Amorphous and crystalline calcium carbonate distribution in the tergite cuticle of moulting Porcellio scaber (Isopoda, Crustacea)

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1. Introduction

Crustaceans possess an exoskeleton (the cuticle) of high mechanical strength that provides support of the body and protection against environmental strains including predation and, in terrestrial species, desiccation. The cuticle consists of an organic matrix composed of chitin–protein fibers and a mineral phase. In the course of growing, it is periodically shed off in a process called moulting. This process includes resorption of mineral and organic material from the old cuticle, the synthesis of the new cuticle before the moult, shedding of the old cuticle, and resorption of mineral from the posterior cuticle. During moulting, crustaceans form large CaCO3 reservoirs that are utilized to mineralize the new cuticle (Greenaway, 1985). An interesting mode to retain calcium during moulting occurs in isopods that are unique in moulting first the posterior and then the anterior half of the body with the border between the fourth and fifth thorax segments (corresponding to the fifth and sixth somites of the pereon). This biphasic moulting allows the animals to reduce the loss of CaCO3 by shifting it between the anterior and poster integument. This mode is generally utilized in terrestrial isopod species (Onisicidea) that developed large CaCO3-deposits within the ecdysial space between the new and old cuticle of the first four anterior sternites (Messner, 1965; Steel, 1993; Strus and Compere, 1996; Ziegler and Miller, 1997; Ziegler et al., 2002, 2004; Fabritius and Ziegler, 2003). Deposition and resorption of the deposits is well studied in the common woodlouse Porcellio scaber (Ziegler, 1996, 1997; Hagedorn and Ziegler, 2002; Ziegler et al., 2002, 2004; Fabritius and Ziegler, 2003). In P. scaber the deposits developed about 1 week before the posterior moult. During this premoult stage calcium and carbonate ions are resorbed from the posterior cuticle into the hemolymph and are then transported into the ecdysial space of the anterior sternites where it forms the deposits. When
the sternal deposits are well developed, additional CaCO$_3$ is stored within the anterior cuticle as indicated by a change in color resulting in a whitish appearance of the anterior dorsal part of the body including the anterior thoracic segments (tergites) (Steel, 1993; Strus and Blejec, 2001). After moulting of the posterior, but before the moulting of the anterior half of the body (intra-moult) calcium and carbonate ions from the deposits and the anterior dorsal integument are transported back to the hemolymph and are used to mineralize the new posterior cuticle (Messner, 1965; Steel, 1993; Ziegler et al., 2007). When the deposits are completely resorbed the animal moults the anterior half of the body and the new anterior cuticle is mineralized in the postmoult stage.

Interestingly, the mineral phase of the sternal CaCO$_3$ deposits of *P. scaber* are completely amorphous with a molar Ca:Mg ratio of 30:1 and a Ca:P ratio of 11.5:1 (Ziegler, 1994, 2002; Becker et al., 2003). The amorphous structure allows a quick dissolution of the deposits since ACC is about 10 times more soluble than calcite (Brecevic and Nielson, 1989). Recent studies on the cuticle of a number of crustaceans has shown that depending on species the cuticle consists of varying amounts of amorphous calcium carbonate, calcite and amorphous calcium phosphate (Al-Sawalmih et al., 2009; Becker et al., 2005; Boßelmann et al., 2007; Neues et al., 2007; Hild et al., 2009; Shechter et al., 2008). This suggests the possibility that ACC is preferentially resorbed from the cuticle over calcite as a result of its high solubility. This hypothesis was supported by a recent analysis of the spatial distribution of calcite and ACC in the terrestrial isopods *Armadillidium vulgare* and *Titanethes albus* employing µ-Raman spectroscopic imaging. It revealed that calcite occurs in the apical region only, while the proximal region of the cuticle is mineralized by ACC, also suggesting preferential resorption of ACC (Hild et al., 2008, 2009). However, direct evidence for changes in the ACC/calcite distribution during the moulting cycle of crustaceans is lacking. In particular it is unknown if a significant amount of calcite is resorbed in addition to ACC.

In an attempt to investigate differential resorption of ACC and calcite and spatial aspects of mineral storage within the cuticle, we have investigated the changes in the distribution and relative amount of ACC and calcite within the anterior and posterior tergites, during the moulting cycle. For this we used a combination of X-ray diffraction analysis, atomic absorption spectroscopy, and µ-Raman spectroscopic imaging. The results suggest that an important function of cuticular ACC is to facilitate the resorption of CaCO$_3$ during the moulting cycle.

