Cribellate thread production as model for spider’s spinneret kinematics

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Abstract
Spider silk attracts researchers from the most diverse fields, such as material science or medicine. However, still little is known about silk aside from its molecular structure and material strength. Spiders produce many different silks and even join several silk types to one functional unit. In cribellate spiders, a complex multi-fibre system with up to six different silks affects the adherence to the prey. The assembly of these cribellate capture threads influences the mechanical properties as each fibre type absorbs forces specifically. For the interplay of fibres, spinnerets have to move spatially and come into contact with each other at specific points in time. However, spinneret kinematics are not well described though highly sophisticated movements are performed which are in no way inferior to the movements of other flexible appendages. We describe here the kinematics for the spinnerets involved in the cribellate spinning process of the grey house spider, Badumna longinqua, as an example of spinneret kinematics in general. With this information, we set a basis for understanding spinneret kinematics in other spinning processes of spiders and additionally provide inspiration for biomimetic multiple fibre spinning.

Keywords Cribellate capture thread · Functional morphology · Biomimetic · Fibre processing · Spigot

Introduction
Spider silk captivates researchers for many decades now, not least because of its extraordinary material properties (Vollrath 2000; Eisoldt et al. 2011). One of the most studied characteristics to uncover the principle behind the high toughness of spider silk is its molecular structure, which enhances toughness by a combination of elastic and crystalline domains in the protein (Termonia 1994; Hayashi et al. 1999; Becker et al. 2003; Mortimer and Holland 2015). However, there is not a single type of spider silk: Spiders possess up to nine different silk glands, each of which produces a specific type of silk with equally distinct properties (Kovoor and Peters 1988; Coddington 1989; Vollrath 1992). These different types of silk are used for various purposes and can even be combined, e.g., to a spider’s web or attachment discs (Foelix 2011). To anchor their threads to a substrate, e.g., spiders specifically combine several fibre types, influencing the mechanical properties of the final attachment disc by this interweaving (Wolff et al. 2015). However, the most complex combination of different silk types to a single thread can be observed in the assembly of capture threads of cribellate spiders. With these threads, cribellate spiders catch their prey without the use of viscous glue. By involving nanofibres, adhesion is achieved by a combination of van der Waals and capillary forces, through which the cuticular waxes of the prey are absorbed (Hawthorn and Opell 2002; Bott et al. 2017). Up to six different types of silk, including thousands of nanofibres (Ø 10–30 nm) and several larger fibres, are connected to complex three-dimensional structures (Peters 1992; Eberhard and Pereira 1993; Opell 2002; Joel et al. 2015; Grannemann et al. 2019). The use of different fibres influences the mechanical properties of the capture threads beyond the molecular properties of their individual
silks, comparable to a fibre-composite (Michalik et al. 2019; Piorkowski et al. 2020).

The silk fibres are extruded from spigots on the spinnerets of a spider. Spinnerets are segmented appendages on the opisthosoma of a spider (Mariano-Martins et al. 2020). They are articulated to the body, and their segments are connected to each other, via movable joints. Most of the spiders possess three pairs of spinnerets: anterior lateral (ALS), posterior median (PMS), and posterior lateral (PLS) spinnerets (Montgomery 1909; Glatz 1972, 1973; Platnick et al. 1991). In addition, cribellate spiders possess a further spinning organ, the cribellum (Cr). It can be described as a transverse and moveable structural unit covered with spigots which is situated anteriorly of the other spinnerets (Blackwall 1839; Bertkau 1882; Montgomery 1909). In contrast, some ancient spiders, such as the Mesothelae, possess a further pair of spinnerets, the anterior median spinnerets, at the corresponding position of the cribellum in cribellate spiders. This fourth pair of spinnerets as well as the cribellum embryologically develop from the same primordia, which implies a homology of both structures, thus, possibly featuring similar manoeuvrability (Pechmann et al. 2010; Foelix 2011). Previous studies of the cribellate capture thread production already proved elaborate manoeuvrability of the spinnerets coupled with a highly coordinated sequence of movements in three-dimensional space to create such complex structures (Joel et al. 2015; Grannemann et al. 2019). The mobility of the spinnerets is achieved by muscles located at different attachment sites (Glatz 1972, 1973; Magalhães et al. 2017). By an increase of haemolymph pressure, some movements can be controlled by a hydraulic mechanism, as described for the movement of legs (Kropf 2013; Labarque et al. 2017). Interestingly, apart from some general descriptions of movements and spinneret morphologies, hardly anything is known about the kinematics of spinnerets. However, the morphology of the spinnerets coupled with their spatial movements is crucial for the accessibility of different spigots extruding the specific fibres and thus the interweaving of fibres (Eberhard 2010).

Though the evolutionary origin of the spinnerets is still open to discussion, several facts hint to originally locomotor appendages which lost their primary function (Shultz 1987; Hilbrant 2008; Pechmann and Prpic 2009). It is conspicuous that spinnerets have a segmentation and complex musculature and thus achieve a flexibility that is in no way inferior to that of the legs (Blackledge et al. 2011). There are a few works on the kinematics of spider legs describing the locomotor system via movement radii, length ratios, and co-ordinates (Ehlers 1939; Weihmann et al., 2010; Foelix 2011). But in contrast to the legs, the spinnerets interact with each other and can create three-dimensional structures. Hence, we aim to analyse the spinneret kinematics using the cribellate spinning process, in which almost all spinnerets are involved in the process and a highly repetitive three-dimensional structure is created as a thread. We investigated the spinneret kinematics of the synanthropic cribellate grey house spider Badumna longinqua (Desidae). By solving the complex choreography and uncovering conserved conditions of the process, we hope not only to unravel the general spinneret kinematics, but also to inspire future biomimetic application for complex small-scale fibre spinning techniques.

