



ARTIMELIA

LEPIDOPTERA IBÉRICA

Biannual Scientific Journal dedicated to research and scientific studies on the Lepidoptera of the Iberian Peninsula.



Artimelia

Lepidoptera Ibérica

Volume 2, Part 1, May 2026

ISSN 3051-6781

Biannual scientific journal dedicated to research and scientific studies on the Lepidoptera of the Iberian Peninsula.

Published on-line by Rede de Estações de Borboletas Noturnas.

Editor

Martin Corley (B.A., Oxon).

E-mail: martin.corley@btinternet.com

Layout Editor: Ana Valadares.

© 2026 Rede de Estações de Borboletas Noturnas.

Front page photo: *Globia algae* (Esper, 1789), Lagoa de Óbidos, 16.vi.2024, and *Friedlanderia cicatricella* (Hübner, 1824), Lagoa de Óbidos, 18.vi.2024 (© Paulo Lemos).

Contents

Editorial	3
Lemos, P. Two wetland-specific Lepidoptera new to Portugal, with notes on ecology, life history and conservation	4
Conde, J., Figueiredo, H., Rodrigues, A. and Boieiro, M. <i>Daphnis nerii</i> (Linnaeus, 1758): registo de um esfingídeo raro em Portugal	31
Revilla, T., Morene, F. and Juana, F. Presencia en España de <i>Willibaldiana culatrae</i> Trematerra, 2014 (Lepidoptera: Tortricidae)	33
Corley, M. <i>Merrifieldia menthae</i> (Chrétien, 1925) (Lepidoptera: Pterophoridae) in Portugal	36
Corley, M. A revision of Portuguese records of the genus <i>Psorosa</i> Zeller, 1846 (Pyralidae: Phycitinae)	41
Ruiz, F. and Garcia, D. <i>Opogona omoscopa</i> (Meyrick, 1893) (Lepidoptera, Tineidae), a new species for Cantabria and new records of <i>Borkhausenia crimnodes</i> Meyrick, 1912 (Lepidoptera, Oecophoridae)	47
Corley, M. <i>Phyllonorycter berberidella</i> Nunes, Laštůvka & Laštůvka, 2026 (Gracillariidae) in Alto Alentejo	56
Corley, M. <i>Depressaria discipunctella</i> Herrich-Schäffer, 1854 in Portugal	59
Guidance for authors	61

Editorial

Articles in this issue of *Artimelia* add one species to the Spanish Lepidoptera list and four to the Portuguese Lepidoptera list, including the first Portuguese record of the sphingid *Daphnis nerii*, for which other records have since been posted on Facebook.

Although this issue is mostly about Portuguese moths, we are very pleased to include two articles on Spanish Microlepidoptera. We would welcome more. We would also like to include short notes and observations and articles of a lighter nature such as memorable moth trapping nights or first experiences with moths.

One vital element in the publication of scientific articles is the peer-review process. This scrutiny helps the editor to decide if an article is worth publishing, or suggests improvements. Without peer-review, relevant literature can be overlooked or other mistakes may be made. Reviewers rarely get the credit they deserve, particularly as they are usually very busy people giving up their valuable time without obvious benefit. I would here like to thank reviewers for this issue including Knud Larsen, Jacques Nel, João Nunes, Jorge Rosete, Martin Townsend, Ana Valadares and José Luis Yela.

Two wetland-specific Lepidoptera new to Portugal, with notes on ecology, life history and conservation

Paulo Lemos^{1,2}

¹ *Independent Researcher, Caldas da Rainha, Portugal.*

² *Escola Superior de Turismo e Tecnologia do Mar; Politécnico de Leiria, Campus 4; Rua do Conhecimento 4; P-2520-614 Peniche; Portugal. email: paulolemos@live.com.pt; <https://orcid.org/0000-0002-3917-0520>*

Abstract

Two wetland-specific moths are documented for the first time in Portugal: *Globia algae* (Esper, 1789) (Lepidoptera: Noctuidae) and *Friedlanderia cicatricella* (Hübner, 1824) (Lepidoptera: Crambidae). Surveys conducted at Lagoa de Óbidos, and later extended to additional localities, provided additional data on their ecology and life histories. Both species are associated with tall helophyte communities characterized by large stands of *Schoenoplectus lacustris* (L.) Palla s.l. The results highlight the importance of dynamically stable micro-sites in wetlands, and support approaches that integrate rural land use with biodiversity conservation.

Keywords: Noctuidae, Crambidae, Apameini, wetlands, host-plants.

Resumo

Duas espécies de lepidópteros de hábitos palustres são documentadas pela primeira vez em Portugal: *Globia algae* (Esper, 1789) (Lepidoptera: Noctuidae) e *Friedlanderia cicatricella* (Hübner, 1824) (Lepidoptera: Crambidae). Levantamentos populacionais realizados na Lagoa de Óbidos e posteriormente alargados a outras localidades, fornecem dados adicionais sobre as suas ecologias e ciclos de vida. Ambas as espécies estão associadas a comunidades de grandes helófitos caracterizados por tufos densos de *Schoenoplectus lacustris* (L.) Palla s.l. Os resultados destacam a importância de microssítios dotados de estabilidade dinâmica em zonas húmidas, apoiando abordagens que integrem as atividades rurais com a conservação da biodiversidade.

Palavras-chave: Noctuidae, Crambidae, Apameini, zonas húmidas, plantas hospedeiras.

Introduction

Knowledge of Lepidoptera in Portugal has been hindered by the cryptic and localized nature of many species. Traditional light-based sampling often fails to detect such populations within dynamic and heterogeneous ecosystems, highlighting the need for more comprehensive and targeted methodological approaches.

Since 2014, several rare populations have been monitored in the remaining natural habitats of central-western Portugal by following their host-plants. The botanical framework supporting these surveys was later strengthened through contributions by the author to the Red List of Vascular Flora of Mainland Portugal, particularly for the Lagoa de Óbidos region (Carapeto et al., 2020; Farminhão et al., 2021).

Subsequent fieldwork revealed several noteworthy findings, identifying the presence of two nocturnal endophytic moths previously unrecorded in the country: *Globia algae* (Esper, 1789) and *Friedlanderia cicatricella* (Hübner, 1824). Both species are associated with tall helophyte communities of the class *Phragmito-Magno-Caricetea* (Landucci et al., 2020), dominated by *Schoenoplectus lacustris* (L.) Palla s.l., which was the only plant on which oviposition was observed.

Given their status as new records for Portugal, their specialized ecology and the scarcity of Iberian records (Farino et al., 2024), these species warrant particular attention, particularly given their tendency toward fragmented distributions and high vulnerability to anthropogenic pressures. Using a multi-approach research framework, this study documents the first national records of these taxa, aims to provide an initial assessment of their populations, and to offer a preliminary evaluation of their conservation status.

Material and methods

Field sessions were conducted across geographically distant municipalities to assess preliminary distribution and abundances. Observations of phenology, ecology, ethology, and host-plant interactions were compared across three Portuguese provinces, and biological material was collected for further research.

Municipalities covered by field sessions:

- Western Coast (Leiria): Alcobaça, Bombarral, Caldas da Rainha, Leiria, Marinha Grande, Nazaré, Óbidos, Peniche.
- Beira Litoral (Coimbra): Figueira da Foz.
- Ribatejo (Santarém): Golegã.
- Setúbal: Santiago do Cacém.

Natural Reserves and protected areas (Rede Natura 2000):

- Reserva Natural do Paul do Boquilobo (UNESCO Biosphere Reserve)
- Reserva Natural das Lagoas de Santo André e da Sancha (SPA; SAC - Natura 2000)
- Reserva Natural Local do Paul de Tornada (Ramsar Site)
- Mata Nacional de Leiria (PGF)
- Dunas de Mira; Gândara e Gafanhas (SAC - Natura 2000)

On-site trapping was employed as an auxiliary method occasionally, utilizing a set of three 10 W mini-projectors (365 nm UVA, unfiltered) mounted on a tripod (Fig. 1) against a white sheet or an aluminum-coated tarpaulin.

Direct observation with high-power handheld flashlights yielded the best results: long-wavelength emitters (deep red light - 660 nm) were used to minimize nocturnal disturbance, while high-power short-wavelength emitters (UVA) were effective for immobilizing fleeing moths, facilitating their identification. For perching insects and larvae, all-around white light flashlight proved to be the most effective tool.

Observations were semi-quantitative and based on repeated standardized field visits, with survey effort controlled by time and area covered per visit in a small pond near Poça do Vau (Lagoa de Óbidos). Sampling sessions were conducted over a two-and-a-half-year period. For the adult stage, surveys were carried out at least four times per week on alternate days during 2024, from May to August, coinciding with the mating seasons of both species and starting at dusk. The most intensive fieldwork took place that year, extending into spring 2025 for the larval stage. In subsequent years, the knowledge gained allowed for less intensive sampling in the water and more efficient surveys from the banks, as a preventive protective measure.

Survey routes ranged from -2 to 50 meters in altitude, primarily across waterlogged areas. Fixed narrow tracks through vegetation were followed, and the systematic or prolonged use of white and UV light was strictly minimized in sensitive areas to reduce environmental impact.

Image documentation was carried out by a Canon EOS 6D with a 100 mm macro lens, employing improvised flash setups with diffusers. Images were later edited in Adobe Lightroom to approximate the field view.

For the host-plants, taxonomic identification was based on the morphological criteria of *Flora Iberica* (Luceño & Jiménez-Mejías, 2008), although adopting the nomenclature of the group (Lansdown, 2020), where *Schoenoplectus lacustris* (L.) Palla s.s. and *S. tabernaemontani* (C.C. Gmel.) Palla are treated as distinct species. However, as noted by the latter author, morphological differentiation is frequently obscured by the presence of the hybrid *Schoenoplectus* × *flevensis* (D.Bakker) Lansdown & Rumsey, which exhibits overlapping characters. In our study, while moth populations were pragmatically considered associated with the parental species, the niche and morphology occasionally overlaps and suggests that host selection may occur across the wider species complex.

Results

Alongside many remarkable riparian and wetland moths, two unexpected discoveries stand out as new records for the country, having likely been overlooked in previous national studies.



Fig. 1 - On-site trapping at Lagoa da Vela left on a bank; close to *S. lacustris* stands, while also inspected directly with handheld flashlights.

Occurring in syntopy, both are associated with aquatic environments, sharing the same main host-plant and an endophytic oviposition, in areas that remain temporarily flooded until the end of the breeding season (from May to late July in western Portugal). In figs. 2 and 3 they are represented by mating pairs, with the females positioned above.

Globia algae (Esper, 1789) - The first Portuguese records of this wetland noctuid originate from Lagoa de Óbidos - Poças do Vau (Óbidos), 29S MD8158, 20.v.2024 (larvae, reared); Reguengo da Parada (Caldas da Rainha), 29S MD8769, 21.v.2024 (larva, reared), where active imagines were also detected days later. Subsequently, records were obtained at two more localities of recognized conservation importance: Reserva da Biosfera do Paul do Boquilobo (Azinhaga), 29S ND4060, 7.vii.2024 (male), and Reserva Natural das Lagoas de Santo André e da Sancha (Santiago do Cacém), 29S NC1614, 10.vii.2024 (male).



Fig. 2 – Mating pair of *Globia algae*.

Friedlanderia cicatricella (Hübner, 1824) - A large crambid associated with marsh and salt-marsh habitats, occupying (often sharing) a similar feeding niche to that of *G. algae*.

Previously mentioned in the Vives Moreno (2014) catalogue without date or locality, this species was not accepted in the national Lepidoptera list because of absence of data (Corley, 2015). The observation at Lagoa de Óbidos, Poças do Vau (Óbidos), 29S MD8158, 18.v.2024, now represents the first documented record for the country (female). It was later recorded in high numbers at the Reserva da Biosfera do Paul do Boquilobo (Azinhaga), 29S ND4060, 6.vi.2024, and sporadically in ditches within floodplains near Chão da Parada (Caldas da Rainha), 29S MD8769, 12.vi.2025. However, it was not detected at Lagoas de Santo André e da Sancha, perhaps because the single survey took place after the seasonal peak and during late-night hours.



Fig. 3 - Mating pair of *Friedlanderia cicatricella*.

To date, three disjunct populations of *G. algae* have been identified in Portugal: the Caldas da Rainha Typhonic Valley (Lagoa de Óbidos and Chão da Parada flood plains), Paul do Boquilobo, and Lagoas de Santo André, which are likely isolated from one another (Fig. 4).

In the coastal regions of the Coimbra district (Dunas de Mira; Gândara e Gafanhas) and Leiria (Mata Nacional de Leiria), surveys produced negative results, with no individuals or interactions recorded during the study period.

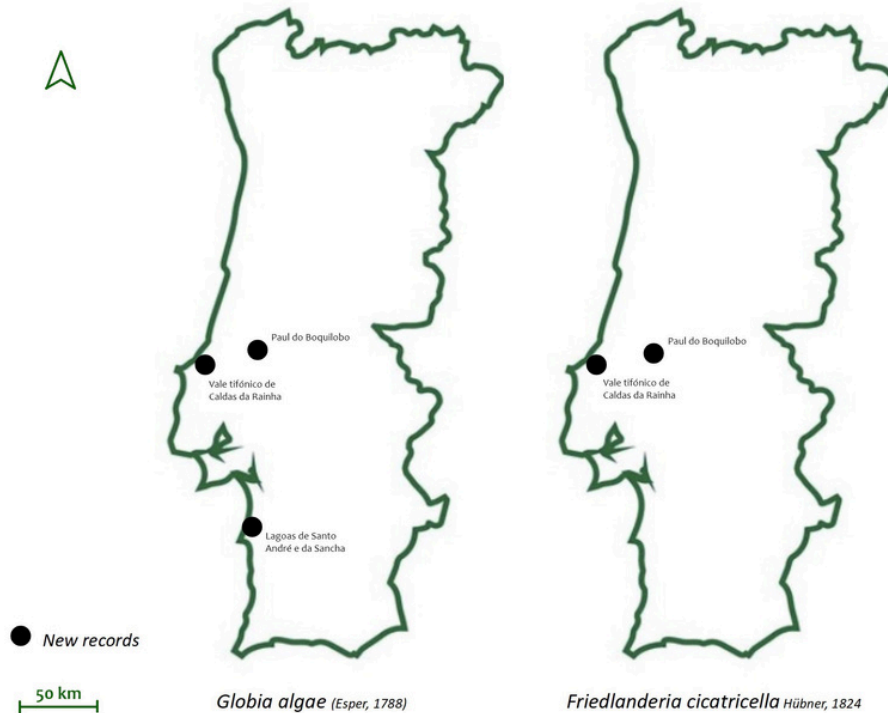


Fig. 4 - Known distribution of *G. algae*, *F. cicatricella* in Portugal (2024).

National distribution maps for the host-plants are available on the Flora-On portal (Flora-On, 2025), which aggregates reliable data from multiple sources.

Due to the erratic flight patterns and high density of individuals in marshy habitat, exhaustive photographic documentation and simultaneous field notation were logistically constrained. Consequently, the hundreds of observed interactions were recorded as real-time visual estimates, with the photographic archive providing a representative subset for taxonomic validation.

A short selection of records is listed at the end of this publication.

Part of the biological material (*G. algae*) is deposited in the José Luis Yela collection (Spain), with additional specimens in the author's private collection (both species).

Discussion

Within the three known disjunct populations of *G. algae* (Caldas da Rainha Typhonic Valley, Paul do Boquilobo and Lagoas de Santo André), the species appears to form metapopulation systems in Paul do Boquilobo and the Caldas da Rainha Typhonic Valley. These are characterized by core populations in a metapopulation context (high density of larvae and breeders annually), residing in small high-quality patches, that likely function as source populations, supporting smaller satellite populations in nearby locations. At Lagoas de Santo André, data are currently insufficient to characterize its population dynamics.

Two micro-sites can be considered hotspots for the species, as core populations:

- A small vestigial pond not far from the large Poça do Vau (Lagoa de Óbidos), dominated by *Schoenoplectus lacustris* (L.) Palla., featuring other tall helophytes in the background. The relative abundance of both moths is significantly higher than at Grande Poça do Vau, located just a few hundred meters away, where dense stands of the hostplant also occur.
- A highly restricted (protected) temporarily flooded area within Reserva da Biosfera do Paul do Boquilobo, featuring large stands of *S. palustris* in deeper areas, closely surrounded by the endangered *Butomus umbellatus* L., several medium-sized nitrophilous alien annuals, and extensive carpets of *Paspalum distichum* L. which seasonally function as a buffer filtering excess nutrients.

In contrast to the Lagoa de Óbidos (largely surrounded by privately owned land), Paul do Boquilobo, and Reserva Natural das Lagoas de Santo André e da Sancha, are subject to active habitat management and hydrological monitoring, outlined in the recent 'Plano de Cogestão', these actions prioritize the conservation of such increasingly rare wetland ecosystems (cf. ICNF, 2024).

The findings highlight the significance of Portuguese marshlands as natural heritage and suggests informed management decisions. Effective conservation depends on an integrated,



Fig. 5a/b

a) Small ancient pond dominated by *S. lacustris*, drying out during the summer (26.iv.2021).

b) Very temporary but exceptionally high water levels were recorded on 12.ii.2026, following the Kristin depression.