### 2. Material and methods

#### 2.1. Animals

*P. scaber* Latreille, 1804, were collected from local biotopes, kept in plastic containers filled with soil and bark and fed with fresh potatoes, carrots and dry oak leaves. The animals were examined daily for the appearance of white calciferous sternal deposits that were used to determine the premoult stages. These animals were placed individually into plastic containers with a diameter of 8 cm containing a disc of wet paper towel and fed as described above. Animals with large but incompletely developed sternal deposits were assigned as being in the mid premoult stage (stage 54 according to Ziegler, 1997) and animals with fully developed sternal deposits and whitish coloration of the anterior dorsal cuticle as being in the late premoult stage. Animals that had shed the posterior cuticle and in which the sternal deposits were still large were denoted as animals being in the early intra-moult stage and those with small sternal deposits as being in the late intra-moult stage. Postmoult animals were used just after the anterior moult (early postmoult) and 1 and 2 days after the anterior moult (1d postmoult and 2d postmoult, respectively). In addition we used animals 6–8 days after moulting. Since in *P. scaber* a true intermoult stage is lacking (Ziegler, 1997) these animals were assigned as being in the early premoult stage rather than intermoult stage.

#### 2.2. Sample preparation

Anterior and posterior tergites were obtained by dissecting animals in 100% methanol. Previous studies have shown that ACC from sternal deposits of *P. scaber* is stable in methanol for at least 1 month (Becker et al., 2003). Subcuticular soft tissue was carefully removed from the tergites and the sample washed first in bidistilled water for 2 s to remove salt residues and then in 100% methanol for at least 5 s to remove the water. Then the samples were air dried at room temperature and stored at −20 °C until further use.

For scanning confocal µ-Raman spectroscopy (SCµ-RS) pieces of the tergites were glued onto polymethylmetacrylate (PMMA) holders using superglue. Tergites were cut sagittally using a Reichert Ultracut ultramicrotome and glass knives to obtain a plane surface. This surface was then polished with a diamond knife by successively cutting and advancing the specimen 15 times each by 70, 40 and 20 nm.

#### 2.3. Field emission scanning electron microscopy

For SEM analysis cuticle samples on PMMA holders were cleaved in the sagittal plane and rotary-shadowed with 3 nm platinum (BAF 100, Balzers) at an angle of 45°. SEM micrographs were recorded with a field emission scanning electron microscope (FES-EM) (Hitachi S-5200) at an acceleration voltage of 4 kV.

#### 2.4. X-ray powder diffractometry

High-resolution X-ray powder diffractometry was carried out in transmission geometry at room temperature (ground samples on Kapton foil) at beamline B2 at HASYLAB/DESY, Hamburg, Germany, at a wavelength of $\lambda = 0.47176$ Å (Knapp et al., 2004a,b). All data were converted to Cu K$_{\alpha1}$ radiation wavelength ($\lambda = 1.54056$ Å) for better comparison. For quantitative Rietveld analyzes (QRA), the cuticle was mixed with crystalline quartz (SiO$_2$) in a 5:1 w:w ratio, followed by thorough grinding. This mixture was subjected to X-ray powder diffraction as described above. The obtained diffractograms were analyzed by the Rietveld software FULLPROF Suite 2005 (Rodríguez-Carvajal, 1990) which gave the relative mass fractions of calcite and quartz, the unit cell parameters and the reflex positions. Thereby, the amount of crystalline calcite in the cuticle was determined. The accuracy of this method is discussed in the literature (Gualtieri, 2000). Due to magnesium replacing calcium in the crystal lattice, the peak intensities and the phase scale factors during Rietveld refinement are affected. Furthermore, minor variations in the intensity occurred due to the biological origin of the samples and the resulting preferred orientation. Therefore, we assumed a relative error of about 5%. The amount of magnesium in calcite was determined by analyzing the peak shifts induced by the ionic substitution of calcium by magnesium using the method of Goldsmith (Goldsmith and Graf, 1958). The unit cell parameters $a$ and $c$ are decreasing and the position of the (104) reflex is shifted to larger angles with increasing content of magnesium.

