Scanning Electron Microscopy of Male Terminalia and Its Application to Species Recognition and Phylogenetic Reconstruction in the Drosophila saltans Group

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Abstract

The Drosophila saltans group consists of five subgroups and 21 species, most of which have been identified only by morphological aspects of the male terminalia revealed by drawings using a camera lucida and a bright-field microscope. However, several species in the group, mainly those included in the saltans subgroup, are difficult to differentiate using only these characteristics. In this study, we used scanning electron microscopy (SEM) to analyze 19 structures of the male terminalia in 10 species from the five saltans subgroups. Among these structures, nine could be identified only through SEM analysis. We aimed to find other characteristics useful for morphological recognition of these species and to use these characteristics for phylogenetic reconstruction. These morphological differences enabled us to effectively distinguish among sibling species. These findings confirmed the monophyly of this group as previously determined in evolutionary studies based on other markers. The single most parsimonious tree (CI = 87 and RI = 90) indicated that the cordata subgroup is the most basal lineage and the saltans subgroup is the most apical lineage, as shown in earlier studies based on morphological data. However, our findings differed somewhat from these studies with respect to the phylogenetic relationships of species in the saltans group indicating that this group is still a puzzle that remains to be deciphered.

Introduction

The saltans group of Drosophila (Diptera, Drosophilidae) is divided into five subgroups on the basis of morphological features [1], [2]. Two of these subgroups (parasaltans and cordata) are found in the Neotropics; the remaining subgroups (elliptica, sturteranti and saltans) are found in both the Neotropics and the Nearctic regions [2].

In addition to these morphological features, several other characteristics of the group have been studied with a focus on the phylogenetic relationships of the species and subgroups. The markers used in these evolutionary studies include morphological characters [3], [4], chromosomal polymorphisms [5], esterase patterns [6], [7], [8], degree of reproductive isolation [5], [9], gene sequence variation [10], [11] and transposable elements [12], [13], [14], [15], [16].

Most of the phylogenetic relationships presented in these earlier studies differ from each other. Molecular analyses using the COL, COI, ITS1, Adh and Xdh genes were expected to be highly informative, but provided different topologies among subgroups and even among species [17]. However, they confirmed the monophyly of the group, which had previously been demonstrated using other markers [3–16].

Within the saltans group, sibling species have been differentiated almost exclusively via analysis of the male terminalia. This structure is considered one of the more variable and rapidly evolving morphological traits in several groups of organisms that exhibit internal fertilization [18], and it has been shown to have considerable taxonomic value for several groups of organisms [19]. The most informative studies on the morphology and taxonomy of the saltans group that are available in the literature were performed by Magalhães and Björnberg [1] and Magalhães [2], who differentiated the species into the five subgroups. In these studies, the structures of the male terminalia were represented in drawings obtained using a camera lucida. However, because the structures of the male terminalia of this group are complex, rich in detail, and similar among species from the same subgroup, identifying species by comparing newly collected specimens with the available terminalia diagrams remains difficult.

The inclusion of SEM in analyses of male terminalia has improved the recognition of different strains, species and subfamilies of several different species (e.g., the screwworms Cochliomyia hominivorax and C. macellaria [20] and the Aphididae family of insects [21]). This methodology has proved to be efficient in studies that integrate morphology and molecular data for
phylogenetic purposes, as in the case of the invasive drosophilid *Zaprionus indanus* [22].

In order to better resolve species diagnostics and phylogenetic relationships, we analysed nineteen male terminalia traits using SEM for species in the *saltans* group. In this paper, we also compared the results of our phylogenetic analysis to other published studies in order to evaluate the potential advantages of using aedeagus fine structure compared to other sources of data.

**Materials and Methods**

**Species and strains**

The species and strains examined in this study are listed in Table 1. Cultures were reared on a banana-agar-medium and housed in the biological laboratory of UNESP/IBILCE at **20°C±1°C**.

**Preparation of the terminalia for analysis**

We used the Kaneshiro technique with some modifications [23] to prepare the terminalia for SEM analysis. The total number of terminalia and aedeagi prepared was approximately 600, and at least 20 terminalia of each strain was analyzed. The prepared parts of the terminalia and aedeagi were placed in modified Karnovsky fixative (2.5% glutaraldehyde, 2.5% formaldehyde in 0.05 M sodium cacodylate buffer, pH 7.2, with 0.001 M CaCl₂) and were maintained at room temperature or in a refrigerator for at least 1 h. These aldehyde fixed samples were immersed in a solution of 1% osmium tetroxide in 0.05 M cacodylate buffer at pH 7.2 for 1 h at room temperature.