Material and methods

Study animals

Specimens of Badumna longinqua (L. Koch, 1867) were collected in public parks in Eastern Australia (Sydney (NSW), Brisbane (QLD)) and kept separately in plastic boxes with a roughened surface at Macquarie University, time-shifted by 12 h. Once a week spiders were fed with Drosophila melanogaster and water was provided once to twice per month by sprinkling the enclosure. Additionally, spiders were exported (permission no. PWS2019-AU-000248) to Aachen (Germany) and kept at elevated room temperature (~26 °C), room humidity (30%), and northern European diurnal rhythm. They were fed with juvenile Acheta domestica or Callosobruchus maculatus. Water was provided as described above.

Video recordings

The spinning apparatus was studied during cribellate thread production using a video camera (Sony Alpha 6300), equipped with a 4× magnification lens of a microscope and stabilized with a magic arm (Manfrotto) at Macquarie University. Videos were recorded with 120 fps under either white light or red light (less disturbance of night-active spiders) with a 12 h shifted day–night cycle. Recordings were performed from different perspectives, if possible from a ventral and lateral view. Video sequences always covered several spinning cycles. Additionally, the general mobility of the spinnerets was recorded using a high-speed microscope (Keyence VW-600C) with the corresponding Software (Keyence VW-9000 Motion Analyzer, Version 1.4.0.0) at a frame rate of 60–250 fps. For this purpose, the spiders were first anaesthetised with diethyl ether and then trapped using minutiae on a preparation dish to immobilise the body and legs.

Scanning electron microscopy

Of the recorded individuals, either the exuvia were collected or living specimens were placed in the freezer (-20 °C) until they were frozen. Frozen specimens were then immediately
transferred to ethanol (70%) and subsequently dehydrated over a series of increasing concentrations of ethanol (80, 90, 95, and 100%). Following, the samples were dried by hexamethyldisilazane (HMDS, ≥ 98%) in increasing concentrations (ethanol: HMDS; 3:1, 1:1, 1:3, 0:1). The concentration step 0:1 was repeated two times. The liquid was then removed, and the samples were air-dried in a fume cupboard. Before examination using a scanning electron microscope (SEM 525 M; Philips AG), samples were sputter-coated with a 10 nm layer of Au (Hummer Technics Inc.).

**Data analysis**

Abstracted models of the spinnerets were created using SEM images (Online Resource 4), video stills, and the software Inkscape (Version 1.1.1, Inkscape Community). Measurements of the length of spinneret legs and bodies were performed on SEM images using a combination of the software Shotcut (Version 20.02.17), GIMP (GNU Image Manipulation Program, Version 2.10.18), and Snipping Tool (Version 10.0.18363, Microsoft Corporation). The resulting data were used as a reference for further distance measurements within the videos. Movements of structures involved in the spinning process could thus be converted into velocities over distances covered within a certain period of time. Velocities were determined on three combing cycles (N = 3) of one individual (n = 1) and listed as means. Data acquisition and statistical analysis were performed using Microsoft® Office Excel® 2020 (Version 16.0.12527.20612, Microsoft Corporation).

**Characterization of the capture thread**

For SEM examinations, capture threads of *B. longinqua* were transferred to sample holders with cut-to-size conductive foil, enabling the thread to hang free between the foil strips. Care was taken to ensure an undamaged and non-stretched transfer of the threads in order to obtain a sample as representative as possible. The samples were examined without sputtering. Cribellate threads without coating charge up quickly in the SEM and can therefore only be recorded with low contrast, but unlike sputtered samples, they retain their original shape and do not collapse (Joel and Baumgartner 2017). Thus, internal fibre types can be identified in the loose mass of nanofibers.

**Preface on kinematics**

Kinematics are also described as the “geometry of motion”. An important aspect of kinematic descriptions includes the degrees of freedom, aside from geometries and velocities to specify all independent possibilities of movement in a physical system. A rigid body in space owns six degrees of freedom along the three axes of an imaginary rectangular cross-axis: Three independent directional movements (translation) and three independent axis rotations (rotation) (Fig. 1a). Any fixation, in a point or plane, reduces the number of degrees of freedom. If several subsystems are connected, e.g., via joints, their individual degrees of freedom are reduced. However, the freedoms of a complete system are the result of an addition of all individual degrees of freedom of each subsystem.

**Results**

**Basic kinematics of spinnerets**

The simplest way of describing a spinneret kinematically is to consider it a rigid body with only one joint at its base. For this abstraction, we considered a possible elastic deformation of the stiff exoskeleton negligible and the simplification to a rigid body appropriate. Otherwise, the number of degrees of freedom of a flexible body would, from a strictly physical point of view, tend to infinity. With this abstraction of a rigid body respecting only one joint, the radius of movement approximately corresponds to a hemisphere (restricted only by the body of the spider) and depends on the total height of the spinneret (Fig. 1b).

