Embryonic development and secondary axis induction in the Brazilian white knee tarantula *Acanthoscurria geniculata*, C. L. Koch, 1841 (Araneae; Mygalomorphae; Theraphosidae)

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Received: 28 November 2019 / Accepted: 29 January 2020 / Published online: 19 February 2020
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Abstract
Tarantulas represent some of the heaviest and most famous spiders. However, there is little information about the embryonic development of these spiders or their relatives (infraorder Mygalomorphae) and time-lapse recording of the embryonic development is entirely missing. I here describe the complete development of the Brazilian white knee tarantula, *Acanthoscurria geniculata*, in fixed and live embryos. The establishment of the blastoderm, the formation, migration and signalling of the cumulus and the shape changes that occur in the segment addition zone are analysed in detail. In addition, I show that there might be differences in the contraction process of early embryos of different theraphosid spider species. A new embryonic reference transcriptome was generated for this study and was used to clone and analyse the expression of several important developmental genes. Finally, I show that embryos of *A. geniculata* are amenable to tissue transplantation and bead insertion experiments. Using these functional approaches, I induced axis duplication in embryos via cumulus transplantation and ectopic activation of BMP signalling. Overall, the mygalomorph spider *A. geniculata* is a useful laboratory system to analyse evolutionary developmental questions, and the availability of such a system will help understanding conserved and divergent aspects of spider/chelicerate development.

Keywords Mygalomorphae · *Acanthoscurria geniculata* · Embryonic development · Axis duplication · Cumulus grafting · Bead transplantation

Introduction

Within the chelicerates, the spiders have become an important model system to study evolutionary, developmental and ecological questions. Furthermore, the diverse field of applications for spider venom and silk have always attracted scientists to study these fascinating animals. Several spider genomes have been published recently allowing to perform functional and evolutionary genomic studies (Kono et al. 2019; Sánchez-Herrero et al. 2019; Sanggaard et al. 2014; Schwager et al. 2017).

Spiders (*Araneae*; around 48000 described species (World Spider Catalog 2019)) can be divided into three suborders; the basally branching segmented Mesothelae, the Mygalomorphae (tarantulas, trapdoor spiders and allies) and the Araneomorphae (“true spiders” including Haplogynae and Entelegynae) (see Fig. 1a).

In recent years, embryonic development has been extensively studied in Araneomorphae (e.g., Akiyama-Oda and Oda 2003; Mittmann and Wolff 2012; Turetzek and Prpic 2016; Wolff and Hilbrant 2011; Yamazaki et al. 2005). Although several classical, and also some recent, studies deal with the embryonic development of mygalomorph spiders and even Mesothelae (e.g., Crome 1963; Pechmann and Prpic 2009; Setton et al. 2019; Yoshikura 1954; Yoshikura 1958), fewer details on the embryonic development are available for these spider suborders. Furthermore, time-lapse recording of the embryonic development has proved incredibly useful for...
analysing araneomorphs (e.g., Mittmann and Wolff 2012; Turetzek and Prpic 2016; Wolff and Hilbrant 2011), but this approach has never been applied to other groups.

There are some important morphological differences between mygalomorph and araneomorph spiders which make it interesting to analyse and compare the development of these morphological characters in these two suborders of spiders. The first major morphological difference is the arrangement of the chelicerae. Until recently, this feature was used to place Mygalomorphae and Araneomorphae into the groups Orthognatha (chelicerae run in parallel to each other) and Labidognatha (chelicerae are in a vertical position opposing each other) (reviewed in e.g., Foelix 2011). As the chelicerae of Mesothelae also run in parallel to each other, it is likely to be the ancestral character of spiders. Other important morphological differences are the absence of the anterior pair of spinnerets in mygalomorph spiders (reviewed in e.g., Foelix 2011, Pechmann et al. 2010) and the presence of two pairs of book lungs in Mygalomorphae and Mesothelae vs. only one pair of book lungs in most Araneomorphae (exception Hypochilidae (e.g., Sharma 2017)). First studies on the embryonic development of tarantulas revealed, that the anterior spinnerets are present as rudimentary structures during the early embryonic development (e.g., Pechmann and Prpic 2009; Setton et al. 2019). It will be interesting to analyse the factors that are responsible for the loss or for the sustainment of the spinnerets during embryonic development. Establishment of a mygalomorph spider model system in the laboratory is thus crucial for the understanding of the evolution and the development of spiders.

Although mygalomorph spider species have a very long generation cycle (most species become sexual reproductive after several years), a large spider culture and the huge amount of eggs per cocoon can at least partially compensate for the long generation time of these animals. In addition, mygalomorph spiders produce very large eggs, which make them amenable to perform classical transplantation experiments (this study).

Here, I provide a detailed analysis of the embryonic development of the tarantula *A. geniculata*. In addition, I show that embryos of *A. geniculata* can be used to perform tissue grafting as well as bead insertion experiments. Applying these techniques, I show that the grafting of the “cumulus” (the organizing centre of early spider embryos) as well as the local activation of the BMP signalling pathway can induce an axis duplication phenotype.

Overall, *A. geniculata* is a promising system that can be used to perform modern and classical embryological studies. The availability of a draft genome assembly (Sanggaard et al. 2014), the presence of transcriptomic data (this work and
Sanggaard et al. 2014) as well as the accessibility to modern molecular biological techniques make this system attractive to analyse a variety of biological questions.

Methods

Spider husbandry

All spiders used for this study originated from a laboratory culture (see Pechmann and Prpic 2009) or were purchased from German breeders. Wild caught animals have not been used. Juveniles of A. geniculata and Brachypelma albopilosum were kept in Drosophila vials and plastic boxes of appropriate size. Adults were kept in glass containers of 20 cm height, 30 cm width and 30 cm depth. Spiders were kept on moist coconut fibres and were fed with Drosophila melanogaster and Gryllus bimaculatus once or twice a week. Adults had access to a water dish. To speed up the development, juveniles were kept in incubators at 27 °C with 10 h light and 14 h dark cycle. Adults were kept in the lab at room temperature (near the window; without direct sun light). As the temperature dropped below 20 °C during the winter, an external heating source (thermo mat, 7 W, Lucky Reptile) was attached to one side of the glass container. Prior to and after mating, adult females were fed more frequently. Cocoons were opened with fine scissors, and eggs were kept within the opened cocoon or were transferred to petri dishes. Cocoons and eggs were incubated in large petri dishes containing a wet paper towel to ensure the desired humidity.

Worldwide, many hobbyists keep mygalomorph spiders as a pet. For this reason, several excellent books are available that describe the breeding and the maintenance of tarantulas in captivity (e.g., Klaas 2013; Schultz and Schultz 2009).

Transcriptome sequencing and assembly

Total RNA from st. 5–14 embryos was extracted using TRIZOL and the Quick-RNA MicroPrep Kit (Zymo Research). In addition total RNA was extracted from fixed st. 5 embryos using the Quick-RNA FFPE Kit (Zymo Research). Both samples were sequenced individually.