The samples fixed in OsO₄ were subsequently washed with distilled water and then treated with increasing concentrations of acetone (30, 50, 70, 90 and 100%) for approximately 10 minutes in each solution, and three washes were performed in the 100% solution. At the end of the dehydration procedure, certain samples were passed through the critical point dried (K550, EMITECH) solution. At the end of the dehydration procedure, certain samples showed the same quality as those were critical-point dried. Other samples were directly mounted to the SEM stub and sputter coated without critical-point drying; these with gold/palladium. Images of the samples were imaged and analyzed using a scanning electron microscope (LEO 435 VPi SEM, Zeiss).

**Phylogenetic analysis**

Nineteen diagnostic features of the terminalia and aedeagus were scored and used to form a data matrix for phylogenetic reconstruction of the *saltans* group (Table 2). All characters were treated as non-additive. A cladistic analysis using equal weights was undertaken with TNT software (Tree Analysis Using New Technology; [24]) including 1,000 iterations and using the following procedures: ratchet, drift, sectorial search, tree fusion and TBR max. Outgroup rooting [25] was implemented with *D. willistoni*. During the analysis, unambiguous optimization was chosen to describe the evolution of the assessed characters.

**Results**

The 19 characteristics of the male terminalia of species from the *saltans* group that were analyzed by SEM are described in Table 2 and illustrated in Figures 1 to 11. The terminology applied for features that had previously been described based on camera lucida drawings was maintained as reported in previous works [1], [2], [26], [27], [28], [29] with certain modifications. The description of the main characteristics of the terminalia and aedeagus analyzed in this study is presented in Table S1 (Supporting Information).

On the basis of the present analysis, we characterized the subgroups and species as follows: *saltans* subgroup: The epandrium (E) is an antero-inferior region that is highly angular, preceding the two recesses similar to curved gloved fingers (Figures 1 A; 2 A; 3 A; 4 A). The hypandrium (H) is elongated with two pairs of extensions, one of which is internal and the other external (Figures 1 A; 2 B, C; 3 B; 4 A, C). *D. prosaltans* (Figure 1): The surstyli (S) are located above the recesses with the form of gloved fingers and just below each side of the epandrium, with a long decasternum (D) between them and with primary and secondary teeth present; the anal plates (AP) exhibit a U-shaped contour (Figure 1A). The aedeagus (A) is characterized by ventral processes (VP) (Figure 1B) similar to curved gloved fingers, with an apical

| Subgroups | Species | Strains | Geographic Origin | Collectors/Identifiers | Date |
|-----------|---------|---------|------------------|------------------------|------|
| saltans   | *D. prosaltans* | ***PBB*** | Barra Bonita/SP/Brazil | Lopes, F.R | 1999 |
|           | *D. prosaltans* | ***PSR*** | Santana do Riacho/MG/Brazil | Vilela, C.R | 1995 |
|           | *D. lusaltans* | **B44 (14045-0891.00)** | Petionville/Haiti | Reed, W. | 1959 |
|           | *D. saltans* | **SAM (14045-0911.00)** | San José/Costa Rica | Wasserman, M. | 1956 |
|           | *D. austsaltans* | ***TUR*** | Turmalina/SP/Brazil | Penariol, L./Madi-Ravazzi, L. | 2008 |
| sturtevanti | *D. sturtevanti* | ***TAQ*** | Taquaritinga/SP/Brazil | Penariol, L./Madi-Ravazzi, L. | 2008 |
|           | *D. sturtevanti* | ***TUR*** | Turmalina/SP/Brazil | Penariol, L./Madi-Ravazzi, L. | 2008 |
|           | *D. dacunhai* | *JD (28.990.28.999)** | Kingston/Jamaica | Mourão; Bicudo | 2008 |
|           | *D. milleri* | **EY (14043-0861.00)** | El Yunque/Puerto Rico | William Reed | 1956 |
| parasaltans | *D. parasaltans* | ***B17-5** | Belém/PA/Brazil | Magalhães, E. | 1973 |
| cordata   | *D. neocordata* | ***CG*** | Campo Grande/MGS/Brazil | Magalhães, E. | 1973 |
| elliptica | *D. emarginata* | ***JD (14042-0841.09)** | Vera Cruz/México | Markow, Th. | 2005 |
| willistoni | *D. willistoni* | ***P9IDW*** | Pindorama/SP/Brazil | Penariol, L./Madi-Ravazzi, L. | 2008 |

* = indicate Department of Zoology of University of São Paulo from Brazil; ** = indicate UC San Diego Drosophila Stock Center ordering number; *** = UNESP/IBILCE/São José do Rio Preto/SP/Brazil.