An example of such simple spinnerets are the posterior median spinnerets of the grey house spider *Badumna longinqua*. They are situated between anterior lateral and posterior lateral spinnerets and consisted each of one segment (Fig. 1e). The segments exhibited a pyramidal shape with a flattened apical site bearing different spigots on it (Fig. 1b). A row of paracribellate spigots, involved in the cribellate spinning process, was situated in the middle of a field of other spigot types, not involved in the process (Fig. 1e). The shafts of these paracribellate spigots originated from a shared base, whereas one further but single paracribellate spigots flanked the minor ampullate spigot (Fig. 1c,d). This dyad was situated at the proximal margin of the field of spigots. The mobility of the posterior median spinnerets was restricted in some directions. On an imaginary x- and y-axis, their position was unalterable. However, the z-axis remained for a directional movement so that the spinnerets could move up and down. Translationally, the posterior median spinnerets possessed therefore one degree of freedom. Additionally, only two rotations were possible: “rolling” to the left and right and “pitching” to back and forth (from posterior to anterior) (Online Resource 1). A “yawing” (a rotation around its own axis), did not occur. As a result, two rotational degrees of freedom, in addition to the translational degrees of freedom, resulted in three degrees of freedom for the posterior median spinnerets in total (Fig. 1b).
Fig. 1 The degrees of freedom of the spinning apparatus of *Badumna longinqua*. 

(a) In three-dimensional space, a freely movable rigid body possesses six degrees of freedom (DOF) due to translational motion along the axes of an imaginary rectangular cross-axis: up/down, left/right, back/forward and the axis rotations: yawing, rolling and pitching. 

(b) A posterior median spinneret (PMS) owns one joint with a restricted number of DOF as the spinneret can only move up and down, roll to the left and right and pitch back and forth. Its radius of movement corresponds to a hemisphere.

(c) On the PMS two different types of paracribellate spigots (PC) occur: One elongated PC and a row of several PC where multiple shafts originate from one base.

(d) The spinning apparatus consists of PMS, anterior lateral (ALS) and posterior lateral (PLS) spinnerets, as well as a pseudo-bipartite cribellum (Cr), containing two separated fields of cribellate spigots (CS) on a common base. Besides several PC, the PMS contain each a minor ampullate spigot (mAP). The PLS contain each a triad of one pseudoflagelliform spigot (PF) flanked by two PC. Spigots involved in the cribellate spinning process are highlighted in black, spigot fields of uninvolved spigots are represented in grey areas.

ALS: anterior lateral spinneret, Cr: cribellum, CS: cribellate spigot, DOF: degrees of freedom, mAP: minor amullate spigot, PC: paracribellate spigot, PF: pseudoflagelliform spigot, PLS: posterior lateral spinneret, PMS: posterior median spinneret.
In addition to the simple movement of a single joint spinneret, as described for the posterior median spinnerets, the posterior lateral spinnerets were both composed of two segments (Fig. 2). The basal segments were long with a constant diameter and articulated to the body via one joint, respectively. The terminal segments were tapered bearing spigots on their inwards directed sides and articulated to the basal segment by a second joint surrounded by an elastic membrane (Fig. 2b). A pseudoflagelliform spigot flanked by two paracribellate spigots was situated apically on each posterior spinneret (Fig. 1e). The posterior lateral spinnerets could either be adducted by an antero-medial alignment or abducted in postero-lateral direction. During complete abduction, the posterior lateral spinnerets were able to incline almost rectangular to the body; Scale bar 500 µm. The PLS consisted of two joints, which created two axes of inclination. Considering a segmented spinneret with two fixed hinge points the movement range of the basal segment would decrease to a hemisphere according to the length of the basal segment while the range of the terminal segment could, in theory, proximally increase represented by an almost spherical range of movement (dotted line). In reality, the range of movement only corresponded to a further hemisphere. An elastic membrane between the segments provided for a mobile positioning of the hinge points. The range of movement could therefore either be further increased or decreased.

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the respective body, thus enabling further positioning by reducing the total length of the spinneret (Fig. 2d). The degrees of freedom of both joints combined, the posterior lateral spinnerets possessed six degrees of freedom (Fig. 2b).

The cribellum, anterior of the other spinnerets, consisted of a flat and oval-shaped base on which two separated and elongated fields of uniformly distributed cribellate spigots were placed (Fig. 1e). The shared base of this pseudo-bipartite cribellum could incline in a posterior or anterior direction and lateral direction to the left or right (Fig. 3a, b). Additionally, the cribellum could be either turned outwards or slightly retracted into the body (Online Resource 1). As described for the other spinnerets, this flexibility was achieved by one joint, enabling one-directional movement and two rotations, thus, creating three degrees of freedom. The radius of movement could be represented by a flattened hemisphere (Fig. 3d). Besides, the two fields of spigots exhibited one further rotational movement independent from their base. During the cribellate spinning process, each field was able to bulge out with their medial sides. We illustrated this further movement by two additional hinge points on the lateral margin of the two fields of spigots, which could incline by a few degrees (Fig. 3c). Therefore, a further degree of freedom is assigned to each half of the cribellum.

**Interplay and limits of motion during the cribellate spinning process**

The cribellate spinning process is a highly conserved sequence of movements that could be subdivided into individual repeating cycles (Fig. 5). Each cycle began with a specific spinneret constellation and proceeded in a highly coordinated choreography with each interplay at a specific point in time (Fig. 5c; Online Resource 3). We defined a cycle by the start of the combing leg at its most anterior position, because the spider always started the cribellate thread production at this position (Fig. 5a, d-f). From this position, the combing leg moved in a posterior direction combing over the spinnerets, changed direction back to anterior, and returned to its initial position (Fig. 5d–f). The return movement was not performed if the spider finished the thread production or was interrupted. As already described above for the basic considerations of the individual structures, all spinnerets, and the cribellum could either be completely adducted or abducted independently or occupy any intermediate position (Fig. 4a, b). In contrast to the observation of

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**Fig. 3** Movements of the cribellum. a In the anaesthetised spider, the cribellum (Cr) inclined to posterior or b anterior, as well as to lateral directions; Scale bar 500 µm. c Due to a fused base, the Cr moved according to one joint with two further angles of inclination enabled by a possible pull out of the right and left spigot fields. d The movement range of the Cr corresponded to a flattened hemisphere

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an individual spinneret, the radii of movement in an interde- 
pended system, consisting of six spinnerets and a cribellum, 
are mutually limited or do even overlap. During cribellate 
thread production, the anterior lateral spinnerets remained 
in an abducted position, thus clearing the way for the move-
ments of the other spinnerets (posterior lateral and posterior 
median spinnerets), the cribellum (Fig. 4b) as well as the 
combing leg bearing the calamistrum.

