Morphological characterization of trichomes show enormous variation in shape, density, and dimensions across the leaves of 14 Solanum species

Sakshi Watts¹ and Rupesh Kariyat¹,²

¹Department of Biology, University of Texas Rio Grande Valley, Edinburg, TX, USA
²School of Earth Environmental and Marine Sciences, University of Texas Rio Grande Valley, Edinburg, TX, USA

Author of Correspondence: Rupesh.kariyat@utrgv.edu
Abstract:

Trichomes are the epidermal appendages commonly observed on plant surfaces including leaves, stem, and fruits. Plant trichomes have been well studied as a structural plant defense designed to protect plants against abiotic and biotic stressors such as UV rays, temperature extremities and herbivores. Trichomes are primarily classified into glandular and non-glandular trichomes, based on the presence or absence of a glandular head. The plant genus *Solanum* is the largest genus of family Solanaceae that houses ~25,000 species of ecological and economic importance have a diverse set of trichomes that vary in density and morphology. However, due to the incomplete and contradictory classification system, trichomes have subjective names and have been largely limited to be grouped into glandular or non-glandular types. Through this study, we did a complete workup to classify and characterize trichomes on both adaxial and abaxial leaf surface of 14 wild and domesticated species of the genus *Solanum*. Using electron microscopy, statistical analyses and artistic rendition, we examined finer details of trichomes and measured their density and dimensions to compile a detailed dataset which can be of use for estimating the variation in trichome types, and their density, with consequences for understanding their functional roles. Our study is the first of its kind that provides us with a better and well-defined classification, density, and dimension analysis to complete the morphological classification of trichomes on both leaf surfaces of a diverse range of members in *Solanum* genus.
Introduction:

Plant surfaces show spectacular variation in the shape, size, location, function, and origin of epidermal projections (Werker, 2000). The most important and well-studied among these are trichomes: unicellular or multicellular appendages (hair-like structures) originating from epidermal cells of various plant parts including leaves, stems and flowers (Oksanen, 2018), and developing outwards (Werker, 2000). Trichomes are distributed almost universally in the plant kingdom and exhibit dramatic variation in their morphology (Seithe and Sullivan, 1990; Dam et al., 1999; Adedeji et al., 2007; Kang et al., 2010; Nurit-Silva and Fatima Agra, 2011; Munien et al., 2015; Talebi et al., 2018) and density (Talebi et al., 2018), both intra (Kang et al., 2010; Munien et al., 2015; Dam et al., 1999) and inter-specifically (Seithe and Sullivan, 1990; Navarro and Oualidi, 1999; Mannethody and Purayidathkandy, 2018; Talebi et al., 2018; Yu et al., 2018), and also among and between related and distant plant families (Kariyat et al., 2018; Deore, 2020; Watts and Kariyat 2021). For example, Munien et al. (2015) found four types of trichomes (glandular, non-glandular dendritic, non-glandular bicellular and non-glandular multicellular) in Withania somnifera (intra-specific); Yu et al. (2018) found a great variation in trichome morphology, dimensions, distribution and density among seven Mentha species (inter-specific); Deore (2020) identified variations in trichomes among 20 species belonging to 12 different plant families and most of the species were reported to have trichomes ranging from unicellular to multicellular, conical to elongated, smooth to grooved, thin to thick walled, and with or without a flat disc at the base. Further, Tsujii et al. (2016) found tremendous variation in trichome leaf dry mass per area in the plant tissue of Metrosideros polymorpha at different elevations (Tsujii et al., 2016). Clearly, within flowering plants, trichomes are both ubiquitous and morphologically diverse.
Trichomes, in general are considered as one of the first line of defenses possessed by plants to protect against abiotic stresses such as UV rays, water loss, temperature extremities (Ehleringer, 1982; Li et al., 2018; Oksanen, 2018) and herbivore damage (Kaur and Kariyat, 2020a; 2020b; Watts and Kariyat, 2021). Moreover, the leaf trichomes can also act as mechanoreceptors for detection of insects on leaf surface as observed in case of Arabidopsis (Zhou et al., 2017). In addition to defense-related functions, trichomes play a role in water usage strategies through maintenance of leaf water content and stomatal traits to name a few. For example, Pan et al. (2021) showed higher percentage increase of leaf water content in epiphytic plants with trichomes (Pan et al., 2021). Additionally, trichomes can play role in translocation and homeostasis of minerals in plants (Blamey et al., 1986; 2015; Li et al., 2021). But the relationships of such traits with trichome types and measurements such as density and dimensions are poorly understood.

Broadly, trichomes can be classified into glandular (presence of glandular head) and non-glandular (absence of glandular head) (Werker, 2000). Both glandular (Tang et al., 2020) and non-glandular (Karabourniotis et al., 2019) trichomes have been well documented to protect plants either by production of chemicals in their glandular heads or by their sturdy structure that assist plants to adapt and/or protect from environmental conditions such as UV radiations and cold stresses. Glandular trichomes deter herbivory by physically entrapping herbivore into sticky exudates (Tingey and Laubengayer, 1981; Neal et al., 1990; Elle et al., 1999; Zalucki et al., 2002), secreting defensive chemical compounds such as proteinase inhibitors (Peiffer et al., 2009), production of volatile organic compounds (Avé et al., 1987; Murungi et al., 2016), or by altering herbivore body odor after providing a sugar rich first meal (Weinhold and Baldwin, 2011). On the other hand, non-glandular trichomes in the Solanum species are mostly spike like structures which deter herbivory primarily by deterring herbivore movement, feeding and oviposition (Corsi & Bottega, 1999; Kennedy, 2003; Løe et
al., 2007; Dalin et al., 2008; Sletvold et al., 2010; Tian et al, 2012; Weigend et al., 2018; Kariyat et al, 2019). Additionally, non-glandular trichomes can cause post-feeding damage to caterpillars by rupturing of caterpillars’ peritrophic membrane (gut lining; Kariyat et al., 2017; Andama et al., 2020). Moreover, some plant species also possess stinging hairs (trichomes with stinging cells which contain irritant fluids) which act as hypodermal syringes and can cause various allergic reactions such as pain, itching, oedema, and visible dermal reactions to mammalian herbivores (Ensikat et al., 2021).

Glandular trichomes are more pliable, so may not cause physical damage (Kariyat 2017; 2019) but can be toxic and can release chemicals to intoxicate herbivores (Hare, 2005) and attract predators of herbivores in association with carcasses of herbivores (LoPresti et al., 2015). Contrary to this, non-glandular trichomes are usually devoid of toxins but their sharp and edgy structure can cause physical damage to herbivores. Thus, while the size and structure of the apical gland can inform about the amount of toxins and their content, knowledge about various types (e.g., unbranched vs stellate) and sizes of non-glandular trichomes can appraise us about their functional significance - either in deterring herbivory or protecting against abiotic stressors (Karabourniotis et al., 2020). More specifically, in case of non-glandular trichomes, we now have multiple lines of evidence on how variation in trichome type and density differentially defend them against insect herbivores (Cho et al., 2017; Kariyat et al., 2017; 2019; Watts and Kariyat, 2021). For instance, Watts and Kariyat (2021) found significant variation in trichome density on abaxial and adaxial leaf surface of 11 Solanaceae species and demonstrated that this variation has functional consequences for caterpillar growth and feeding.

Clearly, a detailed examination and characterization of trichome morphology (even on abaxial and adaxial surfaces) can have multiple benefits, including a reliable and non-contradictory nomenclature of trichomes, understanding the prominent trichome types found
in nature, and their diversity - which later can be explored for defensive functions against different herbivores, and abiotic stressors, with possible implications for our efforts to produce better defended plants for sustainable agriculture (Andama et al., 2020).

Solanaceae is one of the most important plant families consisting of 90 genera and ~25,000 ecologically and economically important species which are found in all habitats ranging from dry deserts to wet tropical rainforest and have growth habits ranging from small ephemeral herbs to large perennial trees (Knapp, 2002). Among all the genera in Solanaceae, the genus Solanum contributes ~75% of all species (Knapp, 2002; Symon, 1981). Solanum genus exhibits tremendous variation and diversity of trichomes; for instance, widely studied and domesticated crops such as tomato (Solanum lycopersicum) and tobacco (Nicotiana tabacum) have different types of glandular trichomes, while wild weeds such as silverleaf nightshade (Solanum elaeagnifolium) and Carolina horsenettle (Solanum carolinense) possess only non-glandular trichomes (Peiffer et al., 2009; Weinhold and Baldwin, 2011; Burrows et al., 2013; Kariyat et al., 2018). A general convention in trichome literature is to reduce the diversity of trichome types by constraining them to just types of glandular and non-glandular, while these types are quite diverse and are often complicated to resolve. Moreover, this basic classification also fails to explain the huge variation among sub-types of glandular and non-glandular trichomes, and consequently potential to explore their function.

Luckwill (1943) was the first to classify trichomes of Lycopersicon into 7 distinct types (4 glandular sub types and 3 non-glandular sub types) of trichomes based on their length, number of stalk and base cells, and the presence or absence of gland. Following Luckwill (1943), Uphof (1962) in a summary for different methods of classification of trichomes concluded that the final classification is still subjective. After that, Payne (1978) provided us with a glossary to name various trichomes or structures found in trichomes to improve trichome nomenclature. Later, Chanrayappa et al. (1992) revised the trichome classification

Downloaded from https://academic.oup.com/aobpla/advance-article/doi/10.1093/aobpla/plab071/6417072 by guest on 05 November 2021
by Luckwill (1943) and the revised classification is used frequently in trichome related studies. While these studies have served as a model for trichome morphology assessment, we used scanning electron microscopy, and artistic rendering along with the previous classical classification systems (Roe, 1971) and the glossary provided by Payne (1978) to characterize and classify trichomes on both adaxial (upperside) and abaxial (lowerside) leaf surface of 14 representative species from Solanum.