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Table 1. Species and strains analysed in this study with geographic origins and collectors/identifiers and date of the collections.
Table 2. Character matrix based on the terminalia and aedeagus of the saltans group species, employed in the cladistic analysis.

| Characteristics/Species | D. willistoni | D. prosaltans | D. lusaltans | D. saltans | D. austrosaltans | D. sturtevanti | D. dacunhai | D. milleri | D. parasaltans | D. neocordata | D. emarginata |
|-------------------------|--------------|---------------|--------------|------------|-----------------|----------------|-------------|------------|----------------|--------------|--------------|
| 1. Fused Ventral Parameres | 0            | 0             | 0            | 0          | 0               | 1              | 1           | 1          | 0              | 0            | 1            |
| 2. Middle Ventral Process | 0            | 0             | 0            | 0          | 1               | 1              | 1           | 1          | 0              | 0            | 0            |
| 3. Scales in the Middle Ventral Process | 0            | 0             | 0            | 0          | 0               | 1              | 1           | 1          | 0              | 0            | 0            |
| 4. Punctiform projection in the Aedeagus Apex | 0            | 0             | 0            | 0          | 0               | 1              | 1           | 1          | 0              | 0            | 0            |
| 5. Ventral Parameres of Epandrium | 0            | 0             | 0            | 0          | 0               | 1              | 1           | 1          | 2              | 0            | 0            |
| 6. Concaved Surtylus | 0            | 1             | 1            | 1          | 1               | 0              | 0           | 0          | 0              | 1            | 1            |
| 7. Apical Crest of Aedeagus | 0            | 1             | 1            | 1          | 1               | 0              | 0           | 0          | 0              | 0            | 0            |
| 8. Apical Crest with Punctiform Projection | 0            | 0             | 0            | 0          | 1               | 0              | 0           | 0          | 0              | 0            | 0            |
| 9. Groove in the Apical Crest | 0            | 0             | 1            | 0          | 0               | 0              | 0           | 0          | 0              | 0            | 0            |
| 10. Bristles in the Aedeagus Apical Crest | 0            | 0             | 0            | 0          | 0               | 0              | 0           | 0          | 0              | 0            | 0            |
| 11. Scales in the Aedeagus Apical Crest | 0            | 1             | 1            | 0          | 1               | 0              | 0           | 0          | 0              | 0            | 0            |
| 12. Aedeagus Cape | 0            | 1             | 1            | 1          | 1               | 0              | 0           | 0          | 1              | 0            | 0            |
| 13. Serrated Edge of Aedeagus Cape | 0            | 1             | 1            | 2          | 1               | 0              | 0           | 0          | 1              | 0            | 0            |
| 14. Dorsal Cleft of Aedeagus | 0            | 0             | 0            | 0          | 0               | 0              | 0           | 0          | 0              | 0            | 0            |
| 15. Frontal Processes of Aedeagus | 0            | 1             | 0            | 1          | 1               | 1              | 1           | 1          | 1              | 0            | 1            |
| 16. Bipartite Aedeagus Apex | 0            | 0             | 0            | 0          | 0               | 0              | 0           | 0          | 0              | 0            | 1            |
| 17. Surtylus Processes | 0            | 0             | 0            | 0          | 0               | 0              | 0           | 0          | 0              | 1            | 0            |
| 18. Sickle-Shaped Processes of Aedeagus | 0            | 0             | 0            | 0          | 0               | 0              | 0           | 0          | 0              | 0            | 1            |
| 19. Long Apodeme | 0            | 1             | 1            | 1          | 1               | 0              | 0           | 0          | 1              | 0            | 0            |

Presence (1), presence with modifications (2) or absence (0). doi:10.1371/journal.pone.0097156.t002
region (AA) (Figure 1A) that is strongly sclerotized and contains a serrated crest covered with scale-like structures (Sc) (Figure 1B–C) that extends to the mid-dorsal region of the aedeagus in the form of a cape (AC) (Figure 1B–D), which is serrated and sclerotized at its edge and displays ventral parameres (VPA) (Figure 1C–D) and ventral prolongations (VPr) (Figure 1B). Additionally, it presents an apodeme (A) without pronounced curvature and triangular ends (Figure 1B). The phallotreme (Ph) is a fissure-like structure located at the apex of the aedeagus that can be observed only in SEM analysis (Figure 1D–E).