The posterior median spinnerets started their spinning cycle adducted and in a medial position perpendicular to 
the body (Fig. 5e). At a velocity of 2.44 mm/s, they first 
moved in posterior direction by a pitching rotation. This 
movement occurred at the same time as the posterior move-
ment of the posterior lateral spinnerets so that the poste-
rior median spinnerets were spatially not restricted. Nev-
evertheless, the posterior movement ended after only a slight 
bending backward to posterior, not using the full possible 
radius of motion. From this position on, the direction of the 
posterior median spinnerets changed to anterior at a veloc-
ity of 11.45 mm/s. Simultaneously, the posterior median 
spinnerets were abducted and thus spread laterally. Due 
to the abduction of the anterior lateral spinnerets and their 
greatest possible clearance, the posterior median spinner-
ets were able to tilt far forward beyond their initial position 
and aligned themselves further anteriorly. With a velocity of 
1.06 mm/s, the posterior median spinnerets finally returned 
posteriorly to their initial adducted position again via a 
change of direction.

The spinning cycle of the posterior lateral spinnerets 
began in a posterior position with the basal segments being 
abducted (Fig. 5d). The posterior lateral spinnerets were 
only slightly tilted backward so that their widest possible 
radius perpendicular to the body was not exploited. Even 
though the terminal segments of the posterior lateral spin-
nerets could theoretically invade the basal parts, e.g., this 
option was not used during the cribellate spinning process. 
By tilting inwards, the terminal segments folded slightly 
towards the basal segment and pointed more and more ante-
riorly. From this position, the posterior lateral spinnerets 
moved in almost uniform motion with 6.1 mm/s in antero-
medial direction. The anterior movement ended as soon as 
the basal segment reached a position approximately perpen-
dicular to the body. At that time, the posterior lateral spin-
nerets approached the posterior median spinnerets closely so 
that their movements were restricted by the posterior median spinnerets anteriorly. The length of the posterior lateral spin-
nerets, in addition to their independent segment movements, 
nevertheless allowed the terminal segments of the posterior 
lateral spinnerets to protrude above the posterior median 
spinnerets with their spigots pointing to the spigots of the 
posterior median spinnerets (Fig. 5c). From this position on, 
the posterior lateral spinnerets changed their direction and 
returned to posterior at a velocity of 3.92 mm/s. The termi-

Fig. 4 Two-dimensional illustration of the spinning apparatus (ven-
tral view). a Most adducted and abducted configurations of posterior 

lateral (PLS), posterior median (PMS), anterior lateral (ALS) spin-

nerets, and cribellum (Cr) represented by overlapping contours of the 

respective spinnerets. b During the cribellate spinning process, the 

a

b

ALS rested in an abducted position while PLS, PMS, and Cr moved 

and occupied different positions shown by transparent contours. ALS: 

anterior lateral spinneret, Cr: cribellum, PLS: posterior lateral spin-

neret, PMS: posterior median spinneret
lateral spinnerets or the cribellum used their complete possible range of motion during the cribellate spinning process. In each cycle, the different spinnerets started at specific positions and moved alternately closer together or further apart. Due to a distinct positioning of the different types of spigots involved, the choreography of the spinnerets, nevertheless, accomplished specific contacts at specific points in time and space. The controlled movement enabled close contact between the pseudoflagelliform spigots on the posterior lateral spinnerets and the minor ampullate spigots on the posterior median spinnerets as well as the paracribellate spigots on both spinnerets.

Fig. 5 Movements of the cribellate spinning process. a During posterior movement of the combing leg (CL), the posterior lateral spinnerets (PLS) moved towards the CL with their terminal segments pointing to anterior; the posterior median spinnerets (PMS) were straightened and followed the movements of the CL; the cribellum (Cr) moved towards posterior. b Just before a change of the CL in anterior direction, the PLS were almost straightened and in an abducted posterior position; the PMS moved towards anterior; the Cr inclined to anterior, as well. c Model of spinneret movements and spigots involved (spigots involved represented by black dots; spigots not involved represented by the whole spinning field in grey): on the PLS a pseudoflagelliform spigot (PF) was flanked by two paracribellate spigots (PC), during abduction the spigots were far away from the other spigots and spinnerets, during adduction the spigots came into close contact with the spigots on the PMS; on the PMS a row of PC and a single PC flanking the minor ampullate spigot (mAP) were situated; on the Cr two separated fields of cribellate spigots (CS) occurred. d Movement velocities of the CL and the PLS during cribellate spinning process: a combing cycle started with the CL at its most anterior position in posterior direction (grey arrow), at the same time the PLS were at a posterior position and moved towards anterior (dark grey arrow). The CL changed direction at its most posterior position returning with a faster movement to its initial position. The PLS returned with a faster movement from its most anterior position in posterior direction, but slowed down and again shortly raised its velocity before the start of a new combing cycle. e Movement velocities of the CL and the PMS: at the start of a combing cycle, the PMS were in a medial position and moved towards posterior. At its most posterior position it changed direction with a fast movement and slowed down until it reached its most anterior direction where it returned with a fast movement in posterior direction and slowed down by reaching its initial position. f Movement velocities of the CL and the Cr: at the start of a combing cycle the Cr was covered by the CL so that a velocity could not be determined (dotted line). From a posterior position, it inclined to anterior with a uniform movement and returned to posterior again by a sudden movement before reaching its initial position, sample size: d–f. (n = 1). (N = 3). CL: combing leg, Cr: cribellum, CS: cribellate spigot, mAP: minor ampullate spigot, PC: paracribellate spigot, PF: pseudoflagelliform spigot, PLS: posterior lateral spinneret, PMS: posterior median spinneret.
Capture thread of *B. longinqua*