Materials and Methods

**Plant Materials:**
A mixture of wild and domesticated species of genus Solanum (14 species in total) were included in the study. We bought seeds of forest bitterberry (*Solanum anguivi*; Product code: Y5SSSOIN), porcupine tomato (*S. pyracanthos*; Product code: Y5SSSOPY), African eggplant (*S. macrocarpon*; Product code: Y5SSSOMC), bittersweet nightshade (*S. dulcamara*; Product code: Y5SSS0DU), lance-leaved nightshade (*S. lanceifolium*; Product code: Y5SSSOLA), potato tree (*S. grandiflorum*; Product code: Y5SSSOGR), tzimbalo (*S. caripense*; Product code: Y5SSSOCA), devil’s fig (*S. asperolanatum*; Product code: Y5SSSOAS), *S. taeniotrichum* (Product code: Y5SSSOTA) from rarepalmseeds.com; tomato (*S. lycopersicum*; Variety: Valley Girl F1) seeds from Johnnyseeds.com; garden huckleberry (*S. melanoceasum*; Brand: Palm Beach Medicinal Herbs), easter eggplant (*S. ovigerum*; Brand: Helens Garden) and Turkish orange eggplant (*S. aethiopicum*; Seller: Seedville USA) from amazon.com, and Aubergine (*S. melongena*; Shikou hybrid Eggplant; Item: 52568-PK- P1) from parkseed.com.

Seeds of all the species were sown in potting mixture (Sunshine professional growing mix: Sun Gro Horticulture Canada Ltd., MA, USA; Tayal et al., 2020) filled trays (12.5” ×7.5” ×2”) and kept in controlled environmental conditions (26°C temperature, ~50% relative
humidity and 16:8 light dark cycle). Germinated seedlings were transplanted in plastic pots (5” × 4” × 4”) with similar soil media and environmental conditions and were watered regularly. For electron microscopy, plants of 4-6 weeks of age post transplanting with at least 10-12 fully developed leaves were used. Young and fully expanded leaves from randomly selected individuals (for each species, sample size varied by treatment, details below) were used for microscopy experiments.

**Desktop Scanning Electron Microscope (DSEM):**

To capture images from both abaxial and adaxial of leaves for trichome morphology (n= 3-11 plants/side/species), dimension measurements (n= 3-11 plants/side/species) and density analysis (n= 3-11 plants/side/species), a desktop Scanning Electron Microscope (SNE-4500 Plus Tabletop; Nanoimages LLC, Pleasanton, California, USA; Watts et al., 2021 in press) was employed. Circular leaf discs (0.63 cm in diameter) of fresh leaf samples (collection method detailed above) were excised from the plants using a hole punch. No chemical treatments (e.g., glutaraldehyde; Kariyat et al., 2017), critical drying or sputter coating was done to the leaf samples, and fresh leaf discs mounted on the aluminum stubs using double-sided carbon tape were directly inserted in the DSEM for scanning and image processing. For more details on operational procedures and methodology of DSEM, see Watts and Kariyat (2021), and Watts et al., (2021) in press.

**Trichome Morphology assessment:**

Fresh leaf samples (n= 3-11 plants/side/species) as described above were used and magnified ranging from 45X to 1000X depending on trichome type and size, to achieve maximum resolution to extract finer details of trichomes. Images of different trichome types from both abaxial and adaxial surface of leaves were captured at different angles in 3-D and later used to classify them. Payne (1978; consisting of glossary for different shapes and structures of trichomes), Roe (1971; consisting of terminology for commonly found Solanum trichomes)
and Werker (2000; glandular trichome characterization based on structure of secretory head) were the major literature used to characterize trichomes post image acquisition.

**Trichome Density Assessment:**

To determine the trichome density from both leaf surfaces of all the species \((n=3-11\text{ plants/side/species})\), sample preparation was done as described above. The images for trichome count were consistently captured at 60X magnification which contains approximately 5.32 mm\(^2\) leaf area measured using ‘Nanoeye’ software linked to DSEM. We calculated the trichome density 1 mm\(^2\) as follows (Chavana et al., 2021; Watts and Kariyat, 2021):

\[
\text{Trichome density (1 mm}^2\text{)} = \frac{\text{Number of trichomes in the image taken at 60X magnification}}{5.32}
\]

**Trichome Dimension Measurements:**

While scanning the leaf samples \((n=3-11\text{ plants/side/species})\), once the image achieved maximum resolution visually, scanning was paused using ‘Nanoeye’ software associated with DSEM, and dimensions of various trichome types were measured by tracing trichomes by straight line in ‘M. tools’ in ‘Nanoeye’ software. For non-glandular trichomes, length of spikes from base to tip was measured, and in case of glandular trichomes, length of trichome from base to tip and diameter of bulb containing glandular secretions was measured using the measurement tool embedded in the software (Figure 7). Magnification was altered among samples depending on trichome type at the best resolution.
**Line art:** The SEM images were used to draw the trichomes manually on paper. The paper was scanned to make a digital copy and the trichomes were traced using a size 4 solid circle brush (Wacom Intuos Pro Digital Graphic Drawing Tablet; Adobe Photoshop). Further, custom brushes in Adobe Photoshop were used to create non-uniform surface of few trichome types. The images were saved into transparent PNG files and inserted into tables with the base of trichomes is on the bottom of the image.

**Analysis**

Our goal was to characterize and document the trichome types in these 14 species, and to define the finer details on individual trichome types, and their dimensions. Using previous publications as a composite reference, we classified the trichome types using images that reflected its most detailed morphological features. The nomenclature of trichomes was decided by following a checklist of features in the order described below: major shape of the trichome; glandular/non-glandular trichome type (italicized); additional minute morphological specifications. Further, additional features have also been added as a separate column to know more details of each trichome type. For morphological representation of trichome types, line art was created for each individual trichome type, by a graphical artist.

Following this, mean trichome density for each trichome type or trichome types as broad groups (glandular; simple non-glandular; stellate non-glandular) was calculated manually from images at 60X magnification. 60X magnification includes 5.32 mm$^2$ of leaf area, and thus to get density of trichomes in 1 mm$^2$ of area, the trichomes density at 60X magnification was divide by a factor of 5.32. The Mean±SE of trichome density has been incorporated in Table 2 and 3. And, to test whether species and trichomes varied across the 14 species of interest, we also ran a Generalized regression analysis with Poisson distribution with species
and types (total, glandular and non-glandular) and their interaction as factors, and trichome number as the response variables. Tukey’s post hoc tests were conducted to examine pairwise comparisons. All analyses were carried out using JMP15 (SAS Inc, Cary, JJ, USA) software and the plots were built using GraphPad Prism (La Jolla, CA, USA). And finally Mean±SE total length and Mean±SE of the diameter of glands on glandular trichomes, and Mean ± SE spike length in case of non-glandular trichomes were measured.

Results

*Trichome Morphology assessment:*

Basic classification of trichomes includes classifying them into glandular and non-glandular types (italicized; Table 2 and 3). Here, detailed classification of trichomes was carried out using previously published works as a composite reference. The terminology used for nomenclature of trichomes is described below in Table 1.

Using above-mentioned terminology, we classified all the trichomes found in our samples. Although, three trichome types viz. stellate non-glandular, simple non-glandular, and glandular type trichomes were the most found in the *Solanum* species in our study, these types have also been further characterized into a numerous sub-types based on minor morphological differences (Figure 1; Table 2 and 3).

Stellate non-glandular trichomes have further been divided into multitangulate, multiradiate, stellate non-glandular hair with subulate rays (2-6 in number) with the presence of pedestal (*Solanum aethiopicum*; adaxial) (Table 2; Serial number 2), porrect-stellate multiradiate non-glandular hair with subulate rays (varying in number) and with short central ray (*S. aethiopicum*; abaxial, *S. anguivi*; abaxial, *S. lanceifolium*; adaxial and abaxial, *S. ovigerum*; abaxial, *S. pyracanthos*; adaxial and abaxial) (Table 2; 9, 45; Table 3; 4, 7, 13, 45, 51),
porrect-stellate multiradiate non-glandular hair with subulate rays (varying in number) with long central ray (S. anguivi; adaxial and abaxial, S. lanceifolium; adaxial, S. ovigerum; adaxial) (Table 2; 8, 10, 39; Table 3; 8), porrect-stellate multi-radiate cruciate non-glandular hair with subulate rays (4 in number) and with short central ray (S. grandiflorum; abaxial) (Table 3; 30), porrect-geminate stellate multi-radiate non-glandular hair with subulate rays (2-16 in number) and short central ray (S. melongena; adaxial and abaxial) (Table 2; 30; Table 3; 34). Moreover, some stellate trichomes also have a pedestal (S. aethiopicum; adaxial, S. anguivi; adaxial and abaxial, S. ovigerum; abaxial, S. pyracanthos; abaxial) (Table 2; 2, 3, 8; Table 3; 7, 8, 45, 51). And some stellate trichomes with only two rays at an angle have been named separately as bifurcated non-glandular hair (S. aethiopicum; adaxial, S. anguivi; adaxial and abaxial, S. lanceifolium; abaxial, S. pyracanthos; abaxial) (Table 2; 3, 7; Table 3; 9, 14, 53). Within each stellate trichome, spike number also varied in almost all the species (Figure 1, Table 2 and 3).