D. lusaltans (Figure 2): The surstyli exhibit morphology similar to a bean seed (Figure 2A, E). The aedeagus is characterized by the presence of an apical crest that is very similar to that of D. prosaltans; however, D. lusaltans displays an apical groove (AG) that isolates two lateral protuberances that end in two frontal processes (Figure 2C). The aedeagus apex has scales and an aedeagus body with a serrated edge (Figure 2D). The ventral processes are longer and thinner and cover the entire length of the ventral region of the aedeagus (Figure 2C). The apodeme is straight and long (Figure 2B–C). D. saltans (Figure 3): The surstyli exhibit morphology similar to a bean seed (Figure 3A–B); (Figure 3B). The aedeagus is characterized by a flattened apical region, with long ventral extensions derived from the insertion of the apodeme that extend through the length of the ventral region of the aedeagus, curving at its end. The ventral parameres of the aedeagus extend from the insertion of the apodeme to the dorsal medial region of the aedeagus (Figure 3C–E). The cape of the aedeagus has a strongly serrated edge that can be observed in detail via SEM (Figure 3B, D, F). The apex of the aedeagus exhibits a punctiform projection in the posterior portion consisting of numerous long setae (Figure 3E, arrowheads). D. austrosaltans (Figure 4): The surstyli show morphology similar to a bean seed (Figure 4A). The hypandrium differs from that of other species of the saltans subgroup species, which have scales on the aedeagus apex, this species displays setae in this region (Figure 3B, E, F). The apex of the aedeagus exhibits a punctiform projection in the posterior portion consisting of numerous long setae (Figure 3E, arrowheads).

Figure 1. Scanning electron micrographs of the terminalia of D. prosaltans (PSR strain). (A) AA = aedeagus apex, AP = anal plate, C = cercus, E = epandrium, H = hypandrium, LP = lateral paramere, S = surstylus, (189 × magnification); (B) AA = aedeagus apex, A = apodeme, AC = aedeagus cape, Sc = scales, VPA = ventral paramere of the aedeagus, VPr = ventral processes (395 × magnification); (C) AA = aedeagus apex, AC = aedeagus cape, (253 × magnification); (D) AC = aedeagus cape, Ph = phallotreme, (376 × magnification); (E) Ph = phallotreme, Sc = scales, (1610 × magnification). doi:10.1371/journal.pone.0097156.g001
The apex of the aedeagus is covered with scales, as observed in *D. parasaltans* and *D. lusaltans* (Figure 4B, D). *sturtevanti* subgroup: The epandrium displays ventral parameres and anal plates forming a U-shaped contour (Figure 5A, 6B, 7A). *D. sturtevanti* (Figure 5): The surstyli are large and concave and are joined by a small decasternum; primary and secondary teeth (T) are present (Figure 5A). The aedeagus is characterized by the presence of fused ventral parameres, where the middle ventral process (MVP) exhibits a pointed apical region and has the appearance of a duck’s beak. The cape of the aedeagus is absent, and the apodeme is short and sinuous (Figure 5C–D). Structures consisting of similar scales (Sc) near the apex of the aedeagus could be visualized only in the SEM analysis and there were fewer of them than observed in *D. dacunhai* and *D. milleri*. No difference in the morphology of the terminalia or aedeagus was observed among the analyzed strains of *D. sturtevanti*. *D. dacunhai* (Figure 6): The surstyli are large and concave and are joined by a small decasternum (Figure 6A–B). The aedeagus is characterized by the presence of fused ventral parameres with the appearance of a duck’s beak (Figure 6C–D); scales are present in the middle ventral process (Figure 6C–D), which is wider and more curved than that of *D. sturtevanti* and presents pointed scales and a groove that has not been described previously (Figure 6C, arrowhead). *D. milleri* (Figure 7): The surstyli are large and concave (Figure 7A). The aedeagus is characterized by fused ventral parameres that combine with the middle ventral process, resulting in a duck-beak-like appearance (Figure 7B), and presents a groove in the upper portion of the middle ventral process (Figure 7C, arrow head), which is covered by scales that are close to the surface of the process (Figure 7B–C). *parasaltans* subgroup: *D. parasaltans* (Figure 8): The epandrium displays ventral parameres similar to those of the *sturtevanti*...
subgroup and exhibits a cercus with many bristles, mainly in the region just above the surstyli (Figure 8A). The surstyli are concave, with a very different morphology; they are characterized by the presence of four to five teeth on the inside edge and six to seven teeth on the outer edge (Figure 8B). The aedeagus has a dorsal part formed by a single structure without recesses and displays extensions projecting laterally that contain spiniform structures. The apodeme is long and straight, not sinuous (Figure 8B), and the ventral parameres are short and thin with two spiniform projections on the apical regions. The cape of the aedeagus presents a serrated edge along its entire dorsal length (Figure 8C).