The site is surrounded by *Eucalyptus* plantations near the large Poça do Vau. Once part of the Lagoa de Óbidos water surface, the pond is now fed by cryptic overland flow (retaining permanent humid soil during summer). After being completely severed by a road, it became a small stronghold in a forgotten roadside corner, only occasionally reached by temporary floods. Despite its reduced size, it hosts high numbers of *Globia algae* and *Friedlanderia cicatricella*.

ecosystem-based approach that explicitly incorporates host-plant ecology and habitat dynamics, as these form the basis for understanding the associated Lepidoptera species.

Broad expanses of tall-helophyte communities still persist in the studied localities, but the specific associations comprising large stands of *S. lacustris* s.l. (crucial for these moths) are actually highly fragmented.

National distribution maps show many coastal occurrence sites for *S. lacustris* s.l., such as the Baixo Mondego or Ria de Aveiro regions (although excluded from this study). However, the plant's presence does not guarantee the moths' occurrence; in Vale Tifónico de Caldas da Rainha (the starting point for the study area), both species are absent from many suitable sites, likely due to anthropogenic pressures disrupting their life cycles. However, instances of discontinuous occupancy across a vast territory and fragmented habitat are possibly due to historical natural factors, such as the establishment of pioneer plants without their associated biodiversity, or the isolation of regions geographically distant from core populations.

I - Dynamic stability across environmental heterogeneity:

The habitat of *G. algae* and *F. cicatricella* may exhibit diverse geological profiles, reflecting the broad substrate tolerance of their host plant. This tolerance ranges from flooded sandstone to heavy clay, with instances where a single nucleus extends across ecotones of both types.

At Lagoa de Óbidos, *S. lacustris* s.l. thrives under slightly brackish influence (ranging from sites with strong to minimal or non-existent tidal action) as well at the intersection of acidic flows with carbonate-rich waters, within ditches, seasonal floodplains and small ponds. At Boquilobo, geologically and edaphically less diverse, the hardness and alkaline influence is even more pronounced due to Tagus alluvial aquifer (cf. Almeida et al., 2000; ICNF, 2024), leading to significant calcium carbonate precipitation at the base of the plant stalks.

Notwithstanding the plants' relative tolerance and the significant phenological plasticity, and despite the escalating pressures that add more variability, a fundamental ecological stability seems to remain essential. This ensures the precise phenological synchrony between host-plants and moths, high nutritional intake for the larvae, still enabling vigorous plants to withstand high densities of stem borers over consecutive years.

In the study area, some of the *Schoenoplectus* s.l. population nuclei may exhibit a reversed phenology, whereas they had been monitored in previous years coinciding with that of the local core populations. This shift has been identified as an important threat in terms of habitat reduction.

A good example of this is a nucleus near Pequena Poça do Vau, despite the absence of obvious physical damage, initiates its vegetative cycle in late summer and reaches maturity in autumn, leaving virtually no organic traces during the rest of the year. When cultivated in a controlled nursery environment, this plant reverts to an almost standard vegetative period (Lemos, P., pers. obs.). The causes of this discrepancy remain unclear, but it is likely related to local environmental conditions such as water chemistry, organic load or turbidity, which are

known to influence helophyte growth and phenology (Clevering, 1998a, b; Coops et al., 1996; Menéndez et al., 2000), potentially creating a 'toxic window' that may force the plants to alter their phenophases.



Fig. 6 – Impact of occasional autumn sheep grazing on *S. lacustris* stands prior to heavy rainfall. Animals follow established tracks, selectively grazing on green, leafy vegetation in humid soil (e.g., the partially grazed *Sparganium erectum* in the center) while leaving approximately 70% of the dry *S. lacustris* stands intact. This grazing results in variable physical damage due to trampling. Poça do Vau (Óbidos, x.2025).

The effect of non-equine livestock on *S. lacustris* appears less significant than on other spreading helophytes, such as *Phragmites australis*, which are known to dominate shallow water levels in the absence of defoliation.

Not every pond with *S. lacustris* provides the necessary conditions to align the life cycles of host-plants and their inquilines. Whether driven by genetics or shaped by stochastic environmental factors, the phenological synchrony between host-plants and these moths is critical to the success of their interaction.

II - Insights on tall-helophyte communities and herbivory:

Favorable habitat for the studied lepidopterans exhibits a certain level of prolonged periodic flooding, the aforementioned tight phenological synchrony between moths and their host-plants, and often a very moderate or well-balanced grazing presence. Moreover, regulated herbivory may safeguard critical riparian ecotones, increasing resilience to severe wildfires.

All known favourable locations for *G. algae* and *F. cicatricella* may lose water for months, having in common the grazing legacy (cattle, horses, and small ruminants) that extends to aquatic environments where rural activities play a role in maintaining riparian biodiversity.



Fig. 7a/b – Recent shot of floodplain meadows in São Martinho do Porto. Three decades ago, the landscape was characterized by more persistent flooding and active grazing, which likely would not have resulted in the wide, monotonous appearance seen today. Currently, land-use promotes drainage via ditches and constant competition between *P. australis* and *S. lacustris*, driven by summer mowing. These macrophytes co-occur, with the latter being seasonally stunted by the dominance of the former. Furthermore, the egg-clutch niches of the target species are removed or severely altered, hence there are no records across this extensive area.

However, during periods of drought and food shortage, livestock can exert disruptive pressure on these ecosystems, even in the absence of chronic overgrazing.

Some examples in the study areas are horse grazing severely consuming stands of *S. tabernaemontani* situated in permanently damp soil, including the dry shoots, or in the same municipality (via Valado dos Frades), cattle may exert a similar impact on *S. lacustris* s.s.; in all cases adding trampling and passage damage, which may also be common to smaller livestock (sheep and goats), as observed in the Óbidos Lagoon (Fig. 6). All ruminants may graze on the green apical halves of the hostplants located peripherally in shallow ditches and banks, particularly as floodwaters recede (a observed phenomenon in the peripheral areas of Paul do Boquilobo).

The egg-clutches of both moths are vulnerable to a range of disturbances, as they remain inside the stem tips for long periods after endophytic laying, from the beginning of the dry season until the following spring eclosion. Eggs remain in dormancy (in Portugal and probably all Mediterranean populations) through much of the summer and following winter (Edelsten, 1907).

In the two identified core populations, *S. lacustris* green shoots annually emerge from the water in late winter, growing rapidly through spring. Later, much like geophytes, they start to senesce as ponds lose water in midsummer, typically almost completely dry in July after fruiting and the egg-laying period. Senescence at this season is advantageous, since it makes the plants less attractive to herbivores on damp ground, especially when forming large monospecific stands (Duncan, 1992). It remains unclear whether this senescence is biologically pre-programmed following peak fruiting in June, or if it is triggered by changes in water parameters just before the pond becomes a damp bed (Clevering, 1998a, b; Coops et al., 1996).

III - Eutrophication and nitrophilous herbs competition:

In theory, the two core populations suggested by this study combine optimal conditions for the interspecific host-moth relationship, despite being vulnerable to significant water level reductions during summer and autumn.

Water depth acts as a limiting factor for *Phragmites australis*, allowing *S. lacustris* s.l. to maintain a competitive advantage in deeper areas (Coops & van der Velde, 1991). However, when water levels drop, the reed can rapidly displace it through superior rhizomatous expansion (Weisner, 1991), a succession often intensified by eutrophication (Clevering, 1998a). Furthermore, the role of herbivory is a decisive factor; while the absence of large herbivores - following rural abandonment in Portugal (Paiva, 1993) - allows *P. australis* to form dense, impenetrable monospecific stands (Vulink, 2001), moderate grazing can act as a beneficial management tool (Bakker et al., 2013). However, this balance is fragile, as excessive grazing and trampling during dry periods can severely compromise *S. lacustris* (Middleton, 1999). Therefore, as the site of Poça do Vau dries and this plant senesces, livestock (cattle and horses in former times, and now small livestock) often shift their grazing pressure towards nearby helophyte species with later phenologies that remain green and nutritious until the end of the season, such as *Phragmites* (Duncan, 1992; Menard et al., 2002).

In the worst-case scenario non-selective, homogeneous industrial-scale forage harvesting in historical wetlands, which replaced former extensive grazing, has proven adverse, as evidenced by the total absence of both lepidopteran species, even where host-plant stands persist (Fig. 6). The egg clutches are taken away with the forage.

Factors associated with nutrient enrichment may also promote the degradation rate of senescent material (Menéndez et al., 2000), potentially compromising the overwintering eggs of the aforementioned moths. The favourable dynamic equilibrium can be difficult to achieve even when science-based management is promoted. It is crucial to ensure both spatial availability and ecosystem variability, guaranteeing that localized niches remain suitable for *G. algae* and *F. cicatricella*.

IV – Forestry



Fig. 8 - Timber harvesting on the banks of the Lagoa de Óbidos during a flood; March 4, 2026.

Intensive afforestation on the periphery of wetlands and sparse shrublands, areas once characterized by a more stable vegetative balance, inevitably triggers silvicultural runoff and relies on harvesting operations currently deemed aggressive to the ecosystem. This is particularly pronounced along the margins and within the drainage basin of the Lagoa de Óbidos, frequently compounded by agricultural practices.

The core population for *G. algae* and *F. cicatricella* in Poças do Vau is currently threatened by a recent *Eucalyptus* plantation that already suppressed a portion of the pond bed. As the grove matures, *Eucalyptus* leaf litter is expected to further degrade water quality through excessive organic pollution.

V – Drought and wildfires

Potentially suitable habitats for *G. algae* and *F. cicatricella* are found in natural floodplains and dune depressions along the Beira Litoral coast. However, surveys conducted in 2024 yielded negative results across the region, most notably at Lagoa da Vela, which retains well-recovered stands of host-plants. The absence of these species possibly dates back to October 2017, when the region was devastated by the combined impact of extreme drought and severe wildfires, which presumably affected the insects during their egg stage. Even today, in areas largely devoid of livestock grazing, the landscape still exhibits lingering evidence of fire damage.

VI- Notes on *Globia algae*

The life cycle of this Euro-Siberian Noctuidae has been documented for over two centuries, primarily by researchers from Northern and Central Europe where it thrives in local, patchy populations. However, its ecological dynamics and phenology in South-western Europe remain poorly studied.

All Portuguese records of *Globia algae* come from wetland environments, with dense stands of *Schoenoplectus lacustris* (L.) Palla. Adults can be found flying over water bodies, sometimes deep (70 cm). They can be active in heavy rain, and males especially may come into contact with the water (unharmed) when in a search for mates.

A variability is known from the species, ranging from pale forms to “sooty brown” (South, 1907). In the studied populations, the vast majority of *G. algae* males are quite dark (grey-brown), corresponding to the description of *ab. fumata* Warren (though not always evident in photographs; Fig. 10). A few exceptions display a mosaic pattern blending female coloration with the typical local male dark greyish-brown tones.

Schoenoplectus lacustris has been mentioned as a host-plant by authors including Edelsten (1907), Bergmann (1954), Kintzl (1966), Bretherton et al. (1983), as well as other large helophytes, in latitudes where *G. algae* is described as more widespread and less rare than *Globia sparganii* (Esper, 1790), both co-existing occasionally (Bergmann, 1954). After a single record from Lagoa de Santo André, in 1997 (Corley, 2004), there are no recent records of *G. sparganii* in Portugal, currently being considered an Endangered (EN) species, according to the Red Book of Invertebrates of Portugal (Boieiro et al. 2023).



Fig. 9 – Chromatic polymorphism in *G. algae*. Upper half: females. Lower half: males (2024).

a) A primary host plant:

In this study, *Schoenoplectus lacustris* (L.) Palla s.s. was identified as the primary host for regional *G. algae*. Contrary to Central and Northern European literature, no life-cycle stages showed a clear affinity for other helophytes, such as *Typha* spp., *Sparganium erectum* L., or *Phragmites australis* (Cav.) Trin ex. Steud, even when in close proximity to *S. lacustris*. The sole exception was *Iris pseudacorus* L.; at Poça do Vau, several larvae in various instars were found on this species, but only in immediate proximity to *S. lacustris* stands (Fig. 10f): young larvae were observed on the leaves, while larger instars were found within the base of the flower stems as reported by Edelsten (1907). Larvae are highly mobile and disperse in marshy environments immediately upon hatching (Fig. 10c), but there were no observations or signs of egg-laying on *Iris*.

Also, egg-bearing stems that detach from the host-plant may disperse in floodwaters, potentially allowing neonate larvae to colonize suitable peripheral plant stands or even other ponds. However, despite the abundance of *Typha* spp. and *Iris* stands on the periphery of core populations with *S. lacustris* (in Lagoa de Óbidos), larvae are consistently absent from surfaces dominated by other helophytes (described in literature as host-plants) other than *S. lacustris*.

Although larval mobility toward helophyte species is documented elsewhere in Europe (Bretherton et al., 1983), only one internal gallery was detected in *Typha latifolia* L., at the Óbidos Lagoon, located within *S. lacustris* stands. The empty gallery likely belonged to *Nonagria typhae* (Thunberg, 1794), often found on this plant genus in nearby ponds. Consequently, *Iris pseudacorus* remains the only confirmed secondary host plant for *G. algae* in Portugal to date (still highlighting that no egg-laying or dedicated imago patrolling was observed around *I. pseudacorus*).

Occasional resting adults, including those expanding their wings or mating after emerging, found on other plants but close to the primary host, are not considered significant.

b) Breeding season:

In Lagoa de Óbidos (2024–2025), the mating season extends from May to late July, peaking approximately two weeks later than in Paul do Boquilobo and becoming infrequent thereafter. The species is rarely recorded by light trapping or direct observations far from the identified core populations, demonstrating a strong affinity to these. The only instance recorded far from host-plant location occurred repeatedly close to a willow tree covered with abundant aphid honeydew, at Pequena Poça do Vau; however, unlike several *Lenisa geminipuncta* (Haworth, 1809), no adults of *G. algae* were observed feeding.

Both sexes exhibit peak activity at dusk, with males searching among *S. lacustris* stems in flooded or saturated conditions. Mating can occur before total darkness, immediately after females emerge, while they are still soft. Females typically remain just over or close to their exit holes (Fig. 2), therefore mating happens right away, low on the stems, approximately 15–30 cm above the water on intertwined stems.

Post-mating, the female climbs higher and may begin oviposition that same night (Fig. 10a), generally over water, and only dispersing when she is lighter. Many mated females were observed ovipositing during the first half of the night, or later.

Shortly after dark, males conceal themselves within the vegetation – though not at great depth – and the majority remain inactive until dawn, eventually seeking daytime refuge deeper within the dense stands. A few females with large abdomens were observed resting quietly late at night at the breeding sites. This suggests they may have emerged long after dark, when males are no longer receptive, or simply that unmated females do not 'call' (release pheromones) late at night. No late-night matings were recorded. Some of the females appearing to be unmated were subsequently detected in mating pairs the following dusk (confirmed through image analysis several times), in close proximity to their previous locations.



Fig. 10 – Life cycle stages *in situ* from Lagoa de Óbidos (except image “e” - Salir do Porto). From left to right: a) Egg-laying inside the stem (15.vi.2024); b) Egg diapause inside the stems (14.x.2024); c) Recently hatched larva (3.iii.2025); d) First instars on the upper half of the stalks (23.iii.2025); e) Full-grown caterpillar inside the stem (21.v.2024); f) Larva on *Iris pseudacorus* (24.v.2024); g) Movements in search of new stems (20.vi.2025); h) Construction of the pupal chamber with an upward exit hole (22.vi.2024); i) Pupa on a submerged stem (21.v.2024); j) Puparium left in a higher position after emergence (20.vi.2024); l) Opened exit hole (20.vi.2024); m) Mating at dusk almost immediately after female emergence (29.vi.2024), with the female on top.

Observations of ovipositing females demonstrate the function of the double hook-shaped apparatus (Figs. 10a, 11).

Perched on the upper vertical half of a cylindrical *Schoenoplectus* stem, often less than half a centimeter in diameter, sometimes on mature green and more often on younger stems (mid-height), she remains firmly anchored by her legs and positions the tip of her abdomen laterally, engaging the 'hook' which clasps the stem's circumference to initiate an active mechanical incision, exploiting the longitudinal 'grain' of the stem (taking advantage of the structural weakness between the epidermal cells, and the stem in *Schoenoplectus* can be notably hard and slippery). She actively creates a slit to lay the eggs inside. This is consistent with what Edelsten (1907) states: "female is provided with special hooks, which enable her to lift the natural folds of the cuticle to deposit the ova".

The process takes several minutes, repeating the slow process in a downward motion on each selected stem, potentially leaving the female vulnerable to certain bat species. Left behind is a line of minute cracks, also described by Edelsten (1907), each sheltering a variable number of inserted ova. Each site remains slightly moist, likely due to a mucilaginous secretion and/or discharged plant fluids. She gradually changes from stem to stem, increasing the distance each time.

c) Egg stage:

The eggs overwinter inside the stems (fig. 10b), which ideally remain upright and can dry very soon without damaging the eggs within. When the flooding season begins, even if the stalks drop into the water, they remain buoyant for some time due to their internal structure rich in air pockets. Many of these remain trapped above the surface in the tangle of dense stands. Most of the stems remain upright long after drying, sometimes until the next year (even after the floods). Winter temperatures probably help to prevent the material from completely decomposing in humidity before the larvae hatch.

At Poças do Vau, some senescent plant material may become submerged during the cold season. However, in the niches where the species is most abundant, the decayed stems generally do not collapse entirely nor remain underwater for prolonged periods (Figs. 5b, 12a). Furthermore, specific environmental factors must ensure that these stems are not washed away by water currents.



