The multi-layered protective cuticle of Collembola: a chemical analysis

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Collembola, also known as springtails, are soil-dwelling arthropods that typically respire through the cuticle. To avoid suffocating in wet conditions, Collembola have evolved a complex, hierarchically nanostructured, cuticle surface that repels water with remarkable efficiency. In order to gain a more profound understanding of the cuticle characteristics, the chemical composition and architecture of the cuticle of Tetrodontophora bielanensis was studied. A stepwise removal of the different cuticle layers enabled controlled access to each layer that could be analysed separately by chemical spectrometry methods and electron microscopy. We found a cuticle composition that consisted of three characteristic layers, namely, a chitin-rich lamellar base structure overlaid by protein-rich nanostructures, and a lipid-rich envelope. The specific functions, composition and biological characteristics of each cuticle layer are discussed with respect to adaptations of Collembola to their soil habitat. It was found that the non-wetting characteristics base on a rather typical arthropod cuticle surface chemistry which confirms the decisive role of the cuticle topography.

1. Introduction

Liquid-repellent, non-fouling and self-cleaning characteristics of natural surfaces receive particular attention in biomimicry research. Unravelling the chemical and morphological origin of the surface properties is crucial to reproduce and translate these characteristics into engineered surfaces [1,2]. The non-wetting and self-cleaning properties of plant surfaces were intensively studied [3–5] over the last 20 years. Plant surfaces are typically decorated by wax crystals, which are rather fragile but regenerate after mechanical destruction [6,7]. Mimicking the needle-like or platelet-shaped crystal structures into artificial surface coatings was widely studied [8–10]; however, the inherent fragility of such micro- and nanostructures limits their durability [11]. Recent studies on the functional morphology of the cuticles of Collembola (springtails) revealed surfaces with higher stability against wear and friction, and outstanding resistance against wetting, even with low-surface-tension liquids [12,13]. While structural features of Collembola cuticles were investigated in detail [14], the impact of the molecular composition on the cuticle characteristics required further elucidation.

Representing more than 8000 species [15], Collembola are the most abundant and widespread arthropods and an important part of the community of soil-dwelling animals [16]. To enable cutaneous respiration [17,18], the cuticle surface needs to be kept clean and dry in the presence of soil microorganisms and water that is often contaminated by surfactants originating from decaying organic matter [19]. Consequently, the collembolan cuticle is adapted to the soil habitat by a robust and repellent surface consisting of nanoscopic, comb-like structures (figure 1a) [14,20]. Polymer replication methods were applied to demonstrate that the particular structures enable a robust non-wetting state, even with low-surface-tension liquids, and, thus, protect the animals against suffocation, even when immersed [21,22].
While the unique surface morphology of collembolan cuticle was recently intensively studied, its chemical composition was not yet comprehensively analysed. A thorough analysis of both structure and composition of the cuticle is, however, needed for reconstituting the non-wetting and non-fouling surface characteristics in the development of bioinspired materials. Earlier studies found the collembolan cuticle to exhibit a layered structure commonly observed for arthropods [23]. Based upon electron micrographs, the authors classified several layers (from inside to outside): an endocuticle, an exocuticle and an epicuticle. The epicuticle was further divided into an internal, a cuticulin, a wax and a cement layer [24–26]. The waxy-like surface coating was suggested to support hydrophobicity under wet conditions and to prevent desiccation under arid conditions [27,28].

As a detailed chemical analysis of each cuticle layer was not yet available, we chose to investigate the cuticle of *Tetrodontophora bielanensis* with respect to the composition of each layer. Morphological characteristics were analysed using electron microscopy, and chemical analyses were performed by time of flight secondary ion mass spectrometry (TOF-SIMS), thin layer chromatography (TLC) and gas chromatography mass spectrometry (GC-MS). Additionally, hydrolysed cuticle samples were analysed by high-performance liquid chromatography (HPLC) and electrospray ionization mass spectrometry (ESI-MS). The combined results of these analyses allowed for establishing a detailed compositional model of the Collembola cuticle.

## 2. Results

The hierarchical structure of the cuticle of *T. bielanensis* is displayed in figure 1a. The scanning electron microscopy (SEM) images show papillose microstructures (designated as secondary granules) covered by a rhombic comb-like mesh exhibiting nanoscopic tubercles (primary granules). (b) TEM sections of the cuticle, revealing the layered structure to consist of (I) the lamellar procuticle (pro), covered by (II) the epicuticle (epi) and (III) a thin envelope (env) as the topmost layer. (Online version in colour.)