Library preparation and sequencing was carried out at the Cologne Center for Genomics (Illumina HiSeq4000). The sequenced reads were subjected to adapter and quality threshold trimming using Trimmomatic 0.33 (Bolger et al. 2014; Trimmomatic 0.33 settings: ILLUMINACLIP:adapter. fa:2:30:10 LEADING:3 TRAILING:3 SLIDINGWINDOW: 4:15 MINLEN:30 where “adapter.fa” contained the adapters used in sequencing). FastQC (Andrews 2010) was used to assess read quality both before and after trimming. There was no difference in sequence quality between the two sequenced samples. The final transcriptome was assembled from both read sets using Trinity 2.6.6 (Grabherr et al. 2011; Trinity 2.6.6 settings: all default options, normalized reads, – CPU 8 –full_cleanup –inchworm_cpu 8 –bflyHeapSpaceMax 15G –bflyCalculateCPU), and the combined assembly can be found on Figshare (https://doi.org/10.6084/m9.figshare.10266314). BioProject accession for the raw reads is PRJNA588224.

The statistics (see Table S2) were calculated using BUSCO v2/3 (Simão et al. 2015), the gVolante server (https://gvolante. riken.jp/analysis.html; Nishimura et al. 2017) and custom scripts that are available upon request.

Gene cloning

CLC Main Workbench 7 (QIAGEN Aarhus A/S) was used to perform a local TBLASTN (Altschul et al. 1990) against the newly assembled transcriptome. For this, protein sequences of known homology were downloaded from FlyBase (Thurmond et al. 2019) and NCBI. Putatively identified genes were reciprocally blasted against the online NCBI nr database using BLASTx to confirm their identity.

cDNA synthesis was carried out using the SuperScript VILO cDNA Synthesis Kit (Thermo Fisher Scientific) or the qScript cDNA Synthesis Kit (Quanta bio). Total RNA or cDNA is available upon request.

PCR amplification and cloning of genes were performed using standard techniques. All genes were cloned into pCRII-TOPO or pCR4-TOPO (Invitrogen). A primer list with corresponding TRINITY_IDs of the genes found in the new embryonic reference transcriptome is provided in the supplemental material (Table S1).

Imaging and image analysis

Most images were recorded using an AxioZoom.V16 (Zeiss; ZEN2 Software), equipped with a movable stage. For live imaging, the membrane of a cell culture dish (lumox dish 35, SARSTEDT) was covered with heptane glue. After the heptane evaporated, the embryos were glued to the membrane and covered with oil (Voltalef H10S or Halocarbon oil 700; SIGMA). Live imaging was performed at room temperature (25 °C ± 2 °C).

The embryos shown in Movie S1, S2, S3, S6 and S9 were imaged via a 45-degree mirror as described in Pechmann 2016.

Helicon Focus (Helicon Soft) was used to generate image stack projections. Fiji (Schindelin et al. 2012) was used to create movies. Created movies were converted and compressed using HandBrake (https://handbrake.fr/). Adobe Photoshop CS5.1 was used to adjust for brightness and contrast.

False color overlays of in situ hybridisation images were generated as described in Pechmann et al. 2017.
**Cumulus grafting**

To extend the time window for cumulus grafting experiments, embryos of the same cocoon were incubated at different temperatures (20 °C, 22 °C, 25 °C, 27 °C). This allowed me to obtain cumulus migration stages (embryonic stage 5, see below) over a time period of 5 days.

For cumulus grafting experiments, 50–100 stage 5 embryos were dechorionated using 2.8% hypochlorite solution (DanKlorix). The dechorionisation step was mandatory for the survival of the embryos. Dechorionated embryos were washed several times with distilled water. The embryos were attached to the membrane of a single cell culture dish (see above) and covered with Halocarbon oil 700 (SIGMA). With the help of SuperFine Vannas Scissors (World Precision Instruments), the vitelline membrane of the donor embryo was cut in a region close to the cumulus. A pulled glass capillary was used to cut out the cumulus of the “donor embryo”. By squeezing the vitelline membrane of the “donor embryo”, it was possible to transfer a small droplet of the perivitelline fluid together with the excised cumulus into the surrounding oil. Using the SuperFine Vannas Scissors, the vitelline membrane of a nearby embryo (the “cumulus acceptor”) was cut at a position that was opposite to the side of the already migrating endogenous cumulus. As the migrating cumulus bulges the overlaying ectoderm in a kind of triangular shape, the direction of the movement of the cumulus was traceable in most embryos. With the help of the glass capillary, the droplet of perivitelline fluid containing the excised cumulus was carefully guided through the oil towards the cut in the vitelline membrane of the “cumulus acceptor”. The droplet within the oil (containing the cumulus of the “donor embryo”) was fused with the perivitelline fluid of the “acceptor embryo” by simultaneously touching both fluids with the glass capillary. After droplet fusion, the “donor cumulus” was carefully inserted into the perivitelline space of the “acceptor embryo” and was moved a bit downward to clamp it between the vitelline membrane and the ectoderm of the “acceptor embryo”.

**Bead transplantation**

Recombinant human BMP4 protein (Thermo Fisher Scientific; PHC9534) was reconstituted in 0.1% bovine serum albumin (BSA). Affi-Gel Blue beads, diameter 150–300 μm (Bio-Rad; #1537301), were washed three times in PBS. Prior to implantation, around 100 washed beads were incubated for 1.5 h (at room temperature) in 5 μl of 100 μg/ml recombinant human BMP4 protein or 0.1% BSA.

Prior to bead transplantation, embryos were prepared as described in the section “cumulus transplantation” and covered with Halocarbon oil 700 (SIGMA). The soaked BMP4 or BSA (as a control) beads were pipetted into the oil, and single beads were separated within the oil using a pulled glass capillary. Using the SuperFine Vannas Scissors, the vitelline membrane of an embryo was cut at a position that was opposite to the side of the already migrating endogenous cumulus. Single beads were guided through the oil towards the cut in the vitelline membrane. With the help of the pulled glass capillary, a small cut was made in the ectoderm of the germ-disc. The bead was inserted into this cut and was pushed below the ectoderm.

**Embryo fixation, in situ hybridisation, durcupan sections and pMad antibody staining**

Embryo fixation was performed as described for *Parasteatoda tepidariorum* (Pechmann et al. 2017) with minor modification. Fixed embryonic material is available upon request.

For the fixation of cumulus – or bead – transplanted embryos, about 3 μl of 10% formaldehyde was injected to the perivitelline space. Excess oil was poured from the lummox dish after an incubation time of 1–2 h. Using a scalpel, the membrane with the attached embryos was cut out from the lummox dish. The embryos (still glued on the membrane) were placed upside-down on the aqueous phase of the fixative (5% formaldehyde in PBS) and were covered with 100% heptane. After 30 min, most of the embryos were washed of the membrane and were transferred to fresh fixative (3 ml PBS, 3 ml 10% formaldehyde and 6 ml heptane). Embryos were fixed on a shaker overnight at room temperature and 50 rpm.

In situ hybridisation was performed as described for *C. salei* (Prpic et al. 2008) with minor modification (no proteinase K treatment; doubling of buffer volumes; duration of washing steps was doubled; final overnight washing step in 15 ml PBST).

Durcupan sections (20 μm cross-sections) of stage 5 embryos and pMAD antibody staining were performed as described in Pechmann et al. 2017. For pMad antibody staining, cumulus or bead transplanted embryos were fixed 15–20 h after transplantation.