Fig. 11 - Details of the female's ovipositor apparatus in action (opposite side in the last photo). Note: they do not represent the same female.

d) Larval stage:

Neonate larvae are observed in large numbers at Lagoa de Óbidos in early March, remaining active in daylight (consistent with positive phototaxis already reported by Eckstein (1920)). Ecdysis was also documented in captivity, with larvae emerging from senescent stalks stored since the previous summer. Larvae are capable of hatching while submerged, subsequently detaching from the stem either underwater or at the surface. First instar larvae are pale brown with a contrasting dark head capsule (Eckstein, 1920; Zilli et al., 2005) and tend to climb and establish themselves within the apical third of young green stems (Fig. 10d). More developed larvae, which are light green with dark marks (Buckler, 1891; Zilli et al., 2005), with “a brown head and a whitish green plate on first ring of the body” (South, 1907), are very prone to change from stem to stem (Eckstein, 1920); (Fig. 10f), frequently observed at night in the water, floating/swimming and moving between obstacles, as well as temporarily submerged clinging to stems while boring into the tissue (making an entry), a phase in which they may be more vulnerable to predation.

Larvae can be found consistently in central and peripheral deep areas of a small pond, but always in standing waterlogged vegetation.

At the core population of Lagoa de Óbidos, they are actually detected in large numbers (100–200 according to reliable evidence of occupation) in late spring, often in water exceeding 70 cm in depth in May–June.

Following the extensive flooding caused by Storm Kristin in March 2026, which merged all the floodplains of the Vau ponds into a single, large, continuous, and deeply inundated water body, small larvae in the subsequent spring, after water levels had receded, hatched and were competing for the few upright plant stems that managed to protrude above the pond surface (fig. 12 a/b).



Fig. 12a/b: Post-Storm Kristin habitat status of the pond from Fig. 5a/b, on 06.iv.2026, with water levels within the seasonal norm. Major threats include allochthonous pollution accumulated during the floods. Population survival is critically dependent on rapid vegetation recovery for larval shelter; however, regrowth appears delayed this year, despite ongoing larval development. This delay, coupled with the turbidity observed after water levels stabilized, suggests that the primary drivers of habitat degradation could be chemical alterations. Currently, larvae are restricted to only two marginal areas of 1 m² each, where high densities of early instars compete for the few dozen available stems. Most plants visible at the pond's periphery are *Scirpoides holoschoenus* (L.) Soják, which do not function as host plants.

e) Pupal stage:

Similar to *F. cicatricella*, the placement of the slightly elongated pupal chamber depends on the pond's water level at the time of construction. As water levels recede throughout the spring, chambers in different stems are found at varying heights, determined by the water surface rather than the distance from the rhizomes. Likely a defense against non-aquatic predators, the chamber is positioned so the pupa stays below the surface, and the still closed exit hole is excavated in the internal tissue until only a thin, transparent membrane remains, as earlier explained by Edelsten (1907) with pupa head upright (Buckler, 1891; South, 1907). The exit hole is consistently placed at or just above the waterline, and the top of the chamber is blocked by plant fragments just over the exit spot (to indicate the exit). This species was observed pupating in thin stems in which it barely fitted.

In contrast to *Phragmataecia castaneae* (Hübner, 1790), the pupa moves clumsily and only slightly along the chamber. Low mobility was already reported in literature (Edelsten, 1907; Zilli et al., 2005).

In contrast to the larval boring effect, the pupal chamber construction can drastically reduce fluid circulation in the stem, somewhat compromising the structure's lifespan, or even cause bending if forced; but generally, not showing signs of decay in the short term, nor lead to the moth failing to emerge (the pupal stage is notably brief, lasting from less than one week to two, likely depending on meteorological conditions). Sometimes, especially in shady conditions and if the water level has dropped significantly, the stem can break next to the exit hole (weak point). If it breaks within a sudden late season flood, the pupa may drown if water enters, or fall out and sink. During the pupal stage (nearly the same as flying period), the summer arrives with partial dormancy of the plant (in Lagoa de Óbidos), and many of the oldest healthy stems lose their vitality, gradually turning yellow or brown, camouflaging all traces of impact left by *G. algae*.

f) Predation:

Confirmed predators include the brown rat *Rattus norvegicus* (Berkenhout, 1769) which swims, climbs the vegetation, and takes larvae and pupae if the water level drops too much; also the great tit *Parus major* Linnaeus, 1758, which bores into stems. Possibly also the red swamp crayfish *Procambarus clarkii* (Girard, 1852), as it infests all aquatic systems where this moth is found but the impact is probably insignificant. Interestingly, the low flight pattern of the males - the lowest observed by the author among noctuids, barely skimming the water's surface - appears to render bat predation quite difficult. While the western swamphen *Porphyrio porphyrio* (Linnaeus, 1758) is a potential destroyer of tall helophytes in deep waters and along banks, no significant host plant damage was confirmed at any studied location. Similarly, signs of predation by smaller Rallidae (which Edelsten (1907) reports as serious predators of the species) were observed sparingly in Natural Reserves and more frequently in the large Poça do Vau, though data remains insufficient to determine if they act as a limiting factor.

Observations revealed a few egg laying females exhibiting significant physical compromise and the appearance of having struggled in the water. This suggests high reproductive resilience following accidents such as predator encounters or emergence failures caused by submerged host stalks, or difficulty climbing to a dry perch. Even in ponds with high densities of aquatic predators (dragonfly larvae, crayfish, and frogs), these damaged or even unable to fly individuals appear to achieve sufficient reproductive success.

g) A metapopulation strategy:

Regional observations suggest that *G. algae* is a highly specialized moth adapted to a metapopulation lifestyle, taking advantage of naturally fragmented niches. This strategy allows for high population densities to offset the scarcity of niches with optimal conditions. Provided that host-plants encounter favourable growth conditions, larvae can infest small flooded areas annually with no serious impact on the plant nucleus.

In the two known core populations, the density of mined stems (including empty galleries) can exceed 60–70% without compromising the habitat. Reproductive behaviour further reinforces site fidelity: by depositing an initial batch of eggs at her emergence site, the female ensures the next generation inhabits a proven, viable environment. Furthermore, evidence from captive specimens indicates a lifespan exceeding one week with gradual oviposition, suggesting significant dispersal potential; the species is not restricted to its natal site and may colonize new habitats that may arise. However, while this metapopulation strategy has historically been effective, the resulting isolation of populations now leaves the species increasingly vulnerable to habitat loss and anthropogenic interference.

VII - Notes on *Friedlanderia cicatricella*

The first Portuguese occurrence of this Crambid has finally been confirmed, in the marshlands of Lagoa de Óbidos. Notably, specimens from this location exhibit a broader chromatic range than those from the Paul do Boquilobo Nature Reserve, especially the females.

Males and females are very active at dusk, mating and ovipositing around the host-plants near or over water, rarely straying from waterlogged areas (depths 5–70 cm). They may fly low, come into contact with the water and even rest on it (Fig. 14g). As the night advances, activity of both sexes tends to stop.

Mating occurs from dusk onwards, often just before emerged females expand their wings, generally in low position, but sometimes higher than *Globia algae*.

Eggs are laid inside the healthy stems, not necessarily upstanding or in high position (very variable), and larvae can be found closer to the base of the plants, or more rarely up to the last third (younger larvae). The larvae (Fig. 14b) are somewhat fragile and clumsy outside the host-plant, and they have never been observed naturally outside the plant. As in *Globia algae*, the position of the pupal chambers inside the stems is more determined by the pond's water surface than by the position of the plant's stem base. In observed cases, the pupa is oriented head-up or, in some instances, horizontally.



Fig. 13 - *Schoenoplectus tabernaemontani* (C. C. Gmel.) Palla; well developed, downstream of a runoff through tidal marsh vegetation. Óbidos Lagoon (Óbidos, x.2025).

Although the pupae are quite mobile inside the pupal chamber, utilizing specialized spines along the body (Schütze, 1931), (Zilli et al., 2005), this mobility is not nearly as advanced as that of *Phragmataecia castaneae*. This behaviour appears to be a defense against predators, rather than other cause.

The oviposition, larvae, and pupal stages were documented in dense stands of *S. lacustris*, in part earlier reported by Reutti (1853) and Schütze (1931). At Lagoa de Óbidos (Bom Sucesso), the larva was also found in a well-developed nucleus of *Schoenoplectus tabernaemontani* (C. C. Gmel.) Palla., downstream of a runoff covered with tidal marsh vegetation (Fig. 12), immediately above the tide line, under strong marine influence.

At Paul do Boquilobo, this species can be locally very numerous, far exceeding *Globia algae*.



Fig. 14 – From left to right in descending order: a) Egg-laying in *S. lacustris* (9.vi.2024). b) Larva in *S. lacustris* (23.v.2024). c) Pupa in *S. lacustris* (28.v.2024). d) Puparium in *S. lacustris* (5.vii.2024). e-h) Males. i-l) Resting females (lower pictures). a-c; e; g-m) Óbidos Lagoon. d;f) Paul do Boquilobo Biosphere Reserve. The set of pictures does not highlight the size difference between males and females.

VIII – Threats

Environmental and Ecological Threats:

Reed Expansion - Reed beds reducing open water areas.

Eutrophication and Pollution - Nutrient enrichment from agricultural runoff and high organic loads.

Phenological Mismatch - Changes in plant life cycles that disrupt synchrony with moth colonies.

Litter Degradation - Accelerated decomposition of senescent material, potentially destroying overwintering eggs.

Management and Human Activity Threats:

Industrial Forestry and Agriculture - Eucalyptus plantations (causing organic pollution) and large-scale forage harvesting.

Hydrological Changes - Wetland drainage and failure of dynamical equilibrium conditions.

Rural Exodus - Abandonment of traditional activities and reduction of beneficial extensive grazing.

Localized Pressures - Overgrazing, harvesting for basketry, and wildfires exacerbated by droughts.

Biotic and Genetic Threats:

Invasive and Weed Species - Presence of harmful species like the red swamp crayfish (*Procambarus clarkii*) and nitrophilous weeds that compete with native flora and increase the organic pollutant load.

Herbivory and Trampling - Impact of wild boar (*Sus scrofa*) on seedlings and banks.

Genetic Risks - Introduction of external host-plant genetic material and proliferation of hybrids due to unnatural environmental changes. These can result in host-plant stands with genetics and possibly phenology divergent from the native.

Habitat Fragmentation - Loss of localized niches and lack of space availability for specific breeding sites.

IX – Conservation

The persistence of both *Globia algae* and *Friedlanderia cicatricella* in Portugal appears to be closely associated with the availability of small wetland microhabitats dominated by *Schoenoplectus lacustris* that remain structurally persistent under fluctuating hydrological conditions. In the studied sites, these habitats supported high densities of larvae and breeding adults, suggesting that they function as core population nuclei within a fragmented landscape.

Observations of the life cycle indicate that eggs are deposited inside the stems and remain in diapause for extended periods, including during the dry season and winter. As a result, disturbances that remove or damage senescent stems, such as harvesting, trampling, or

vegetation clearance, may directly affect overwintering stages and reduce local recruitment. This highlights the potential sensitivity of these species to small-scale habitat disturbance.

The study also suggests that a balance between hydrological variability and habitat stability is important for maintaining suitable conditions. Temporary flooding, followed by gradual drying, appears to support both host-plant development and larval survival. Similarly, moderate grazing pressure may contribute to maintaining habitat heterogeneity, although excessive grazing during dry periods could negatively impact host-plant structure and reproductive success.

The absence of the target species from apparently suitable habitats indicates that the presence of the host plant alone is not sufficient. This suggests that additional factors, such as phenological synchrony, water quality, and disturbance regimes, may play a key role in determining habitat suitability.

Given the localized nature of the observed populations and their apparent dependence on specific microhabitat conditions, conservation efforts may benefit from focusing on the maintenance of small-scale habitat mosaics. Protecting areas where dense stands of *Schoenoplectus lacustris* coincide with stable hydrological conditions could be particularly important for ensuring long-term population persistence.

Although the present study is based on a limited number of sites, it provides initial evidence that these species may be vulnerable to habitat fragmentation and changes in land use. Further research incorporating quantitative monitoring would be valuable to better assess population trends and refine conservation strategies.

Mitigation strategies should be defined at a local scale and evaluated on a case-by-case basis. The precautionary principle (Carraro & Siniscalco, 1993) should guide wetland management, particularly when enhancing habitat availability or reinforcing host-plant populations. It is recommended to prioritize the protection of autochthonous plant communities and prevent the introduction of external genetic material, while remaining vigilant regarding the proliferation of phenotypic forms or hybrids that may arise from unnatural environmental changes.

Legal Protection and Land Management:

Niche Protection - Establish formal protection and restoration by implementing micro-reserves or long-term local management agreements.

Private Land Agreements - Collaborate with large-scale forage and grazing farms to secure specific micro-reserves in small flooded areas that meet the species' environmental requirements.

Habitat and Resource Regulation:

Harvesting Restrictions - Prohibit or strictly avoid the traditional harvesting of *S. lacustris* (common club-rush) within primary breeding hotspots to maintain habitat structural integrity.

Herbivory Control - Implement rigorous monitoring and mitigation protocols to prevent livestock overgrazing in sensitive ecological zones, utilizing seasonal fencing where necessary.

Ecological Restoration:

Helophyte Recovery - Prioritize the restoration of degraded wetland plant communities with *Schoenoplectus lacustris*, focusing on preserving local genetic resources.

Host-plant Expansion - Target the strategic expansion of the primary larval host-plant into large, high-density stands.

Research and Monitoring:

Applied Research - Promote and expand scientific studies on wetland ecology in Portugal to bridge current knowledge gaps regarding associated biodiversity.

Impact Assessment - Establish long-term monitoring programs to evaluate the success of restoration efforts.

Selection of relevant records

Globia algae (Esper, 1789)

Poça do Vau, Óbidos, 20.v.2024 (handheld flashlight).

Several last instar larvae on *Schoenoplectus lacustris*.

Bouro, Reguendo da Parada (Caldas da Rainha), 21.v.2024 (handheld flashlight).

Last instar larva on *Schoenoplectus lacustris* (daylight observation) .

Poça do Vau, Óbidos, 22.v.2024 (handheld flashlight).

Two females on *Schoenoplectus lacustris* (handheld flashlight).

Poça do Vau, Óbidos, 23.v.2024 (handheld flashlight).

Larva moving on stalk and over water (swimming), within *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 12.vi.2024 (handheld flashlight).

Mating on *Iris pseudacorus*, on *Schoenoplectus lacustris* with exit hole (female's) at a distance of 1 centimeter.

Poça do Vau, Óbidos, 12.vi.2024 (handheld flashlight).

Mating pair on dry leaves of *Typha* sp. (hiding place?) of a female, probably a virgin photographed in the previous morning on *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 15.vi.2024 (handheld flashlight).
Female, laying eggs on green *Schoenoplectus lacustris* stalks.

Poça do Vau, Óbidos, 15.vi.2024 (handheld flashlight).
Larva on an *Iris pseudacorus* stalk, in a pond with *Schoenoplectus lacustris*.

Paul do Boquilobo, Azinhaga, 5.vii.2024 (daytime observation).
Pupal chamber and puparia on *Schoenoplectus lacustris*.

Paul do Boquilobo, Azinhaga, 7.vii.2024 (handheld flashlight).
Male on patrol near *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 25.vii.2024 (handheld flashlight).
Male on *Schoenoplectus lacustris* (the latest record of the species on the West Coast so far).

Lagoas de Santo André, Santiago do Cacém, 10.vii.2024 (handheld flashlight).
Male patrolling the shore with *Phragmites australis* and *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 3.iii.2025 (daytime observation).
Neonate larva in *Schoenoplectus lacustris* on immerse litter.

Poça do Vau, Óbidos, 23.iii.2025 (daytime observation)
Small larva inside the tip of the *Schoenoplectus lacustris* stalk.

Friedlanderia cicatricella (Hübner, 1824):

Poça do Vau, Óbidos, 18.v.2024 (handheld flashlight).
Female laying eggs on *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 23.v.2024.
Male patrolling on *Schoenoplectus lacustris* (handheld flashlight).

Poça do Vau, Óbidos, 23.v.2024 (daylight observation).
Last instar larva on *Schoenoplectus lacustris* (reared afterwards, emerged on 28.vi.2024).

Poça do Vau, Óbidos, 28.v.2024 (handheld flashlight).
Female pupa on *Schoenoplectus lacustris* (reared afterwards, emerged on 8.vi.2024).

Poça do Vau, Óbidos, 18.vi.2024, 21.27 WEST (handheld flashlight).
Mating couple on *Schoenoplectus lacustris*; tender female expanding wings during the process.

Paul do Boquilobo, Azinhaga, 5.vi.2024 (daytime observation and handheld flashlight).
Several imagines of both sexes, larvae, a dead larva, and puparia (daytime observation), on *Schoenoplectus lacustris*.

Bouro, Reguendo da Parada (Caldas da Rainha), 12.vi.2025 (handheld flashlight).

Mating couple on *Schoenoplectus lacustris*.

Poça do Vau, Óbidos, 17.vii.2024 (handheld flashlight).

Male on *Schoenoplectus lacustris* (the latest record of the species on the West Coast).