The capture thread of *B. longinqua*, created during the spinning process, consisted of two separate strands, which were placed either parallel or detached from each other in the web (Fig. 6). The separation resulted from the bilateral arrangement of the spinning apparatus together with the bipartite cribellum, which was particularly evident in the recordings of the spinning process (Fig. 6b). Each of the strands incorporated a single axial fibre (af), an undulating fibre (uf) and a mat of cribellate fibres (cf), which was alternately denser and more loosely packed (Fig. 6a).

**Discussion**

Spinnerets move with astonishing manoeuvrability. We studied the kinematics of spinnerets and revealed the spatio-temporal movement possibilities during the cribellate spinning process. The general kinematics of spinnerets, besides, indicate their close relationship to the locomotor system and provide information for other silk-spinning processes. In addition to a transfer to other spinning processes in spiders, this information could also serve as biomimetic inspiration for small-scale fibre assembly.

**Transfer of our model to other spinning systems**

The movements observed in the spinnerets of *Badumna longinqua* confirmed a theoretical model of spinneret kinematics very nicely: the range of movement of a one-joint spinneret, such as the posterior median spinnerets, is restricted to a space determined by the length of the spinneret and limited by the body on which the spinneret rests. The movements of the cribellum strongly resemble the flexible movements of a spinneret. A possible evolution from an ancestrally fourth pair of spinnerets has already been suggested due to the different variants of a cribellum (Pechmann et al. 2010). In some species, it has been described that the cribellum can be either undivided with a continuous field of spigots (e.g. in *Uloborus plumipes*) or split into two (pseudo-bipartite; e.g. in *Badumna longinqua*) to several fields of spigots (e.g. in *Dresserus*) (Griswold et al. 2005; Alfaro et al. 2018). We observed that a cribellum possesses almost the same number of translational and rotational movements as a single posterior median spinneret. By an additional bulging of the spigot fields, the cribellum of *B. longinqua* even exhibited a further movement axis on each half of the cribellum. Restrictions due to the merged base of the cribellum are at least partially reduced by this additional mobility. Such a movement has not been described before and could be achieved by either internal musculature or an increase of haemolymph pressure. A multi-joint spinneret, such as the 2-joint posterior lateral spinneret in *B. longinqua* and its formation. The thread consisted of two strands, which could be either a closely parallel or b separate. a Each strand (left and right half) contained one axial fibre (af), one undulating fibre (uf) and many cribellate fibres (cf) formed into a mat; Scale bar 200 µm. b The halves of the cribellate thread (CT) were produced separately right from the beginning. The emergence of individual fibres could be seen at the posterior lateral (PLS) and posterior median (PMS) spinnerets. The final thread structure became visible after processing by the combing leg (CL). af: axial fibre, cf: cribellate fibres, CL: combing leg, CT: cribellate thread, PLS: posterior lateral spinneret, PMS: posterior median spinneret, uf: undulating fibre
spinneret, effects further positioning as several inclinations achieve more combinations of movements. It can be thus concluded that the type and number of joints directly affect the flexibility of a spinneret. However, 2-joint spinnerets are not the most flexible spinnerets that exist: depending on the species, a posterior lateral spinneret possesses up to 4 (e.g. in *Nemesia*), in *Mesothelae* (e.g. in *Liphistius*), even up to 12 segments with the corresponding number of joints (Marples 1967; Glatz 1973). Besides, most of the anterior lateral spinnerets possess 2 to 3 segments (in *Liphistius* again many more) (Marples 1967; Glatz 1972, 1973). The agility of the spinnerets can, therefore, be refined.

Nevertheless, a limiting factor of spinneret movements is a mutual interference in the overall system (i.e. the spinning apparatus). The spinnerets are grouped very close to each other. If certain action ranges of different spinnerets overlap, space can only be occupied if it is released by another spinneret. The extent to which a restriction is expressed depends on the movement of the adjacent spinnerets at the same time: the further, e.g. the posterior median spinnerets tilt anteriorly, the higher the manoeuvrability of the posterior lateral spinnerets. By protrusion, as possible for the posterior lateral spinnerets to protrude above the posterior median spinnerets, ranges of motion may even overlap without competing. The length of the segments is, therefore, another crucial factor for spinneret movements. During the cribellate spinning process, the abduction of the anterior lateral spinnerets enables free movement of the other spinnerets by clearing possible spatial positions. Due to its flat nature and anterior position in front of the other spinnerets, the cribellum as well is not visibly restricted during the cribellate spinning process. However, during the production of the attachment discs, e.g., the movements of the anterior lateral spinnerets can be restricted anteriorly by the presence of the cribellum (Wolff et al. 2019). As there is hardly any information on spinneret kinematics for other spinning processes, further restrictions caused by spinnerets cannot be excluded. Even a difference in the size proportions of the spinnerets in different species could render certain movements more problematic (Eberhard 2010). However, the general kinematics may already be estimated very well based on the models described here.