Calcium and magnesium contents were determined by atomic absorption spectroscopy (AAS) in a Unicam 939 instrument after extraction of calcium from the cuticle by suprapure HCl (average from two determinations). Typically the variation between the
two measurements was 0.2 wt.% or less. The content of calcium carbonate was computed from the calcium content as w(CaCO$_3$) = w(Ca) M(CaCO$_3$)/M(Ca). In contrast to our earlier studies on crustacean cuticles (Becker et al., 2005; Boßelmann et al., 2007; Neues et al., 2007), no thermogravimetric data could be recorded due to the small amount of sample. Thus, the amount of ACC was estimated as difference between the total amount of calcium carbonate and the amount of crystalline calcite obtained by Rietveld refinement. The content of amorphous calcium phosphate (ACP) could not be determined due to the small amount of sample and was neglected.

2.5. Raman spectroscopy and imaging

A confocal Raman microscope (WITec, Ulm, Germany) equipped with a NdYag laser (wavelength of 532 nm) for excitation and a Nikon 100× (NA = 0.95) objective were used to locate calcite, ACC and organic matrix within the tergite cuticle by recording single Raman spectra and Raman spectral images. As reference material for the Raman measurements, calcite was precipitated from a solution containing 24 mmol L$^{-1}$ Ca$^{2+}$ and 24 mmol L$^{-1}$ CO$_2$ at pH 8.6. Amorphous calcium carbonate without any organic additives was synthesized by the method described by Faatz et al. (2004). Purified chitin from crab-cuticles (Sigma–Aldrich) and bovine serum albumin (BSA) (VWR International) served as chitin and protein reference materials, respectively. The Raman spectra of the standards were recorded with integration times of 5 s.

Sagittally cut and polished surfaces of the cuticles were prepared and analyzed in single imaging and in the Raman spectral imaging mode. To get high resolution Raman images, the samples were scanned point-by-point and line-by-line, and Raman spectra between 0 and 3750 cm$^{-1}$ were recorded with an integration time of 250 ms at every pixel (Pixel density: 10 spectra at 1 μm). Images were obtained analyzing the integral of specific bands above background employing the WITecProject software (Version 1_86, WITec, Ulm, Germany) (Schmidt et al., 2005). To determine the distribution of calcite, carbonate in total, and organic material we used the peak areas for the lattice vibration at 280 cm$^{-1}$, for CO$_2$ stretching vibration, and that of the band at 2800–3020 cm$^{-1}$, which is specific for CH$_3$ stretching vibration, respectively. All images were color-coded in a way that higher values of the integral sum which correlates to a higher amount of a specific component appear brighter. For the different materials color tables were chosen as following: calcite (red), carbonate (orange), and organic components (green).

The overall Raman spectrum for a tergite cuticle of P. scaber was obtained by averaging spectra collected along a 10 μm wide area across the polished surface of the sample. These spectra were used to calculate the ACC/calcite ratio. The spectra were fitted by a linear combination of normalized ACC ($S_{ACC}$), calcite ($S_{calcite}$) and chitin ($S_{chitin}$) reference spectra.

$$S_{sample} = a \cdot S_{ACC} + b \cdot S_{calcite} + c \cdot S_{chitin}$$

The ratio a/b of the coefficients obtained for $S_{ACC}$ and $S_{calcite}$ respectively, was used to present the ACC/calcite ratio.

To get high resolution Raman line scans, the samples were collected point-by-point across the polished surface of the sample. Raman spectra between 0 and 3750 cm$^{-1}$ were recorded with an integration time of 1 s at every pixel (Pixel density: 10 pixel/μm).

The line scans were obtained in the same way as Raman images by analyzing the integral of specific bands above background employing the WITecProject software. For each moulting stage tergites of 2–4 animals were used. For each tergite we recorded 2–3 line scans at three different locations.

3. Results

3.1. Field emission scanning electron microscopy (FESEM)

The ultrastructure of the mineralized tergite cuticle from intermoult animals was already described in previously (Hild et al., 2008). Here we therefore just show FESEM micrographs of selected moulting stages (Figs. 1 and 2). During late premoult the cuticle of the anterior tergites resembles those in intermoult, except, that just proximal of the unmineralized membranous layer a new layer appears that contains numerous spherules with diameters of about 200 nm that are interconnected by strands of an organic matrix (Fig. 1A and B). The spherules are composed of small granules about 20 nm in size. This layer and considerable amounts of mineral within the anterior cuticle are resorbed before the cuticle is shed (see Raman spectroscopic images below). The tergites of the posterior exuviae are about 6 μm thick in late premoult (Fig. 1C). The exocuticle appears intact. In particular the distal part of the exuviae including an outer smooth layer (Hild et al., 2008) is still fully mineralized (Fig. 1D). Fibrous structures within the proximal regions of the exocuticle and the distal region of the endocuticle indicate partial demineralization (Fig. 1C and E). In the proximal part of the endocuticle the organic and inorganic material is almost completely resorbed leaving only a thin degradation zone of smooth appearance (Fig. 1C).