**cordata** subgroup: *D. neocordata* (Figure 9): The epandrium does not have ventral processes but exhibits long and short bristles in the basal region. The antero-inferior region of the epandrium is situated at nearly a right angle and presents anal plates forming a U-shaped contour (Figure 9A-B). The surstyli are concave formed by cuticular plates that are inserted into five to six primary teeth and six long bristles; large processes are present (SPr) along with protruding structures with the appearance of gloved fingers, which are diagnostic characteristics of the species (Figure 9A-B). The apodeme of the aedeagus is short and thick (Figure 9A). The frontal processes (FrP) at the bottom of the apex are similar to claws (Figure 9C-E). The ventral parameres of the aedeagus are long and thin and bifurcate in the middle region, giving rise to cuticular prolongations that extend to the ventral region of the body of the aedeagus and end in the sickle-shaped processes (SSP). The surstylus processes and the sickle-shaped processes are characteristic of this species (Figure 9A-D). *elliptica* subgroup: *D. emarginata* (Figure 10): The hypandrium is elongated and thin (Figure 10A). The epandrium has two pairs of cuticular hooks on the inside edge of its angular lower region (Figure 10A). The aedeagus is extremely large compared with the other analyzed species of the *saltans* group and appears sickle-like. The central axis of the aedeagus has a slit in its dorsal region (FD) and ends in a hook (BH) that branches into two pointed ends (Figs. 10B-C). Its apodeme is wide and short (Figure 10B), and its lateral processes are large and long and follow almost the entire length of the central axis (Figure 10B, arrow heads). The fused ventral paramere is similar to that of the *sturtevanti* subgroup species, and the dorsal slit is covered by plates with serrated edges, which is another feature that was detailed only in the SEM analysis (Figure 10B-C). *D. willistoni* (outgroup) (Figure 11): The hypandrium is short and wide with a quadrangular morphology (Figure 11A, B). The surstyli are concave and are formed by two
Figure 5. Scanning electron micrographs of *D. sturtevanti* terminalia (TUR strain). (A) C = cercus, E = epandrium, VP = ventral paramere, S = surstylus (400 × magnification); (B) AA = aedeagus apex, VP = ventral paramere (747 × magnification); (C) AA = aedeagus apex, MVPr = middle ventral process, (817 × magnification); (D) MVPr = middle ventral process, VPA = ventral paramere of the aedeagus, Sc = scales, (578 × magnification). doi:10.1371/journal.pone.0097156.g005

Figure 6. Scanning electron micrographs of *D. dacunhai* terminalia. (A) S = surstylus, VP = ventral paramere, E = epandrium, (256 × magnification); (B) AA = aedeagus apex, C = cercus, E = epandrium, S = surstylus, (350 × magnification); (C) AA = aedeagus apex, E = epandrium, groove on the MVPr (arrow heads), (420 × magnification); (D) MVPr = middle ventral process, Sc = scales, (582 × magnification). doi:10.1371/journal.pone.0097156.g006
parts, with a larger apical part that contains primary teeth and a smaller part that is fused to the larger part and contains secondary teeth (Figure 11B–D). The dorsal portion of the aedeagus is formed by a plate that extends on both its sides (Figure 11C, arrowhead). In the dorsal portion of this plate, there is a tubular projection that extends along the aedeagus body and ends in two frontal projections (Figure 11 C, F). The ventral parameres of the aedeagus are short and thick, and the apodeme is short and thin (Figure 11 E).

Among the described features, nine could be observed only through SEM analysis: 1. the scales on the middle ventral process in *D. dacunhai* and *D. milleri* (Figure 6C–D); 2. the punctiform projection on the aedeagus apex in the three species of the *sturtevanti* subgroup (Figure 5C–D, Figure 6C–D); 3. the apical crest with the punctiform projection in *D. austrosaltans* (Figure 4B); 4. the groove on the apical crest in *D. lusaltans* (Figure 2C–D); 5. the bristles on the apical crest of the aedeagus in *D. saltans* (Figure 3E–F); 6. the scales on the apical crest of the aedeagus in *D. prosaltans, D. lusaltans* and *D. austrosaltans* (Figure 1C–E; 2D; Figure 4B, D); 7. the serrated edge of the aedeagus cape in all species of the *saltans* subgroup and *D. parasaltans* (parsaltans subgroup) (Figure 1C–D; Figure 2D; Figure 3B, C, D, F; Figure 4D–E; Figure 8C); 8. the dorsal cleft of the aedeagus; and 9. the bipartite aedeagus apex in *D. emarginata* (Figure 10C) (table 2).