**Results and discussion**

**A. geniculata as a model system in the lab**

*A. geniculata* is a huge and colourful tarantula (body length of around 8 cm and a leg span of around 20 cm; see Fig. 1b and Fig. S1a,b) from Brazil. The spider should be handled with care as the behavioural character can vary from shy to very aggressive, and urticating hairs are regularly used as a defence mechanism.

Regarding the long generation time of Mygalomorphs, it is challenging to analyse evolutionary developmental questions in this group of spiders. However, *A. geniculata* females
produce a huge number of eggs/cocoon (between 1000 and 2000), and the spiders can be maintained and raised in the lab throughout the year. In addition, transcriptomic data (this work and Sanggaard et al. 2014), as well as a draft assembly of the sequenced genome of *A. geniculata*, are available (Sanggaard et al. 2014), and in situ hybridisations were previously established in embryos of this spider species (Pechmann and Prpic 2009). Finally, the big size of the embryos makes them amenable to transplantation experiments (this work). These facts compensate for the long generation time of this spider species and allow the analysis of evolutionary developmental questions.

Depending on temperature and food supply, females reach adulthood after 3–5 years. In contrast to the males, which reach adulthood after 3–4 years and only live for an additional year, the females can reach an age of more than 10 years. During that time the females moult regularly (on average once per year) to regenerate lost hairs or appendages. As transferred sperm is lost during the moultng process, the females have to be mated again. However, a spider culture of around 30 individuals with 10–20 adult females and males of different age ensures that several cocoons can be obtained every year.

After their final moult, male spiders show a clear sexual dimorphism (Fig. S1a,b). The distal part of the pedipalp has developed into a copulatory organ, the bulb (Fig S1c,d). Prior to copulation, males produce a specialized web that is used to attach a drop of sperm. This drop of sperm is then actively taken up into the bulb. In addition to the pedipalpal bulb, the first walking leg has developed a tibial apophysis (Fig. S1e). During mating, the male (spider to the right in Fig. 1b) uses the tibial apophysis to lift up the female. This enables the male to reach the genital opening of the female on the ventral side of the opisthosoma and to insert the bulb of the pedipalp at the right angle to transfer the stored sperm. The female is able to store the sperm for several months in the receptaculum seminis (Fig. S1f).

A complicated mating dance precedes the copulation, and it can take minutes to hours for the final step of sperm to transfer. Males that are stimulated by touching the silk or an exuvium of a female spider will start to send out species-specific vibrations. Females that are willing to mate will answer violently by tapping the ground (like a drum roll). This behaviour can also be used to check whether a female has reached adulthood.

After mating, females were regularly fed to ensure egg production, and cocoons were observed 10.5 weeks after mating (time between mating and cocoon production of four spider females was monitored). The timing of cocoon production may vary depending on the season, feeding status of the female, temperature and humidity. The cocoon production itself takes several days and involves the establishment of a giant bowl-shaped web (see Fig. 1c). The final cocoon has a ball-like appearance (Fig. 1d,e) and is protected by the female. The size of the cocoon and the number of eggs depend on the size of the female and food supply. One of the cocoons obtained for this study contained around 2000 eggs (Fig. 1d–f).

**Embryonic development of *A. geniculata***

Several publications deal with the description of the development of Araneomorphae spider species (Mittmann and Wolff 2012; Turetzek and Prpic 2016; Wolff and Hilbrant 2011). Unfortunately, the embryonic staging varies across the different publications (e.g., 21 embryonic stages for *Cupiennius salei* vs. 14 embryonic stages for *P. tepidariorum* or hours after egg lay in *Pholcus phalangioides*). This fact makes it difficult to compare embryonic stages of different spider species. Nevertheless, stage 1 (early cleavage stages before contraction), stage 5 (cumulus migration), stage 8 (segmented germ band before appendage outgrowth) and stage 9 (appendages start to develop) are embryonic stages that are common to all spider species studied so far. These stages should be used as a landmark for staging spider species and have been used in this study to compare and align the development of *A. geniculata* embryos to other spider species, like *C. salei* or *P. tepidariorum* (Fig. 1a).

**Stage 1–3: energid cleavages, contraction and blastoderm**

Freshly laid eggs are very fragile and surrounded by a fluid that is absorbed during the first hour of development (Foelix 2011). This absorbance process leads to the formation of the chorion that surrounds the egg and most likely protects the eggs against desiccation and bacterial infections (e.g., Makover et al. 2019). The chorion is irregularly distributed on the surface of the vitelline membrane and can be recognized as a whitish and granulated material (see Fig. 2 and Movies S1, S2). In contrast to many other spider species (e.g., *P. tepidariorum*), the chorion of *A. geniculata* eggs is quite transparent, and it is not necessary to remove the chorion or to immerge the eggs in oil to stage the embryos.

Eggs that were removed a few hours after egg deposition did not survive. Only around 24 h after egg deposition it was possible to carefully remove single eggs from the cocoon and to analyse the very early steps of embryogenesis (see below).

In contrast, most of the eggs of a cocoon that was opened 3 days after egg lay could easily be separated from each other. These embryos were still at stage 1 of development, and it was possible to observe the cortical periplasm that was divided into ovoid or polygonal fields (dotted line in Fig. 2a).

During stage 1, gravity is leading to an irregular distribution of larger and denser yolk granules within the egg. As a result, the upper third of the stage 1 pre-contraction egg is more transparent than the lower two thirds of the egg (see brackets in Fig. 2b and the dotted line in Fig. S3a, Movies S1 and S2). This phenomenon has been described for other spider species as well, and it has been suggested that this is
common to large eggs (Holm 1940). However, the subdivision of a region of more densely packed yolk granules seems not to be connected to any polarity of the egg as the process can be reverted as soon as one rotates the egg upside-down (own observation; Holm 1952). In addition to *A. geniculata*, I also had access to very early embryonic stages of *Brachypelma albopilosum*, another theraphosid spider species. Time-lapse imaging of *B. albopilosum* embryos that were at the beginning of stage 1 (around 24 h after egg deposition) revealed that the establishment of these cortical polygonal fields occurred several hours after egg lay (see Movie S2 and Movie S3). The nature and the content of the cortical polygonal fields are not well described. However, these fields also develop in other spider species, and it has been suggested that these fields are required for the cellularization process that occurs at the end of stage 1 (Balbiani 1873; reviewed in Holm 1940).

Nuclei with surrounding cytoplasm (so-called cleavage energids) are the characteristics of the embryonic stage 1. The energids start to divide in the centre of the egg and reach the periphery of the egg at the end of stage 1 (Fig. 2b; Fig. S2a; Movies S1, S2 and S4). In spiders, cytoplasmic strands connect the cleavage energids to the cortical fields (reviewed in Schwager et al. 2015), and these strands seem to be required to bring the energids from the centre towards the surface of the egg. If this is an active (via contraction of the strands) or passive (flexible strands under tension) process is unknown.