Acknowledgments

I would like to express my sincere gratitude to everyone who helped with the data collection and inspired me in the preparation of this paper. Special thanks go to Martin Corley for challenging me to publish as an author several years ago, and to the editorial team for their patience and professional support in processing such a lengthy manuscript. I would also like to thank Fernando Pereira (ICNF) for his hospitality and for his previous tip regarding the club-rush population in Paul do Boquilobo. I am also grateful to Martin, José Luis Yela and Pedro Pires for their valuable insights and for difficult-to-obtain publications. Finally, I wish to thank the anonymous landowners whose field work and dedication to traditional farming and extensive land management in Lagoa de Óbidos continue to sustain this unique biodiversity hotspot.

References

- Almeida, C., Mendonça, J.J.L., Jesus, M.R. & Gomes, A.J. 2000. *Sistemas Aquíferos de Portugal Continental*. 350 pp. Instituto da Água, Lisboa.
- Bakker, J.P., Esselink, P., Dijkema, K.S., van Duin, W.E. & de Jong, D.J. 2013. Grazing and large herbivores in wetland ecosystems. In D. Batzer & A. Baldwin (eds): *Wetland habitats of North America: Ecology and conservation*: 51–66. University of California Press, Berkeley.
- Bergmann, A. 1954. *Die Großschmetterlinge Mitteleuropas*. Band 4/1, Eulen: Verbreitung, Formen und Lebensgemeinschaften. Urania-Verlag, Jena.
- Boieiro, M., Ceia, H., Caramujo, M.J., Cardoso, P., Garcia Pereira, P., Pires, D. & Rego, C. (eds.) 2023. *Livro Vermelho dos Invertebrados de Portugal Continental*. 468 pp. FCIências.ID & ICNF I.P., Lisboa.
- Bretherton, R.F., Goater, B. & Lorimer, R.I. 1983. Noctuidae: Cuculliinae to Hypeninae. In J. Heath (ed.): *The Moths and Butterflies of Great Britain and Ireland*, **10**: 36–413. Harley Books, Colchester.
- Buckler, W. 1891. *The Larvae of the British Butterflies and Moths*. Vol. 4 (The Mottled Willow to the Small Angle Shades). 115 pp. + plates. Ray Society, London.
- Carapeto, A., Francisco, A., Pereira, P. & Porto, M. 2020. *Lista Vermelha da Flora Vasculare de Portugal Continental*. 374 pp. Sociedade Portuguesa de Botânica, PHYTOS & ICNF, Lisboa.
- Carraro, C. & Siniscalco, D. 1993. *The Precautionary Principle in Environmental Policy*. Edward Elgar Publishing, Cheltenham.
- Clevering, O.A. 1998a. An investigation into the effects of nitrogen on growth and morphology of helophytic species (*Phragmites australis*, *Scirpus lacustris*). *Aquatic Botany*, **60**: 185–201.
- Clevering, O.A. 1998b. Effects of nitrogen, phosphorus and potassium supply on the growth of *Phragmites australis* and *Scirpus lacustris*. *Aquatic Botany*, **61** (2): 153–167.
- Coops, H. & van der Velde, G. 1991. Differences in competence between *Phragmites australis* and *Schoenoplectus lacustris* in a depth gradient. *Freshwater Biology*, **26** (1): 71–82.
- Coops, H., van den Brink, F.W.B. & van der Velde, G. 1996. Growth and morphological responses of four helophyte species in an experimental water-depth gradient. *Aquatic Botany*, **54** (1): 11–24.
- Corley, M.F.V. 2004. Provisional list of the Lepidoptera of Lagoa de Santo André, Baixo Alentejo, Portugal. *SHILAP Revista de lepidopterologia*, **32** (126): 105–138.
- Corley, M.F.V. 2015. *Lepidoptera of Continental Portugal. A fully revised list*. 288 pp. Martin Corley, Faringdon.
- Duncan, P. 1992. *Horses and Grasses: The Nutritional Ecology of Free-ranging Horses and Their Impact on the Vegetation*. Springer-Verlag, New York.
- Eckstein, K. 1920. *Die Schmetterlinge Deutschlands mit besonderer Berücksichtigung ihrer Biologie*. Band 3: Die eulenartigen Falter. 96 pp. + plates. K. G. Lutz, Stuttgart.
- Edelsten, H.M. 1907. The life-history of *Archanara algae*. *The Entomologist's Record and Journal of Variation*, **19**: 7–10.
- Farino, T., Ortiz, A.S. & Yela, J.L. 2024. Segundo registro ibérico de *Archanara neurica* (Hübner, 1808) (Lepidoptera: Noctuidae: Noctuinae: Xylenini: Apameina): implicaciones biogeográficas. *Arquivos Entomológicos*, **30**: 153–158.
- Farminhão, J. (ed.) 2021. *Sítios de Interesse Botânico de Portugal Continental: Tomo II (Coleção Botânica em Português, Vol. 5)*. Imprensa Nacional-Casa da Moeda, Lisboa.
- Flora-On. 2025. *Flora de Portugal Interativa*. Sociedade Portuguesa de Botânica. Available at: <https://flora-on.pt/>.
- ICNF. 2024. *Plano de Cogestão da Reserva Natural do Paul do Boquilobo 2024-2027*. 85 pp. Instituto da Conservação da Natureza e das Florestas, Lisboa.
- Kintzl, W. 1966. *Archanara algae* - Funde in *Scirpus lacustris* (Lep., Noctuidae). *Entomologische Zeitschrift*, **76** (7): 78–79.
- Klika, J. & Novák, V. 1941. *Die Pflanzengesellschaften des Sumpf- und Feuchtgebietes Mitteleuropas*. J. G. Calve Verlag, Prague.
- Landucci, F., Šumberová, K., Tichý, L., Chytrý, M., et al. 2020. Classification of the European marsh vegetation (*Phragmito-Magnocaricetea*) to the association level. *Applied Vegetation Science*, **23** (2): 297–316.
- Lansdown, R.V. & Rumsey, F. 2020. *Schoenoplectus × flevensis* (*S. lacustris* × *S. tabernaemontani*, Cyperaceae) in Britain. *British & Irish Botany*, **2** (3): 190–203.

- Luceño, M. & Jiménez-Mejías, P. 2008. Cyperaceae. In S. Castroviejo (ed.): *Flora Iberica*, **18**: 109–250. Real Jardín Botánico, CSIC, Madrid.
- Menard, C., Duncan, P., Fleurance, G., Georges, J.Y. & Lila, M. 2002. Comparative foraging and nutrition of horses and cattle in European wetlands. *Journal of Applied Ecology*, **39** (1): 120–133.
- Menéndez, M., Martínez, M., Hernández, O. & Comín, F.A. 2000. Main factors controlling aquatic macrophyte decomposition in a shallow Mediterranean coastal lagoon. *Hydrobiologia*, **435** (1): 153–164.
- Middleton, B.A. 1999. *Wetland Restoration: Flood Pulsing and Disturbance Dynamics*. John Wiley & Sons, New York: 135–150.
- Paiva, J. 1993. *A flora e a vegetação da Reserva Natural do Paul de Arzila*. 66 pp. Serviço Nacional de Parques, Reservas e Conservação da Natureza, Lisboa.
- Reutti, C. 1853. *Uebersicht der Lepidopteren-Fauna des Grossherzogthum's Baden*. 216 pp. Herder'sche Verlagsbuchhandlung, Freiburg im Breisgau.
- Schütze, K.T. 1931. *Die Biologie der Kleinschmetterlinge unter besonderer Berücksichtigung ihrer Nährpflanzen e Erscheinungszeiten*. Verlag des Internationalen Entomologischen Vereins e.V., Frankfurt am Main.
- South, R. 1907. *The Moths of the British Isles*. First Series. Frederick Warne & Co., London & New York.
- Vives Moreno, A. 2014. *Catálogo sistemático y sinonímico de los Lepidoptera de la Península Ibérica, de Ceuta, de Melilla y de las Islas Azores y Canarias*. 1184 pp. SHILAP Revista de lepidopterología, Madrid.
- Vulink, J.T. 2001. *Hungry herds: Management of temperate lowland wetlands by grazing*. 382 pp. University of Groningen, Groningen.
- Weisner, S.E.B. 1991. Within-lake patterns in depth penetration of emergent macrophyte communities. *Freshwater Biology*, **26** (1): 133–142.
- Zilli, A., Ronkay, L. & Fibiger, M. 2005. *Noctuidae Europaeae, Vol. 8: Apameini*. Apollo Books, Stenstrup.

***Daphnis nerii* (Linnaeus, 1758): registo de um esfingídeo raro em Portugal**

José Conde ^(1,*), Hugo Figueiredo, António Rodrigues & Mário Boeiro ⁽²⁾

⁽¹⁾Centro de Interpretação da Serra da Estrela (CISE), Rua Visconde de Molelos, 6270-423 Seia, Portugal.

email: joseconde@cise.pt; <https://orcid.org/0000-0003-2981-7076>

⁽²⁾Centro de Ecologia, Evolução e Alterações Ambientais (cE3c) & CHANGE - Instituto para as Alterações Globais e Sustentabilidade, Universidade dos Açores, Angra do Heroísmo, Portugal. <https://orcid.org/0000-0002-9087-091X>

Resumo

Um exemplar de *Daphnis nerii* (Linnaeus, 1758) foi observado em Seia (Serra da Estrela), a 26 de setembro de 2014, provavelmente atraído por luz artificial. Este corresponde ao primeiro registo confirmado da espécie para Portugal.

Abstract

A specimen of *Daphnis nerii* (Linnaeus, 1758) was observed in Seia (Serra da Estrela, Portugal) on 26 September 2014, probably attracted to artificial light. This represents the first confirmed record of the species for Portugal.

A 26.xi.2014, um indivíduo fêmea da espécie *Daphnis nerii* (Linnaeus, 1758) (Lepidoptera: Sphingidae) foi recolhido ao início da noite no Estádio Municipal de Seia, provavelmente atraído pela iluminação do complexo desportivo, tendo sido entregue a técnicos do Centro de Interpretação da Serra da Estrela, em Seia. A espécie é um heterócero de grandes dimensões (90-110 mm de envergadura) originário da região Afrotropical e da metade ocidental da região Oriental (Pérez De-Gregório *et al.*, 2001), tendo sido detetada a sua presença num período de temperaturas anormalmente elevadas para a época do ano, resultante da prevalência de ventos do quadrante sul, com origem no Norte de África. Salienta-se que no mesmo período, também nesta cidade, foram efetuados registos envolvendo exemplares de outras espécies de macroheteróceros migradores pouco comuns na região, nomeadamente *Acherontia atropos* (Linnaeus, 1758) e *Hippotion celerio* (Linnaeus, 1758), tendo a sua presença chamado a atenção de residentes da localidade.

Daphnis nerii caracteriza-se pela sua capacidade de efetuar amplos movimentos migratórios, podendo, exemplares procedentes de África, alcançar vastas áreas do continente Europeu, desde o Báltico, a Norte, ao Cáucaso, a Leste, sobretudo em anos mais quentes (Pérez De-Gregório *et al.*, 2001). Durante os voos migratórios, estes heteróceros deslocam-se à noite, sendo frequentemente atraídos por luzes artificiais. Na Península Ibérica, os registos desta espécie são escassos, ainda que repartidos por amplas áreas da geografia peninsular, em particular ao longo das zonas mais litorais (Pérez De-Gregório *et al.*, 2001; Montagud-Alario & Engra, 2017). Curiosamente, foram já observadas lagartas desta espécie em Espanha (Montagud-Alario & Engra, 2017). Segundo Corley (2015), embora *Daphnis nerii* figure na lista dos lepidópteros de Portugal de Karsholt & Razowski (1996), não há registos publicados para o país.

Esta observação reveste-se de especial interesse, uma vez que constitui a primeira citação indubitável da espécie para Portugal e um dos poucos registos conhecidos no interior peninsular. Por fim, dado que o exemplar referido exibia evidentes danos no abdómen, após se proceder ao seu registo fotográfico (Figura 1), tomou-se a opção de o conservar na coleção do Centro de Interpretação da Serra da Estrela.

Referências

- Corley, M.F.V. 2015. *Lepidoptera of Continental Portugal. A fully revised list*. 288 pp. Martin Corley, Faringdon.
- Karsholt, O. & Razowski, J. 1996. *The Lepidoptera of Europe: A distributional Checklist*. 380 pp. Apollo Books.
- Montagud-Alario, S. & Engra, M.A. 2017. Nuevo registro de *Daphnis nerii* en la Península Ibérica (Lepidoptera: Sphingidae). *Boletín de la Sociedad Entomológica Aragonesa*, **60**: 369-371.
- Pérez De-Gregório, J.J., Muñoz, J. & Rondós, M. 2001. *Atlas fotográfico de los lepidópteros macroheteróceros ibero-baleares 2*. 210 pp. Argania Editio, Barcelona.



Figura 1. Exemplar de *Daphnis nerii*, recolhido em Seia, a 26 de novembro de 2014.
(Fotografia: © Hugo Figueiredo)

Presencia en España de *Willibaldiana culatrae* Trematerra, 2014 (Lepidoptera: Tortricidae)

Txema Revilla ^{1,4}, Francisco Morente ² & Fernando de Juana ³

¹ c/ Simón Otxandategi, 122. E-48640 Berango (Vizcaya) (ESPAÑA / SPAIN). e-mail: txema.revilla@gmail.com

² c/ Camino de La Zubia, 21, 2ºC. E-18006 Granada (ESPAÑA / SPAIN). e-mail: paleohistoria@gmail.com

³ c/ Adriano VI 7A, 6º izqda. E-01008 Vitoria-Gasteiz (Alava) (ESPAÑA / SPAIN). e-mail: fdejuana@euskalnet.net

⁴ Autor para la correspondencia / Corresponding author

Resumen

Se cita *Willibaldiana culatrae* Trematerra, 2014 (Lepidoptera: Tortricidae) por primera vez para España, y segunda para Europa, gracias a la captura de 4 ejemplares en septiembre de 2022 en la depresión de Baza (Granada). Se aportan también fotografías del adulto y su genitalia, así como un mapa con la ubicación en la península ibérica de las localidades en la que fueron capturados los ejemplares.

Palabras clave: Lepidoptera, Tortricidae, *Willibaldiana culatrae*, primera cita, Granada, España.

Abstract

Occurrence of *Willibaldiana culatrae* Trematerra, 2014 (Lepidoptera: Tortricidae). *Willibaldiana culatrae* Trematerra, 2014 (Lepidoptera: Tortricidae) is reported for the first time in Spain, only the second record for Europe, by the capture of four specimens in September 2022 in the depression of Baza (Granada). Photographs of the adult and its genitalia are also provided, as well as a map showing the current distribution on the Iberian Peninsula.

Key words: Lepidoptera, Tortricidae, *Willibaldiana culatrae*, new record, Granada, Spain.

Introducción

Como resultado de los muestreos sistemáticos que se llevan realizando en la provincia de Granada en los últimos años, se ha encontrado una interesante especie de Tortricidae que resulta ser nueva para Granada y España; se trata de *Willibaldiana culatrae* Trematerra, 2014.

El género *Willibaldiana* Larsen, 2013 (Lepidoptera: Tortricidae, Olethreutinae, Eucosmini) contenía inicialmente dos especies *Willibaldiana paasi* Larsen, 2013 y *Willibaldiana schmitzi* Larsen, 2013 ambas originarias de las Islas Canarias (Fuerteventura), a las que se añadió una tercera *Willibaldiana culatrae* Trematerra, 2014, del suroeste de la península ibérica.

Esta tercera especie *W. culatrae*, fue descrita de Ilha da Culatra, Algarve (Portugal); Trematerra (2014) indica que el hábitat donde se localizó esta especie es una zona baja y arenosa con escasa vegetación de dunas, con especies como *Lotus* L. y *Medicago* L. entre otras.

En Granada, el biotopo donde se ha encontrado *W. culatrae* es un criptohumedal enclavado en un área con una elevada salinidad del suelo, donde abunda una flora halófila y marítima, a pesar de estar a más de un centenar de kilómetros del litoral. Entre otras, destacan por su abundancia, las salsolas, *Salicornia fruticosa* (L.) L. y dos especies de *Limonium* Mill. que son endémicas de esta zona.

Material y método

Los ejemplares fueron atraídos por trampas de luz fluorescente UV + actínica de 8W, alimentadas por baterías de 12V.

Para la identificación de la especie se han tenido en cuenta tanto los caracteres morfológicos externos, como la estructura genital. La preparación de los órganos genitales se ha efectuado siguiendo el método de Robinson (1976).

Asimismo, se ha utilizado el programa Adobe Photoshop© para el retoque fotográfico.

Material estudiado

Granada: Cúllar (depresión de Baza), 4 ♀ ♀, 840 m, 2-ix-2022, Fr. Morente leg. prep. gen. FdJ-1675 (Figs. 1 y 2).

Resultado

Se trata de la primera cita de esta especie para Granada y España, y la segunda para la fauna europea, ampliando su distribución a zonas halófilas del interior de Andalucía (mapa Fig. 3).

Agradecimientos

No podemos terminar esta nota sin el expreso agradecimiento a Martin Corley por su inestimable y desinteresada ayuda y a la Consejería de Agricultura, Ganadería, Pesca y Desarrollo Sostenible de la Junta de Andalucía la concesión de las autorizaciones necesarias para poder realizar estos estudios y trabajo de campo.