![Figure 1.](http://rsif.royalsocietypublishing.org/fig1a.png) **Figure 1.** (a) SEM studies of the cuticular morphology of *T. bielanensis*. SEM images showing papillose microstructures (secondary granules), covered by a rhombic comb-like mesh exhibiting nanoscopic tubercles (primary granules). (b) TEM sections of the cuticle, revealing the layered structure to consist of (I) the lamellar procuticle (pro), covered by (II) the epicuticle (epi) and (III) a thin envelope (env) as the topmost layer. (Online version in colour.)
of the measurements (figure 3b) showed lipids, such as fatty acids ($m/z = 255; 279; 281; 283$) and higher molecular weight aliphatic compounds ($m/z = 489; 561; 575; 603; 631$). Hexane/dichloromethane extracts of the examined animal sample were applied to glass surfaces and also analysed by TOF-SIMS. The resulting data (electronic supplementary material, figure S1) were similar to those of the untreated cuticle surface, demonstrating the relevance of the extracted sample. As TOF-SIMS only allows for the detection of fragments, TLC was also performed to obtain further insights into the cuticle lipid composition using a standard lipid mixture, were detected (electronic supplementary material, figure S2). The intensity of the TLC spots revealed that fatty acids, esters and terpenes dominated the extracts of both sources. The extract of the complete animals contained a higher amount of steroids, fatty acids and esters. This might be due to lower amounts of lipids retained in the moulted cuticle extract.

GC-MS analysis was performed to identify the lipids as detected by TLC (electronic supplementary material, figures S3–S9). For the moulted cuticle extract, steroids, fatty acids and one terpene were detected. The detected steroids were identified as cholesterol and desmosterol, the fatty acids as palmitic and stearic acid and the terpene as lycopane (table 1). In case of the complete animal extract, steroids, fatty acids, esters and terpenes could be detected, which was in accordance with the TLC measurements. The steroids again were identified as cholesterol and desmosterol, in addition the fatty acids palmitic acid, stearic acid, oleic acid and linoleic acid. The esters were identified as linolenyl myristate and linoleyl palmitate, which are common wax esters. The terpenes were in line with the results from [29] identified as lycopan, lycopene and lycopadien (table 1). The extract of moulted cuticles contained fewer substances, which might be due to lower amounts of lipids retained.

Extraction of the outermost lipid layer of freshly shed cuticles left the epicuticle and the procuticle behind for further analysis. The epicuticle was subsequently dissolved in a 2.5 M KOH solution for hydrolysis of proteins (figure 4). Figure 4b depicts the destruction of the comb-like structure after 0.5 h incubation time. After 24 h, the protein-rich epicuticle was completely dissolved. In order to verify the proteinaceous nature of the nanoscopic surface features, the solution was further analysed by HPLC. Table 2 summarizes the amino acid composition and the particular concentrations of the moulted cuticle extract. Glycine represents more than 50% of the detected amino acids. Furthermore, the sample showed a high amount of tyrosine and serine residues. For comparison, the amino acid compositions of structural proteins, such as fibroin (Bombyx mori), collagen (Periplaneta americana) and resilin (Schistocerca gregaria) [30–32] are included in table 2. A common feature of structural proteins, glycine was the dominating amino acid in all proteins. Therefore, the epicuticular structures of the collembolan cuticle are concluded to also consist of structural proteins. Additionally, decellularized (TritonX treated) animals were analysed by amino acid analysis. The results clearly differed (the glycine amount was reduced to 35%, the amounts of alanine and serine increased) determining the decellularization of animals by TritonX to be unsuitable in identifying cuticle proteins.

For protein identification, the epicuticular structures were exposed to 8 M urea, 6 M guanidine hydrochloride, guanidinium thiocyanate (TriFast) and 20% sodium dodecyl sulfate (SDS) for at least one week at 60°C. However, these treatments did not alter the epicuticular pattern (figure 4a), indicating a robust protein structure. Interestingly, the common covalent cross-links, such as disulfide bonds or dityrosine amino acids residues, were not observed in the samples by UV–Vis spectroscopy (electronic supplementary material, figure S10), suggesting a non-covalent association.

After KOH hydrolysis, the remaining procuticle skeletons (figure 4c) were further hydrolysed by 6 M HCl and analysed by ESI-MS (electronic supplementary material, figure S11) that confirmed the presence of chitin. Additionally, SEM (figure 4c) and Cryo-SEM (figure 5) studies revealed a regular distribution of pore channels within the chitin skeleton. The pore channels showed different sizes and morphologies. Most channels were small with a diameter of about 200 nm and vertically interpenetrated the entire procuticle. Some pore channels appeared to be thicker (diameter of about 2 μm) but branched into smaller channels close to the epicuticle.