At the end of stage 1, the energids have reached the surface of the egg, and each energid fuses with several polygonal cortex fields (see Fig. 2b; Fig. S2a; Movies S1, S2 and S4). As soon as the energids have invaded the cortex, polygonal cortex fields and energids divide synchronously (see Movie S1). This was in contrast to *B. albopilosum* embryos in which the cleavage energids reached the surface of the egg only in the lower two third of the egg (in the region of the densely packed yolk granules; see Fig. S3a,b and Movie S2). Thereafter, the upper, energid free cortex, collapsed on
top of energids that arrested at the surface of the densely packed yolk mass (see Fig. S3b,c and Movie S2).

As the polygonal cortical fields are very fragile structures, single or groups of fields were regularly damaged while handling stage 1 embryos. However, this damage of the cortex allowed me to see below the cortex, and I was able to image the movement of the energids along the connecting strands during the contraction phase in *B. albopilosum* embryos (see Fig. S3e,e' and Movies S4 and S5).

At stage 2, blastodermal cells are large and irregularly distributed (Fig. 2c,d; Fig. 3a; Movie S1, S2, S4, S7). In addition, not every polygonal field of the cortex has been occupied by an energid (dashed line in Fig. 2d and boxed region in Fig. S4a–d). Asynchronous cell divisions lead to stage 3 embryos that show a relatively regular distribution of small blastodermal cells (Fig. 2e; Fig. 3b; Movie S1 and S7). At the same time, contraction of cortical fields leads to the formation of big white patches that can be observed throughout the blastoderm (the development of the white patch marked by dashed line in Fig. 2e is shown in Fig. S4).

At stage 3, some cells appear whiter than others, and this difference in colour is sometimes connected to a different cell fate. Movie S6 shows a group of cells of white appearance that start to become migratory during stage 5 of development.

**Stage 4–5: primary thickening, germ-disc and cumulus migration**

During stage 4, hundreds of cells accumulate and form a white disc of around 500 μm in diameter (Fig. 3e; Movie S7 and

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**Fig. 3**  Embryonic development of *A. geniculata* (st. 2 until hatching; top view). Stills from Movie S7 (a–j) and Movie S10 (k–o). a Embryo at late stage 2 showing irregularly distributed blastoderm cells. b A regular blastoderm has formed that is interspersed with white spots (compare to Fig. 2e). c At stage 4 many cells aggregate and form the “primary thickening” that seems to be the source for the developing cumulus and for most of the cells that gastrulate during stage 5. d and e The cumulus (cu) appears at early stage 5 and migrates from the region of the primary thickening towards the rim of the “germ-disc” (please note that in tarantula embryos a distinct germ-disc is not visible via bright field imaging). f The dorsal field (df) gets established and extensive morphogenetic rearrangements are leading to the formation of the axially symmetric germ band. The region of the primary thickening has developed into the caudal lobe (cl). g A clearly segmented germ-band is present at stage 8. h–l Embryonic development proceeds. Appendages start to grow out and elongate to their final length, posterior segments are added and dorsal closure is completed. See text and Fig. 5 and Fig. 6 for more details. m,n Ventral closure and the pigmentation of the egg tooth (arrow in m), bristles and of the “false pincer” of the chelicera (arrowheads in n) occurs at embryonic stage 14. o The postembryo hatches from the egg. Abbreviations: st. 5e, stage 5 early; st. 5l, stage 5 late; cu, cumulus; df, dorsal field; II-14, walking legs; pp, pedipalp; ch, chelicera. Scale bar is 500 μm.
S8). A similar cell accumulation is visible in a variety of spider species and was named primitive plate or primary thickening in former publications (e.g., Holm 1952; Wolff and Hilbrant 2011). In the following, I refer to this accumulation of cells as the primary thickening.

The primary thickening is the characteristic structure of a stage 4 embryo, and it seems to be the main source of cells that gastrulate and spread over the embryo during embryonic stage 5. In addition the migratory cumulus arises in the centre of the primary thickening (Fig. 3d; Movies S7 and S8).

While a distinct radially symmetric germ-disc is visible at stages 3–5 in spider species like P. tepidariorum or Zygella x-notata (Chaw et al. 2007; Mittmann and Wolff 2012), no distinct germ-disc is visible in tarantula embryos. This is similar to C. salei in which a germ-disc is first visible at stage 5 when gastrulating cells only invade the embryonic half of the egg leading to a distinct border (termed the equator) between the embryonic and ab-embryonic half of the egg (Wolff and Hilbrant 2011). Also in A. geniculata, gastrulating cells seem to primarily invade the embryonic half of the egg (the germ-disc), and an equator-like rim of the disc is visible at stage 5 of development (dotted line in Fig. 4b”). Physical sectioning of stage 5 embryos revealed that while the upper, embryonic germ-disc is multi-layered, the lower ab-embryonic half of the egg is single-layered (Fig. 4a–a’). Further evidence for the presence of a germ-disc, albeit not visible with bright field imaging, is the fact that the cumulus migrates only until it has reached the equator (Fig. 2f,g; Fig. 3d,e; Movies S1 and S7-S9).

In spiders, the cumulus is a group of mesenchymal cells that function as a migratory cell cluster to break the radial symmetry of the germ-disc. The cumulus is likely an evolutionary novelty of chelicerates, and it has been observed in a variety of chelicerate species (reviewed in Schwager et al. 2015 and Hilbrant et al. 2012). In spiders, it has been shown that the cumulus is required to induce (via the activation of the BMP signalling pathway, see below) the so-called dorsal field (Akiyama-Oda and Oda 2006; Oda and Akiyama-Oda 2008). In A. geniculata, the cumulus is about 250 μm in diameter and consists of hundreds of cells (Fig. 2f,g; Fig. 3d,e; Fig. 4a–c).

As in P. tepidariorum, the cumulus expresses the cumulus marker fascin (Akiyama-Oda and Oda 2010), and it activates the BMP signalling pathway (Akiyama-Oda and Oda 2006) in overlaying ectodermal cells (Fig. 4b,c). A. geniculata fascin expression is also visible in single cells and in clusters of cells (Fig. 4c’) throughout the germ-disc. It is likely that these cells are migrating gastrulating endodermal cells as it was shown for fascin positive cells in P. tepidariorum (Akiyama-Oda and Oda 2016).

Stage 6–8: dorsal field and germ-band formation

In P. tepidariorum the gene orthodenticle (otd) is a good marker for anterior tissue identity, and it is expressed all around the anterior rim of the germ-disc (e.g., Akiyama-Oda and Oda 2003; Pechmann et al. 2009). When the cumulus reaches the rim of the germ-disc, it initiates the opening of the disc via the establishment of the dorsal field. This is also reflected by the expression of otd, which forms an “open” circle (Akiyama-Oda and Oda 2003; Pechmann et al. 2009). Also, in A. geniculata otd is expressed as a faint “open” circle at stage 6 of embryonic development (Fig. 4d,e), and the cumulus seems to be required to induce the dorsal field (Fig. 3f; Fig. 4d”; Movie S7). At the beginning of stage 6, the cells of the cumulus lose contact to each other, and they spread over the dorsal field of the embryo (best visible in Movies S1 and S9). So far, the final fate of the cumulus cells is still unclear. Labelling experiments and molecular data suggest that cumulus cells as well as cells within the dorsal field possess an endodermal cell fate (Akiyama-Oda and Oda 2003; Edgar et al. 2015; Feitosa et al. 2017; Holm 1952), and the spreading of the cumulus cells over this area in A. geniculata supports this idea. However, the recent finding that twist (a marker for mesodermal cells) is also expressed within the developing cumulus in P. tepidariorum suggests a mesendodermal nature of the cumulus and indicates that the situation might be more complicated (Pechmann et al. 2017).