**MOTORIC AND SENSORY CONTROL OF SPINNERET KINEMATICS**

Besides geometrical descriptions and positions of the spinnerets, the velocity of movements represents an important value to define the kinematics of spinnerets. The velocity of motions during the cribellate spinning process is astonishingly high. During the production of attachment discs in cribellate spiders, the maximum velocity of the anterior lateral spinnerets ranges from 0.6 (e.g. in *Hickmania troglodytes*) to 34 mm/s (e.g. in *Oecobius navus*) (Wolff 2020). Maximum velocities of the spinnerets and the cribellum during cribellate thread production between 11 and 18 mm/s in *B. longinqua* are in the mid-range and seem to be species-specific. We assume the maximum velocity might be restricted to the coordination and manoeuvrability of the spinnerets as well as the combing leg. This raises the question of how muscles move with such a high velocity and how this is controlled (Barth 2020). Usually, muscles of spiders are described to fatigue rapidly after a few seconds of exertion (Bristowe 1932). There are different types of muscle fibres: one that contracts rapidly but fatigues quickly, two which contract slower but for a longer time, and one that remains contracted permanently (Maier et al. 1987; Paul et al. 1991). The long-lasting and fast contraction during the cribellate spinning process could hint to the use of the latter. An increased energetic demand for muscle contraction during spinning could be satisfied by the evolution of respiratory structures as in different lineages within the Araneomorphae the tracheal system shifted towards the spinning apparatus (Ramirez et al. 2020). It has already been described that the spinnerets and their segments can move independently from each other or perform synchronous movements operated by a combination of extrinsic and intrinsic muscles (Whitehead and Rempel 1959; Glatz 1973; Grannemann et al. 2019). Additionally, the cycle always starts at a specific position. This could hint to a constitutional program (e.g. a cascade) of nervous information controlling the movement sequences. It has been described that during dragline production, the spinneret movements and silk release are controlled by several sensilla (mechanoreceptors) which give information about the forces while pulling out the silk and to the orientation of the fibre in space (Gorb and Barth 1996). Similar sensory control of the spinneret movements could be possible by a combination of sensilla on the spinnerets or the calamistrum controlling the cribellate thread production (Foelix and Jung 1978). Furthermore, muscles not only control the movements of a spinneret but define the motion axes of the joints (translation and rotation) according to their length and attachment sites (Glatz 1973; Magalhães et al. 2017). Inside the cribellum, muscles attach anteriorly and posteriorly and thus already indicate possible directions of movement (Glatz 1972). However, a further study would be needed to specify the dynamics of the cribellum as well as the spinnerets.

**THE SPINNERET MOVEMENTS DURING THE CRIBELLATE SPINNING PROCESS**

The cribellate spinning process is a rhythmic interplay of different spatio-temporal factors leading to a combination of cribellate, axial, and possibly other fibres, such as undulating and supporting fibres (Peters 1984; Joel et al. 2015; Granemann et al. 2019; Michalik et al. 2019). This combination of different types of silk affects the mechanical properties of...
a cribellate capture thread (Michalik et al. 2019; Piorkowski et al. 2020). The spinning sequence is highly conserved, not only in *B. longinqua* but also in other cribellate spiders, such as *Uloborus plumipes* or *Kukulcania hibernalis* (Joel et al. 2015, 2016, Grannemann et al. 2019). In *B. longinqua*, the movements are all synchronous between the pairs of spinnerets. In other spiders, such as in *K. hibernalis*, the spinnerets can also move asynchronously during cribellate thread production creating another thread geometry (Grannemann et al. 2019). Aside from the movements, the morphology of the spinnerets is crucial for the accessibility of spigots involved (Eberhard 2010; Koebley et al. 2017). The spinnerets can be brought into close contact due to their adapted shape, as also described for *Polenecia producta* (Kovoor and Peters 1988). Besides, the fields of spigots on the terminal segments of the spinnerets are mostly inwards directed, as in the posterior median and posterior lateral spinnerets of *B. longinqua*. During the cribellate spinning process, the apical position of the triad of pseudoflagelliform spigots and two flanking paracribellate spigots on the posterior lateral spinnerets ensures an enlarged radius of movement during an abduction. Thus, the spinnerets function as an extension for the spigot range in this position. During adduction and their most anterior position, however, the spigots of the posterior lateral spinnerets come into close contact with the spigots of the posterior median spinnerets. The posterior median spinnerets on the other hand can approach the cribellum by inclining anteriorly and come into close contact with the cribellum. For the approach of the different structures during the cribellate spinning process, the flexibility of the spinnerets, however, is not fully exploited: only necessary movements are executed and theoretically possible ranges of movements are not completely used. By reducing to essential movements in combination with the required spigot distribution, the spinning apparatus of a cribellate spider can thus achieve different process sequences with one configuration of spinnerets. This not only affects the capture thread production but all the different silk-spinning processes a spider performs.

**Biomimetic inspiration**

Studying biological small-scale fibre processing, such as the cribellate thread production, could offer biomimetic inspiration to engineers. Biomimetics, in general, deals with the technical implementation of principles of biological systems. By emphasising necessary conditions which essentially stand for the spinneret movements in the cribellate spinning process, we aimed to find out which conditions can and cannot be neglected for future technical abstractions. Interestingly, it is the spider itself that reduces the scope of movement and processes, as in the cribellate spinning process not all movements of the spinnerets are executed—in comparison to other spinning processes. The course of movement of each spinneret thus takes place along a defined and limited path. This also reduces the possibilities of potential kinematic conditions. For technical implementation, this insight means two things: on one hand, the complexity of the cribellate spinning process takes us to the limits of visualising three-dimensional movements within time. However, this is also a result of the fact that the spinning apparatus compromises due to the compatibility of various process options. On the other hand, the process can be reduced to necessary components of a single function, e.g. the cribellate spinning process. This facilitates biomimetic abstraction by offering scope for optimisation and could inspire the technical fibre spinning of complex systems.