Some characteristics were shared by different subgroups, whereas others were subgroup-specific or species-specific. Five features were specific to the *saltans* subgroup (the apical crest of the aedeagus, the apical crest with a punctiform projection, the groove on the apical crest, the bristles on the aedeagus apical crest and the scales on the aedeagus apical crest); two were specific to the *sturtevanti* subgroup (the middle ventral process, the scales on the middle ventral process and the punctiform projection on the aedeagus apex magnification); two were specific to the species *D. neocordata*, representing the *cordata* subgroup (the surstylus processes and sickle-shaped processes of the aedeagus); and two were specific to the species *D. emarginata*, representing the *elliptica* subgroup (the dorsal cleft of the aedeagus and the bipartite aedeagus apex).
Drosophila parasaltans, representing the parasaltans subgroup, did not show subgroup-specific or species-specific features (Table 2).

Species-specific features were observed in D. lusaltans (the groove on the apical crest), D. saltans (the bristles on the aedeagus apical crest), D. neocordata (the surstylus processes and sickle-shaped processes of the aedeagus) and D. emarginata (the dorsal cleft of the aedeagus and the bipartite aedeagus apex) (Table 2).

Subgroup-specific features that are shared among species were found in the saltans subgroup (the aedeagus cape, serrated edge of the aedeagus cape, frontal processes of the aedeagus and long apodeme) and the sturtevanti subgroup (the fused ventral parameres, ventral parameres of the epandrium and concave surstyli), which were the only subgroups for which more than one species was available for analysis (Table 2).

Features shared between subgroups were observed in the D. sturtevanti subgroup and D. parasaltans (the ventral parameres of the epandrium and concave surstyli); the D. sturtevanti subgroup and D. emarginata (fused ventral parameres); the saltans subgroup and D. parasaltans (the aedeagus cape, serrated edge of the aedeagus cape and long apodeme); and the D. saltans subgroup and D. neocordata (the frontal processes of the aedeagus; Table 2).

The nineteen characters evaluated in detail via SEM were used to construct a matrix for phylogenetic analysis of the species of the saltans group, which is shown in Table 2. Cladistic analysis using only the data obtained in this study recovered a single most parsimonious tree with a length of 25 steps (CI = 87; RI = 90) and recovered the monophyly of the saltans group. The consensus tree indicated that the cordata subgroup is the most basal lineage and the saltans subgroup is the most apical lineage (Figure 12).

Discussion

The application of SEM to the study of male terminalia in the Drosophila saltans group provided tools for improving the recognition of species and for developing a phylogenetic tree based on new morphological markers. Nine of the 19 structures selected for analysis were described for the first time in this study, whereas the remaining structures were assessed by detailing known structures that were previously analyzed via light microscope.

Some of our findings distinguished sibling species in the saltans and in the sturtevanti subgroups more precisely. In the saltans subgroup, a species-specific structure allowed differentiation of D. saltans from D. prosaltans, D. austrosaltans and D. lusaltans.
Figure 9. Scanning electron micrographs of *D. neocordata* terminalia. (A) A = apodeme, C = cercus, FrP = frontal process, SPr = surstylus processes, SSP = sickle-shaped processes, H = hypandrium, VPA = ventral paramere of the aedeagus, (161 \times \text{magnification}); (B) S = surstylus, SPr = surstylus processes, (578 \times \text{magnification}); (C) FrP = frontal process, H = hypandrium, (248 \times \text{magnification}); (D) FrP = frontal process, SSP = sickle-shaped processes, (318 \times \text{magnification}); (E) C = cercus, E = epandrium, H = hypandrium, epandrium bristles (arrow heads), (202 \times \text{magnification}).

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Figure 10. Scanning electron micrographs of *D. emarginata* terminalia. (A). C = cercus, E = epandrium, S = surstylus, T = primary and secondary teeth, H = hypandrium, cuticular hook of the epandrium (arrow heads), (141 \times \text{magnification}); (B) AA = aedeagus apex, VPA = ventral paramere of the aedeagus, lateral processes of the aedeagus (arrow heads), (130 \times \text{magnification}); (C) AA = aedeagus apex, BH = bipartite hook, DC = dorsal cleft, SP = serrated plates, (309 \times \text{magnification}).

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lusaltans and D. austrosaltans also had species-specific characters. In the sturtevanti subgroup, by the same criterion, D. sturtevanti was differentiated from D. dacunhai and D. milleri. D. parasaltans, D. neocordata and D. emarginata, the unique available species representing the subgroups parasaltans, cordata and elliptica, respectively, presented characters that allowed differentiation among them and from the species in the subgroups saltans and sturtevanti.