It has been shown that convergent extension mechanisms are the main driver to form an axially symmetric spider embryo out of a radially symmetric germ-disc (Kanayama et al. 2011). In A. geniculata, the differences between a stage 6 and a stage 7 embryo are only subtle (stage 7 is more triangular shaped, see Movie S7 and the respective figure legend), and it is a smooth transition to form the segmented stage 8 embryo (Fig. 3f,g). Stage 8 is completed when prosomal segments are clearly segmented and prosomal appendage buds are about to form (Fig. 3g; Fig. 5a–d). The expression of the segment polarity gene engrailed (en) reveals that prosomal segments are well established at stage 8 of embryonic development (Fig. 7a).

Stages 9–11: appendage formation, inversion and a switch in the segment addition zone

Embryonic stage 9 is characterized by the outgrowth of the prosomal appendages (Fig. 3h; Fig. 5e–i). At the beginning of stage 9, broad blocks of segments showing continuous en expression are sequentially added from the segment addition zone (saz) (Fig. 7e). At the end of stage 9, the first sign of the ventral sulcus (ventral opening of the embryo which initiates the process of inversion (described in e.g., Crome 1963; Wolff and Hilbrant 2011)) is visible (Fig. 5e,h,i), and en stripes of the opisthosomal segments 1–3 are split into a left and a right stripe per segment (Fig. 7b). Only at the anterior end of the saz, continuous, semi-circular expression of en was still detectable (o4 segment in Fig. 7b). In spiders that belong to the Entelegynae (like C. salei or
P. tepidariorum), inversion (the ventral splitting of the embryo into a left and a right body half) is a process that is relatively uniform along the anterior-posterior body axis. In these spiders, the ventral sulcus is widest in the median region of the germ-band, and it converges pointedly towards the posterior of the embryo (e.g., Mittmann and Wolff 2012; Wolff and Hilbrant 2011). This is in contrast to the inversion process of A. geniculata in which the ventral sulcus broadens during embryonic stages 9 and 10 and is widest in the second opisthosomal segment and converges pointedly towards the anterior of the embryo (Fig. 5i–k; Fig. 7c,f,g; Fig. 10a). At the end of stage 10 the opisthosoma has a horseshoe-like appearance (Fig. 5k,l). This horseshoe-like split of left and right side of the opisthosoma has been noted in other mygalomorph and chelicerate species before (reviewed in Setton et al. 2019).

Connected to the described shape changes within the opisthosoma, there seems to be a switch in the segmentation process as well. While complete and continuous segments are added one after the other from the posteriorly located sz at embryonic stages 8 and 9 (Fig. 7e), segments seem to be added in a ring and wave like fashion to the left and right body half during embryonic stage 10 (Fig. 5k,l; Fig. 7c,g; Fig. 10a; Fig. S7c). The ring- and wave-like addition of segments of stage 10 A. geniculata embryos is also evident in the expression of segmentation genes. It was shown that in the sz of P. tepidariorum, caudal (cad) might be required for the activation and maintenance of the expression of the pair-rule gene even-skipped (eve) (Schönauer et al. 2016). Also in A. geniculata cad and eve seem to be expressed in overlapping domains within the sz (Fig. 7i and k). Furthermore, similar to P. tepidariorum, A. geniculata eve also shows a wave like
cyclic expression (compare Fig. 7j,k) in the saz. In addition, the pair rule-gene *hairy* (*h*) is also expressed in ring-like domains within the newly formed segments (Fig. 7l). Wave- and ring-like addition of segments from the saz has also been observed in arthropods like myriapods (e.g., Chipman and Akam 2008; Janssen et al. 2011). This suggests that this kind of genetic patterning during segment addition may represent an ancestral feature of arthropod segmentation. Between stage 10 and stage 11 the posterior end of the spider embryo passes through another shape change. Similar to what has been described for the posterior end of Mesothelae (Yoshikura 1954), other mygalomorph spiders like *Atypus karschi* (Crome 1963; Yoshikura 1958) as well as for representatives of haplogyne spiders like *P. phalangioides* or *Segestria bavarica* (e.g., Crome 1963; Holm 1940; Turetzek and Prpic 2016; Wolff and Hilbrant 2011), the segment...
addition zone starts bulging outwards, and a tube-shaped structure grows into an anterior direction (Fig. 5m–o; Fig. 6d; Fig. 10a; Fig. S7d). This tube-like structure has been named the “post-abdomen” (Holm 1940) and the most posterior segments are added sequentially from this tube. This “downturned” opisthosoma is a plesiomorphic character of Arachnopulmonata (reviewed in Setton et al. 2019; Turetzek and Prpic 2016).

In A. geniculata and B. albopilosum embryos, the “post-abdomen” is never as prominent as in A. karschi, P. phalangioides or S. bavarica. However, as the “post-abdomen” is present in all big branches of the Araneae, patterning the posterior segments within a tube-like structure, is likely to resemble the ancestral state of posterior segment addition within spiders (reviewed in Turetzek and Prpic 2016; Wolff and Hilbrant 2011).

Going from stage 8 to stage 10, the anterior of the tarantula embryo also changes its morphology. The precheliceral region is very uniform during stages 8 and 9 (Fig. 5c–i; Fig. 7a,b). As soon as the stomodeum (at stage 10) and the labrum (at stage 11) are established, the pre-cheliceral region gets divided into a left and right pre-cheliceral lobe (Fig. 5j,k; Fig. 6c). At stage 10 and 11, small indentations within the pre-cheliceral lobes and along the ventral neuro-ectodermal tissue (Fig. 5j,k) indicate that neurogenesis has started. In addition to the establishment of the bi-lobed labrum also the appendage buds on the opisthosomal segments 2–5 (Fig. 5m–o; Fig. 6a,d; Fig. S7d) have developed at stage 11. These appendage buds will develop into the respiratory system (the book lungs) and the spinnerets. While the book lungs will develop from the buds on opisthosomal segments 2 and 3, the spinnerets will develop from the bud within spiders (reviewed in Turetzek and Prpic 2016; Wolff and Hilbrant 2011).

During stage 12.2, dorsal closure is at an advanced stage, and mainly the dorsal region of the prosoma has to close (dotted line in Fig. 6k). In the opisthosoma, the heart tube is already visible. The two precheliceral lobes come together and fuse along the dorsal midline (compare dotted line in Fig. 6d and e). In the anterior and all along the dorsal rim of the germ-band, some tissue is detectable that is growing dorsally to close the back (arrows in Fig. 6f,g).

Stage 12: dorsal closure

The main characteristic of stage 12 is the process of dorsal closure. However, as many morphogenetic events take place during stage 12, I have divided this stage into two sub-stages (stages 12.1 and 12.2).