Simple non-glandular trichomes have also been further sub-divided as osteolate (S. dulcamara; abaxial) (Table 3; 57), subulate (S. aethiopicum; adaxial and abaxial, S. anguivi; adaxial and abaxial, S. lycopersicum; abaxial, S. macrocarpon; adaxial and abaxial, S. melanocerasum; adaxial, S. grandiflorum; adaxial and abaxial, S. melongena; adaxial and abaxial, S. taeniotrichum; abaxial, S. caripense; adaxial and abaxial, S. ovigerum; adaxial, S. pyracanthos; adaxial, S. dulcamara; adaxial) (Table 2; 1, 6, 20, 22, 25, 31, 36, 37, 40; Table 3; 3, 6, 18, 23, 27, 33, 39, 40, 43), falcate (S. grandiflorum; abaxial, S. ovigerum; abaxial) (Table 3; 32, 44), setiform (S. grandiflorum; adaxial) (Table 2; 29), crescent (S. melanocerasum; abaxial) (Table 3; 24), attenuate (S. lycopersicum; adaxial and abaxial) (Table 2; 17, 18; Table 3; 19, 20), hooked (S. aethiopicum; abaxial; S. lycopersicum; adaxial and abaxial, S. melongena; abaxial, S. caripense; abaxial; S. pyracanthos; abaxial) (Table 2; 3, 16; Table 3; 18, 33, 40, 54). And most of these trichome types were smooth (all species)
(Figure 1 and 2; Table 2 and 3) while few others were pustulated (S. macrocarpon; adaxial and abaxial, S. dulcamara; abaxial) (Table 2; 20; Table 3; 23, 57). Similar to stellate non-glandular trichomes, some simple non-glandular trichomes have pedestal (S. macrocarpon; adaxial and abaxial, S. grandiflorum; adaxial, S. melongena; adaxial, S. asperolanatum; adaxial, (Table 2; 20, 25, 29, 31, 41; Table 3; 23, 27) while others do not have pedestal. The third most found trichome type was glandular type which has been further characterized based on the presence of a globular head (all species) (Figure 1 and 2; Table 2 and 3) or small glandular tip (S. aethiopicum; abaxial, S. lanceifolium; abaxial, S. lycopersicum; adaxial and abaxial, S. macrocarpon; adaxial, S. melanocerasum; adaxial and abaxial, S. grandiflorum; adaxial, S. taeniotrichum; adaxial and abaxial, S. asperolanatum; adaxial and abaxial, S. dulcamara; abaxial) (Figure 2 and 3; Table 2; 13, 19, 23, 28, 33; Table 3; 1, 10, 17, 20, 25, 36, 37, 56). Further globular head can be large (Table 2; 5, 11, 12, 14, 21, 24, 26, 32, 41; Table 3; 2, 5, 12, 15, 16, 22, 26, 28, 31, 35, 42, 48, 52) or small (Table 2; 4, 5, 34, 35, 38, 44, 48; Table 3; 38, 46, 55, the characterization made based on comparative visual observations (Figure 2 and 3; Table 2 and 3). Globular headed trichomes can vary in shape as was observed in case of S. lanceifolium (doliform globular head; adaxial; Table 2; 9). Additionally, globular headed glandular trichomes having distinct four-celled head have also been observed (Table 2; 5; Table 3; 5, 15, 16, 42). Similar to simple non-glandular trichomes, glandular trichomes were also found in various major shapes such as attenuate (S. aethiopicum; abaxial, S. lanceifolium; abaxial, S. lycopersicum; adaxial and abaxial, S. macrocarpon; adaxial, S. grandiflorum; adaxial, S. dulcamara; abaxial) (Table 2; 13, 19, 28; Table 3; 1, 10, 20, 57), acuminate (S. lycopersicum; adaxial) (Table 2; 15), subulate (S. lycopersicum; abaxial, S. melanocerasum; adaxial and abaxial, S. taeniotrichum; adaxial and abaxial) (Table 2; 23, 33; Table 3; 17, 25, 36, 37), hooked (S. lycopersicum; abaxial) (Table 3; 17). In case of non-glandular hair with attenuate non-glandular branches and one gland-
tipped branch in *S. asperolanatum*, trichome has one branch with a glandular tip making it both glandular and non-glandular type trichome but named as non-glandular because of a greater number of rays being non-glandular (Table 3; 50).

And based on previous literature (Werker, 2000) and since these trichome lack a clear distinction of glandular head, ovoid (*S. grandiflorum;* adaxial and adaxial, *S. asperolanatum;* adaxial and abaxial) (Figure 1, table 2; 27, 42; Table 3; 29), mammilla (*S. lycopersicum;* abaxial) (Table 3; 21) and verrucate (*S. lancefolium;* abaxial) (Table 3; 11) trichomes were characterized as non-glandular. Although DSEM microscope used in the study had numerous benefits including no sample preparation or the use of critical chemicals and machinery, and faster image processing (critical point dryers and sputter coaters; Kariyat et al., 2017), it also had some drawbacks. One of them was that we were unable to further classify the glandular trichomes based on the number of their head cells as SEM images lacked those details. And, since fresh leaf samples were used for the study, few glandular trichomes with pliable heads burst on their encounter with vacuum of the machine while scanning, due to the lack of critical point drying and sputter coating. For detailed results of species, trichome type, their density, and dimensions with pictorial representation of each trichome type on adaxial leaf surface see table 2, and on abaxial leaf surface, see table 3.

**Density Measurements:**

Consistent with morphological diversity and variation among and within each species, density of trichomes also varied across species (Watts and Kariyat, 2021; Figure 2 and 3; Tables 2 and 3). Oddly, in some of the species, while acquiring images for density count at 60X, we did not observe any trichomes, but while zooming in on different leaf samples at a higher magnification, we observed few trichome types although they were quite rare. These include attenuate basilatus glandular hair with small glandular tip on adaxial leaf surface of *S.*
macrocarpon (Table 2; Serial number 19), attenuate basilatus glandular hair with small glandular tip; mamilla non-glandular hair on abaxial leaf surface of *S. lycopersicum* (Table 3; 20, 21), subulate glandular hair with multicellular jointed stalk and small glandular tip on abaxial leaf surface of *S. melanocerasum* (Table 3; 24), subulate non-glandular hair with multiseriate base and tall pedestal on abaxial leaf surface of *S. grandiflorum* (Table 3; 27), hooked subulate non-glandular hair on abaxial leaf surface of *S. caripense* (Table 3; 40), and falcate non-glandular hair on abaxial leaf surface of *S. ovigerum* (Table 3; 44).

The trichome types with highest trichome density include glandular hair with large globular head, single stalk cell and no neck cell on adaxial leaf surface of *S. asperolanatum* (Figure 3, Table 2; 41), and porrect-stellate multiradiate non-glandular hair with subulate rays (5-10 in number) and with short central ray on abaxial leaf surface of *S. aethiopicum* (Figure 2; Table 3; 4). Although the variation was huge, we found that some trichome types had higher trichome density than the other types in each species. For example, in *S. anguivi* (abaxial) porrect-stellate multi-radiate non-glandular hair with subulate rays (5-10 in number) with short/long central ray and pedestal (density: 13.00±1.56; Table 3; 7, 8 and 9) had considerably higher trichome density than the other two trichome types (Table 3; 5 and 6) (Figure 2). Similarly, were observed that in almost all the species in which one/two trichome types dominated over the other. In addition to this, occasionally it was difficult at 60X to distinguish between some trichome types, so the density of some trichome types have been compiled. For example, manual counting of both glandular hair with large quadricellular globular head and single stalk cell in case of abaxial leaf surface of *S. lycopersicum* resulted into total density of 0.94±0.19 (Table 3; 15 and 16). Such cases were observed for all three major trichome types (stellate non-glandular, simple non-glandular and glandular trichomes) (Table 2 and 3). Additionally, we found significant variation in trichome numbers (total, glandular and non-glandular; at 60X magnification) among species (Generalized Regression;
p ≤ 0.0001) and interaction of species with trichome type (total, glandular and non-glandular) (Generalized Regression; p ≤ 0.0001) (Figure 4, 5 and 6), but the variation was non-significant between trichome types (Generalized Regression; p= 0.6971).

**Trichome Dimensions**

Similar to density, dimensions of each trichome type also varied across species and location (Figure 7; Table 2 and 3). Non-glandular trichome types- subulate non-glandular hair with multiseriate base and tall pedestal on abaxial leaf surface of *S. grandiflorum* (Table 3; Serial number 27) was the longest in dimensions, and the shortest non-glandular trichome was subulate non-glandular hair with multicellular jointed stalk and multicellular base on abaxial leaf surface of *S. ovigerum* (Table 3; 43). Longest glandular trichome type was subulate glandular hair with multicellular jointed stalk, multicellular base, distinct subsidiary cells, and small glandular tip on adaxial leaf surface of *S. melanocerasum* (Table 2; 23), and the shortest glandular trichome type was glandular hair with large globular head on abaxial leaf surface of *S. melongena* (Table 3; 35). The glandular trichome with largest head was glandular hair with large quadricellular globular head and single stalk cell on abaxial leaf surface of *S. lycopersicum* (Table 3; 16). The glandular trichome with smallest glandular tip was hooked subulate glandular hair with multicellular jointed stalk and small glandular tip on abaxial leaf surface of *S. lycopersicum* (Table 3; 17). Dimensions of some trichome types seemed comparable and thus dimension data for few trichome types was collected as one type (Stellate and bifid trichomes; all globular glandular trichomes; glandular trichomes with small tip; simple trichomes).

Although the stellate trichomes had almost consistent spike length within a species or/and leaf surface, but central ray of stellate trichomes varied (long/short) resulting into subdividing them into stellate trichomes with short or long central ray. Simple trichomes had the
most variation resulting them being both the shortest and the longest trichome found among species in the study. Further, in general, glandular trichomes with globular heads were shorter than glandular trichomes with small tip on the top. Additionally, glandular trichomes with globular heads had greater diameter of their glandular heads than glandular trichomes with small tip on the top. Due to the high species numbers, processing of multiple samples, and incredible trichome diversity, we could not acquire dimensions of some trichome types (for example, osteolate non-glandular hair with multicellular stalk on abaxial leaf surface of *S. dulcamara*; Table 3; 57).