Bibliografía

- Larsen, K. 2013. A new genus and two new species of Tortricidae (Lepidoptera) from the Canary Islands. *Phegea*, **41** (3): 50-54.
- Robinson, G. S. 1976. The preparation of slides of Lepidoptera genitalia with special reference to the microlepidoptera. *Entomologist's Gazette*, **27**: 127-132.
- Trematerra, P. 2014. *Lobesia arzilae* sp. n. and *Willibaldiana culatrae* sp. n. new species from Portugal (Lepidoptera: Tortricidae: Olethreutinae). *Journal of Entomological and Acarological Research*, **46** (2): 66-69.

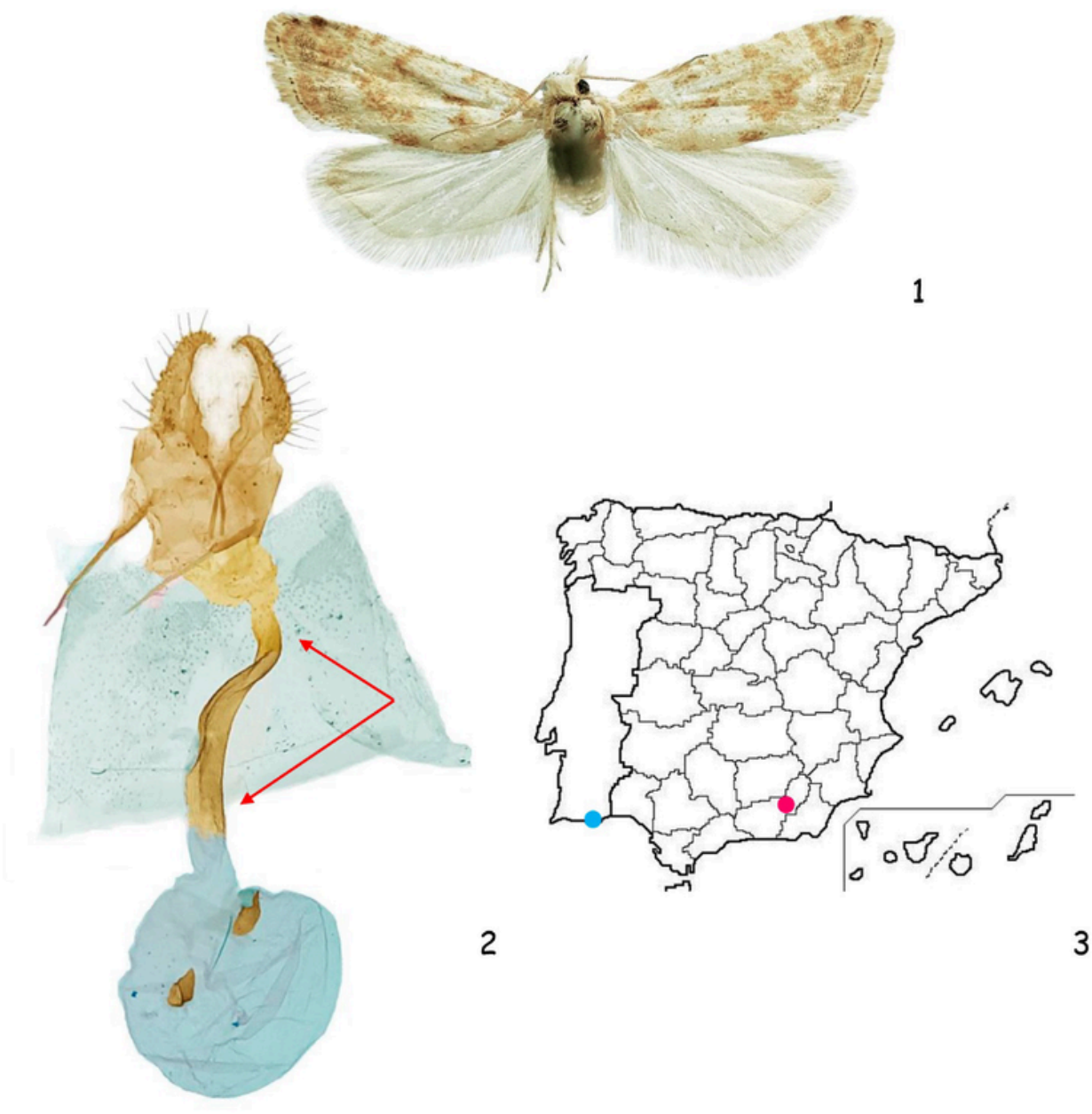


Fig. 1: Adulto de *Willibaldiana culatrae*, Fig. 2: genitalia ♀, Fig. 3: punto azul Isla de Cádiz (Algarve), punto rojo Cúllar (Granada).

***Merrifieldia menthae* (Chrétien, 1925) (Lepidoptera: Pterophoridae) in Portugal**

Martin Corley

Pucketty Farm Cottage, Faringdon, Oxfordshire, SN7 8JP, U.K. email: martin.corley@btinternet.com;
<https://orcid.org/0000-0003-4240-8007>

Introduction

I have been aware for some years that the name *Merrifieldia tridactyla* (Linnaeus, 1758) has been applied to more than one taxon in Portugal. Regrettably no great effort was made to collect specimens of this group. However, it was suspected that a large *Merrifieldia* associated with *Mentha suaveolens* Ehrh. in wet places was different from *M. tridactyla* from dry limestone sites. Although based on very limited material, the *Mentha*-associated species clearly belongs to *Merrifieldia menthae* (Chrétien, 1925). The moth from limestone sites in Estremadura, Ribatejo and Beira Litoral requires further investigation, but is apparently not *M. tridactyla*. Other records attributed to *M. tridactyla* cannot be assigned to that species with any certainty.

In *Merrifieldia* the valvae are asymmetrical, each bearing a sacculus of complex structure, usually with three sections, an inflated elongate basal section, a median section usually narrower and shorter than the basal section and a terminal usually slender section, often needle-pointed and especially on the left valva, curved. The median and terminal sections are often referred to as the saccular process (e.g. Skyva & Elsner, 2007). Bigot & Picard (1989) figured the genitalia of all nine French species of *Merrifieldia*, with *M. menthae* added by Nel (1991).

Abbreviations

ALG Algarve
BA Beira Alta
E Estremadura
Gen. prep. Genitalia preparation
leg. legit = collected by
MNHN Muséum National d'Histoire Naturelle, Paris
R Ribatejo
TM Trás-os-Montes

***Merrifieldia menthae* (Chrétien, 1925)**

Alucita baliodactyla menthae Chrétien, 1925: 243.

Male lectotype in MNHN: San Ildefonso, Segovia, Spain, 29.vi.1902, leg. P. Chrétien. Also recorded from mainland France (Alpes-Maritimes), Corsica and Morocco (Nel, 1991).

Material examined:

Caldas de Manteigas, Serra da Estrela, Beira Alta, 9.ix.2001, M. Corley P6326, gen. prep. 1667 male. Published as *M. tridactyla* in Corley et al. (2012) and Corley (2015).

Caldas de Manteigas, Serra da Estrela, Beira Alta, 17.vii.2007, M. Corley P8631, gen. prep. 2909 male.

Moimenta, Vinhais, Trás-os-Montes, 8.vii.2009, M. Corley, gen. prep. 3304 male, as *M. tridactyla* in Corley et al. (2012).

Vilarinho, E. of Parâmio, Bragança, Trás-os-Montes, 8.vii.2009, M. Corley P9140.

Methods. All specimens were netted in afternoon sunshine, resting on or flying over *Mentha suaveolens* Ehrh. Dissections were made following the methods detailed by Robinson (1976).

Description of Portuguese specimens

Wingspan 23–24 mm. (Fig. 1). Head pale lemon yellow; antennae light fuscous, not ringed. Thorax pale lemon yellow. Forewings creamy white, costa narrowly edged dark brown in middle, fringes very pale grey; hindwings light grey.

Male genitalia

Fig. 2. Middle section of left saccular process with a conspicuous thorn. The left saccular process has terminal section about 1.3 times length of middle section, while this ratio for the right saccular process is 1.0. The equivalent figures for *M. tridactyla* are approximately 0.7 and 0.65.

Female genitalia

No females have been collected in Portugal, but Nel et al. (2025) illustrate the female genitalia of *M. menthae*.



Fig. 1. *Merrifieldia menthae* (Chrétien, 1925). Vilarinho, E. of Parâmio, Bragança, Trás-os-Montes, Portugal, 8.vii.2009, leg. M. Corley, P9140.

Bionomics

In Portugal *M. menthae* has only been found resting on and flying around *Mentha suaveolens* Ehrh. in July and September. Nel (1991) mentions *Mentha sylvestris* L. and *M. rotundifolia* Sole as host-plants. Both are synonyms of *Mentha longifolia* (L.) L.



Fig. 2. *Merrifieldia menthae* (Chrétien, 1925) – male genitalia. Moimenta, Trás-os-Montes, 8.vii.2009, leg. M. Corley., gen. prep. 3304.

Discussion

Alucita baliodactla menthae was described by Pierre Chrétien from San Ildefonso (Segovia) in Spain.

The two 'standard' works covering European Pterophoridae, Arenberger (1995) and Gielis (1996) treat *menthae* as a synonym of *M. tridactyla*, but it should be noted that both authors were inclined to doubt the value of the numerous species described by French lepidopterists mainly in the 1980s and 1990s. These species were based on differences in host-plants together with small differences in larvae and in genitalia. Arenberger (1995) and Gielis (1996) do not always agree on which species should be recognised and which should be treated as synonyms. Vives (2014) also treats *menthae* as a synonym of *tridactyla*. Skyva & Elsner (2007) briefly

mention *M. menthae*, stating that it is currently considered to be a junior synonym of *M. tridactyla*, but they point out that it shows differences from *M. tridactyla* in external characters and in female genitalia.

More recently DNA barcodes have indicated that many of the French species (including all the *Merrifieldia* species) that were synonymised by both Gielis (1996) and Arenberger (1995), are valid species (Nel, pers. comm.). In Portugal barcode studies showed that records of *Merrifieldia malacodactylus* (Zeller, 1847) in Portugal actually refer to *Merrifieldia garrigae* Bigot & Picard, 1989 (Corley et al., 2024). In Nel et al. (2025) DNA barcoding revealed that a *Merrifieldia* from France is a different species from *M. tridactyla* and that this species also occurs in more or less wet places in Portugal, resulting in its description as *Merrifieldia thibaulti* Nel, Corley & Varenne, 2025. The saccular processes of *menthae* and of *thibaulti* were compared by Nel et al. (2025).

At present there is no DNA barcode available for *M. menthae*, but genitalia characters indicate that this is clearly different from *M. tridactyla*.

Arenberger (1995) considered *M. tridactyla* to be very variable, both in habitus and in male genitalia. He illustrated male genitalia from Germany and left and right valvae of a specimen from Albarracin, Spain and another from the Middle Atlas Mountains of Morocco. The illustrated differences between the saccular processes are such that these must represent three different taxa, but further research is needed to discover if these taxa already have names.

As a result of Arenberger's broad concept of *M. tridactyla*, Portuguese specimens were identified as that species, but can now be considered as definite or possible misidentifications. Some of these are here recognised as *M. menthae*, but unfortunately genitalia preparations have not been retained of all such specimens. Published records include those from Cabecico da Vinha, Freixiosa, Miranda do Douro, Trás-os-Montes (Corley et al., 2012) and from Valverde, Ribatejo (Corley et al., 2020). Both of these are females with retained genitalia preparations, but more work is needed to clarify the identity of females previously named as *M. tridactyla*.

To summarise, Portuguese records attributed to *M. tridactyla* may belong to *M. menthae* (recorded from BA and TM), *M. thibaulti* (recorded from BA and TM) or an as yet un-named species from limestone habitats in R, E and BL. The distributions of all three species may well be wider. Currently there is no certainty that *M. tridactyla* occurs in Portugal. Further collections of this complex are needed.

Acknowledgement

I am most grateful to Mike Dale for the photograph of the male genitalia and to Jacques Nel and João Nunes for helpful comments.

References

- Arenberger, E. 1995. *Pterophoridae I*. In Amsel, H.G. (Ed.) *Microlepidoptera Palaearctica* **9**: i-xxv, 1-258, Plates 1–153. G. Braun, Karlsruhe.
- Bigot, L. & Picard, J. 1989. Remarques sur les Pterophoridae français du genre *Merrifieldia*: *M. neli* et *M. garrigae*, espèces nouvelles. *Alexanor*, 1988, **15** (7): Suppl. 25–40.
- Chrétien, P. 1925. La légende de *Graëllsia isabellae*. Appendice – *L'Amateur de Papillons* **2** (16): 241–247.
- Corley, M.F.V. 2015. Lepidoptera of Continental Portugal. A fully revised list. 288 pp. Martin Corley, Faringdon.
- Corley, M.F.V., Cardoso, J.P., Dale, M.J., Marabuto, E., Maravalhas, E., Pires, P. 2012. New and interesting Portuguese Lepidoptera records from 2010. (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **40** (157): 5-21.
- Corley, M.F.V., Fabião, J.L., Lemos, P., Nunes, J., & Rosete, J. 2024. New and interesting Portuguese Lepidoptera records from 2023 (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **52** (208): 761–786. <https://doi.org/10.57065/shilap.1018>
- Corley, M.F.V., Nunes, J., Rosete J., Terry, R. & Ferrerira, S. 2020. New and interesting Portuguese Lepidoptera records from 2019 (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **48** (192): 609-641.
- Gielis, C. 1996. *Pterophoridae*. In Huemer, P., Karsholt, O. & L. Lyneborg |(Eds.). *Microlepidoptera of Europe* **1**: 1–222. Stenstrup (Apollo Books).
- Nel, J. 1991. Deuxième note sur les Ptérophores de la Corse. *Stenoptilia cyrnea* n. sp. et *Merrifieldia moulignieri* n. sp. (Lepidoptera Pterophoridae). *Alexanor* **17** (3): 167–182.
- Nel, J., Corley, M., Varenne, Th. & Ferreira, S. 2025. Description de *Merrifieldia thibaulti* sp. n., découverte en France et au Portugal (Lepidoptera, Pterophoridae). *Revue de l'Association Roussillonnaise d'Entomologie*, **118**: 247–251.
- Robinson, G.S., 1976. The preparation of slides of Lepidoptera genitalia with special reference to the microlepidoptera. *Entomologist's Gazette*, **27**: 127–132.
- Skyva, J. & Elsner, G. 2007. A new species of *Merrifieldia* Tutt from Slovenia (Pterophoridae). *Nota lepidopterologica*, **30**: 115–119.
- Vives Moreno, A., 2014. Catálogo sistemático y sinonímico de los Lepidoptera de la Península Ibérica, de Ceuta, de Melilla y de las Islas Azores, Baleares, Canarias, Madeira y Salvajes (Insecta: Lepidoptera.): 1184 pp. *Suplemento de SHILAP Revista de Lepidopterología*, Improitalia, Madrid.

A revision of Portuguese records of the genus *Psorosa* Zeller, 1846 (Pyralidae: Phycitinae)

Martin Corley

Pucketty Farm Cottage, Faringdon, Oxfordshire, SN7 8JP, U.K. email: martin.corley@btinternet.com;
<https://orcid.org/0000-0003-4240-8007>

Introduction

The Portuguese checklist (Corley, 2015a) was a synthesis of published information on the Portuguese Lepidoptera fauna together with the results of examination of recent and historical collections, all subject to critical evaluation. Some of the province records were unpublished, but many of these were later made available on GBIF (Corley & Afonso, 2021).

Knowledge of the Lepidoptera of southern Europe has always lagged behind that of more northern parts of the continent, so it was inevitable that a work of this sort would be found to contain errors. Most often these result from poor understanding of particular species or genera at the time of publication. Improved knowledge of many species comes from new papers clarifying the taxonomy of difficult groups or from the results of DNA barcoding, but also from collection of new specimens. When such new information becomes available it can present an opportunity for a revision updating knowledge of particular species or genera. In this paper the genus *Psorosa* Zeller, 1846 in Portugal is re-examined.

According to GlobIZ (2003–2026) the genus *Psorosa* has 26 known species, and occurs in Europe, Asia extending east to India and China and Africa, reaching South Africa. Very little is known of the bionomics of *Psorosa* species. One host-plant is recorded for *P. mediterranea*.

Corley (2015a) listed *Psorosa dahliella* (Treitschke, 1832) and *P. mediterranea* Amsel, 1953 for Portugal but excluded *P. nucleolella* (Möschler, 1866). In 2018 *P. ferrugatella* (Turati, 1924) was discovered in Algarve (Corley et al., 2019). Before 2018 *P. dahliella* was considered to be a species that could be distinguished without resorting to examination of genitalia. After the discovery of *P. ferrugatella*, this was no longer the case. As a result it became necessary to re-examine earlier records of *Psorosa*. Unfortunately several records of *P. dahliella* were based on specimens that were identified in the field but not collected.

Abbreviations

AAL Alto Alentejo
ALG Algarve
BA Beira Alta
BAL Baixo Alentejo
BB Beira Baixa
BL Beira Litoral
Coll. Collection of

DL Douro Litoral

GP genitalia preparation

Leg. Legit – collected by

NHMW Natural History Museum, Vienna, Austria

TM Trás-os-Montes

ZMCP Zoological Museum of the University of Coimbra, Portugal

***Psorosa mediterranea* Amsel, 1953 (Fig. 1)**

In Corley (2015a) this species was only recorded from Trás-os-Montes but subsequently was recorded from Beira Litoral, Beira Baixa, Beira Alta and Douro Litoral (Corley et al., 2019, 2020). Externally it does not closely resemble *P. dahliella* or *P. ferrugatella*. There are few permanent genitalia preparations of this species, but some records were confirmed by genitalia examination without the preparations being retained.