General features of stage 12 embryos are the fully elongated and segmented prosomal appendages (Fig. 6e–l). Furthermore, the chelicera is now divided into a proximal base and a distal fang (Fig. 6h), and endites are growing from the coxae of the pedipalps and the walking legs (arrow heads in Fig. 6f). These endites have been described before, and the endites on the walking legs might represent a kind of ground pattern of the arachnid post-cheliceral appendage (Pechmann and Prpic 2009). In comparison to the endites of the walking legs, the endites on the pedipalps are bigger and will develop into the gnathendite that is required for food processing. At the end of embryonic development, no endite-like structures will be visible at the proximal base of the walking legs. Similar to the situation observed in the spinnerets, only the endite of the pedipalp expresses Distal-less and does not degenerate during embryonic development (Pechmann and Prpic 2009). Finally, brain development has also progressed further, and the anterior margin is a very prominent structure (Fig. 6f–h) during stage 12.

Stage 13: ventral closure

During stage 13, the ventral sulcus closes. Dorsal closure led to the formation of the prosomal shield and the heart tube (Fig. 6n,o). At the rim of the prosoma, a prominent ridge is visible (arrowhead in Fig. 6n and p). The fang of the chelicera is now very pointed (arrow in Fig. 6p). While the posterior spinnerets have grown to their final size, the anterior rudimentary bud of the spinneret is greatly reduced (ras in Fig. 6p). As opisthosomal segments 2–5 are greatly enlarged, the segments can be recognized as big “yolk sacs” (ys in Fig. 6n).
Stage 14: petiolus, cuticle secretion, pigmentation and muscle twitching

Ventral closure brings the left and right spinneret buds into a ventral position. The spinnerets are now directly anterior to the anus (Fig. 6q). Median spinnerets are visible (Fig. 6q). The petiolus has formed out of the first opisthosomal segment (pt in Fig. 6s). As the yolk is now mainly located in the opisthosoma, a flat prosoma has developed. In the centre of the prosomal shield, an indentation, the apodeme (a muscle attachment point), has formed (arrowhead in Fig. 6q,t). Cuticle secretion has started and the egg tooth gets secreted from the base of the pedipalp (Movie S7 and S10; arrows in Fig. 6q,r,t; Fig. S6a,b).

During stage 14, a dorsally and a ventrally located tooth (the so-called false pincer (Galiano 1996)) are also secreted from the distal tip of the fang of the chelicera (Fig. S5). The bristles of the appendages and the false pincer of the chelicera
Fig. 6 Embryonic development of A. geniculata (stages 11–14) a-d At stage 11 the sz is pointing towards the anterior (arrowhead in d). The ventral sulcus has a triangular shape with its narrowest point in the cheliceral segment. The labrum has started to form. Appendage buds on opisthosomal segments 2–5 are clearly visible. e-h Dorsal closure starts at stage 12.1 (see arrows in f and g). The left and right side of the opisthosoma fuse along the dorsal midline (compare dotted line in d and e). Endites are visible at the base of the appendages (arrowheads in f). The brain is framed by the anterior margin. The chelicera is clearly divided into the base and the fang (see asterisk in h). i–l At stage 12.2 dorsal closure is at an advanced stage (dotted line in k indicates the dorsal region that has not closed yet). The two head lobes start to fuse along the midline (arrow in l), and the chelicere of the left and right body half start to close over the labrum (l). During stage 12, only the appendage buds on the second and third (the book lungs) and the fifth (the posterior spinneret) opisthosomal segment grow and develop further. The rudimentary bud on o4 degenerates until the end of embryonic development. m–p Ventral closure starts at stage 13. The labrum sits below the cheliceres. Dorsal closure has led to the formation of the prosomal shield and the heart. The rim of the prosomal shield is pronounced (arrow head in n). The opisthosoma is divided into big “yolk sacks”. The fang of the chelicera is very pointed (arrowhead in p). q–t Stage 14. Ventral closure has led to the formation of a flat prosoma. The egg tooth (arrows in q, r and t) is visible and the petiolus has formed. The posterior spinnerets are in a ventral posterior position, close to the anus. Median spinnerets have formed. The apodeme is visible in the centre of the prosomal shield (arrowhead in q and t). All embryos were fixed and stained with the fluorescent nuclear dye Sytox green. Abbreviations: at, anal tubercle; am, anterior margin; bl, book lung; ch, chelicera; h, heart; l1–l4, walking legs; lr, labrum; ms, median spinneret; o1–9, opisthosomal segments 1–9; pcl, pre-cheliceral lobe; pp, pedipalp; ps, posterior spinneret; psh, prosomal shield; pt, peti- lus; ras, rudimentary anterior spinneret; sz, segment addition zone; st, stomodaeum; te, telson; ter, tergite; vs, ventral sulcus; yolk sack. Scale bar is 500 μm

Fig. 7 Marker gene analysis in embryos of A. geniculata. a–h Segmental expression of the segment polarity gene engrailed at stages 8–12. The double headed arrow indicates the width of the ventral sulcus in c and g. i Posterior expression of cad. j–l Wave like posterior expression of the pair-rule genes eve and h are indicated by the arrows in j, k and l. Segmental expression of eve in the posterior region of the ventral sulcus is indicated by the arrowheads in j and k. a–d, i and m are flat mounted embryos with false colour overlays of the indicated gene expression pattern. Abbreviations: bl02, book lung on opisthosomal segment 2; bl03, book lung on opisthosomal segment 3; ch, chelicera; l1–l4, walking legs; lr, labrum; o1–9, opisthosomal segments 1–9; pcl, pre-cheliceral region; pcl, pre-cheliceral lobe; pp, pedipalp; ps05, posterior spinneret on opisthosomal segment 5; ras, rudimentary anterior spinneret on opisthosomal segment 4; sz, segment addition zone; st, stomodaeum; vs, ventral sulcus. Scale bar is 500 μm

Postembryo

About 25 days after egg lay (analysed cocoons: n = 3; cocoons/embryos were incubated at 25 °C), embryonic development is completed, and the postembryo hatches from the egg (Movie S10; Movie S11; Fig. S6d–i). Hobbyists know this stage as “eggs with legs”. The hatching process itself is a relatively quick process (see Movie S11), and it takes only 30 min for the postembryo to completely hatch from the egg. In many spiders, hatching from the egg is connected to the first moulting from the embryonic cuticle (e.g., Mittmann and Wolff 2012; Wolff and Hilbrant 2011). I could not observe any embryonic cuticle in A. geniculata. Only the egg tooth (Fig. S6a–c) detached from the cuticle during the hatching process and was found at random positions within the stripped off eggshell or on the cuticle of the hatching. At the beginning, the postembryo is shiny and mostly unpigmented (Fig. S6d–f) but is able to move its legs (see Movie S11). Some postembryos already feed on non-hatched eggs (not shown). A few days later, the cuticle of the 1st instar larva develops under the postembryonic cuticle. As the postembryonic cuticle is transparent, one can easily see the pigmented hairs and eyes through the cuticle (Fig. S6g–i). In addition, the fang of the chelicera has developed and projects into the dorsal tooth of the false pincer of the postembryonic
1st and 2nd instar larva

After moulting from the postembryonic cuticle, the first instar larva is the first stage that is able to actively walk around. This stage takes about 5 weeks (analysed cocoons: \( n = 2 \); cocoons/1st instar larva were incubated at 25 °C). During this stage the larva is still consuming the rest of the yolk that is present within the opisthosoma. The spinnerets are now completely segmented. Internal organs like the silk glands are developing, and a few days after moulting into the first instar larvae, the spinnerets are able to produce first silk strands. At the end of this larval stage, the cuticle of the 2nd instar larva is secreted underneath the cuticle of the 1st instar larva. Urticating hairs are developing, and as soon as these hairs are getting pigmented, a circular spot is visible on the opisthosoma (Fig. S6k).