In summary, the chemical analyses revealed a multi-layered cuticle structure, consisting of a topmost lipid-rich envelope, a protein-rich epicuticle and a chitin-rich procuticle (figure 6). The homogeneous lipid-rich layer, covering the entire surface, encompassed aliphatic hydrocarbons such as steroids, triglycerides, fatty acids, wax esters and terpenes. The characteristic morphological surface features of the cuticle consisted of a protein-rich layer, dominated by a glycerine-rich structural protein. The inner cuticle layer consisted of a lamellar chitin skeleton that was interpenetrated by numerous pore channels.
3. Discussion

Our presented study revealed that the cuticle of *T. bielanensis*, showing the characteristic collembolan ornamentations, consisted of a cross-sectional layer structure known from arthropods, namely, a chitin-rich procuticle, a chitin-free but protein-rich epicuticle and an envelope composed of a lipid mixture. The reported results confirm earlier investigations on basic features of the collembolan cuticle [24–26] and on cuticle differentiation [33].

The chitin-rich procuticle forms the basis of the exoskeleton and provides rigidity and mechanical protection to the body [23]. The numerous nanostructured pore channels enable material transport [25,34]. Moreover, these channels may indicate a preliminary stage of a tracheal system that particularly...
allows *T. bielanensis*, the European giant springtail, to respire by gaseous diffusion through the cuticle surface. Nevertheless, the channels do not interpenetrate the entire cuticle, which can be considered a physiological advantage in preventing high transpiration rates and penetration of microorganisms.

The protein-rich epicuticle covers the chitin layer. The epicuticle contains the distinctive surface features of Collembola in displaying a comb-like mesh covering the entire body. Cavities inside the epicuticular mesh allow for respiration under wet conditions and preventing suffocation in temporarily rain-flooded habitats. In a recent study, the durability of these epicuticular structures was demonstrated by sand blast experiments [12]. We found that proteins with high amounts of glycine, tyrosine and serine formed the epicuticular structures. The composition of the amino acid mixture resembled that of known structural proteins such as fibroin, collagen or resilin [30–32], which can often combine stiffness and toughness [35]. Thus, it can be reasonably assumed that the durability of patterned epicuticle of *T. bielanensis* to withstand wear in soil habitats results from epicuticular protein structures.

As the outermost layer, a lipid mixture of fatty acids, wax esters and terpenes envelopes the epicuticular structures, forming the first protective barrier of the animal. Wax esters support the non-wetting properties due to their hydrophobic characteristics [6]. The thin lipid layer enables gas exchange, but hardly protects against transpiration and desiccation [36]. Therefore, Collembola, with some exceptions, depend on humid surroundings as given in soil habitats. The composition of the amino acid mixture resembled that of known structural proteins such as fibroin, collagen or resilin [30–32], which can often combine stiffness and toughness [35]. Thus, it can be reasonably assumed that the durability of patterned epicuticle of *T. bielanensis* to withstand wear in soil habitats results from epicuticular protein structures.

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The unique hierarchical topography of the layered cuticle reflects the adaptation of Collembola to their soil habitats, enabling the permeability to gases, minimizing liquid wettability and fouling by microorganisms and resisting abrasion. Those remarkable cuticle characteristics were found to be achieved by a rather typical arthropod cuticle surface chemistry. Thus, our data highlight the importance of topographical features of the cuticle for its functional characteristics.

### 4. Material and methods

#### 4.1. Animals

*Tetrodontophora bielanensis* species were collected in the wooded mountains of Saxony near Dresden, southeastern Germany. The animals were kept as laboratory colonies in large Petri dishes using soil, litter, decaying wood and moss from their original habitat as substrate and food source. The substrate was wetted regularly to maintain humid conditions. *Tetrodontophora bielanensis* species survived up to six months in captivity, preferably at 12–14°C, which was maintained by storage in wine cooler board. During this time, animals were collected regularly for experiments. Additionally, moulted cuticles of the animals were collected from the Petri dishes and used as cuticle samples. Moultng occurred at three to four week intervals.

#### 4.2. Electron imaging

SEM studies were performed using a XL30 ESEM-FEG microscope (Philips) in the usual HighVac mode at voltages of 5 kV. Cryo-SEM studies were performed by a Zeiss Supra 40VP with an Emitech K1250× cryo transfer device in the usual HighVac mode at voltages of 5 kV. The animals were prepared by freezing.
and subsequent air-drying without any fixation. Samples were coated with a 15 nm gold layer (BALZERS SCD 050 Sputter Coater) to avoid surface charging effects. TEM studies were carried out using an EM 912 Omega (Carl Zeiss SMT). The samples were fixed, stained and subsequently sliced into ultrathin sections as described in detail by Helbig et al. [12].

4.3. Sample preparation for chemical analysis

4.3.1. Lipid layer analysis

Moulted cuticles (30–40 specimens) were collected, washed in distilled water, air-dried and subsequently extracted in a hexane/dichloromethane (1 : 1) solvent mixture for 30 min at 60°C. The extract was concentrated by solvent evaporation. Complete animals (20 specimens) were shock frozen, air-dried for 24 h at room temperature (RT) and extracted under the same conditions. The hexane/dichloromethane extracts were used for TLC and GC-MS analysis.