Records confirmed by genitalia examination: **DL**: Maragoutos, Lousada, 25.viii.2022, leg. J. Nunes and F. Gil (Nunes & Gil, 2023). **TM**: Arnal, Serra de Alvão, 2.ix.2002, leg. M. Corley (P6609), Corley GP 1772 male (Corley et al., 2006); Vilarinho das Azenhas, Tua Valley, 25.vii.2006, leg. M. Corley (P8189), Corley GP 6119 male (Corley et al., 2014). **M**: Fradelos, 3.ix.2020, leg. J. Nunes, Corley GP6015 male.

Other records accepted here, although without retained genitalia preparations: **BL**: Quinta do Canal, Bizarreiro, 14.ix.2019, leg. M. Corley, J. Rosete, J. Nunes and S. Ferreira (Corley et al., 2020). **BB**: Segura, 30.ix.2018, leg. M. Corley and S. Ferreira (Corley et al., 2019). **BA**: Algodres, Faia Brava, 15.ix.2018, leg. C. Silva, E. Jesus and J. Nunes. **TM**: Amieiro, Tua Valley, 28.viii.2013, leg. M. Corley (Corley et al., 2014); Guadramil, 20.ix.2019, leg. M. Corley, J. Nunes and S. Ferreira. This is the only species of the genus with known host-plant: *Polygonum equisetiforme* Sm. (Huertas Dionisio, 2007). According to Flora-on this plant is not present in some of the localities from which *P. mediterranea* is known. Other host-plants are therefore likely. Accepted records indicate that the species flies from July to September.



Fig. 1. *Psorosa mediterranea* Amsel, 1953. Arnal, Serra de Alvão, Trás-os-Montes, 2.ix.2002, leg. M. Corley.

***Psorosa dahliella* (Treitschke, 1832) (Fig. 2.)**

Monteiro & Carvalho (1984) recorded this species from Sapais de Castro Marim in Algarve. This probably refers to *P. ferrugatella* as there are confirmed records of that species from the locality. Corley (2015a) lists *P. dahliella* from Algarve, Alto Alentejo, Beira Alta and Trás-os-Montes, the last three provinces based on records published in Corley et al. (2015). More recently it was also recorded from Baixo Alentejo (Marabuto, 2018) and Douro Litoral (Corley et al., 2018), but this last record was later found to be a misidentification of *Pempelia genistella* (Duponchel, 1836) (Nunes, 2023).

Confirmed records: **ALG**: Algoz, 22.ix.1995, leg. M. Corley (P3404), Corley GP 6120 female (Corley et al., 2000). **AAL**: São Julião Igreja, Serra de São Mamede, 27.ix.1999, leg. M. Corley (P5433), Corley GP 1458 female (Corley et al., 2015b).

Records from BAL and BA are currently unconfirmed as also are Algarve records from Santa Catarina (Carvalho & Corley, 1995) and Sagres (Corley et al., 2000).

September is the only month with confirmed records.



Fig. 2. *Psorosa dahliella* (Treitschke, 1832). São Julião Igreja, Serra de São Mamede, Alto Alentejo, 27.ix.1999, leg. M. Corley.

***Psorosa ferrugatella* (Turati, 1924) (Fig. 3)**

This species was described from Libya. Leraut (2014) mentions it from Tunisia and figures female genitalia. This figure allowed the recognition of *P. ferrugatella* as a Portuguese species (Corley et al., 2019). Slamka (2019) mentions the new Portuguese record and also an old record from Spain, Granada, 14.vi.1914, leg. Knitschke, in coll. NHMW.

Confirmed records: **ALG**: Moncarapacho, 23.iv.1979, leg. A. Contente, in Passos de Carvalho coll., Corley GP JPC98-70; Praia Verde, 20.iv.1992, leg. M. Corley (two specimens, P1101, P1104) Corley GP 289 male, 290 male (Carvalho & Corley, 1995, as *P. dahliella*); Ludo, 6.v.1995, leg. M. Corley (P2838), Corley GP 709 female (Corley et al., 2000 as *P. dahliella*); Sapais de Castro Marim, 18.v.2003, leg. M. Corley, GP 1906 female; Sapais de Castro Marim, 2.x.2018, leg. M. Corley and

S. Ferreira, Corley GP 5514 female (Corley et al., 2019). **TM**: Amieiro, Tua Valley, 10.ix.2006, leg. M. Corley (P8299), Corley GP 6118 male (Corley et al., 2015b, as *P. dahliella*).

Records indicate that the species flies in April and May and again in September and October.

The presence of this mainly North African species in Trás-os-Montes is surprising, but the Tua Valley fauna (Corley, 2025) also includes other species with strongly southern distribution in the Iberian Peninsula, such as *Horisme scorteata* (Staudinger, 1901) and *Idaea nigrolineata* (Chrétien, 1911).



Fig. 3. *Psorosa ferrugatella* (Turati, 1924). Sapais de Castro Marim, 2.x.2018, leg. M. Corley and S. Ferreira.

Excluded species: *Psorosa nucleolella* (Möschler, 1866)

Mendes (1904) recorded *P. nucleolella* from Sao Fiel, Beira Baixa. An unlabelled specimen under this name in the National Collection (ZMCP) is presumably this specimen. It is *P. mediterranea* (Corley GP 3837 female) (Corley, 2015b).

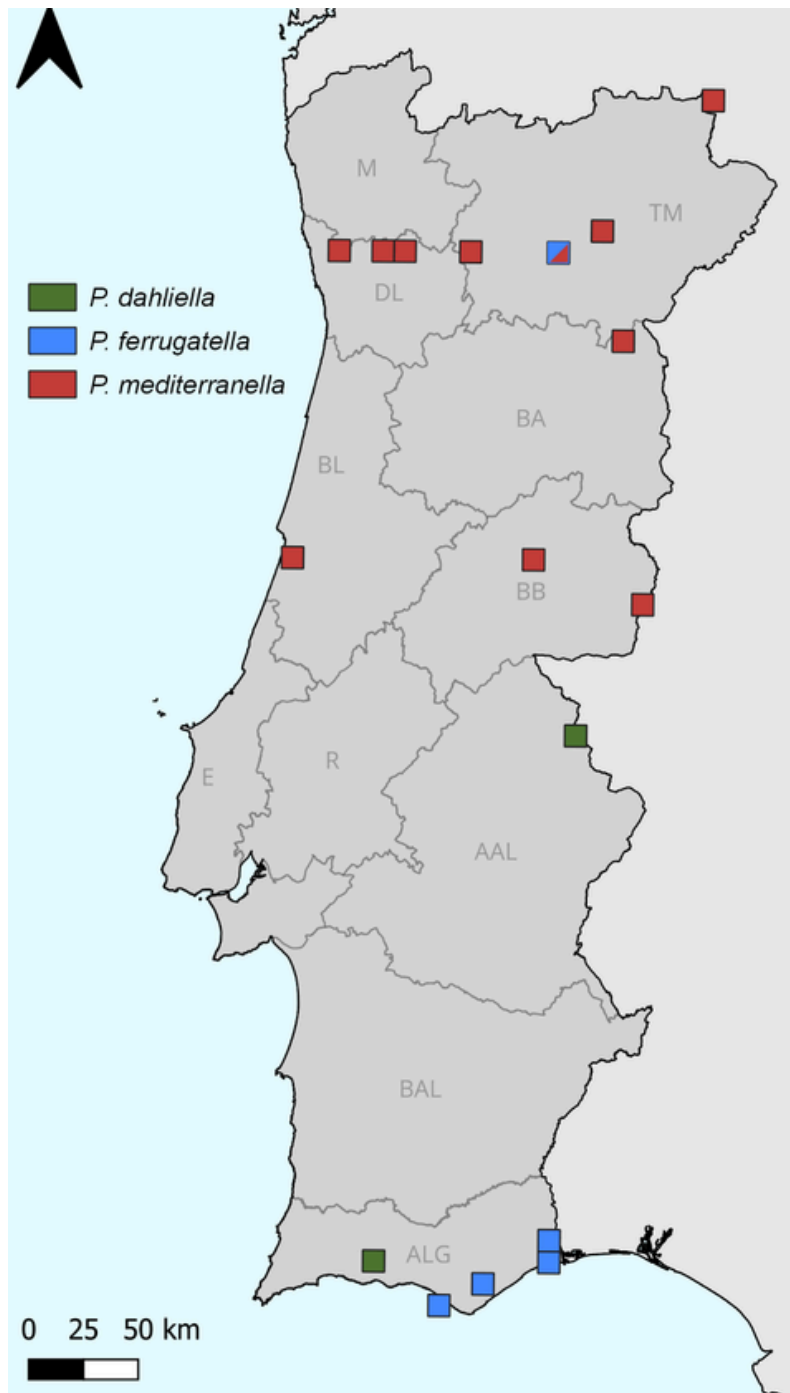


Fig. 4. The distribution of *Psorosa* species in Portugal.

Acknowledgement

I am most grateful to João Nunes for providing the map and for a thorough review of this paper.

References

- Carvalho, J. Passos de & Corley, M.F.V. 1995. Additions to the Lepidoptera of Algarve, Portugal. *SHILAP Revista de lepidopterología*, **23** (91), 191-230.
- Corley, M.F.V., 2015a. *Lepidoptera of Continental Portugal. A fully revised list*. 1–288 pp. Martin Corley, Faringdon.
- Corley, M.F.V., 2015b. The Lepidoptera collections of deceased Portuguese entomologists. *Entomologist's Gazette* **66**: 25–49.
- Corley, M. & Afonso, B. 2021. Portuguese Lepidoptera records compiled by Martin Corley's database. Version 1.5. CIBIO (Research Center in Biodiversity and Genetic Resources) Portugal. Occurrence dataset <https://doi.org/10.15468/ca4xt8> Accessed via GBIF.org on 20 February 2026.
- Corley, M.F.V., Gardiner, A.J., Cleere, N. & Wallis, P.D. 2000. Further additions to the Lepidoptera of Algarve, Portugal (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **28** (111), 245-319.
- Corley, M.F.V., Maravalhas, E. & Passos de Carvalho, J. (2006). Miscellaneous additions to the Lepidoptera of Portugal (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **34** (136): 407-427.
- Corley, M.F.V., Rosete, J., Romão, F., Dale, M.J., Marabuto, E., Maravalhas, E., & Pires, P., 2015. New and interesting Portuguese Lepidoptera records from 2014. (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología* **43** (172): 583–613.
- Corley, M.F.V., Ferreira, S., Grundy, D., Nunes, J., Pires, P. & Rosete, J. 2018. New and interesting Portuguese Lepidoptera records from 2017 (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **46** (184): 551–576.
- Corley, M.F.V., Nunes, J., Rosete, J. & Ferreira, S. 2019. New and interesting Portuguese Lepidoptera records from 2018 (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **47** (188): 611–630.
- Corley, M.F.V., Nunes, J., Rosete, J., Terry, R. & Ferrerira, S. 2020. New and interesting Portuguese Lepidoptera records from 2019 (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **48** (192): 609–641.
- Corley, M.F.V., Rosete, J., Marabuto, E., Maravalhas, E. & Pires, P., 2014. New and interesting Portuguese Lepidoptera records from 2013. (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **42** (168): 587-613.
- Corley, M. 2025. The Lepidoptera of the Tua Valley, Portugal 2006–2014. *Artimelia – Lepidoptera Ibérica*, **1** (1): 4–42.
- Flora-On. 2025. *Flora de Portugal Interativa*. Sociedade Portuguesa de Botânica. Available at: <https://flora-on.pt/>. Accessed 20 February 2026.
- GlobIZ, 2003-2026. Global Information System on Pyraloidea. www.globiz.pyraloidea.org Accessed 20 February 2026.
- Huertas Dionisio, M. 2007. Lepidópteros de los Espacios Naturales Protegidos del Litoral de Huelva (Micro y Macrolepidóptera). *Sociedad Andaluza de Entomología*. Monográfico 2. 251 pp.
- Leraut, P. 2014. *Moths of Europe* **4**. *Pyralids* 2. 1–441. NAP Editions, Verrières-le-Buisson.
- Marabuto, E. 2018. Butterfly and moth diversity in Serpa (Baixo Alentejo, Portugal): an advance in a yet poorly surveyed region (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **46** (183): 371–410.
- Mendes, C. de Azevedo, 1904. Lepidopteros de Portugal. II. Lepidopteros da região de S. Fiel (Beira Baixa). *Brotéria*, **3**: 223–254.
- Monteiro, T. & Carvalho, J. Passos de, 1984. Lepidópteros do Algarve – *Anais da Faculdade de Ciências de Porto*, **64**: 95-219.
- Nunes, J. 2023. As borboletas noturnas de Lousada – Inventário. *Lucanus – Revista de Ambiente e Sociedade*, **7**: 78-119.
- Nunes, J. & Gil, F. 2023. The moths of Lousada, Portugal (Insecta: Lepidoptera). *Boletín de la Sociedad Entomológica Aragonesa (S.E.A.)*, **72**: 71–80.
- Slamka, F., 2019. *Pyraloidea of Europe (Lepidoptera)*, **4**. *Phycitinae* – Part1. 432 pp. František Slamka, Bratislava.

***Opogona omoscopa* (Meyrick, 1893) (Lepidoptera, Tineidae), a new species for Cantabria and new records of *Borkhausenia crimnodes* Meyrick, 1912 (Lepidoptera, Oecophoridae)**

Francisco Eduardo Oliver Ruiz¹ & Daniel Oliver García²

¹ Ceballos 8, P.3, 3D Suances, Cantabria. España. email: eduardooliverruiz@gmail.com;
<https://orcid.org/0009-0007-3463-8319>

² Marqués de la Valdavia 13, P.A, 1C Collado Villalba, Madrid, España. email: danielpiracucu@gmail.com;
<https://orcid.org/0009-0008-6600-7367>

Abstract

This paper documents the presence in Cantabria of a new species: *Opogona omoscopa* (Meyrick, 1893), as well as *Borkhausenia crimnodes* Meyrick, 1912, a species previously cited in the region (Oliver Ruiz & Oliver García, 2025), providing new records that confirm its geographical expansion from the area from which it is presumed to have been initially introduced.

Key Words: Lepidoptera, Tineidae, Oecophoridae, alien species, dispersal, Cantabria, Spain.

Resumen

En el presente trabajo se documenta la presencia en Cantabria de una nueva especie: *Opogona omoscopa* (Meyrick, 1893), así como de *Borkhausenia crimnodes* Meyrick, 1912, especie previamente citada en la región (Oliver Ruiz & Oliver García, 2025), aportando nuevos registros que confirman su expansión geográfica desde el área donde se presume tuvo lugar su introducción inicial.

Palabras clave: Lepidoptera, Tineidae, Oecophoridae, especie alóctona, dispersión, Cantabria, España.

Introduction

In the current context of globalisation and increasing mobility of goods and people, the presence of alien species in various biotopes is an increasingly frequent phenomenon, resulting from both anthropogenic and natural biological dispersion processes. In this contribution to the knowledge of the Lepidoptera of Cantabria, two species with these characteristics are presented. The presence of both may be linked to human activity. *Opogona omoscopa* (Meyrick, 1893) is a species of uncertain origin (Bylli, 2009). It is speculated that the species may be native to the Atlantic island of Saint Helena and has spread throughout the world through human trade (EPPO, 2010; GBIF, 2024), with sightings from Australia and New Zealand, through the Hawaiian Islands, South Africa, Central and North America (Landcare Research, 2024) and several countries in Europe (Sterling et al., 2009). There are confirmed sightings in the Canary Islands (Gaedike & Falck, 2019), Madeira and the Azores (Gaedike & Karsholt, 2001) and in the Iberian Peninsula, sightings in Portugal (Corley et al., 2018), **in Catalonia (Requena et al., 2022)** and a vague mention in Gibraltar (EPPO, 2010). As for *Borkhausenia crimnodes* Meyrick, 1912, we already reported its presence in Cantabria in 2024 (Oliver Ruiz & Oliver García, 2024), and to test the theory that the species may have arrived in shipments of wood from the southern hemisphere

via the Sniace company (a cellulose factory near the town of Torrelavega), in 2025 we set up a series of light traps for sampling in different locations more or less close to its site.

Materials and methods

Two female *Opogona omoscopia* were obtained on 9.vii.2025 in Suances (UTM 30TVP1408). After genital dissection of both specimens identification was confirmed by Martin Corley. For *Borkhausenienia crimnodes*, sampling was carried out between 29.vi.2025 and 14.ix.2025. Light traps with different wavelength spectra were used. Most frequently, 400W mercury vapour lamps were used, sometimes supplemented with 125W black mercury lamps. Morphological, genital and distribution observations and capture data for *Opogona omoscopia* are included, as well as data confirming the expansion of *Borkhausenienia crimnodes* in the north of the Iberian Peninsula. When necessary, genital dissections were performed using the standard procedure of macerating abdomens in potassium hydroxide and subsequently preparing the genitalia between slides and coverslips (Yela, 1992). Microscopic and macro photographs were taken with a Canon 3500 camera, which were enhanced with photographic software and are stored together with the specimens in the authors' private collection. At the time of writing this report, it is assumed that all the literature relating to this species in Cantabria has been consulted. The geographical coordinates are given in the MGRS system, WGS84 datum.