It is interesting to compare the morphological data on the aedeagus with data on the reproductive isolation of the same species [29], [30], [31]. In the present phylogenetic analysis, D. saltans differed from D. lusaltans, D. austrosaltans and D. prosaltans. However, data on reproductive isolation [29] indicated that D. austrosaltans is completely reproductively isolated from these species at the premating level. In contrast, D. prosaltans, D. saltans and D. lusaltans are not completely reproductively isolated from each other, as D. lusaltans can produce fertile F1 offspring with D. prosaltans and with D. saltans. Additionally, D. prosaltans can be intercrossed with D. saltans yielding fertile F1 females and males in one direction and fertile females and sterile males in the other direction. The sturtevanti subgroup, whose species show great similarity with regard to the male terminalia produced a low number of hybrid progeny in the intercrosses of D. milleri with D. sturtevanti or D. magalhaesi, generating a phylogenetic sequence in which D. magalhaesi was the most basal species, followed, in sequence, by D. milleri, D. sturtevanti and D. dacunhai [31]. In the tree proposed in the present paper, D. milleri and D. dacunhai were considered sister taxa. These incongruent results are compatible with the finding that premating mechanisms (probably sexual isolation) are predominantly involved in the reproductive isolation of species in the saltans group [29], [31].

In general, phylogenetic relationships within the saltans group based on different markers have been found to be incongruent, but monophyly of the group has always been recovered [4], [7], [17]. The same occurred in the present study. However, comparison of the evolutionary relationships based on SEM analysis with previous data revealed high consistency with the sequential order of the subgroups obtained using morphological and biochemical data [2][3]. According to the results of these studies, the cordata
and elliptica subgroups are the most basal, followed by startecendi and parasaltans, with the saltans subgroup being the most derived.

Unlike, as observed regarding the reproductive isolation already mentioned, comparison of the present phylogenetic data with other studies using different markers indicated only partial concordance. For example, one molecular phylogenetic study using the genes that code for ADH, ITS1, COI and COII [10] produced phylogenetic relationships that were predominantly incongruent relative to those of Magalhães [1], Throckmorton and Magalhães [2] and the present SEM data. O’Grady et al. [10] suggested that unresolved branching patterns were due to the relatively recent divergence of these species and conflicting information from each locus. In both O’Grady et al. [10] and the present study, Drosophila parasaltans and D. australis were considered to be closer to each other than to D. saltans.

When our cladogram was compared with that of Yassin [4], which was based on morphological markers, the main difference observed was related to the basal group, which was the cordata clade in the present work and the startecendi subgroup in Yassin’s study. In both studies, the parasaltans and saltans subgroups appeared as sister clades. Yassin [4] suggested that the discrepancies in molecular phylogenies in the saltans group may have resulted from a characteristic shift of codon bias in Sophophorans from the New World. Yassin [32] found that different morphological characteristics provide varying signals at different phylogenetic scales. Therefore, molecular and morphological data must be employed to better understand the taxonomy and phylogeny of these organisms.

The analysis of datasets based on phylogenetic relationships determined using different markers for species in the saltans group suggests that increased taxon sampling and identification of more informative markers are necessary to better resolve the phylogeny of this group. The present results indicate that SEM characteristics may be useful in the species recognizing and useful as a marker for attempting to clarify the evolutionary history of the group.

Supporting Information

Table S1 Description of the structures of the terminalia and the aedeagus analysed by SEM (Scanning Electron Microscopy).

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Author Contributions

Conceived and designed the experiments: LMR TAJ5. Analyzed the data: TAJ5 LMR FBN HEMCB. Contributed reagents/materials/analysis tools: LMR HEMCB. Wrote the paper: TAJ5 LMR HEMCB.