An example of the posterior cuticle during early intramoult is shown in Fig. 2. The new exocuticle appears already well mineralized (Fig. 2A and B). Higher magnification reveals a granular texture of the smooth layer and more proximal regions of the exocuticle (Fig. 2B).

The endocuticle is not yet fully synthesized. Most regions of the endocuticle that are present are already mineralized to some extent (Fig. 2A and C). However, still fibrous structures are visible and pore canals are devoid of mineral (Fig. 2C). Only an about 0.5 μm thick proximal layer consists of unmineralized fibers of the organic matrix (Fig. 2D).

3.2. The ACC/calcite ratio

A typical X-ray powder diffractogram and the superimposed diffractogram calculated according to the Rietveld refinement are shown for a sample containing 5.66 mg of the tergal cuticle mixed with 0.63 mg quartz (Fig. 3). The diffractograms indicate the presence of calcite within the samples and the virtual absence of crystalline calcium phosphate. The amount of crystalline magnesium calcite determined according to the Rietveld refinement are given for the anterior and posterior tergite cuticle of each moulting stage (Table 1). Small shifts in the peak position are due to magnesium replacing calcium within the calcite crystal lattice. The magnesium concentration within the crystal lattice as determined by the Goldsmith method was between 0.1 and 0.2 wt.%. AAS indicates that the amount of magnesium not in Mg-calcite is about five times higher than that within the calcite lattice. In all moulting stages, AAS of anterior and posterior cuticle samples yielded calcium concentrations that exceed the amount of calcium expected from the amount of calcite within the samples (Table 1), indicating that a significant amount of calcium occurs in the form of ACC or amorphous calcium phosphate (ACP). Because of the small sample size (see Section 2) we did not distinguish between ACP and ACC by performing TG analysis. For the sake of simplicity, we assumed that all non-calcite calcium would occur in the form of ACC, although we know that in the early premoult stage the cuticle of the tergites contains 12 wt.% ACP (Becker et al., 2005). Because of changes in the mass of the cuticle between moulting stages due to resorption/
secretion of mineral and organic material only mass-invariant values can be compared. Therefore we calculated virtual ACC/calcite ratios for each moulting stage (Fig. 4). The resorption of mineral from the posterior cuticle is accompanied by a reduction of the ACC/calcite ratio within the posterior tergites by 1.08 units from early premoult to late premoult. Concomitant to this...
decrease the ratio in the anterior tergites increases by 0.74. Until posterior ecdysis the ACC/calcite value within the posterior cuticle decreases further by 0.15 units to the value within the exuviae (Fig. 4A). Between the posterior moulting and the time of collection of the posterior exuviae in early intramoult, some of the mineral within the anterior tergites is already resorbed for

Fig. 2. Field emission scanning electron micrographs of the mineralizing posterior cuticle of *P. scaber* during early intramoult. (A) Overview, showing mineralised exo (ex) and partly mineralized endocuticle (en) containing many pore canals. The well-mineralized exocuticle and its distal smooth layer (sl) have a granular texture (B). (C) Detail of the transition zone between exo and endocuticle. Empty pore canals (pc) and filaments (arrows) of the organic matrix are apparent. (D) Proximal region of the endocuticle. The very proximal organic matrix (om) appears still unmineralized.
mineralization of the new posterior cuticle, resulting in a small decrease in the ACC/calcite value by 0.37 units (Fig. 4B). In late intramoult the mineralising ACC/calcite ratio of the new posterior cuticle increases to a value 1.66 units higher than in the exuviae and even 0.43 units higher than in the early premoult control stage. Surprisingly, the ACC/calcite ratio within the posterior tergites decreases by 0.78 units from late intramoult to early postmoult. The mineralization of the new posterior tergites is accompanied by a decrease of the ACC/calcite ratio in the anterior tergites by 1.4 units from late premoult to the value within the anterior exuviae. During further mineralization of the posterior cuticle the ACC/calcite ratio increases by 1.1 units within 1 day from early postmoult, followed by a decrease of 0.25 units during the next day. One day after the anterior moult the ACC/calcite ratio of the mineralising anterior tergite cuticle is 0.61 units larger than during early premoult and increases further by 0.69 units during the next day.