Abbreviations

Fig: Images, graphs and maps represented in the text.

mm: millimetres

UTM: Universal Transverse Mercator

MGRS: Military Grid Reference System

KOH: Potassium hydroxide

Results and Discussion



Figure 1. *Opogona omoscopa*, female. Suances, Cantabria, Spain, 9.ix.2025.



Figure 2. *Opogona omoscopa*, enlarged detail of the wings. Suances, Cantabria, Spain, 9.ix.2025.



Figure 3. *Opogona omoscopia*, gynopygium. Suances, Cantabria, Spain, 9.vii.2025.



Figure 4. Anal papillae and apophyses of *Opogona omoscopia*. Suances, Cantabria, Spain. 9.ix.2025.

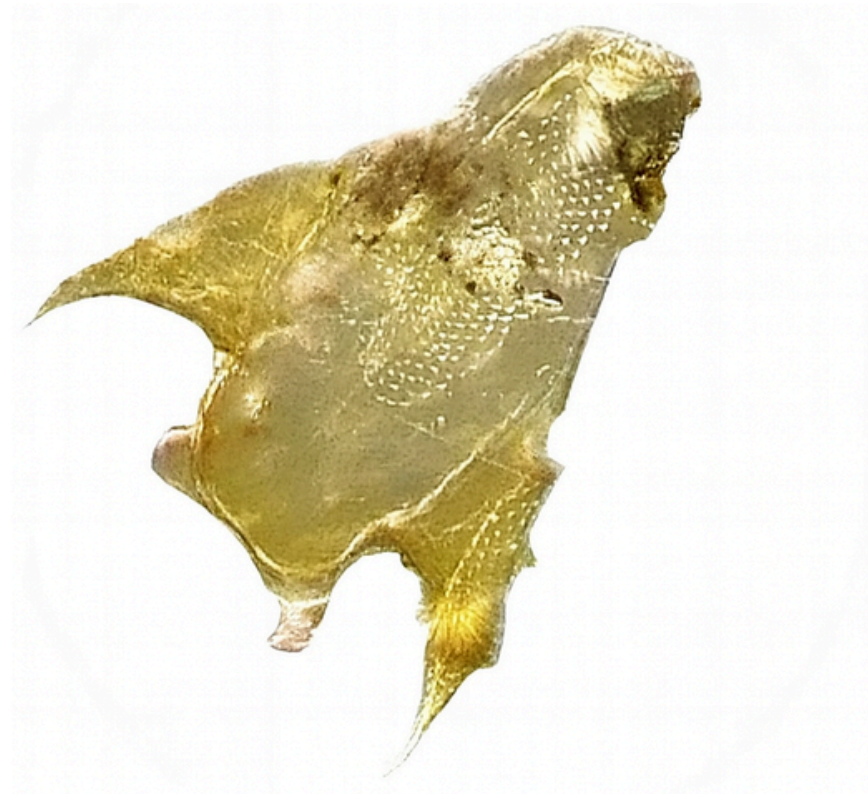


Figure 5. *Opogona omoscopia*, signum. Suances, Cantabria, Spain, 9.vii.2025.



Figure 6. *Borkhausenia crimnodes*. Cortiguera, Cantabria, Spain, 22. viii.2025.

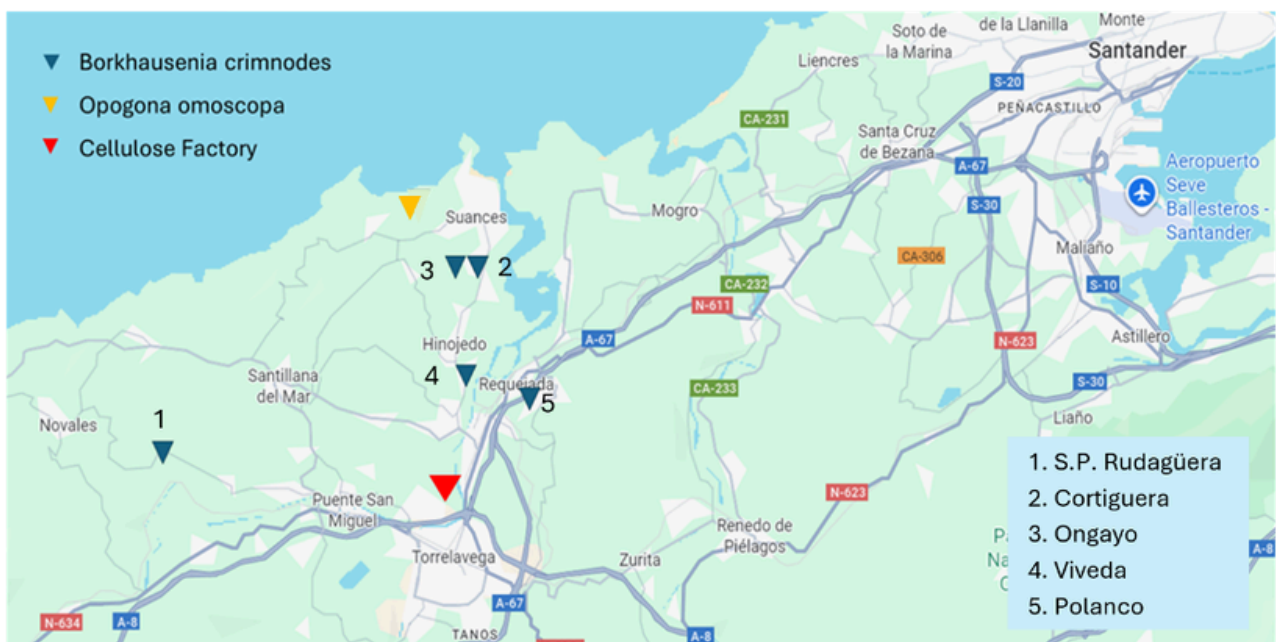


Figure 7. Distribution of *Opogona omoscopa* and *Borkhausenia crimnodes* in Cantabria, Spain.

Locations:

Opogona omoscopa:

Suances 30TVP1408: 9.ix.2025: 2 specimens.

Borkhausenia crimnodes:

1. San Pedro de Rudagüera, 30TVP0602: 29.vi.2025: 2 specimens; 16.vii.2025: 2 specimens.
2. Cortiguera, 30TVP1506: 10.vii.2025: 6 specimens; 16.vii.2025: 6 specimens; 29.vii.2025: 19 specimens; 12.viii.2025: 22 specimens; 22.viii.2025: 5 specimens; 2.ix.2025: 3 specimens.
3. Ongayo, 30TVP1406: 4.vii.2025: 4 specimens; 15.vii.2025: 6 specimens; 22.vii.2025: 13 specimens; 8.viii.2025: 16 specimens; 23. viii.2025: 8 specimens; 3.ix.2025: 2 specimens.
4. Viveda, 30TVP1404: 12.vii.2025: 1 specimen; 30.vii.2025: 11 specimens; 12.viii.2025: 17 specimens; 31.viii.2025: 6 specimens.
5. Polanco, 30TVP1803: 20.vii.2025: 15 specimens; 18.viii.2025: 9 specimens; 8.ix.2025: 1 specimen.

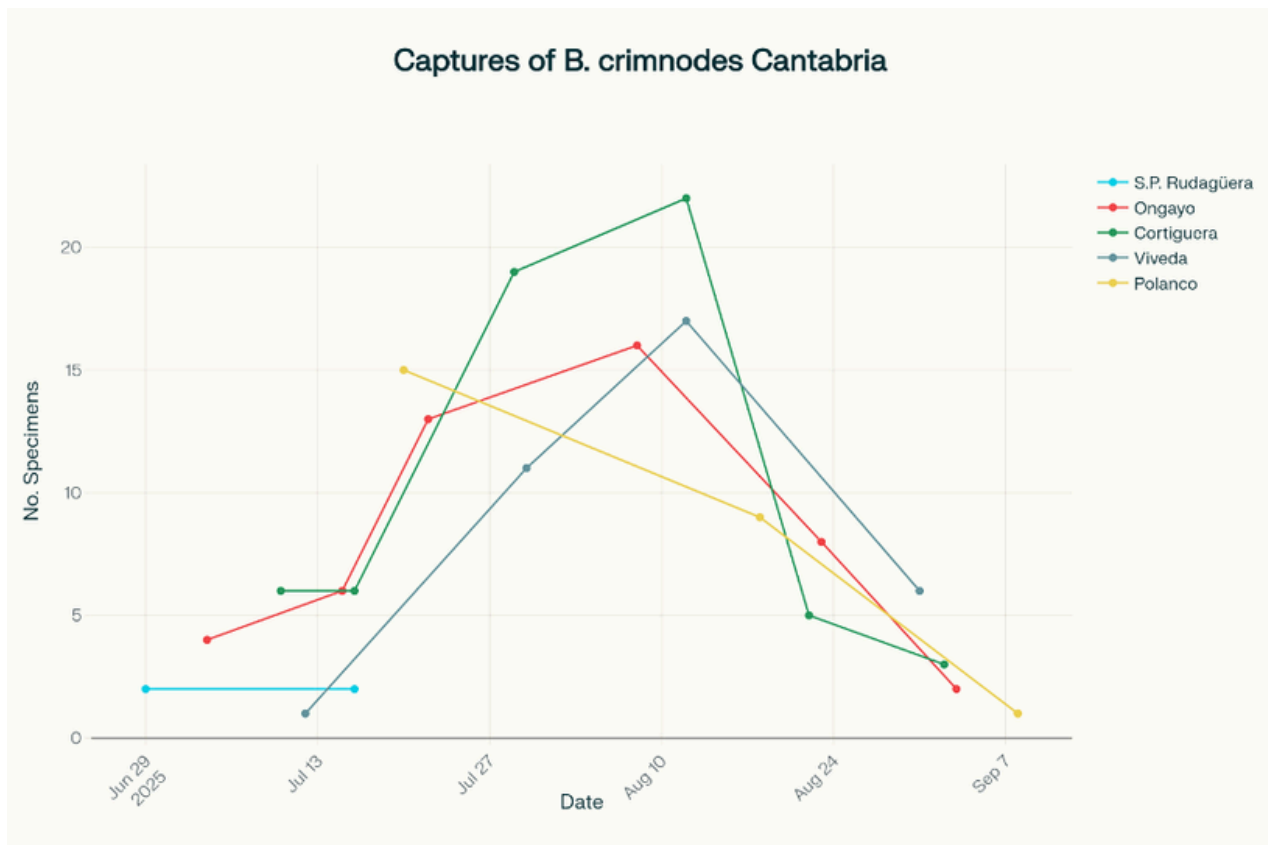


Figure 8. Density plot showing peak occurrence of *Borkhausenia crimnodes*.

1. *Opogona omoscopa* (Meyrick, 1893) (Fig: 1)

The caterpillars of this species feed on leaf litter and decaying fruit, especially on palm trees, and can perforate the crown of palms, but they are generally considered secondary pests attracted to decaying plant tissue (EPPO, 2010). Moore (1959) reported that the pupation period lasted 25 days during June in Lisarow, Australia. The time required from oviposition to adult emergence was 10 to 12 weeks during the warmer months in the same location, extending to

four months during the colder periods. Moore (1959) found no evidence of parasitism during his studies, but he did report that the larvae were infected by an entomopathogenic fungus of the *Rhizocybe vermicularis* type. In New Zealand, it is found in all types of natural and artificial biotopes (Landcare Research, 2024). The two specimens (females) from Suances have a wingspan of 16 mm and 19.5 mm respectively, and in New Zealand they are given an average size of between 7 and 11.5 mm (Landcare Research, 2024), which differs greatly from the size observed in the Cantabrian specimens. Lepiforum shows a photo of a specimen from the Azores with a wingspan of 20 mm, which is closer to the measurements taken on our specimens (Lepiforum e.V, 2006-2025). The wings are greyish with yellowish reflections under direct light; under indirect light, they show muted grey-brown tones. The forewings are elongated and relatively narrow with a slightly pointed apex, while the hindwings are narrower than the forewings, with a rounded apex and a slightly concave distal margin (Fig. 2). The most notable feature of its *gynopygium* (Fig. 3) is that the *signum* (Fig. 5) on the *corpus bursae* is irregular and occupies a large part of it, with a peculiar and distinctive shape. Other characteristics are that the *apophyses* are long and very thin (Fig. 4).

2. *Borkhausenia crimnodes* Meyrick, 1912 (Fig: 6)

The larval food source of *Borkhausenia crimnodes* is unknown, although it is probably dry plant matter or fungi (Corley et al., 2017). This type of larval behaviour facilitates its passive transport through human trade and subsequent establishment in altered habitats. In the previous study (Oliver Ruiz & Oliver García, 2024), we only had access to one male specimen with a wingspan of 13 mm, but now that we have more specimens, we have been able to take average measurements of between 9 and 13 mm. The most common size is 10 mm. As can be seen, both species have similar feeding habits.

Relationship between the two species

The detection of *Opogona omoscopia* in Cantabria extends its Iberian range to the north of the peninsula, confirming its expansion, probably through human activity. Its discovery in heavily modified coastal environments is consistent with previous European records (EPPO, 2010). The high density of *Borkhausenia crimnodes* specimens around the Sniace factory supports its local establishment, possibly linked to past importation of wood from the southern hemisphere, as probably occurred in Portugal (Corley et al., 2008). A total of 194 specimens were recorded in locations near the former Sniace factory, with captures reaching double figures at some sampling points. The phenology observed suggests an annual generation with a peak between late July and early August. It is more than likely that the species has been expanding for years and is now inhabiting many more locations in the area. A map showing the location of the species mentioned is provided (Fig. 7), as well as a graph explaining and visualising the above (Fig. 8). Both species may benefit from habitats altered by human activities, such as *Eucalyptus* plantations, abandoned sites, nurseries, etc.

Acknowledgements

To the Directorate-General for Biodiversity, Environment and Climate Change of the Government of Cantabria for granting permission to carry out sampling in 2025 and capture the material studied. To Martin Corley, for his invaluable help in identifying the specimens.

References

- Billi, F. 2009. *Opogona omoscopa* (Meyrick, 1893), parasite occasionnel des palmiers, espèce nouvelle pour la France (Lep. Tineidae). *Oreina*, **6**: 1–3.
- Corley, M.F.V., Marabuto, E., Maravalhas, E., Pires, P. & Cardoso, J.P. 2008. New and interesting Portuguese Lepidoptera records from 2007 (Insecta: Lepidoptera). *SHILAP Revista de Lepidopterología*, **36**(143): 283–300.
- Corley, M.F.V., Ferreira, S., Lvovsky, A.L., & Rosete, J. 2017. *Borkhausenia crimnodes* Meyrick, 1912 (Lepidoptera, Oecophoridae), a southern hemisphere species resident in Portugal. *Nota Lepidopterologica*, **40**(1): 15–24. DOI 10.3897/nl.40.10938.
- Corley, M.F.V., Rosete, J., Gonçalves, A.R., Mata, V., Nunes, J. & Pires, P. 2018. New and interesting Portuguese Lepidoptera records from 2016 (Insecta: Lepidoptera). *SHILAP Revista de Lepidopterología*, **46**(181): 33–56.
- EPPO 2010. First report of *Opogona omoscopa* in France. *European and Mediterranean Plant Protection Organization*. <https://gd.eppo.int>
- Gaedike, R. & Falck, P. 2019. Tineoidea (Meessiidae, Tineidae) y Glyphipterigidae: Acrolepiinae de las Islas Canarias, España (Insecta: Lepidoptera). *SHILAP Revista de Lepidopterología*, **47**(187): 507–517.
- Gaedike, R. & Karsholt, O. 2001. Contribution to the Lepidoptera fauna of the Madeira Islands. Part 2. Tineidae, Acrolepiidae, Epermeniidae. *Beitrage zur Entomologie*, **51**: 161–213.
- GBIF. 2024. *Opogona omoscopa*. *Global Biodiversity Information Facility*. <https://www.gbif.org>
- Landcare Research. 2024. Moths and Butterflies of New Zealand: *Opogona omoscopa* species file. <https://www.landcareresearch.co.nz>
- Lepiforum.V. 2006–2024. https://lepiforum.org/wiki/page/Opogona_omoscopa.14.xi.2025.
- Meyrick, E. 1893. Descriptions of Australian Micro-Lepidoptera. *Proceedings of the Linnean Society of New South Wales*, **7**: 435–487.
- Moore, K. M. 1959. Observations on Some Australian Forest Insects. *The Australian Zoologist*, **12**(4):337-350.
- Oliver Ruiz, F.E. & Oliver García, D. 2025. *Borkhausenia crimnodes* Meyrick, 1912 (Lepidoptera, Oecophoridae), a new species for Cantabria and Spain. *Artimelia – Lepidoptera Ibérica*, **12**: 45–52.
- Requena, E. & Pérez De-Gregorio, J. J. 2022. Noves dades sobre la fauna de microlepidòpters de Catalunya (Lepidoptera: Nepticulidae, Adelidae, Tischeriidae, Psychidae, Tineidae, Bucculatricidae, Gracillariidae, Ypsolophidae, Lyonetiidae, Douglasiidae, Autostichidae, Oecophoridae, Batrachedridae, Momphidae). *Butlletí de la Societat Catalana de Lepidopterologia*, **113**: 25–33.
- Yela, J.L., 1992. *Los Noctuidos (Lepidoptera) de la Alcarria (España Central), y su relación con las principales formaciones vegetales de porte arbóreo*. Ministerio de Agricultura, Pesca y Alimentación, Dirección General de Sanidad de la Producción Agraria, Subdirección General de Sanidad Vegetal. Madrid, 509 pp.