**Discussion**

In this study, we examined the trichome characteristics including their nomenclature, density, and dimensions on both adaxial and abaxial leaf surface of 14 *Solanum* species using scanning electron microscopy on fresh leaf samples. We found that *Solanum* genus consists of numerous trichome types which vary not only among the species, but also within each species and between adaxial and abaxial leaf surfaces. We also found that three trichome types are most common in *Solanum*: stellate non-glandular, simple non-glandular and glandular trichomes. Broadly, all trichomes have been characterized into glandular and non-glandular trichomes, but it is not fair, because both glandular and non-glandular trichomes can further be classified into various types based on their shape, size, number of cells, basal cells, neck cells, etc. Besides, it was not certain whether ovoid, verrucate and mammila trichomes are glandular or non-glandular because of their unusual and perplexing structure but we confirm that they can be characterized as non-glandular trichomes because of their lack of distinction of a clear glandular head.
Trichomes, in general, have been proven to be an excellent phenotypic trait for finding evolutionary and taxonomic relationships among species (Cantino, 1990; Khosroshahi and Salmaki, 2019). For example, *Phlomis* genus has characteristic multi-nodal branched trichomes and can be considered as synapomorphy for this group, but *Phlomoides* lacks this feature and thus, this feature most likely represents a plesiomorphy in the genus (Khosroshahi and Salmaki, 2019); and separation of African and Asian *Leucas spp* were made more explicit by the differences of capitate trichomes with definite morphology and absence of non-glandular trichomes with one cell and more than three cells (Mannethody and Purayidathkandy, 2018). Our study characterized the finer details of trichome morphology, and it can be of an aid in exploring phylogenetic and taxonomic relationships among the members of genus *Solanum*, and their relationship with members of other genus of Solanaceae family and among other plant families. Additionally, trichomes act as excellent cell differentiation models (Hülskamp, 2004) and provided with the diversity from this study, it can be explored how differentiation in trichome cells result into production of glandular/non-glandular of various shapes and sizes.

Defense against herbivores is one of the major functions of trichomes, and both glandular and non-glandular trichomes have been well documented to deter herbivore movement and feeding (Karabourniotis et al., 2020; Kariyat et al., 2013; 2017; 2019). Morphology, density and dimensions relationships of sub-types of trichomes can be employed to find correlations between trichome characteristics with herbivore feeding intensity and behavior (Li et al., 2020). For instance, Watts and Kariyat (2021) found that significantly higher trichome density on abaxial leaf surface than adaxial leaf surface of Solanaceae species resulted into delayed feeding and lower mass gain of tobacco hornworm caterpillars (*Manduca sexta*; Lepidoptera: Sphingidae); Kariyat et al. (2017) found damage done to peritrophic membrane (gut-lining) of *M. sexta* caterpillars after feeding on stellate trichomes of horsenettle...
Further, since the damage done by non-glandular trichomes is primarily because of their structure and many herbivores mow the trichomes off the plant surface before feeding (Kariyat et al., 2017; 2019; Kaur & Kariyat, 2020b), we speculate that trichomes with a greater number of spikes, and spikes with higher dimensions can result into higher negative impacts on herbivore feeding, but warrants closer examination (Medeiros & Moreira, 2005; Andama et al., 2020). The variation in trichome types, density, dimensions, and their functional consequences (Watts and Kariyat, 2021) between abaxial and adaxial leaf surfaces also warrants detailed exploration, a reason why we examined these differences in detail (Tables 2 and 3). Contrary to non-glandular trichomes, glandular trichomes are the secretory structures and contain various types of chemicals in their head cells and those chemicals have been found to trigger different defense related pathways against herbivores (Tissier, 2012), and some glandular trichome types have been found to play more prominent roles than the others. For example, type VI (as named by Luckwill, 1943; glandular trichomes with quadricellular head) trichomes of *Lycopersicon* genus were found to contain chemicals possessing insecticidal properties against lepidopteran larvae (Lin et al., 1987). Thus, knowledge of glandular trichome density and dimensions such as length and diameter can help us know the plant parts with higher trichome density, amount of chemicals possessed by trichomes and if the trichomes are tall enough to act against herbivores with its structural features along with chemical defense. Additionally, expanding on this study, histochemistry and volatile collection of various glandular secretions can be done, and anti-herbivore chemicals can be identified (Muravnik et al., 2019), and can lead to further functional assessment and classification, an area that we are currently exploring. Trichomes have been of great importance against herbivores as a defense trait, and thus, have been incorporated in integrated pest management of insect pests of various crops of economic importance including potato (*Solanum tuberosum*; Solanaceae), cotton (*Gossypium spp.*;
Malvaceae), cowpea (*Vigna unguiculata*; Fabaceae), to name a few. For instance, *hooked* non-glandular trichomes of *Phaseolus vulgaris* (Fabaceae) entrap pests such as black bean aphid (*Aphis fabae*; Hemiptera: Aphididae) and green stink bug (*Nezara viridula*; Hemiptera: Pentatomidae) (Rebora et al., 2020). Trichomes also play multiple roles in plants including leaf water uptake and protection from UV light, in addition to defense against herbivores (Sack and Buckley, 2020). For example, Li et al., 2021, showed that among three trichome types viz. non-glandular trichomes, linear glandular trichomes and glandular trichomes, only non-glandular trichomes of Sunflower (*Helianthus annuus*) were found to accumulate and translocate Zinc, an important micronutrient for plants, an potential area of research to be explored for mineral and nutrient uptake to enhance crop yield through trichomes. Clearly, trichome characteristics has the potential to be explored in relation plant ecophysiological functions such as water use efficiency (Thitz et al., 2017), UV protection and mineral uptake (Feng et al., 2021).

Taken together, this study documented the variation in trichome types, their density, and dimensions in representative *Solanum* species. So far, the major reasons which withheld the detailed classification and nomenclature of trichomes were (1) the conventional approach to classify trichomes as just glandular or non-glandular trichomes failed to document their morphotypes and function, (2) the requirement of expensive machinery and skilled professionals, required to process samples across species and families, and (3) the need of intensive and detailed workflow for the closer examination of images to extract all the data on morphology, density and dimensions. By overcoming these challenges, we show the variation in trichome traits within a subset of the genus *Solanum* and encourage more detailed examination across various plant families.
Conclusions:

Overall, the study provides unusually fine details and morphological characterization of trichomes of a mixture of wild and domesticated, annual, and perennial, food crops and weed species of genus *Solanum* which can act as referral source for further studies of most of trichome related parameters and their relationships with biotic and abiotic stresses (Tian, 2012). Since *Solanum* is the largest genus of one of the major angiosperm family viz. Solanaceae, exploring trichome diversity by considering fourteen species from various groups (for example, *S. macrocarpon* and *S. lycopersicum* are cultivated species, while *S. anguivi* and *S. pyracanthos* are wild species) of the family provided us with an updated data source of trichome characteristics with such details that never has been done before. Further, future directions in trichome studies can be focused on understanding variability and organ development while studying gene expression simultaneously using trichomes as a model. Moreover, trichomes are also known to play role in multi-trophic interactions in ecosystem (Weinhold and Baldwin, 2011) and thus, each trichome type can be explored for its potential in strengthening plants’ defenses.
References

Adedeji O, Ajuwon OY, Babawale OO. 2007. Foliar epidermal Studies, organographic distribution and taxonomic importance Of trichomes in the family Solanaceae. *International Journal of Botany* 3: 276–282.

Andama JB, Mujiono K, Hojo Y, Shinya T, Galis I. 2020. Nonglandular silicified trichomes are essential for rice defense against chewing herbivores. *Plant, Cell & Environment* 43: 2019–2032.

Avé DA, Gregory P, Tingey WM. 1987. Aphid repellent sesquiterpenes In glandular trichomes of Solanum berthaultii and S. tuberosum. *Entomologia Experimentalis et Applicata* 44: 131–138.

Blamey FP, Joyce DC, Edwards DG, Asher CJ. 1986. Role of trichomes in sunflower tolerance to manganese toxicity. *Plant and Soil* 91: 171–180.

Blamey P, Hernandez-Soriano M, Cheng M, Tang C, Paterson D, Lombi E, Wang WH, Scheckel K, Kopittke PM. 2015. Synchrotron-based techniques shed light on mechanisms of Plant sensitivity and tolerance to High manganese in the root environment. *Plant Physiology*.

Burrows GE, White RG, Harper JD, Heady RD, Stanton RA, Zhu X, Wu H, Lemerle D. 2013. Intrusive trichome bases in the leaves Of SILVERLEAF nightshade (Solanum elaeagnifolium; Solanaceae) do not facilitate fluorescent tracer uptake. *American Journal of Botany* 100: 2307–2317.

Cantino PD. 1990. The phylogenetic significance of stomata and trichomes in the Labiatae and Verbenaceae. *Journal of the Arnold Arboretum*.

Channarayappa C, Shivashankar G, Muniyappa V, Frist RH. 1992. Resistance of lycopersicon species to Bemisia tabaci, a tomato leaf curl virus vector. *Canadian Journal of Botany* 70: 2184–2192.

Chavana J, Singh S, Vazquez A, Christoffersen B, Racelis A, Kariyat RR. 2021. Local adaptation to continuous mowing makes the noxious weed Solanum elaeagnifolium a superweed candidate by improving fitness and defense traits. *Scientific Reports* 11.

Cho K-S, Kwon M, Cho J-H, Im J-S, Park Y-E, Hong S-Y, Hwang I-T, Kang J-H. 2017. Characterization of trichome morphology and aphid resistance in cultivated and wild species of potato. *Horticulture, Environment, and Biotechnology* 58: 450–457.

CORSI G, Bottega S. 1999. Glandular hairs of Salvia officinalis: New data on morphology, localization and histochemistry in relation to function. *Annals of Botany* 84: 657–664.

Dalin P, Ågren J, Björkman C, Huttunen P, Kärkkäinen K. 2008. Leaf trichome formation and plant resistance to herbivory. *Induced Plant Resistance to Herbivory*: 89–105.

Deore CR. 2020. Morphologicalvariation of trichomes in some common species of angiosperm and their systematic enumeration useful for taxonomic significance.
Ehleringer J. 1982. The influence of water stress and temperature on leaf pubescence development in Encelia Farinosa. American Journal of Botany 69: 670.

Elle E, van Dam NM, Hare JD. 1999. Cost of glandular trichomes, a "Resistance" character in Datura Wrightii REGEL (SOLANACEAE). Evolution 53: 22.