In both the posterior and anterior tergites the ACC/calcite ratio decreases within 4–6 days from the second day of postmoult to early intramoult by 0.50 and 1.30 units, respectively.

### 3.3. Raman imaging

Fig. 5A and B present integral spectra calculated from the individual spectra obtained during a scan over a sagittal microtome polished surface of tergite cuticle in early premoult and the posterior exuviae, respectively. Comparing these spectra with the reference spectra for synthetic ACC, purified chitin, BSA and synthetic calcite (Fig. 5C–F) reveal that in the exuviae the signal for calcite...
at 280 cm\(^{-1}\) is larger and the broad band between 100 and 300 cm\(^{-1}\) that is typical for presence of ACC, is smaller than in the cuticle from an early premoult animal. The organic matrix consists of chitin and protein.

Micro Raman spectroscopic images of calcite, carbonate in total, and organic material reveal changes in the distribution of these components during the moulting cycle of \textit{P. scaber} (Fig. 6). From these images it is apparent that during mineral resorption in the posterior sternal cuticle the amount of calcium carbonate and of organic material decreases from the early premoult to that in the exuviae collected in early intramoult. Concomitant to these changes the thickness of the cuticle decreases (Fig. 6A). The changes in the calcite distribution are small and may result from intraspecific variations between the animals. In the cuticle of the anterior tergites there appears to be no change in the distribution of CaCO\(_3\), calcite and organic material between early premoult and mid premoult, however, in late premoult an additional layer of non-calcitic CaCO\(_3\) that also contains organic material appears at the base of the cuticle, clearly below the non-mineralized membranous layer (Fig. 6B). In contrast to the membranous layer that is composed of little protein and large amounts of chitin the additional mineralized layer contains high amounts of proteins and less chitin (data not shown). In early intramoult this additional CaCO\(_3\) layer has disappeared. During the mineralization of the new

**Fig. 5.** Averaged Raman spectra of (A) the anterior tergite during early premoult, and (B) the anterior exuviae obtained from Raman spectroscopic images, and reference spectra of (C) synthetic ACC, (D) purified chitin from crab cuticles, (E) BSA as a protein standard and (F) synthetic calcite.

**Fig. 6.** Raman spectroscopic images recorded from sagittally cleaved and microtome-polished surfaces of posterior (A) and anterior (B) tergite cuticles of \textit{P. scaber} during various stages of the moulting cycle. The images provide the distribution of carbonate (orange), calcite (red) and organic matrix (green). Calcite is located in the exocuticle, whereas calcium carbonate occurs within the whole endocuticle. The epicuticle and the membranous layer are devoid of mineral. 1dpost: 1 day after postmoult, 2dpost: 2 days after postmoult, eintra: early intramoult, epost: early postmoult, epre: early premoult, lpre: late premoult, mpre: mid premoult. (C) Raman spectroscopic images recorded from sagittally cleaved and microtome-polished surfaces of the posterior tergite cuticle of \textit{P. scaber} sampled 1 h after the posterior moult. Carbonate (orange) occurs within an about 4 \(\mu\)m thick layer of the exocuticle. Calcite (red) formation has already started within a thin layer at the distal side of the exocuticle. Only a thin layer of the organic matrix (green) at the proximal side of the developing cuticle appears unmineralized.
posterior cuticle the amount of CaCO₃ and organic material increase in the posterior tergites from late intramoult to the stage 2 days after the anterior moult and further within the next 4–6 days to the early premoult stage (Fig. 6A). Concomitant to the mineralization of the posterior cuticle the amount of CaCO₃ and organic material decrease in the anterior tergite cuticle from late premoult until ecdysis of the anterior exuviae that were collected during early postmoult (Fig. 6B). The new anterior cuticle is already well mineralized 1 and 2 days after the posterior moult, and the amount of calcium carbonate and organic material increases further during the next 4–6 days until early premoult.