***Phyllonorycter berberidella* Nunes, Laštůvka & Laštůvka, 2026 (Gracillariidae) in Alto Alentejo**

Martin Corley

Pucketty Farm Cottage, Faringdon, Oxfordshire, SN7 8JP, U.K. email: martin.corley@btinternet.com;

<https://orcid.org/0000-0003-4240-8007>

Between 1995 and 2003 I visited Parque Natural da Serra de São Mamede a number of times. This was on the recommendation of José Passos de Carvalho. My previous entomological activities in Portugal had been confined to Algarve. He thought that I should extend my activities to other parts of the country. He proposed PN Serra de São Mamede because there were no Lepidoptera records from the Parque Natural. He informed me that Serra de São Mamede was a locality where some plants and animals had their most southerly Portuguese records, while others had their most northerly records. From a base at Escusa most of my work was in the northern half of the Parque, around Castelo de Vide and Marvão, south as far as the high point of São Mamede itself. We also stayed a few times at Hortas de Baixo at the southern extremity of the Parque. The area between São Mamede and Esperança appeared to me to have little interest and was neglected for the first few years. After a day spent with a local botanist, I became aware that there were localities with interesting plants in this area and that these could harbour interesting moths.

On 19.iv.2000 I found several plants which I was able to identify as *Genista berberidea* Lange. The site (39.214620N, 7.261614W) was about 1 km south of Besteiros, on moist slightly peaty soil on a slope between a track and a small stream at an altitude of 450 m. Aware that several *Genista* species are host to particular microlepidoptera, I examined these plants and found a single upperside leaf-mine of a *Phyllonorycter* species. I also netted a single female of a *Phyllonorycter*. On a later visit on 6.xi.2000 another leaf-mine was found, but no moth was reared from either of these leaf-mines. Further visits to the site on 1.ix.2001 and 20.v.2003 failed to produce any more mines or moths. On the 2003 visit, I had the impression that there were fewer plants of the *Genista*, perhaps because brambles (*Rubus*) were becoming more dominant at the site.

It was known at the time that there were several species of *Phyllonorycter* associated with various species of *Genista* and related genera in the tribe Genisteae of Fabaceae. Laštůvka & Laštůvka (2006) reviewed the European species of this group, describing 12 new species. None of these species was recorded from *Genista berberidea*, nor did the single female from Besteiros closely resemble any of the described species, suggesting that this was an undescribed species. As females in the group are not safely distinguishable by genitalia characters, it was hoped that further specimens would eventually be found.

On 24.iii.2017 Sónia Ferreira and I visited a site by Rio Ferreira at Ponte do Couce, Valongo, Douro Litoral. On a steep slope above a track east of the river there was a number of plants of *Genista berberidea* bordering a very small stream. A quick look at some of these plants did not reveal any leaf-mines. Subsequently I alerted João Nunes to the presence of these plants and the possibility that they could harbour an undescribed *Phyllonorycter*. As his home at Valongo was only a few kilometres from this site and because I knew that he had considerable expertise in rearing *Phyllonorycter* species, I hoped he would be able to find leaf-mines on the *Genista*. In the spring of 2024 he did indeed find leaf-mines and was able to rear adults, with further mines collected in autumn 2024 and spring 2025. This resulted in the description of *Phyllonorycter berberidella* Nunes, Laštůvka & Laštůvka, 2026 (Nunes et al., 2026).

After the description of the new *Phyllonorycter* I looked out my single specimen (numbered P5648) intending to compare the external appearance of the moth with illustrations in Nunes et al. (2026). Unfortunately there had been some insect damage to some of the specimens in the store box and the *Phyllonorycter* had been entirely consumed.

According to Flora-on and Castroviejo (1999) *Genista berberidea* is a rare and threatened species of the north-west of the Iberian Peninsula, designated Vulnerable on the Portuguese Red-list of Flowering Plants. The Flora-on distribution map shows 12 hectads (considered extinct in one) between Serra do Caramulo and Serra de Arga. The locality in PN Serra de São Mamede is therefore surprising, being much the furthest inland and about 160 km south-east of the Serra do Caramulo population. As it is far outside the known distribution area of the species, the legend on this species in Flora-on does not match the location of the Besteiros population. However, the legend describes what is known of the species and therefore could not cover what is not known. The flora of Serra de Sao Mamede is not particularly well known. From my own plant records there are many plant species in the area which are not shown on the distribution maps in Flora-on. As with *Genista berberidea*, some of these also show significant disjunction from other populations.

When I visited the site in 2003, I feared that *G. berberidea* was already in decline. There is a strong possibility that it is now extinct there. However there is also the possibility that there could be additional populations in the area, but any such discovery would require extensive field work.

References

- Flora-On: Flora de Portugal Interactiva. 2026. Sociedade Portuguesa de Botânica. www.flora-on.pt. Accessed 28 February 2026.
- Castroviejo, S. 1999. *Flora iberica. Plantas vasculares de la Península Ibérica e Islas Baleares. Vol. VII(I). Leguminosae (partim)*. Real Jardín Botánico, CSIC, Madrid. 578 pp.
- Laštůvka, A. & Laštůvka, Z. 2006. The European *Phyllonorycter* species feeding on the plants of the tribe Genisteeae (Fabaceae), with descriptions of twelve new species (Lepidoptera: Gracillariidae). *Acta universitatis agriculturae et silviculturae Mendelianae Brunensis*, **54** (5), 65–84. <https://doi.org/10.11118/actaun200654050065>
- Nunes, J., Laštůvka, A., Hansson, C. & Laštůvka, Z. 2026. A new species of *Phyllonorycter* Hübner (Lepidoptera: Gracillariidae) associated with *Genista berberidea* Lange (Fabaceae), and a new species of *Achrysocharoides* Girault (Hymenoptera: Eulophidae) associated with both, from Portugal. *Zootaxa*, **5760** (1): 139–150. <https://doi.org/10.11646/zootaxa.5760.1.7>

Depressaria discipunctella Herrich-Schäffer, 1854 in Portugal

Martin Corley

Pucketty Farm Cottage, Faringdon, Oxfordshire, SN7 8JP, U.K. email: martin.corley@btinternet.com;
<https://orcid.org/0000-0003-4240-8007>

In 1998 José Passos de Carvalho asked me to try to name part of his backlog of unidentified Portuguese Microlepidoptera. Among those I was able to name was a single *Depressaria* specimen which I identified as *Depressaria discipunctella* Herrich-Schäffer, 1854. It had been collected at his mother's house at São Romão, São Brás de Alportel in Algarve on 19.vi.1976. The record was published in Corley et al. (2000). This remains the only Portuguese record of this species.

In 2017 a new species *Depressaria junnilaineni* Buchner, 2017 was described from Greece and other Mediterranean countries, with many paratypes from Spain. The new species had been overlooked due to the relatively small differences in its genitalia from those of *D. discipunctella*. The presence of the new species in Spain cast doubt on the identity of the Portuguese specimen. This could not be checked, because the specimen and the genitalia preparation were in the Passos de Carvalho collection which was not then available for study.

In Buchner & Corley (2024) *D. discipunctella* was listed for Portugal in the Distribution Table in spite of the reservations about its identity.

The Passos de Carvalho collection is now in the Museu Nacional de História Natural e da Ciência, Universidade de Lisboa (MNHNC). The specimen and microscope slide (fig. 1) have the museum number MNHNCENT0076400.

The genitalia preparation (fig. 2) (Corley gen. prep. JPC 98-14) has now been re-examined and the original identification confirmed.

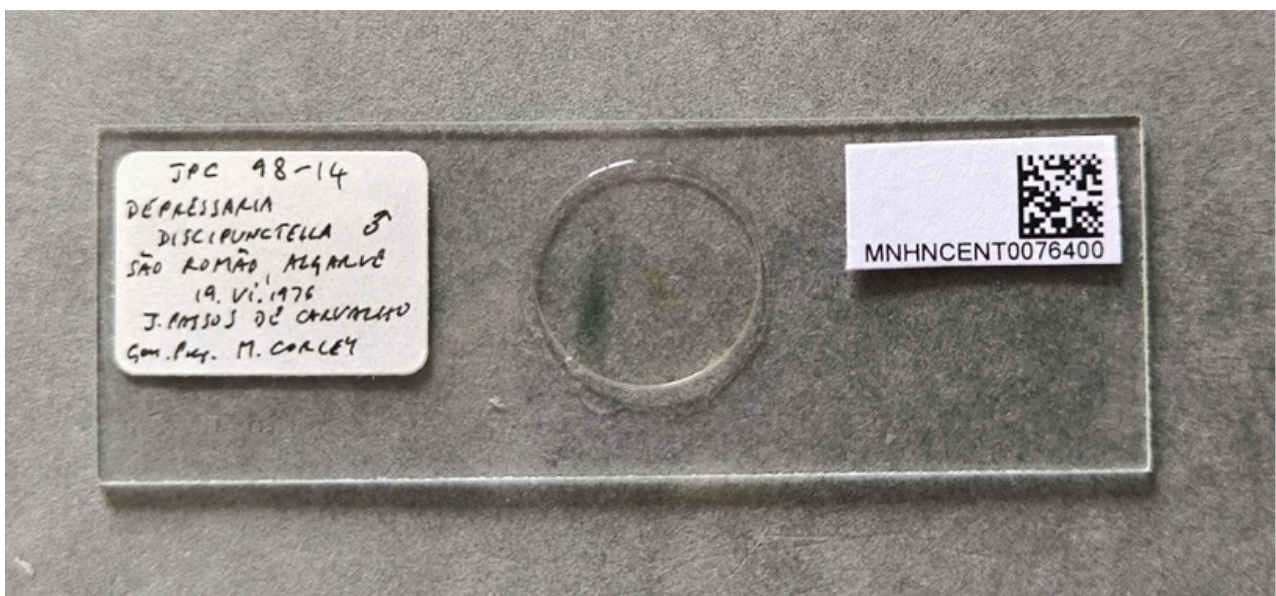


Fig. 1. M. Corley gen. prep. JPC 98-14.

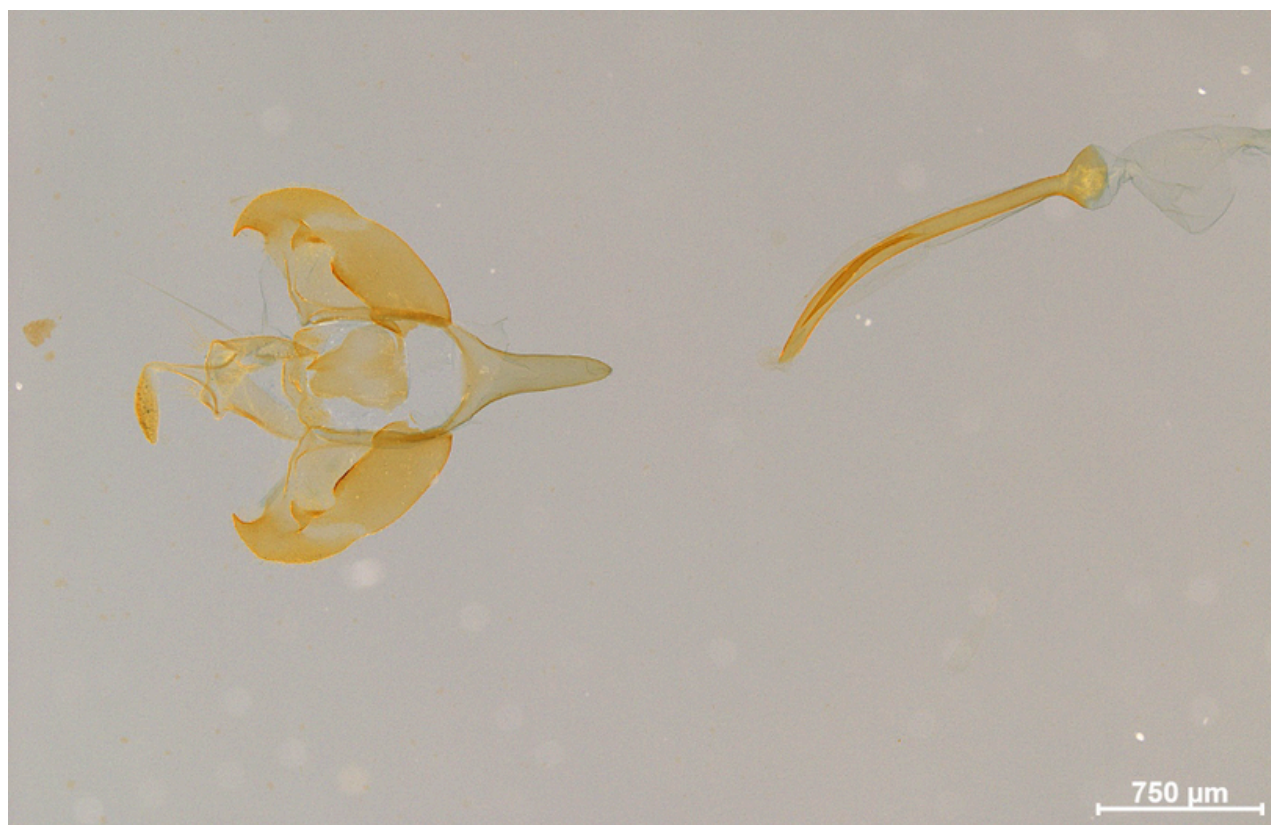


Fig. 2. *Depressaria discipunctella* Herrich-Schäffer, 1854. Male genitalia, São Romão, São Brás de Alportel, Algarve, 19.vi.1976, leg. J. Passos de Carvalho. M. Corley gen. prep. JPC 98-14.

Acknowledgement

I am most grateful to Roberto Keller for sending me the photo of the genitalia preparation and for permission to reproduce this here; my thanks also to Tatiana Moreira for facilitating this contact.

References

- Buchner, P. 2017. *Depressaria junnilaineni*, a new species from the *veneficella* species-group (Depressariidae, Lepidoptera) from the West Palaearctic, with additional information on the rare species *D. pentheri* and *D. erzurumella*. *Centre for Entomological Studies, Miscellaneous Papers*, **166**: 1–19.
- Buchner, P. & Corley, M. 2024. Depressariidae. In Karsholt, O. & Mutanen, M. (Eds.): *Microlepidoptera of Europe*, 10. Brill, Leiden. 605 pp.
- Corley, M.F.V., Gardiner, A.J., Cleere, N. & Wallis, P.D. 2000. Further additions to the Lepidoptera of Algarve, Portugal (Insecta: Lepidoptera). *SHILAP Revista de lepidopterología*, **28** (111): 245–319.

Guidance for authors

Articles should be submitted to the editor as Word files. They must be original work not published elsewhere.

Longer articles should be written in English, with a Portuguese or Spanish abstract. These papers will be peer-reviewed by one or more referees.

Short articles may be in English, Spanish or Portuguese, but English is preferred. They may be reviewed by a referee or not at the discretion of the editor.

Longer articles should have this format:

- Title
- Author(s)
- Address and e-mail address of author(s)
- Orcid number of author(s) (if available not obligatory).
- Abstract
- Key words
- Resumo or Resumen
- Key words
- Introduction
- Materials and Methods
- Abbreviations
- Results
- Discussion
- Acknowledgements
- References

Dates should appear in the format 6.vii.2024 (day.month.year).

The date of a record should be the actual date for daytime records, but records from overnight work should be the date on which the light was set, not the date when trap contents were examined.

References in the text should be given in the form (Vivaldi, 2014), or for two authors (Vivaldi & Beethoven, 2024), for more than two authors (Vivaldi *et al.*, 2024).

In the References papers should be cited in the form:

Vivaldi, A. 2024. A quartet of new moths from eastern Portugal. *Lepidoptera fantastica*, **25** (2): 100–105.

Vivaldi, A. & Beethoven, L. 2024. Moonlight moth records. *Lepidoptera fantastica*, **25** (3): 180–197.

Vivaldi, A., Liszt, F. & Boccherini, L. 2024. Evidence that moths respond to classical music. *Lepidoptera fantastica*, **25** (3): 200–206.

Books should be cited:

Corley, M.F.V. 2015. *Lepidoptera of Continental Portugal. A fully revised list*. 288 pp. Martin Corley, Faringdon.

Book that is part of a series:

Huemer, P. & Karsholt, O. 1999. Gelechiidae I (Gelechiinae: Teleiodini. Gelechiini). In P. Huemer, O. Karsholt & L. Lyneborg (eds): *Microlepidoptera of Europe* **3**: 1–356.

Internet sites:

BOLD. 2007–2026. The Barcode of Life Data System.
<http://www.v4.boldsystems.org/index.php>

Lepiforum e.V. 2006–2026.

https://lepiforum.org/wiki/page/Lymantria_dispar

If available doi numbers should be included at the end of the reference:

Zlatkov, B., & Huemer, P. 2023. *Eucosma subvittana* (Staudinger 1892) stat. rev., a Mediterranean species resurrected by DNA barcodes and morphology (Lepidoptera, Tortricidae), *Zootaxa*, **5361** (4): 451-462.
<https://doi.org/10.11646/zootaxa.5361.4.1>