Ensikat H-J, Wessely H, Engeser M, Weigend M. 2021. Distribution, ecology, chemistry and toxicology of Plant Stinging Hairs. Toxins 13: 141.

Eyvazadeh Kehroshahi E, Salmaki Y. 2019. Evolution of trichome types and its systematic significance in the Genus Phlomoides (lamioideae–lamiaceae). Nordic Journal of Botany 37.

Feng L-jie, An W-jing, Di LUI, Wang K-jie, Liang W-hong. 2021. Progress in Research of Rice Trichome Related Genes. Biotechnology Bulletin.

Grafius EJ, Douches DS. 2008. The present and future role of insect-resistant genetically modified potato cultivars in IPM. Integration of Insect-Resistant Genetically Modified Crops within IPM Programs: 195–221.

Hare JD. 2005. Biological activity of acyl glucose esters from Datura wrightii glandular trichomes against three native insect herbivores. Journal of Chemical Ecology 31: 1475–1491.

Hülskamp M. 2004. Plant trichomes: A model for cell differentiation. Nature Reviews Molecular Cell Biology 5: 471–480.

Kang J-H, Liu G, Shi F, Jones AD, Beaudry RM, Howe GA. 2010. The tomato Odorless-2 Mutant is defective in trichome-based production of diverse specialized metabolites and broad-spectrum resistance to insect herbivores. Plant Physiology 154: 262–272.

Karabourniotis G, Liakopoulos G, Nikolopoulos D, Bresta P. 2019. Protective and defensive roles of non-glandular trichomes against multiple stresses: Structure–function coordination. Journal of Forestry Research 31: 1–12.

Kariyat RR, Balogh CM, Moraski RP, De Moraes CM, Mescher MC, Stephenson AG. 2013. Constitutive and herbivore- induced structural defenses are compromised by inbreeding in Solanum carolinense (Solanaceae). American Journal of Botany 100: 1014–1021.

Kariyat RR, Hardison SB, Ryan AB, Stephenson AG, De Moraes CM, Mescher MC. 2018. Leaf trichomes affect caterpillar feeding in an instar-specific manner. Communicative & Integrative Biology 11: 1–6.

Kariyat RR, Raya CE, Chavana J, Cantu J, Guzman G, Sasidharan L. 2019. Feeding on glandular and non-glandular leaf trichomes negatively affect growth and development in tobacco hornworm (Manduca sexta) caterpillars. Arthropod-Plant Interactions 13: 321–333.

Kariyat RR, Smith JD, Stephenson AG, De Moraes CM, Mescher MC. 2017. Non-glandular trichomes of Solanum carolinense deter feeding by Manduca sexta caterpillars and
cause damage to the gut peritrophic matrix. *Proceedings of the Royal Society B: Biological Sciences* **284**: 20162323.

**Kaur I, Kariyat RR. 2020a.** Eating barbed wire: Direct and indirect defensive roles of non-glandular trichomes. *Plant, Cell & Environment* **43**: 2015–2018.

**Kaur J, Kariyat R. 2020b.** Role of trichomes in plant stress biology. *Evolutionary Ecology of Plant-Herbivore Interaction*: 15–35.

**Kennedy GG. 2003.** Tomato, pests, parasitoids, and predators: tritrophic interactions involving the genus Lycopersicon. *Annual Review of Entomology* **48**: 51–72.

**Knapp S. 2002.** Tobacco to tomatoes: a phylogenetic perspective on fruit diversity in the Solanaceae. *Journal of Experimental Botany* **53**: 2001–2022.

**Li C, Wu J, Blamey FP, Wang L, Zhou L, Paterson DJ, van der Ent A, Fernández V, Lombi E, Wang Y, Kopittke PM. 2021.** Non-glandular trichomes of sunflower are important in the absorption and translocation of foliar-applied Zn. *Journal of Experimental Botany* **72**: 5079–5092.

**Li S, Tosens T, Harley PC, Jiang Y, Kanagendran A, Grosberg M, Jaamets K, Niinemets Ü. 2018.** Glandular trichomes as a barrier against atmospheric oxidative stress: Relationships with ozone uptake, leaf damage, and emission of lox products across a diverse set of species. *Plant, Cell & Environment* **41**: 1263–1277.

**Li Y, Lei L, Luo R, Li C, Luo C. 2020.** Morphological structures of bamboo (Bambusa Emeiensis) shoot shells and trichomes and functions in response to herbivory. *Journal of Plant Growth Regulation*.

**Lin SY, Trumble JT, Kumamoto J. 1987.** Activity of volatile compounds in glandular trichomes of Lycopersicon species against two insect herbivores. *Journal of Chemical Ecology* **13**: 837–850.

**LoPresti EF, Pearse IS, Charles GK. 2015.** The siren song of a sticky plant: columbines provision mutualist arthropods by attracting and killing passerby insects. *Ecology* **96**: 2862–2869.

**Luckwill LC. 1943.** The genus Lycopersicon: an historical, biological, and taxonomic survey of the wild and cultivated tomatoes. Aberdeen, Scotland: Aberdeen University Press.

**Løe G, Toräng P, Gaudeul M, Ågren J. 2007.** Trichome production and spatiotemporal variation in herbivory in the perennial herb Arabidopsis lyrata. *Oikos* **116**: 134–142.

**Mannethody S, Purayidathkandy S. 2018.** Trichome micromorphology and its systematic significance in Asian Leucas (lamiaceae). *Flora* **242**: 70–78.

**Medeiros L, Moreira GR. 2005.** Larval feeding behavior of Gratiana spadicea (klug) (Coleoptera: Chrysomelidae: Cassidinae) on its host plant, Solanum sisymbriifolium lamarck (Solanaceae): Interaction with trichomes. *The Coleopterists Bulletin* **59**: 339–350.
Mehdi Talebi S, Ghorbani Nohooji M, Yarmohammadi M, Azizi N, Matsyura A. 2018. Trichomes morphology and density analysis in some "Nepeta" species of Iran. *Mediterranean Botany* **39**: 51–62.

Miyazaki J, Stiller WN, Wilson L.J. 2017. Sources of plant resistance to thrips: A potential core component in cotton IPM. *Entomologia Experimentalis et Applicata* **162**: 30–40.

Munien P, Naidoo Y, Naidoo G. 2015. Micromorphology, histochemistry and ultrastructure of the foliar trichomes of Withania somnifera (L.) Dunal (Solanaceae). *Planta* **242**: 1107–1122.

Muravnik LE, Kostina OV, Mosina AA. 2019. Glandular trichomes of the leaves in three Doryicum species (Senecioneae, Asteraceae): morphology, histochemistry, and ultrastructure. *Protoplasma* **256**: 789–803.

Murungi LK, Kirwa H, Salifu D, Torto B. 2016. Opposing roles of foliar and glandular trichome volatile components in cultivated nightshade interaction with a specialist herbivore. *PLOS ONE* **11**.

Navarro T, El Oualidi J. 1999. Trichome morphology in Teucrium L. (Labiatae). a taxonomic review. *Anales del Jardín Botánico de Madrid* **57**.

Neal JJ, Tingey WM, Steffens JC. 1990. Sucrose esters of carboxylic acids in glandular trichomes of Solanum berthaultii deter settling and probing by green peach aphid. *Journal of Chemical Ecology* **16**: 487–497.

Nurit-Silva K, De Fátima Agra M. 2011. Leaf epidermal characters of Solanum Sect. Polytrichum (Solanaceae) as taxonomic evidence. *Microscopy Research and Technique* **74**: 1186–1191.

Oghiakhe S, Jackai LEN, Makanjuola WA, Hodgson CJ. 1992. Morphology, distribution, and the role of trichomes in cowpea (Vigna unguiculata) resistance to the Legume Pod borer, Maruca testulalis (Lepidoptera: Pyralidae). *Bulletin of Entomological Research* **82**: 499–505.

Oksanen E. 2018. Trichomes form an important first line of defence against adverse environment-new evidence for ozone stress mitigation. *Plant, Cell & Environment* **41**: 1497–1499.

Pan Z-L, Guo W, Zhang Y-J, Schreel JDM, Gao J-Y, Li Y-P, Yang S-J. 2021. Leaf trichomes of Dendrobium species (epiphytic orchids) in relation to foliar water uptake, leaf surface wettability, and water balance. *Environmental and Experimental Botany* **190**: 104568.

Payne WW. 1978. A glossary of plant hair terminology. *Brittonia* **30**: 239.

Peiffer M, Tooker JF, Luthe DS, Felton GW. 2009. Plants on early alert: Glandular trichomes as sensors for insect herbivores. *New Phytologist* **184**: 644–656.

Rebora M, Salerno G, Piersanti S, Gorb E, Gorb S. 2020. Entrapment of Bradysia paupera (Diptera: Sciaridae) by Phaseolus vulgaris (Fabaceae) plant leaf. *Arthropod-Plant Interactions* **14**: 499–509.
Roe KE. 1971. Terminology of hairs in the genus Solanum. *TAXON* **20**: 501–508.

Sack L, Buckley TN. 2020. Trait multi-functionality in plant stress response. *Integrative and Comparative Biology* **60**: 98–112.

Seithe A, Sullivan JR. 1990. Hair morphology and systematics of Physalis (Solanaceae). *Plant Systematics and Evolution* **170**: 193–204.

Sletvold N, Huttenen P, Handley R, Kärkkäinen K, Ågren J. 2010. Cost of trichome production and resistance to a specialist insect herbivore in Arabidopsis lyrata. *Evolutionary Ecology* **24**: 1307–1319.

Symon DE. 1981. A revision of the genus Solanum in Australia. *Journal of Adelaide Botanic Gardens*.

Tang T, Li C-H, Li D-S, Jing S-X, Hua J, Luo S-H, Liu Y, Li S-H. 2020. Peltate glandular trichomes of Colquhounia vestita harbor diterpenoid acids that contribute to plant adaptation to UV radiation and cold stresses. *Phytochemistry* **172**: 112285.