**Conclusion**

The movements of the spinning apparatus in the cribellate spider *B. longinqua* reveal how specifically the morphological features of a spider’s spinning apparatus can be used to accomplish specific functions. By describing the kinematics of random movements and the regular sequences of movements during cribellate capture thread production, spatial constraints as well as free spaces could be identified. During the cribellate spinning process, multi-joint spinnerets are precisely coordinated for the intertwaving of a characteristic multi-fibre system and only appropriate movements are executed in the process. The versatility of spinneret kinematics, combined in a single spinning apparatus, could thus also provide inspiration for the complex spinning of technical fibre systems.

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**Availability of data and materials** Three selected video recordings of the spinneret movements during cribellate thread production and random movements of an anaesthetised *Badumna longinqua* as well as two...
SEM images of the spinning apparatus are provided as supplemental material accessible in the online version of this publication.

Compliance with ethical approval

Conflict of interest The authors have no conflicts of interest to declare that are relevant to the content of this article.

Ethics approval All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. Export of spiders was kindly granted by the Department of Agriculture, Water and the Environment (Australian Government, permission no. PWS2019-AU-000248).

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References

Alfaro RE, Griswold CE, Miller KB (2018) The ontogeny of the spinning apparatus of *Tengella perjuga* (Araneae: Zoopsidae). Invertebr Biol 137:187–204
Barth FG (2020) A spider in motion: facets of sensory guidance. J Comp Physiol A. https://doi.org/10.1007/s00359-020-01449-z
Becker N, Oroudjev E, Mutz S, Cleveland JP, Hansma PK, Hayashi CY, Makarov DE, Hansma HG (2003) Molecular nanosprings in spider capture-silk threads. Nat Mater 2(4):278
Bertkau P (1882) Über das Cribellum und Calamistrum: ein Beitrag zur Histologie. Biologie und Systematik der Spinnen vol 48. Arch Naturgesch
Blackledge TA, Kuntner M, Aagnarsson I (2011) The form and function of spider orb webs: evolution from silk to ecosystems. In: Adv Insect Phys, vol 41. Elsevier, pp 175–262
Blackwall J (1839) XIII. On the Number and Structure of the Mammule employed by Spiders in the Process of Spinning. Trans Linn Soc Lond 18:219–224
Bott RA, Baumgartner W, Braunig P, Menzel F, Joel AC (2017) Adhesion enhancement of cribellate capture threads by epicular waxes of the insect prey sheds new light on spider web evolution. Proc R Soc B Biol Sci 284:6. https://doi.org/10.1098/rspb.2017.0363
Bristowe WS (1932) The Liphistiiid Spiders. In: Proc Zool Soc Lond, vol 4. Wiley Online Library, pp 1015–1057
Coddington JA (1989) Spinneret silk pigot morphology: evidence for the monophony of orbweaving spiders, Cyrtophorinae (Araneidae), and the group Theridiidae plus Nesticidae. J Arachnol 17(1):71–95
Eberhard WG (2010) Possible functional significance of pigot placement on the spinnerets of spiders. J Arachnol 38:407–414
Eberhard W, Pereira F (1993) Ultrastructure of cribellate silk of nine species in eight families and possible taxonomic implications (Araneae: Amaurobiidae, Deinopidae, Desidae, Dictynidae, Filistatidae, Hypochilidae, Stiphidiidae, Tengellidae). J Arachnol 21(3):161–174
Ehlers M (1939) UntersuchungenüüberFormenaktiverLokomotionsbeispielSniventieren. Zool Jb, System 72:1–374
Eisoldt L, Smith A, Scheibel T (2011) Decoding the secrets of spider silk. Mater Today 14:80–86
Foelix R (2011) Biology of spiders, 3rd edn. OUP, USA
Foelix RF, Jung H (1978) Some anatomical aspects of *Hypochilus thorelli* with special reference to the calamistrum and cribellum. Symp Zool Soc London 42:417–422
Glatz L (1972) Der spinnapparathalopiespinnen (Arachnida, Araneae). Z Morphol Tiere 72:1–25
Glatz L (1973) Der spinnapparat der orthognatha (Arachnida, Araneae). Z Morphol Tiere 75:1–50
Gorb SN, Barth FG (1996) A new mechanosensory organ on the anterior spinnerets of the spider *Cupiennius salei* (Araneae, Ctenidae). Zoomorphology 116:7–14
Grannemann CC, Meyer M, Reinhardt M, Ramírez MJ, Herberstein ME, Joel A-C (2019) Small behavioral adaptations enable more effective prey capture by producing 3D-structured spider threads. Sci Rep 9:1–10
Griswold CE, Ramirez MJ, Coddington JA, Platnick NI (2005) Atlas of phylogenetic data for entelegyne spiders (Araneae: Araneomorphae: Entelegynae) with comments on their phylogeny. Proc Calif Acad Sci 56:1–324
Hawthorn AC, Opell BD (2002) Evolution of adhesive mechanisms in cribellar spider prey capture thread: evidence for van der Waals and hygroscopic forces. Biol J Linn Soc Lond 77(1):1–8
Hayashi CY, Shipley NH, Lewis RV (1999) Hypotheses that correlate the sequence, structure, and mechanical properties of spider silk proteins. Int J Biol Macromol 24(2–3):271–275
Hilibrant M (2008) Development and evolution of the spider silk producing system. Dissertation, Universität zu Köln, http://kups.ub.uni-koeln.de/id/eprint/2795
Joel A-C, Baumgartner W (2017) Nanofibre production in spiders without electric charge. J Exp Biol 220(12):2243–2249
Joel A-C, Kappel P, Adamova H, Baumgartner W, Scholz I (2015) Cribellate thread production in spiders: complex processing of nano-fibres into a functional capture thread. Arthropod Struct Dev 44:568–573
Koebel SR, Vollrath F, Schniepp HC (2017) Toughness-enhancing metastructure in the recluse spider’s looped ribbon silk. Mater Horizons 4:377–382
Kovoor J, Peters HM (1988) The spinning apparatus of *Poleneca producta* (Araneae, Uloboridae): structure and histochemistry. Zoomorphology 108:47–59
Kropf C (2013) Hydraulic system of locomotion. In Spider ecophysiology, Springer, Berlin, Heidelberg 43–56
Labarque FM, Wolff JO, Michalik P, Griswold CE, Ramírez MJ (2017) The evolution and function of spider feet (Araneae: Araneomorphae: Entelegynae) with comments on their phylogeny. Proc Zool Soc London 181:308–341
Magalhães I, Ravelo A, Scioscia C, Peretti A, Michalik P, Ramírez MJ (2017) Recluse spiders produce flattened silk rapidly using a highly modified, self-sufficient spinning apparatus. J Zool 303:27–35
Maier L, Root TM, Seyfarth E-A (1987) Heterogeneity of spider leg muscles: histochemistry and electrophysiology of identified fibres in the claw levator. J Comp Physiol B 157:285–294
Mariano-Martins P, Lo Man Hung N, Torres TT (2020) Evolution of Spiders and Silk Spinning: mini review of morphology, evolution, and development of spider’s spinnerets. Front Ecol Evol 8:109
Marles B (1967) The spinnerets and epiandrous glands of spiders. Zool J Linn Soc 46:209–212