4.3.2. Amino acid analysis

Extracted moulted cuticles were treated directly with 2.5 M KOH solution for 0.5–24 h at RT [42]. Extracted animal samples were decellularized to remove cellular proteins by placing animals in 0.5% TritonX (Sigma-Aldrich) solution for one week. Afterwards, the decellularized animals were washed in distilled water several times to remove TritonX. The decellularized animals were treated with 2.5 M KOH solution for 0.5–24 h at RT. The KOH solution with the hydrolysed cuticle proteins were used for HPLC analysis.

4.3.3. Chitin analysis

Remaining cuticle samples treated with 2.5 M KOH were further hydrolysed in 6 M HCl for 4 days at 80°C. The obtained samples were filtered with 0.4-mm filter and freeze dried. The solid remnant was dissolved in methanol for ESI-MS analysis.

4.4. Time of flight secondary ion mass spectrometry

For TOF-SIMS analysis, animals were shock frozen and air-dried for 24 h at RT. For extract analysis, animal samples were prepared as described in §4.3.1. The extract was coated on glass surfaces and analysed. Measurements were performed using an ION TOF TOF-SIMS V instrument equipped with a Bi liquid metal ion gun. Analysis was carried out as described in detail by Nygren et al. [43]. All image analyses were performed within the Ion-Tof Ion image software (v. 3.1, Ion-Tof, GmbH, Münster, Germany).

4.5. Thin layer chromatography

Hexane/dichloromethane extracts of moulted cuticles and complete animals were prepared as described in §4.3.1. A mixture of 30 mg lipids in 2 ml of dichloromethane was used as a lipid standard, containing phosphatidylcholine, cholesterol, glyceryltriol-eate, palmitic acid, stearyl palmitate and cholesteryl palmitate. TLC was performed using TLC sheets with a 0.2 mm silica gel layer (ALUGRAM, Machery-Nagel). Three solvent systems were used stepwise on one TLC plate for lipid separation (adapted from [44]):

(1) methanol : chloroform : acetic acid (50 : 30 : 16),
(2) hexane : diethyl ether : acetic acid (70 : 2 : 0.2) and
(3) hexane.

Figure 4. Separation and analysis of the procuticle and epicuticle. (a) Cuticle treated with 8 M urea, 6 M guanidine hydrochloride, 20% SDS and guanidinium thiocyanate for one week at 60°C without changes in the epicuticular structures. (b) Cuticle samples treated with 2.5 M KOH solution for 0.5 h for dissolving the cuticle structures. The KOH solution was used for HPLC analysis. (c) Further treatment of the cuticle with 2.5 M KOH led to complete dissolution of the epicuticular proteins uncovering the chitin skeleton. (Online version in colour.)

Figure 5. Cryo-SEM image of the cuticle with visible pore channels.
Figure 6. Model of the multi-layered cuticle of Collembola. (Online version in colour.)

Table 2. Detected amino acids by HPLC of the moulted cuticle KOH hydrolysate and from decellularized animals. Amino acid analyses of fibroin (B. mori), collagen (P. americana) and resilin (S. gregaria) are included for comparison.

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4.6. Gas chromatography mass spectrometry
Hexane/dichloromethane extracts of moulted cuticles and complete animals were prepared as described in §4.3.1. Measurements were performed using an Agilent Technologies 6890 N GC System equipped with a 5973 Mass Selective Detector. GC separation was conducted with a temperature programme from 110°C (1 min) to 340°C (30 min) at a rate of 10°C min⁻¹, respectively. One microlitre of the sample was injected for analysis. The mass spectrometer was operated in the electron impact ionization mode with ionization energy of 70 eV. Data evaluation was performed using LIPID MAPS Lipidomics Gateway database.

4.7. High-performance liquid chromatography
KOH hydrolysates of extracted moulted cuticles and extracted animals were prepared as described in §4.3.2 and used for amino acid analysis. HPLC analysis was performed by amino
acid sample derivatization using o-phthalaldehyde for primary amino acids and 9-fluorenylmethyl chloroformate for secondary amino acids [46]. Measurements were performed by analytical HPLC Zorbax Eclipse-AAA column (4.6 x 150 mm, 3.5 μm, Agilent Technologies, USA) for 40 min and a flow rate of 0.8 ml min⁻¹ for the analytical column. A linear gradient of 0.1 M phosphate buffer (pH7.8) and acetonitrile/methanol/water 45/45/10 was used as the mobile phase. A two-pump system (Agilent Technologies 1100 Series, USA) equipped with a UV/Vis detector/spectrophotometer in line with fluorescence detector both having a 1 cm path length cell was used for analysis of the amino acid composition.

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