Tayal M, Somavat P, Rodriguez I, Thomas T, Christoffersen B, Kariyat R. 2020. Polyphenol-rich purple corn pericarp extract adversely impacts herbivore growth and development. *Insects* **11**: 98.

Thitz P, Possen BJHM, Oksanen E, Mehtätalo L, Virjamo V, Vapaavuori E. 2017. Production of glandular trichomes responds to water stress and temperature in silver Birch (Betula pendula) leaves. *Canadian Journal of Forest Research* **47**: 1075–1081.

Tian D, Tooker J, Peiffer M, Chung SH, Felton GW. 2012. Role of trichomes in defense against herbivores: Comparison of herbivore response to woolly and hairless trichome mutants in tomato (Solanum lycopersicum). *Planta* **236**: 1053–1066.

Tingey WM, Laubengayer JE. 1981. Defense against the green peach aphid and potato leafhopper by glandular trichomes of Solanum berthaultii123. *Journal of Economic Entomology* **74**: 721–725.

Tissier A. 2012. Trichome Specific Expression: Promoters and Their Applications. INTECH Open Access Publisher.

Tsujii Y, Onoda Y, Izuno A, Isagi Y, Kitayama K. 2016. A quantitative analysis of phenotypic variations of Metrosideros polymorpha within and across populations along environmental gradients on Mauna Loa, Hawaii. *Oecologia* **180**: 1049–1059.

Uphof JC. 1962. *Plant Hairs*. Berlin, Berlin: Gebr. Borntraeger.

van Dam NM, Hare JD, Elle E. 1999. Inheritance and distribution of trichome phenotypes in Datura wrightii. *Journal of Heredity* **90**: 220–227.

Watts S, Kaur I, Singh S, Jiminez B, Chavana J, Kariyat R. Desktop scanning electron microscopy in plant-insect interactions research: A fast and effective way to capture electron micrographs with minimal sample preparation. *Biology Methods and Protocols*.
Watts S, Kariyat R. 2021. Picking sides: Feeding on the abaxial leaf surface is costly for caterpillars. *Planta* **253**.

Weigend M, Mustafa A, Ensikat H-J. 2018. Calcium phosphate in plant trichomes: The overlooked biomineral. *Planta* **247**: 277–285.

Weinhold A, Baldwin IT. 2011. Trichome-derived o-acyl sugars are a first meal for caterpillars that tags them for predation. *Proceedings of the National Academy of Sciences* **108**: 7855–7859.

Werker E. 2000. Trichome diversity and development. *Advances in Botanical Research*: 1–35.

Yu X, Liang C, Fang H, Qi X, Li W, Shang Q. 2018. Variation of trichome morphology and essential OIL composition of seven Mentha species. *Biochemical Systematics and Ecology* **79**: 30–36.

Zalucki MP, Clarke AR, Malcolm SB. 2002. Ecology and behavior of first instar larval lepidoptera. *Annual Review of Entomology* **47**: 361–393.

Zhou LH, Liu SB, Wang PF, Lu TJ, Xu F, Genin GM, Pickard BG. 2017. The Arabidopsis Trichome is an active mechanosensory Switch. *Plant, Cell & Environment* **40**: 611–621.
Figure legends:

Figure 1. The three major trichome types found in the study includes (A-E) Stellate non-glandular trichomes, (F-L) Simple non-glandular trichomes, and (M-U) glandular trichomes. Figure shows scanning electron microscopic images of (A) porrect-stellate multi-radiate non-glandular hair with subulate rays (2-7 in number) and with short central ray on adaxial leaf surface of *Solanum lanceifolium* (B) porrect-stellate multi-radiate non-glandular hair with subulate rays (3-7 in number) and with long central ray on adaxial leaf surface of *S. ovigerum* (C) bifurcated basilatus non-glandular hair with subulate rays (one shorter than the other) on adaxial leaf surface of *S. anguivi* (D) multitangulate multiradiate stellate non-glandular hair with subulate rays (2-6 in number) with the presence of pedestal on adaxial leaf surface of *S. aethiopicum* (E) porrect-geminate stellate multi-radiate non-glandular hair with subulate rays (2-16 in number) and short central ray on abaxial leaf surface of *S. melongena* (F) osteolate non-glandular hair with multicellular stalk on abaxial leaf surface of *S. dulcamara* (G) subulate basilatus non-glandular trichome with distinct subsidiary cells on adaxial leaf surface of *S. aethiopicum* (H) subulate basilatus non-glandular trichome with distinct subsidiary cells on adaxial leaf surface of *S. aethiopicum* (H) subulate basilatus non-glandular hair with multicellular jointed stalk and multicellular base on adaxial leaf surface of *S. caripense* (I) hooked subulate non-glandular hair on abaxial leaf surface of *S. lycopersicum* (J) subulate non-glandular hair with multiseriate base and tall pedestal on *S. grandiflorum* (K) subulate non-glandular hair with multicellular jointed stalk, multicellular base and distinct subsidiary cells on adaxial leaf surface of *S. melanocerasum* (L) subulate non-glandular hair with pulvinate base and a pedestal on adaxial leaf surface of *S. macrocarpon* (M) glandular hair with single stalk and neck cell and large quadricellular globular head on adaxial leaf surface of *S. anguivi* (N) glandular hair with single stalk and neck cell and large doliform globular head on adaxial leaf surface of *S. lanceifolium* (O) glandular hair with large quadricellular globular head and single stalk cell on abaxial leaf surface of *S. lycopersicum* (P) glandular hair with large globular head and single stalk cell on abaxial leaf surface of *S. macrocarpon* (Q) glandular hair with large globular head and single stalk cell on abaxial leaf surface of *S. grandiflorum* (R) subulate basilatus glandular hair with multicellular jointed stalk and small glandular tip on abaxial leaf surface of *S. taeniotrichum* (S) subulate glandular hair with multicellular jointed stalk and small glandular tip on adaxial leaf surface of *S. taeniotrichum* (T) glandular hair with small globular head on adaxial leaf surface of *S. ovigerum*.

Figure 2. Scanning Electron Microscopic (SEM) images captured at 60X magnification of abaxial (A-G) and adaxial (H-N) leaf surface of (A, H) *Solanum aethiopicum* (B, I) *Solanum anguivi* (C, J) *Solanum lanceifolium* (D, K) *Solanum melongena* (E, L) *Solanum pyracanthos* (F, M) *Solanum ovigerum* (G, N) *Solanum grandiflorum* with the presence of stellate non-glandular trichomes as one of the major trichome types.

Figure 3: Scanning Electron Microscopic (SEM) images captured at 60X magnification of abaxial (A-G) and adaxial (H-N) leaf surface of (A, H) *Solanum macrocarpon* (B, I) *Solanum melanocerasum* (C, J) *Solanum asperalanatum* (D, K) *Solanum taeniotrichum* (E, L) *Solanum caripense* (F, M) *Solanum dulcamara* and (G, N) *Solanum lycopersicum*, with stellate non-glandular trichome absent as a major trichome type.
Figure 4. Significant variation in total trichome density (Generalized regression; p= <0.0001) among 14 Solanum species.

Figure 5. Significant variation in total glandular trichome density (Generalized regression; p= <0.0001) among 14 Solanum species.

Figure 6. Significant variation in total non-glandular trichome density (Generalized regression; p= <0.0001), among 14 Solanum species.

Figure 7. Dimension measurement of different trichome types on (A) adaxial leaf surface of Solanum anguivi at 100X (B) adaxial leaf surface of S. aperolanatum at 45X (C) abaxial leaf surface of S. melongena at 150 X (D) abaxial leaf surface of S. grandiflorum at 60X, by tracing the trichome projections (length of hair and diameter of bulb in case of glandular trichomes, and length of hair/spike in case of non-glandular trichomes) using the ‘Nanoeye’ software.
Table 1. List of terminology used to define trichomes in the study.

| Terminology        | Definition (compiled from previously published literature)                                      |
|--------------------|------------------------------------------------------------------------------------------------|
| Attenuate          | Long and gradually tapering                                                                   |
| Base               | Lowermost part of the trichome                                                                |
| Basilatus          | Emerging from a broad base                                                                    |
| Bifurcated         | Divided into two branches                                                                     |
| Brevicolate        | Short-necked                                                                                  |
| Compound           | Having multiple rays                                                                          |
| Cruciate           | Shaped as a cross with four equal arms                                                        |
| Doliform           | Barrel-shaped                                                                                 |
| Falcate            | Sickle-shaped                                                                                 |
| Glandular          | Has secretory/excretory function                                                              |
| Hooked             | Bent/incurved apex shaped                                                                    |
| Head               | Has an enlarged terminal portion                                                              |
| Jointed            | Presence of apparent articulation                                                             |
| Mamilla            | Nipple-shaped projection                                                                       |
| Multiradiate       | Multi-rayed                                                                                   |
| Multitangulate     | Rays at many angles                                                                           |
| Muticus            | No pointed tip                                                                                |
| Neck               | Middle cell of a uniseriate glandular hair                                                    |
| Non-glandular      | Without secretory/excretory function                                                           |
| Osteolate          | Thighbone shaped with many cells having swollen ends                                           |
| Ovoid              | Egg-shaped with attachment at larger end                                                       |
| Pedestal           | Raised base to which hairs are attached                                                       |
| Porrect-geminate   | Similar to porrect stellate but consisting of two whorls of rays                              |
| Term               | Description                                      |
|--------------------|--------------------------------------------------|
| one over the other | Resemblance with porrect rays of cacti with multiple horizontal rays and a central ray |
| Porrect-stellate   | Resemblance with porrect rays of cacti with multiple horizontal rays and a central ray |
| Pulvinate          | Swollen base                                     |
| Pustulated         | Blistered surface                                |
| Quadricellular     | Four cells                                       |
| Sessile            | Without stalks (for glands)                      |
| Setiform           | Bristle-shaped                                   |
| Simple             | Unbranched                                       |
| Smooth             | Without any surface irregularities               |
| Stalk              | Supporting part of a hair                        |
| Stellate           | Star-shaped                                      |
| Subsidiary cells   | Neighboring base cells                           |
| Subulate           | Awl-shaped                                       |
| Tufted/penicillate | Branched from a base                             |
| Uniseriate         | Single rows/columns of cells                     |
| Verrucate          | Warty shaped                                     |
Table 2. Detailed morphological characterization (Major shape and additional features) of trichomes on adaxial leaf surface along with pictorial representation, density, and dimensions of trichomes in fourteen *Solanum* species in the study (‘very rare’ in density table indicate the absence of certain trichome types while counting trichomes at 60X, however the trichome was present very rarely in very few images which were not at 60X magnification; Blank values in density and dimensions table indicate that the density of that particular trichome types has already been included in the density of a broader trichome type, such as density of all glandular trichomes, or density of simple non-glandular trichome types or density of all stellate trichome types as one number rather than individual density of all sub-types of these trichomes; ‘no data’ in dimension table indicate the lack of dimensions of that trichome; Asterisks in dimensions table indicate the multiplication sign showing the length of trichome multiplied by width of gland of trichome, in case of glandular trichomes).