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Michalik P, Piorkowski D, Blackledge TA, Ramírez MJ (2019) Functional trade-offs in cribellate silk mediated by spinning behaviour. Sci Rep 9:9092
Montgomery THJ (1909) On the spinnerets, cribellum, colulus, tracheae and lung books of Araneads. Proc Acad Nat Sci Philadelphia 61:299–320
Motimer B, Holland C (2015) The use of spider silk as a biomaterial. In: Advances in Silk Science and Technology. Elsevier, pp 233–260
Opell BD (2002) How spider anatomy and thread configuration shape the stickiness of cribellar prey capture threads. J Arachnol 30:10–20
Paul R, Zahler S, Werner R, Markl J (1991) Adaptation of an open circulatory system to the oxidative capacity of different muscle cell types. Sci Nat 78:134–135
Pechmann M, Prpic N-M (2009) Appendage patterning in the South American bird spider Acanthoscurria geniculata (Araneae: Mygalomorphae). Dev Genes Evol 219:189–198
Pechmann M, Khadjeh S, Sprenger F, Prpic N-M (2010) Patterning mechanisms and morphological diversity of spider appendages and their importance for spider evolution. Arthropod Struct Dev 39:453–467
Peters H (1984) The spinning apparatus of Uloboridae in relation to the structure and construction of capture threads (Arachnida, Aranea) Zoomorphology 104:96–104
Peters HM (1992) On the spinning apparatus and the structure of the capture threads of Deinopis subrufus (Araneae, Deinopidae) Zoomorphology 112:27–37
Piorkowski D, Blackledge TA, Liao C-P, Joel A-C, Weissbach M, Wu C-L, Tso I-M (2020) Uncoiling springs promote mechanical functionality of spider cribellate silk. J Exp Biol 223:1–11. https://doi.org/10.1242/jeb.215269
Platnick NI, Coddington JA, Forster RR, Griswold CE (1991) Spinneret morphology and the phylogeny of halopigyne spiders (Araneae, Araneomorphae). Am Mus Novit 3016:1–73
Ramírez MJ, Magalhaes IL, Derkarabetian S, Ledford J, Griswold CE, Wood HM, Hedin M (2020) Sequence-capture phylogenomics of true spiders reveals convergent evolution of respiratory systems. Syst Biol 0(0):1–7
Shultz JW (1987) The origin of the spinning apparatus in spiders. Biol Rev 62:89–113
Termonia Y (1994) Molecular modeling of spider silk elasticity. Macromolecules 27:7378–7381
Vollrath F (1992) Spider webs and silks. Sci Am 266:70–77
Vollrath F (2000) Strength and structure of spiders’ silks. Reviews in Mol Biotechnol 74:67–83
Weihmann T, Karner M, Full RJ, Blickhan R (2010) Jumping kinematics in the wandering spider Cupiennius salei. J Comp Physiol A 196:421–438
Whitehead W, Rempel J (1959) A study of the musculature of the black widow spider, Latrodectus mactans (Fabr.). Can J Zool 37:831–870
Wolff JO (2020) Evolutionary kinematics of spinneret movements for rapid silk thread anchorage in spiders. J Comp Physiol. https://doi.org/10.1007/s00359-020-01453-3
Wolff JO, Grawe I, Wirth M, Karstedt A, Gorb SN (2015) Spider’s super-glue: thread anchors are composite adhesives with synergistic hierarchical organization. Soft Matter 11:2394–2403
Wolff JO, Paterno GB, Liprandi D, Ramírez MJ, Bosia F, Van Der Meijden A, Michalik P, Smith HM, Jones BR, Ravelo AM, Pugno N, Herberstein ME (2019) Evolution of aerial spider webs coincided with repeated structural optimization of silk anchorages. Evolution 73:2122–2134

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