References

1. Magalhães LE, Bjornberg AJ (1957) Estudo da genitália masculina de Drosophila, Drosophilidae, Diptera. Braz J Biol 16: 273–280.
2. Magalhães LE (1962) Notes on the taxonomy, morphology and distribution of saltans group of Drosophila, with description of four new species. UT Publications 6205: 135–154.
3. Throckmorton LH, Magalhães LE (1962) Changes with evolution of pteridine accumulations in species of the saltans group of genus Drosophila. UT Publication 6205: 489–503.
4. Yassin A (2009) Phylogenetic relationships among species subgroups in the saltans saltans group (Diptera: Drosophilidae): can morphology solve a molecular conflict? Zool Res 30: 225–232.
5. Bicudo HEMC (1973a) Reproductive isolation in the saltans group of Drosophila. Genetica 44: 313–329.
6. Nascimento AP, Bicudo HEMC (2002) Esterase patterns and phylogenetic relationships of Drosophila species in the saltans subgroup (saltans group). Genetica 114: 41–51.
7. Nascimento AP, Bicudo HEMC (2006) Further study on the esterase patterns of sibling species in the Drosophila saltans subgroup (saltans group): intraspecific and interspecific variations in the development. Genetica 126: 263–276.
8. Bernardo AA, Bicudo HEMC (2009) Variability of esterase patterns in adult flies of the saltans species group of Drosophila (subgenus Sophophora). Genetica 137: 111–124.
9. Bicudo HEMC (1973b) Chromosomal polymorphism in the saltans group of Drosophila. Genetica 44: 520–552.
10. O’Grady PM, Clark JB, Kidwell MG (1998) Phylogeny of the Drosophila saltans species group based on combined analysis of nuclear and mitochondrial DNA sequences. Mol Biol Evol 15: 656–664.
11. Rodriguez-Trelles F, Tarrio R, Ayala FJ (1999) Molecular evolution and phylogeny of the Drosophila saltans species group inferred from the XdH gene. Mol Phylogenet Evol 13: 110–121.
12. Clark JB, Altheide TK, Schllosser MJ, Kidwell MG (1995) Molecular evolution of P transposable elements in genus Drosophila. I. The saltans and allistius species groups. Mol Biol Evol 12: 902–913.
13. Clark JB, Kidwell MG (1997) Phylogenetic perspective on P transposable elements evolution in Drosophila. Proc Nail Acad Sci USA 94: 11428–11433.
14. Silva CJ, Kidwell MG (2008) Horizontal transfer and selection in the evolution of P elements. Mol Biol Evol 17: 1542–1557.
15. Castro JP, Carareto CMA (2004a) P elements in the saltans group of Drosophila: a new evaluation of their distribution and number of genomic insertion sites. J Mol Evol 52: 383–387.
16. Castro JP, Carareto CMA (2004b) Canonical P elements are transcriptionally active in the saltans group of Drosophila. J Mol Evol 59: 31–40.
17. O’Grady PM, Kidwell MG (2002) Phylogeny of the Subgenus Sophophora (Diptera: Drosophilidae) Based on Combined Analysis of Nuclear and Mitochondrial Sequences. Mol Phylogenet Evol 22: 442–453.
18. Evans JP, Gasparini C, Hollow GI, Rammarine I, Pitcher TE, et al. (2011) Intraspecific evidence from guppies for correlated patterns of male and female genital trait diversification. Proc R Soc B 278: 2611–2620.
19. Jamjoom MB, Shalaby IM (2006) The Contribution of Electron Microscopic Studies to the Taxonomy and Biology of Parasitic Trematodes. World J Zool 2: 64–81.
20. Leite ACR (1995) Ultrastructure of the male terminalia of Coeliochroma hominivorax and C. macellaria. Mem Inst Oswaldo Cruz 90: 473–481.
21. Wiczerek K, Plachno BJ, Sweitack PA (2012) Comparative morphology of male genitalia of Aphididae (Insecta, Hemiptera) part 2. Zoo morphology 131: 303–324.
22. Yassin A, Capi P, Madi-Ravazzi L, Ogereau D, David JR (2008) DNA barcode discovers two cryptic species and two geographic radiations in the invasive drosophilid, Zaprionus indistinctus. Mol Ecol Resour 8:491–501.
23. Torres FR, Madi-Ravazzi L (2006) On the use of open or closed traps in the capture of drosophilids. Biota Neotropica 8: 47–53.
24. Globoff PA, Farris JS, Nixon K (2003) TNT: Tree analysis using New Technology. Version 1.0, version Beta test v. 0.2. Program and documentation. Available: http://www.zmuc.dk/public/phylip/tnt/. Accessed 26 December 2013.
25. Nixon KC, Carpenter JM (1993) On outgroups. Cladistics 9: 415–426.
26. Mourão CA, Bicudo HEMC (1967) Duas novas espécies de Drosophila do grupo saltans (Drosophilidae, Diptera). Pap Avulsos Zool 20: 123–134.
27. Vilela CR (1983) A revision of the Drosophila repleta (Drosophilidae, Diptera) species group (Diptera, Drosophilidae). Rev Bras Entomol 27:1–114.
28. Grimaldi DA (1990) A phylogenetic revised classification of genera in Drosophila (Diptera: Drosophilidae). Bull Am Mus Nat Hist 197: 1–139.
29. Bacll G, Vilela CR, Escher SA, Saura A (2004) The Drosophilidae (Diptera) of Fennoscandia and Denmark. Fauna Entomologica Scadinavica 39: 1–362.
30. Bicudo HEMC (1975) Reproductive isolation in the saltans group of Drosophila. I. The saltans subgroup. Genetica 44: 313–329.
31. Bicudo HEMC (1976) Reproductive isolation in Drosophila parasaltans. Revista Brasileira de Genétic 1: 11–28.
32. Bicudo HEMC (1979) Reproductive isolation in the saltans group of Drosophila. IV The startecendi subgroup. Braz J Genet 2: 247–250.