In an attempt to analyze an early mineralization step we imaged the new posterior cuticle just 1 h after the posterior moult (Fig. 6C). In this stage a more than 4 µm thick mineralized layer has been already formed. This layer contains calcium carbonate, however, only the distal 200–300 nm in the form of calcite. The proximal part of the mineralized layer contains ACC in a rather patchy distribution. The Raman image of the organic matrix indicates that the whole cuticle is only little thicker than the mineralized layer. Not mineralized are the thin distal epicuticle and a less than 1 µm thick proximal layer (Fig. 6C).

Fig. 7 gives an example for the estimation of the ACC/calcite ratio from the SCµ-RSI data of tergite cuticles. The ratios for both the posterior and anterior cuticle the amount of CaCO₃ and organic material increase in the posterior tergites from late intramoult to the stage 2 days after the anterior moult and further within the next 4–6 days to the early premoult stage (Fig. 6A). Concomitant to the mineralization of the posterior cuticle the amount of CaCO₃ and organic material decrease in the anterior tergite cuticle from late premoult until ecdysis of the anterior exuviae that were collected during early postmoult (Fig. 6B). The new anterior cuticle is already well mineralized 1 and 2 days after the posterior moult, and the amount of calcium carbonate and organic material increases further during the next 4–6 days until early premoult.

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Fig. 7 gives an example for the estimation of the ACC/calcite ratio from the SCµ-RSI data of tergite cuticles. The ratios for both the posterior and anterior cuticles from the early premoult stage (epre) with the tergite of the exuviae (eintra) reveals that the thickness of the calcite-rich exocuticle is equal for both samples whereas the total thickness of the mineralized part of the tergites is reduced in the exuviae. This indicates that mainly ACC is resorbed.

Fig. 8. Line scans across sagittal planes of the posterior tergite cuticle of P. scaber. Comparing a posterior tergite from the early premoult stage (epre) with the tergite of the exuviae (eintra) reveals that the thickness of the calcite-rich exocuticle is equal for both samples whereas the total thickness of the mineralized part of the tergites is reduced in the exuviae. This indicates that mainly ACC is resorbed.

Fig. 9. Mean values for the thickness of the calcite-containing exocuticle and that of the endocuticle were determined from line scans recorded from tergites of P. scaber at the different moulting stages. For both posterior (A) and anterior (B) samples the thickness of the exocuticle shows only small variations. Variations in the thickness of the endocuticle can be attributed to the resorption of and mineralization with ACC. The numbers on top of the error bars denote the mean values. The numbers in parentheses below the x-axis give the number of animals before and the number of records after the slash. 1dpost: 1 day after postmoult, 2dpost: 2 days after postmoult, eintra: early intramoult, epre: early premoult, lpre: late premoult, mpre: mid premoult. Error bars give the standard deviation.
after the anterior moult the values obtained by quantitative SC
the anterior moult and of the new anterior tergites from 1 to 2 days
new posterior tergites from late intramoult to the second day after
individual animals. It appears that during the mineralization of the
postmoult (exuviae), with the exception of the values in late pre-
(exuviae) and in the anterior tergites from early premoult to early
postmoult, epost, early postmoult, epre: early premoult, lpre: late premoult, mpre:
mid premoult. Error bars give the standard deviation.

Fig. 10. Mean values for the thickness of the integral intensity of the Raman counts
of the band for calcite lattice vibration at 280 cm⁻¹ within the exocuticle, and that
of the band for carbonate stretching vibration between 1070 and 1090 cm⁻¹ within
the endocuticle, as determined from line scans recorded from posterior (A) and
anterior (B) tergites of P. scaber at different moulting stages. The amount of calcite is
nearly constant within the various moulting stages. Large changes related to
moulting occur in the amount of ACC within the endocuticle. Numbers on top of the
error bars denote the mean values. The numbers in parentheses below the x-axis
give the number of animals before and the number of records after the slash. 1dpost: 1 day after postmoult, 2dpost: 2 days after postmoult, intra: early
intramoult, epost, early postmoult, epre: early premoult, lpre: late premoult, mpre:
mid premoult. Error bars give the standard deviation.

(1) ACC resorption. The spatial location at the proximal side of the cuti-
no indication of ACC crystallization within the cuticle during min-
Calcite can be excluded during mineral resorption from the same
anterior tergite. An assumption confirmed directly by SC-
ential resorption of ACC over calcite. An increase in the amount of
the underlining epithelial cells. This involves lowering of the mean
of the latter are probably the shorter intervals for mineral resorp-
section 4 for further evaluation of differences obtained by the two
methods.