Moulting from the first instar larval, cuticle is leading to a small spiderling (the second instar larva; Fig. S6l). This little spider is now able to feed on small insects like *Drosophila* and will cannibalize on its siblings.

Axis duplication via cumulus grafts

Within the arthropods, axis patterning via a migrating signalling centre (the cumulus) seems to be an evolutionary novelty of chelicerates, and the presence of a cumulus in myriapods is still debated (reviewed in Hilbrant et al. 2012). As shown in this and several other studies, the cumulus of spider embryos migrates from the centre of the germ-disc towards the periphery of the disc and activates the BMP signalling pathway in ectodermal cells that are close to the cumulus (Fig. 4b, e.g. Akiyama-Oda and Oda 2006, 2003; Hilbrant et al. 2012; Oda and Akiyama-Oda 2008; Pechmann et al. 2017). This activation of the BMP signalling pathway in a subset of germ-disc cells is essential to break the radial symmetry of the disc and to initiate the formation of the dorsal field and the bilateral symmetry of the spider embryo (reviewed in Hilbrant et al. 2012; McGregor et al. 2008; Oda et al. 2019; Schwager et al. 2015). The importance of the cumulus and its signalling activity has been demonstrated in a variety of studies. First, the removal of the cumulus via extirpation, laser ablation or through genetic manipulation resulted in spider embryos that failed to break the radial symmetry of the germ-disc and failed to establish a dorsoventral body axis (Holm 1952; Oda et al. 2019; Pechmann et al. 2017). Second, inhibiting the activation of the BMP signalling pathway also inhibited the formation of an axially symmetric spider embryo (Akiyama-Oda and Oda 2006). Finally, the grafting of the cumulus led to the induction of a secondary body axis by inducing an ectopic dorsal field (Holm 1952; Oda et al. 2019). However, cumulus grafting experiments were so far only performed in araneomorph spider species (the funnel-web spider *Agelena labyrinthica* (Holm 1952) and the jumping spider *Hasarius adansoni* (Oda et al. 2019)), and functional analysis of the cumulus in basally branching spiders is entirely missing.

I performed cumulus grafting in about 50 *A. geniculata* embryos at mid to late stage 5. The ectopic cumulus was always placed within the germ-disc opposite to the endogenous cumulus, which was already migrating towards the rim of the disc (see Fig. 8a; Fig. 10b; Movie S12). The cumulus material from the donor embryo was clamped between the germ-disc and the vitelline membrane of the acceptor embryo. For technical reasons, cumuli were not transplanted below the ectoderm of the germ-disc. Nevertheless, the cumulus of the donor embryo regularly fused with the ectoderm of the germ-disc of the acceptor embryo (see Movie S12). As cumulus material from the donor embryo sometimes detached from the germ-disc of the acceptor embryo, not all analysed embryos showed an axis duplication phenotype. In addition, I failed to fix all of the cumulus-grafted embryos. In the end, I obtained 12 embryos that showed a complete or partial axis duplication phenotype. Figure 8 shows three examples of complete (Fig. 8a–e) or partial (Fig. 8g–j,l,m) axis duplication events after cumulus grafting. Embryos of the same cocoon that were not manipulated did not show any axis duplication (\( n > 100 \)).

Some cumulus-grafted embryos were tested for ectopic BMP pathway activity. In these embryos, 15–20 h after cumulus grafting, a strong ectopic pMad signal was detectable in the region around the cumulus graft (Fig. 8f).

In his comprehensive work, Holm already showed a huge variety of axis duplication phenotypes that can occur if cumulus material is placed at different regions within the germ-disc (Holm 1952). Also in my grafting experiments, I observed many different axis duplication phenotypes. Depending on the timing, the position within the germ-disc and the amount of grafted cumulus material, twinning is more or less perfect. A kind of perfect twinning event is depicted in Movie S12 and in the schematics shown in Fig. 10b. Here, the grafted cumulus material is big enough and fuses at the right time with the germ-disc of the acceptor embryo. This leads to an ectopic dorsal-field, which is very similar in size and shape to the endogenous dorsal field. As a result, convergent extension mechanisms drive the formation of two embryos that share one posterior pole (see Fig. 10b).

In spiders, only the prosomal segments are laid down during germ-disc stage and posterior segments are added sequentially via the posteriorly located segment addition zone (reviewed in Hemmi et al. 2018; Kauflholz and Turetzek 2018; McGregor et al. 2008). A perfect twinning event leads to the formation of two opisthosomata that share the saz at early germ band stages (see stages 8–10 in Fig. 10 and the drawings of Holm (Holm 1952)). However, as a separation event between the two opisthosomata seems to occur during advanced embryonic
development (see schematics in Fig. 10b and the embryo shown in Fig. 8c–e). Mis-patterning of one of the duplicated opisthosomata was frequently observed (see Fig. 8d,e). In the twinned embryo shown in Movie S12 and Fig. 8a–e, it is likely that only one opisthosoma duplicate took over most portions of the saz. As a result, only one opisthosoma duplicate showed a wild type-like morphology, whereas the other duplicate was slightly deformed (marked with an asterisk in Fig. 8d,e).

Fusion of prosomal segments and imperfect twinning of embryos was probably due to the fact that grafted cumulus material was not big enough or that the fusion of the grafted material did take place at the wrong place and time. In these cases the developing embryos shared a single wild type looking opisthosoma (Fig. 8g–j,l,m; schematics in Fig. 10c).

The results show that not only in true spiders like A. labyrinthica (Holm 1952) and H. adansoni (Oda et al. 2019) but also in the basally branching tarantula A. geniculata, cumulus grafting is leading to an axis duplication phenotype. In spiders, the cumulus is the source of secreted Decapentaplegic protein (Dpp; homologous to the vertebrate

Axis duplication via local activation of the BMP signalling pathway

In spiders, the cumulus is the source of secreted Decapentaplegic protein (Dpp; homologous to the vertebrate...
BMP2/4 protein), and the knockdown of dpp or the loss of the cumulus results in embryos that fail to break the radial symmetry of the germ-disc (Akiyama-Oda and Oda 2003; Akiyama-Oda and Oda 2006; Pechmann et al. 2017).