| Species    | Serial number | Trichome types (Major shape) | Additional features | Line art of morphology | Density (Average±Standard error; trichome number/mm² leaf area) | Dimensions (in µm; Length of non-glandular trichome; Length*width of gland in case of glandular trichome; Average±Standard error) |
|------------|---------------|------------------------------|---------------------|------------------------|-----------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------|
|            |               |                              |                     |                        |                                                                 |                                                                                                                                |

Downloaded from https://academic.oup.com/aobpla/advance-article/doi/10.1093/aobpla/plab071/6417072 by guest on 05 November 2021
1. subulate subulate basilatus non-glandular trichome with distinct subsidiary cells

2. multitangu late multitiradiate stellate non-glandular hair with subulate rays (2-6 in number) with the presence of pedestal

3. bifurcated non-glandular hair with subulate rays with the presence of pedestal

uniseri ate- P)

|   |   |   |   |
|---|---|---|---|
|   | S | 4.73±1.02 | 153.71±15.67 |
|   | C | 1.06±0.73 | 185.08±7.29 |

Downloaded from https://academic.oup.com/aobpla/advance-article/doi/10.1093/aobpla/plab071/6417072 by guest on 05 November 2021
4. **glandular** hair with small globular head on the top  
   - **S**  
   - Size: $4.88 \pm 1.40$  
   - Atrichous: $65.25 \pm 5.58 \times 25.08 \pm 1.88$

5. Forest bitterberry (**Solanum anguivi**)  
   - **glandular** hair with single stalk and neck cell and large quadricellular globular head  
   - **S**  
   - Size: $3.50 \pm 0.60$  
   - Atrichous: $59.58 \pm 3.13 \times 27.11 \pm 1.69$

6. **subulate** basilatus **non-glandular** hair  
   - **S**  
   - Size: $2.40 \pm 0.90$  
   - Atrichous: $93.14 \pm 9$

7. **bifurcated** basilatus **non-glandular** hair with subulate rays (one shorter than the other)  
   - **C**  
   - Size: see #8  
   - Atrichous: see #8
8. Porrect-stellate multi-radiate non-glandular hair with subulate rays (3-10 in number) with long central ray and pedestal

C 5.92±0.96 133.01±5.39 (#7 + #8) (#7 and #8)

((Lance-leaved nightshade
Solanum lancefolium))

9. Porrect-stellate multi-radiate non-glandular hair with subulate rays (2-7 in number) and short central ray

C 0.72±0.14 193.75±10.22 (#9 + #10) (#9 and #10)

10. Porrect-stellate multi-radiate non-glandular hair with subulate rays (2-7 in number) and long central ray

C see #9 see #9
11. **glandular hair** with single stalk and neck cell and large doliform globular head

12. **glandular hair** with large globular head

13. **attenuate glandular hair** with small glandular tip

14. **glandular hair** with large globular head

15. **acuminate glandular hair** with bicellular stalk and small glandular tip

---

Garden tomato (*Solanum lycopersicum*)

|    |    |     |    |
|----|----|-----|----|
| 11 | S  | see #12 | see #12 |
| 12 | S  | 1.45±0.21 (#11 + #12) | 66.46±7.19*28.64±2.67 (#11 and #12) |
| 13 | S  | 0.55±0.15 | 63.55±6.35*9.85±1.25 |
| 14 | S  | 0.44±0.11 | 58.74±3.18*25.38±2.23 |
| 15 | S  | 0.05±0.05 | 327±93.30 |
16. hooked non-glandular hair

\[
S = 0.97 \pm 0.22, \quad 218.21 \pm 18.24
\]

17. attenuate non-glandular hair with multicellular jointed stalk and multicellular base

\[
S = 0.17 \pm 0.05, \quad 327.33 \pm 93.30
\]

18. attenuate non-glandular hair with jointed multicellular stalk

\[
S = 0.12 \pm 0.06, \quad 436.33 \pm 44.63
\]

19. African eggplant (Solanum macrocarpon) attenuate basilatus glandular hair with small glandular tip

very rare no data
20. subulate non-glandular hair with pulvinate base and a pedestal

21. glandular hair with multicellular large globular head and single stalk cell

Huckleberry
(Solanum melanocerasum)

22. subulate non-glandular hair with multicellular jointed stalk, multicellular ar base, and distinct subsidiary cells

23. subulate glandular hair with multicellular jointed stalk, multicellular ar base, distinct subsidiary cells and

|   | Description                                                                 | P    | S    |
|---|------------------------------------------------------------------------------|------|------|
| 20| subulate non-glandular                                                       | 0.22±0.09 | 288 |
| 21| glandular                                                                   | 1.88±0.36 | 72.05±6.99*24.23±1.55 |
| 22| subulate non-glandular                                                       | 0.23±0.07 | 421.42±43.01 |
| 23| subulate glandular                                                           | 0.02±0.02 | 461.8±26.03*20.68±3.67 |
24. *glandular* hair with large globular head, single stalk cell and no neck cell

Potato tree (*Solanum grandiflorum*)

25. subulate non-*glandular* hair with multiseriate base and tall pedestal

26. *glandular* hair with large globular head and single stalk cell

27. ovoid sessile non-*glandular* hair

28. attenuate basilatus *glandular* hair with small glandular tip

| Type          | Count | Size        | Measurement         |
|---------------|-------|-------------|---------------------|
| *glandular*   |       | 0.77±0.18   | 78.81±3.11*32.37±1.49 |
| Potato tree   |       | 0.47±0.47   | 583.83±121.39       |
| *glandular*   |       | 14.57±6.48  | 80.55±12.72*32.21±4.34 |
| *ovoid*       |       | 0.19±0.19   | no data             |
| *attenuate*   |       | 0.19±0.19   | 256*25.2            |
29. **setiform**
   - non-glandular hair with multicellular base and a pedestal
   - Eggplant (*Solanum melongena*)
   - \( S \) 0.94±0.75 549.25±96.85

30. **porrect-geminate stellate multiradiate non-glandular hair with subulate rays (2-16 in number) and short central ray**
   - C 8.34±1.26 168±7.46

31. **subulate non-glandular hair with pulvinate base and pedestal**
   - S 0.19±0.19 64.4±14

32. **glandular hair with large globular head**
   - S 0.41±0.20 47.3±0.9*13.1±0.85
| Species                        | Type Description                                                                 | S   | Value 1   | Value 2   | Value 3   |
|-------------------------------|-----------------------------------------------------------------------------------|-----|-----------|-----------|-----------|
| *Solanum taeniotrichum*       | subulate glandular hair with multicellular jointed stalk and small glandular tip | S   | 7.20±0.76 | 385±48.1±2| 3.8       |
|                               |                                                                                   |     |           |           |           |
| *Pepino lloron*               | glandular hair with small globular head                                           | S   | 0.44±0.35 | no data   |           |
| *(Solanum caripense)*         |                                                                                   |     |           |           |           |
|                               | glandular hair with small globular head                                           | S   | 0.75±0.11 | 52.3±32.2 | ±12.5     |
|                               |                                                                                   |     |           |           |           |
|                               | subulate basilatus non-glandular hair with multicellular jointed stalk and multi  | S   | 0.88±0.19 | 431.44±73.48 |           |
|                               | cellular base                                                                      |     |           |           |           |
|                               |                                                                                   |     |           |           |           |
|                               | subulate non-glandular hair with multicellular jointed stalk                       | S   | 0.12±0.06 | 287±107   |           |
Easter white eggplant (*Solanum ovigerum*)

38. *glandular* hair with small globular head

39. *porrect-stellate multi-radiate non-glandular* hair with subulate rays (3-7 in number) and with long central ray

40. *subulate non-glandular* hair with pulvinate base

*Solanum asperolanatum*

41. *glandular* hair with large globular head, single stalk cell and no neck cell
42. ovoid sessile non-glandular hair

43. tufted/penisicillate non-glandular hair with multiseriate long pedestal

Porcupine tomato (*Solanum pyracanthos*)

44. glandular hair with small globular head

45. porrect-stellate multi-radiate non-glandular hair with subulate rays (2-12 in number) and short central ray

46. subulate non-glandular hair with multicellular stalk and base cells
Bittersweet nightshade (Solanum dulcamara) 