Fig. 8 shows two examples from Raman spectroscopic line scans
for carbonate and calcite across the sagittal planes of anterior ter-
and in the anterior tergites from early premoult to early
postmoult (exuviae), with the exception of the values in late pre-
moiunt for the posterior tergites and in late intramoult for the ante-
ior tergites. These differences between the values obtained by the
two methods may be explained by intraspecific variations between
individual animals. It appears that during the mineralization of the
new posterior tergites from late intramoult to the second day after
the anterior moult and of the new anterior tergites from 1 to 2 days
after the anterior moult the values obtained by quantitative SCµ-
RSI analysis lead, in almost all cases, to considerably lower values
than combined QRA and AAS. Intraspecific variations cannot be
completely discounted for this difference. However, because of the
similarity of the trend during the moulting cycle, it appears likely that the difference in the values is due to the application of
different methods. In fact statistical analysis of the pooled data
from tergites sampled 1 and 2 days after the anterior moult re-
vealed a significant difference (two tailed unpaired t-test, 
\( p < 0.01 \)) between the means ± SD (n) of 2.26 ± 0.30 (4) and
1.47 ± 0.36 (8) analyzed by QRA/AAS and SCµ-RSI, respectively.
See the Section 4 for further evaluation of differences obtained by the two methods.

4. Discussion

The results of the present study indicate that ACC within the
Cuticle of terrestrial isopods play an important role during re-
peated storage and recycling of CaCO₃ within the integument. For
the first time we show by combined QRA and AAS and SCµ-RSI that
most of the amorphous calcium carbonate (ACC) is resorbed from
the cuticle, whereas calcite remains in the old cuticle and is shed
during moulting. Furthermore we show that in the isopod exocuti-
CLE, ACC functions as a precursor phase for calcite during
biomineralization.

4.1. Resorption and storage of ACC

The decrease in the ACC/calcite ratio within the posterior and
anterior tergites during the stages of mineral resorption, as re-
vealed by combined QRA and AAS are in accordance with a prefer-
ential resorption of ACC over calcite. An increase in the amount of
calcite can be excluded during mineral resorption from the same
tergite. An assumption confirmed directly by SCµ-RSI, which show
no indication of ACC crystallization within the cuticle during min-
eral resorption. The spatial location at the proximal side of the cuti-
cle facilitates the resorption of ACC occurring by active processes
of the underlying epithelial cells. This involves lowering of the mean
pH and an increase in the expression of calcium transport mech-
isms (Ziegler, 2008, 2002; Ziegler et al., 2004; Hagedorn and
Ziegler, 2002). It appears that ACC within the cuticle has at least
two functions: (1) spatial modulation of the mechanical properties
within the cuticle from distal towards the proximal layers (Neues
et al., 2007; Hild et al., 2008) and (2) facilitation of mineral resorp-
tion from the posterior and anterior cuticle during the biphasic
moult cycle of isopods (present study). The adaptive advantages of
the latter are probably the shorter intervals for mineral resorp-
tion in pre and intramoult in which the cuticle is not fully miner-
alized and cannot provide its full support and protective functions. In particular this aspect is important in isopods that rely
on a thick cuticle in avoiding predation by rolling into a sphere (Schmalfuss, 1984). These animals have a higher ACC/calcite ratio

