang Liu

Data type: video (.mov file)
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