47. Subulate basilatus non-glandular hair with multicellular stalk 

|   |   |   |   |   |
|---|---|---|---|---|
| S |   |   |   |   |
| 0.52±0.10 | 158±14 |

48. Glandular hair with small globular head 

|   |   |   |   |   |
|---|---|---|---|---|
| S |   |   |   |   |
| 0.19±0.11 | no data |
Table 3. Detailed morphological characterization (Major shape and additional features) of trichomes on adaxial leaf surface along with pictorial representation, density, and dimensions of trichomes in fourteen *Solanum* species in the study (‘very rare’ in density table indicate the absence of certain trichome types while counting trichomes at 60X, however the trichome was present very rarely in very few images which were not at 60X magnification; Blank values in density and dimensions table indicate that the density of that particular trichome types has already been included in the density of a broader trichome type, such as density of all glandular trichomes, or density of simple non-glandular trichome types or density of all stellate trichome types as one number rather than individual density of all sub-types of these trichomes; ‘no data’ in dimension table indicate the lack of dimensions of that trichome; Asterisks in dimensions table indicate the multiplication sign showing the length of trichome multiplied by width of gland of trichome, in case of glandular trichomes).

| Species  | Serial number | Trichome types  | Additional features | Line art of morphology | Density (Average±Standard error; trichome number/mm² leaf area) | Dimensions (in µm; Length in case of non-glandular trichome; Length*width of gland in case of glandular trichome; Average±Standard error) |
|----------|---------------|-----------------|--------------------|------------------------|-----------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Simple smooth uniseriate-S; Compound smooth uniseriate-C; Simple pustulate uniser | | | | | | |

Downloaded from https://academic.oup.com/aobpla/advance-article/doi/10.1093/aobpla/plab071/6417072 by guest on 05 November 2021
Ethiopian eggplant
(*Solanum aethiopicum*)

1. attenuate glandular hair with small glandular tip
   
   ![Illustration](image1)
   
   $$S$$
   
   $0.86 \pm 0.86$
   
   $101.24 \pm 20.42^*$$
   
   $13.66 \pm 1.74$

2. glandular hair with large globular head on the top
   
   ![Illustration](image2)
   
   $$S$$
   
   $0.52 \pm 0.20$
   
   $52.06 \pm 3.83^*$$
   
   $17.11 \pm 0.82$

3. subulate hooked non-glandular hair with pulvinate base
   
   ![Illustration](image3)
   
   $$S$$
   
   $0.82 \pm 0.79$
   
   $88.55 \pm 10.79$

4. porrect-stellate multiradiate non-glandular hair with subulate rays (5-10 in number) and short central ray
   
   ![Illustration](image4)
   
   $$C$$
   
   $15.26 \pm 4.60$
   
   $241.82 \pm 6.36$

5. glandular hair with single stalk and neck cell and quadricellular globular
   
   ![Illustration](image5)
   
   $$S$$
   
   $0.72 \pm 0.21$
   
   $46.6 \pm 5.1^*$$
   
   $23.35 \pm 0.25$
6. subulate non-glandular hair with pulvinate base

7. porrect-stellate multi-radiate non-glandular hair with subulate rays (5-10 in number) and short central ray and pedestal

8. porrect-stellate multi-radiate non-glandular hair with subulate rays (5-10 in number) with long central ray and pedestal

9. bifurcated basilatus non-glandular hair with subulate rays (one shorter than
Lance-leaved nightshade (Solanum lanceifolium)

10. attenuate glandular hair with small globular tip
   S
   see #12
   no data

11. verrucate non-glandular hair
   S
   see #12
   no data

12. glandular hair with large globular head
   S
   6.04±0.35
   40.88±1.22*23.
   (#10, #11 and #12)
   79±1.23

13. porrect-stellate multi-radiate non-glandular hair with subulate rays (2-6 in number) and with short central ray
   C
   3.38±0.63
   156.28±11.76
   (#13+ #14)
   (#13 and #14)

14. bifurcated basilatus non-glandular hair with subulate rays (one arm reduced)
   S
   see #13
   see #13
|   | Description                                                                 | Letter | S      | #15    | #16    | #17    |
|---|------------------------------------------------------------------------------|--------|--------|--------|--------|--------|
| 15 | *glandular* hair with large quadricellular globular head and single stalk cell | S      | 0.94±0.19 | 42.25±4.69*23. (15 + 16) | 32*3.07 |
| 16 | *glandular* hair with large quadricellular globular head and single stalk cell | S      | see #15  | 158±21.38*66. 57±3.45 |
| 17 | hooked subulate *glandular* hair with multicellular jointed stalk and small glandular tip | S      | 0.30±0.22 | 378.5±29.23*3. 50 |
| 18 | hooked subulate *non-glandular* hair | S      | 5.4±1.03 | 204.34±16.40 |
| 19 | attenuate *non-glandular* hair with jointed multicellular stalk | S      | 0.35±0.10 | 884.38±84.30 |
| No. | Description                                                                 | Type | Abundance     | Measurement                        |
|-----|-----------------------------------------------------------------------------|------|---------------|------------------------------------|
| 20  | attenuate basilatus glandular hair with small glandular tip                 | S    | very rare     | 57.45±8.14*7.5 5±0.91              |
| 21  | mamilla non-glandular hair                                                  | S    | very rare     | no data                            |
| 22  | glandular hair with large globular head and single stalk cell               | S    | 3.03±0.58     | 72.83±6.79*31.69±1.38              |
| 23  | subulate non-glandular hair with pulvinate base and a pedestal              | P    | 0.02±0.02     | 301.66±21.6                       |
| 24  | crescent non-glandular hair with multicellular jointed stalk                | S    | 0.18±0.05     | 312±38.83                         |

Gboma (Solanum macrocarpon)

Huckleberry (Solanum melanocerasum)
25. **subulate glandular** hair with multicellular jointed stalk and small glandular tip

![Image](323x628)

| S | very rare | 337.5±39.63*2 |
|---|-----------|---------------|
|   |           | 4.85±2.7      |

26. **glandular** hair with large globular head, single stalk cell and no neck cell

![Image](268x542)

| S | 0.81±0.12 | 73.46±4.52*35.09±2.16 |
|---|-----------|------------------------|

27. **subulate non-glandular** hair with multiseriate base and tall pedestal

**Potato tree (Solanum grandiflorum)**

![Image](320x381)

| S | very rare | 1194±395.5 |
|---|-----------|------------|

28. **glandular** hair with large globular head and single stalk cell

![Image](295x278)

| S | 10.15±2.25 (#28 + #29 + #31) | 283.2±180.46* ( #28, #29 and #31) |
|---|-----------------------------|----------------------------------|
|   | 37.9±7.26 ( #28, #29 and #31) |

29. **ovoid sessile non-glandular** hair

![Image](316x171)

| S | see #28 | see #28 |
|---|---------|---------|

30. porrect-stellate multiradiate cruciate non-glandular hair with subulate rays (4 in number) and short central ray

31. glandular hair with large globular head, single stalk cell and no neck cell

32. falcate non-glandular hair with pulvinate base

Eggplant (Solanum melongena)

33. hooked subulate non-glandular hair with a pulvinate base
34. porrect-geminate stellate multi-radiate *non-glandular* hair with subulate rays (2-16 in number) and short central ray

35. glandular hair with large globular head

*Solanum taeniotricum*

36. subulate basilatus glandular hair with multicellular jointed stalk and small glandular tip

37. subulate basilatus glandular hair with multicellular jointed stalk, multicellular base and small glandular tip

---

34. C

10.12±1.96

211±8.9

---

35. S

1.48±0.33

44.3±3.86*16.9 ±0.94

---

36. S

8.21±0.76

344±62*21.8±2

(#36+ #37) .33 (#36 and #37)

---

37. S

see #36

see #36
| No. | Hair Type                          | Common Name | Description                                                                 | Length (μm) | Width (μm) |
|-----|-----------------------------------|-------------|------------------------------------------------------------------------------|-------------|------------|
| 38. | glandular hair with small globular head |            |                                                                              | 0.44±0.27   | 51.9±2.55*24.4±8.83 |
| 39. | subulate non-glandular hair        |            |                                                                              | 0.06±0.06   | 1010       |
| 40. | hooked subulate non-glandular hair | Pepino lloron (Solanum caripense) | Very rare, no data                                                           |             |            |
| 41. | subulate non-glandular hair        |            |                                                                              | 0.66±0.11   | 318±58.5   |
| 42. | glandular hair with large quadricellular globular head | Easter white eggplant (Solanum ovigerum) |                                                                              | 0.66±0.22   | 53±8*22.9±4.65 |
| 43. | subulate non-glandular hair        |            |                                                                              | 0.23±0.12   | 63.7±6.35  |
44. falcate non-glandular hair

45. porrect-stellate multi-radiate non-glandular hair with subulate rays (2-12 in number) and with short central ray and pedestal

46. glandular hair with small globular head

47. ovoid sessile non-glandular hair

48. glandular hair with large globular head, single stalk cell and no neck cell
49. **porrect-stellate multi-radiate**
   **non-glandular** hair with subulate rays (4-5 in number) and with short central ray

![Image](image1)

C | 0.12±0.12 | 127±20.4

50. **non-glandular** hair with attenuate non-glandular branches and one gland-tipped branch

![Image](image2)

C | 0.06±0.06 | 201*39.4

51. **Porcupine tomato** *(Solanum pyracanthos)*

**porrect-stellate multi-radiate**
**non-glandular** hair with subulate rays (2-12 in number) and with short central ray and pedestal

![Image](image3)

C | 7.08±1.50 | 265±8.5 (#51 and #53)

### Footnotes:
*(#51+ #53)*
52. *glandular* hair with large globular head

S

0.53±0.24  55±5.2*30.1±3.29

53. *bifurcated non-glandular* hair with subulate rays with pulvinate multicellular base

C

see #51

see #51

54. *hooked non-glandular* hair

S

0.25±0.14  126±6.5
|   |   |   |   |   |   |
|---|---|---|---|---|---|
|   | Bittersweet nightshade (*Solanum dulcamara*) | 55. | glandular hair with small globular head | S | 3.2±0.33 86.7±5.25*28±1.65 |
| 56. | attenuate basilatus glandular hair with small glandular tip | S | 0.04±0.04 no data |
| 57. | osteolate non-glandular hair with multicellular stalk | P | 0.07±0.05 no data |
Figure 4

$p \leq 0.0001$
Figure 5

\[ p \leq 0.0001 \]
Figure 6

$p \leq 0.0001$