\( \text{ACC from the endocuticle is resorbed and that calcite within the}
\text{exocuticle remains more or less unchanged. The difference be-
\text{tween the total thickness of the mineralized part of the tergites}
\text{determined from the carbonate signal and the exocuticle thickness}
\text{will result in the thickness of endocuticle where only ACC can be
found. From line scans like those in Fig. 8 we plotted mean values for}
\text{the thickness of the calcite containing exocuticle and that of the}
\text{endocuticle (Fig. 9). To quantify the amount of calcite and ACC in}
\text{the exocuticle and the endocuticle, the total amount of the Raman}
\text{counts of the integral intensity of the band for calcite lattice vibra-
\text{tion at 280 cm⁻¹ in the exocuticle region and the total amount of}
\text{the Raman counts of the band for the carbonate stretching vibra-
\text{tion in the endocuticle are determined (Fig. 10). It is important}
\text{to note that differences between the stages reflect also moult-inde-
pendent differences in cuticle thickness and mineral content in}
\text{addition to moult related changes. Nevertheless, the plots show}
\text{that the thickness and the calcite content within the exocuticle,}
\text{vary only within a small range and do not follow the large mostly}
\text{moult related changes in the thickness and ACC content of the}
\text{endocuticle (Figs. 9 and 10). Therefore, the variations of the values}
\text{within the exocuticle are likely due to intraspecific variations}
\text{rather than moult related events. This gives indication that during}
\text{mineral resorption apparently no calcite is dissolved and also that}
\text{no ACC is transformed to calcite. Furthermore, it appears that the}
calcite layer is quickly reformed within less than 1 day. The thick-
ess and ACC content of the endocuticle show strong changes dur-
ing the moulting cycle that appear to be mostly but not always due
to resorption and deposition of ACC. However, the overall trend
shows that the thickness of the endocuticle and the content of
ACC decrease when mineral is resorbed, and increase during min-
eralization (Figs. 9 and 10).}
and larger calcium stores than those with thinner cuticles, in order to accelerate cuticle mineralization after ecdysis (Becker et al., 2005; Neues et al., 2007; Hild et al., 2008). Resorption of ACC from the cuticle during premoult was also reported for the cuticle of the crayfish Cherax quadricarinatus (Shechter et al., 2008). Crayfish resorb calcium carbonate from the cuticle to weaken the cuticle before ecdysis (Wheatly and Gannon, 1995). Most of this calcium is excreted, but a part of it is stored in gastrocoels that also consist of ACC (Shechter et al., 2008).

The location below the membranous layer suggests that the transient layer inside of the anterior tergite was secreted by the hypodermal epithelium of the tergites. Disappearance of this layer just after posterior ecdysis indicates that it serves as a transitory reservoir for the mineralization of the new posterior cuticle. Its composition of ACC-microspherules and a proteinaceous organic matrix reminds to the sternal ACC deposits, a transitory reservoir for the mineralization of the new posterior cuticle just 1 h after the posterior moult supports our finding that ACC precedes calcite within the exocuticle. However, because of a thin layer of calcite in the most distal region of the cuticle we cannot exclude a direct precipitation of calcite without a preceding amorphous phase in that very distal region. A precursor function of ACC for calcite formation in the crustacean cuticle was previously suggested by Dillaman et al. (2005) based on the solubility of cuticular calcium carbonate just after its deposition in the cuticle of the dorso-carpal scutum of Callinectes sapidus.

4.2. Methodical aspects

As already mentioned in the Section 3, differences in the ACC/calcite ratio obtained when using the combination of QRA and AAS at one hand and SCµ-RII on the other may be due to variations between individuals. Intraspecific variations can be high and only a small number of tergites were analyzed by SCµ-RIII, whereas for the combined QRA/AAS study the pooled tergites of 6–15 animals were analyzed for each moulting stage. In addition, since the ACC/calcite ratio is changing quickly during mineral resorption and deposition, differences between individuals also reflect variations in the time of animal preparation within a specific moulting stage. However, when the mineralization is almost complete, differences due to intraspecific variations or time of sampling should be rather low. Furthermore, the significant difference obtained when the two methods are compared, using data from tergites sampled 1 or 2 days after the anterior moult, argues against variations between the specimens as the reason for the difference. An overestimation of the ACC/calcite value using QRA and AAS because of the ACP content of 12% within the cuticle was neglected in this study can by no means explain the differences between the methods. This suggests that QRA/AAS and SCµ-RIII differ in the sensitivity to specific properties of the calcium carbonate phases in the tergites of these moulting stages. The decrease of the ACC/calcite ratio from the second day of the postmoult to the early premoult stage in both the anterior and posterior tergites as measured by QRA/AAS can be explained by conversion of a part of the X-ray amorphous CaCO₃ into calcite. It appears likely that SCµ-RIII does not confirm this result because the sensitivity of Raman spectroscopy to detect calcite-like lattice vibrations is higher than the sensitivity of XRD to detect calcite-like short-range order in crystallising nanoparticles converting from ACC precursors. This interpretation, however, is speculative and requires further experimental support.

Acknowledgments

We thank the Deutsche Forschungsgemeinschaft (DFG) for financial support within the Priority Program “Principles of Biomaterialization” SPP 1117 (IZ 368/4-3 and Ep 22/16-3) and HASYLAB at DESY for generous allocation of beamtime.
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