Regarding the functional analysis of the cumulus, only loss of function experiments has been performed so far, and the final proof that Dpp is sufficient to induce the establishment of the dorsal field is still missing. It has been shown that arthropod Dpp is able to activate the BMP signalling pathway in vertebrate species and vice versa (Padgett et al. 1993; Sampath et al. 1993). For this reason, I have transplanted human BMP4-soaked agarose beads to the rim of the germ-disc of stage 5 A. geniculata spider embryos. To better mimic the function of the endogenous cumulus, beads were transplanted below the ectoderm of the disc. As a control, BSA-soaked beads were transplanted to embryos of the same cocoon. Embryos were fixed 15–20 h after bead transplantation, and the activation of the BMP signalling pathway was analysed.

Fig. 9 Axis duplication via local BMP pathway activation. a,a’ A BSA-soaked agarose bead is not able to locally activate the BMP signalling pathway. pMad signal is only visible in the region of the dorsal field (df). a’’ Magnified view on the region of the transplanted bead. b,e Embryos develop normal and show no axis duplication. d,d’ An agarose bead soaked with human BMP4 is able to locally induce the BMP signalling pathway. d” Magnified view on the region of the transplanted bead. f-h Three embryos in which a single BMP4-soaked bead (f,g) or multiple BMP4-soaked beads (h) have been transplanted. Nuclear pMad signal is detectable in a broad region around the transplanted beads. e,e’,j In the shown embryos, a BMP4-soaked bead was transplanted to the germ-disc (opposite to the cumulus). The local activation of the BMP signalling pathway did lead to axes duplication phenotypes during further embryonic development. a,a’,d,d”,f-h Bright field (the agarose bead is visible as a blue spot). a”,b,e,i,e” Nuclear Sytox green staining (the agarose bead is visible as a black spot). e,e”,j Flat mounted embryos. Same embryo in e (whole mount) and e’ (flat mount). Abbreviations: ch, chelicera; df, dorsal field; f14, fused 14; 11-14, walking legs; pp, pedipalp; saz, segment addition zone. Scale bar is 500 μm.

### Table 1 Analysis of the activated BMP signalling pathway in cumulus-grafted and bead-transplanted embryos

| Embryos          | pMad signal around the endogenous cumulus/in the endogenously induced dorsal field | pMad signal around the grafted cumulus/in the ectopically induced dorsal field | pMad signal around the transplanted bead | Number of analysed embryos |
|------------------|----------------------------------------------------------------------------------|---------------------------------------------------------------------------------|------------------------------------------|---------------------------|
| Embryos with ectopic cumulus graft | 10                                                                               | 9                                                                              | n/a                                      | 10                        |
| BSA bead-transplanted embryos       | 29                                                                               | n/a                                                                            | 0                                        | 29                        |
| BMP4 bead-transplanted embryos     | 32                                                                               | n/a                                                                            | 16                                       | 32                        |

Embryos were analysed for nuclear localized pMad 15–20 h after cumulus grafting or bead transplantation, respectively. Please note, only embryos in which the ectopic cumulus graft fused to the germ-disc were used for the analysis.
segment addition zone occurs (compare to the embryo shown in Somewhere between stage 10 and stage 12, a separation event of the are leading to a double-headed embryo that shares the posterior pole. Convergent extension mechanisms (indicated by the arrows at stage 6) grafted cumulus (gra-cu) is able to induce a secondary body axis. the germ-disc has the potential to become the anterior of the embryo, a hypothetical situation of an ideal twinning event. As the complete rim of sulcus (white area in st. 9 – and appendages grow out from the germ-band (st. 8 and 9). The ventral convergent extension mechanisms (indicated by the arrows at stage 6) are the rim of the germ-disc and is inducing the dorsal-field (df). Extensive type development. At the end of stage 5, the cumulus (cu) has shifted to

via pMad antibody staining. In 100% of the control embryos, no pMad signal was detectable around the transplanted bead (n = 29; see Table 1; Fig. 9a–a’). In contrast, the transplantation of BMP4-soaked beads led to a detectable pMad signal in 50% of the analysed embryos (n = 32; see Table 1; Fig. 9d–d’,f–h). This result shows that human BMP4 is able to locally induce the BMP signalling pathway in the germ-disc of A. geniculata germ-disc stage embryos. However, the direct comparison between cumulus-grafted and BMP4 bead-transplanted embryos showed that an ectopic cumulus graft (a strong pMad signal around the cumulus graft was visible in 90% of the embryos) is more reliably inducing the BMP signalling pathway in manipulated embryos (see Table 1).

Nevertheless, although a pMad signal in late stage 5/early stage 6 embryos was only detectable in 50% of the BMP4 bead-transplanted embryos, 85% of the transplanted embryos did show an axis duplication phenotype during later stages of development (n = 40; e.g., see Fig. 9e,e’,i–j). The remaining 15% of the BMP4 bead-transplanted embryos showed a wild type or an unspecific phenotype (not shown). This indicates that the BMP signalling pathway around the transplanted bead was only slightly activated (below the detection level via pMad antibody staining) in a large portion of the embryos. Control, BSA-soaked beads (e.g., Fig. 9b,c) did never induce an axis duplication phenotype (n = 38; 36 embryos were wild type-like; 2 embryos showed an unspecific phenotype).

In contrast to the cumulus transplantation (see Fig. 8a–e), I never observed a “perfect” axis duplication phenotype upon BMP4 bead transplantation. Most embryos showed an axis duplication event within the prosoma of the embryo (Fig. 9e,e’,i–j). As a result, fusion of prosomal appendages was frequently observed (Fig. 9e, e’,i–j; Fig. 10c). One of the strongest phenotypes upon BMP4 bead transplantation is shown in Movie S13. In this embryo almost the entire prosoma was duplicated and only the central part of the fourth walking leg segment was fused.

Overall, these results are the final proof that the local activation of the BMP signalling pathway within the germ-disc is sufficient to induce a secondary body axis in spider embryos.

Conclusions and future directions

A. geniculata is a promising model system that can be used to answer a variety of evolutionary developmental questions. It can be reared in the lab throughout the year, genomic and transcriptomic sources are available (this work; Sanggaard et al. 2014), and the embryos are amenable to standard molecular biological techniques like in situ hybridisation and antibody staining (this work; Pechmann and Prpic 2009). Future studies will show if it is possible to use embryonic RNA interference to analyse gene functions. The observation
that RNA probes are cross hybridising across tarantula genera (Fig. S7) might help to compare gene expression pattern across mygalomorph species.

It will be interesting to see whether anterior as well as posterior segmentation mechanisms are conserved between mygalomorph and araneomorph spider species. Also the timing of the establishment of the cumulus seems to be different between mygalomorph and many araneomorph spider species, and it will be interesting to see whether the same set of genes is expressed in with the cumuli of distantly related spiders.

In this study I have demonstrated that A. geniculata embryos are well suited to perform grafting and transplantation experiments. In future studies, especially the bead transplantation experiments will allow analysing the local activation of different signalling pathways at different regions within the developing spider embryo.

Acknowledgements I thank Siegfried Roth and Matthew Benton for critical discussions and comments on the manuscript. A special thanks goes to Nathan Kenny for assembling and analysing the transcriptome. I also thank the two reviewers for their constructive comments and suggestions.

Author contributions MP designed the study, performed the experiments, analysed the data and wrote the manuscript.

Funding information Open Access funding provided by Projekt DEAL. This work has been funded by the Deutsche Forschungsgemeinschaft (DFG grant PE 2075/1–2).

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