The timing of female genital mutilation and the role of contralateral palpal insertions in the spider *Cyclosa argenteoalba*

Kensuke Nakata

Kyoto Women’s University, Kitahiyoshi-cho 35, Higashiyama-ku, Kyoto 605-8501, Japan

KN, 0000-0002-1207-3448

Competing interests

The author declares no competing interests.

Received: 7 July 2017
Accepted: 25 October 2017

Subject Category:
Biology (whole organism)

Subject Areas:
behaviour

Keywords:
female genital mutilation, mating behaviour, paternity, spider, sexual conflict

Author for correspondence:
Kensuke Nakata
e-mail: nakatake@kyoto-wu.ac.jp

Some male spiders exhibit female genital mutilation behaviour (FGM) by removing the female genital appendage (scape) to control the mating frequency of females. Female spiders have two, i.e. right and left, genital openings connected with separate spermathecae into which males transfer sperm successively using one pedipalp (secondary genitalia) at a time. Thus, males must complete at least two palpal insertions to fill both spermathecae, before FGM. The present study examined whether (i) scape removal is only associated with the second palpal insertion (one-action hypothesis) or (ii) two contralateral palpal insertions facilitate FGM, with each insertion cutting the basal part of the scape halfway (two-actions hypothesis). Experiments in which females were replaced after a male had made the first insertion did not support the one-action hypothesis, because scapes remained intact after the newly introduced virgin females received their first palpal insertion, which was the second insertion by the males. In comparison, mating experiments using two half-eunuchs (i.e. one of the palps of each male had been manually removed, forcing them to fill female spermatheca on one side only) supported the two-actions hypothesis. FGM was more frequent in females that received two contralateral palpal insertions than in females that received ipsilateral insertions.

1. Background

In the majority of taxa, a female often mates with multiple males [1,2]. A female that has more than one reproductive partner might benefit from higher fecundity and improved genetic quality and/or diversity of her offspring [3,4]. However, this action reduces male fitness; thus, to secure paternity, males have developed various strategies to inhibit females from mating with...
multiple males, including guarding females [5], injecting chemical substances to lower female re-mating tendency [6] and mating plugs [7].

Another way to prevent females from mating multiply is through female genital mutilation (FGM) by males. For example, FGM has been described in two orb-web spiders, *Cyclosa argenteoalba* [8] and *Larinia jeskovi* [9]. In these species, virgin females have a small projection, called a scape, on their external genitalia (epigynum) [10]. The scape is essential for successful mating. The male must ‘clasp’ the scape with its genitalia (pedipalp) to position itself appropriately for palpal insertion [11]. In FGM, the scape is removed from the epigynum during mating, preventing the success of subsequent genital coupling. Thus, mutilated females cannot mate with additional males. FGM is a powerful way in which males can inhibit females from mating with multiple partners, and thus ensure full male paternity. As a result, FGM contrasts with other ways to secure paternity that are more costly to males in terms of energy expenditure and mating opportunity. For instance, mate guarding might reduce the glycogen reserves of males in stream-dwelling isopods [12]. Some male spiders form mating plugs by breaking their pedipalps and leaving breakages in the copulatory openings of females [13,14]. Male *Nephilengys malabarensis* also detach their entire pedipalp, which continues sending sperm into a female. Such an emasculated and, thus, lightened male can guard the female from other males more efficiently [15–17]. Other males sacrifice themselves as food to females in copula, to send more sperm while being eaten [18]. All these males lose all their future mating opportunities. Male garter snakes adjust the size of mating plugs according to the size of the females [19], suggesting that the production of these plugs is costly. Nevertheless, males often fail to monopolize their partnered female [20–22]; thus, monandry is relatively rare [23]. Thus, it is important to understand FGM to elucidate the evolution of monandry/polyandry and evolutionary outcomes of male–male competition over paternity and male–female conflict over the number of times a female copulates.

*Cyclosa argenteoalba* is a diurnal spider that occurs in Japan, Korea and China [24]. This species builds vertical orb webs. The male spider has two pedipalps (i.e. right and left), while the females of entelegyne spiders, including *C. argenteoalba*, have two copulatory openings, each of which is connected to separate spermatheca. On mating, a male makes a mating thread and sends courtship signals by tapping and jerking the thread with its legs. When a female accepts courtship, the male inserts one of its pedipalps into one of the genital openings of the female. Thus, at least two palpal insertions are required to fill both spermatheca with sperm, which is considered to lower the risk of insufficient sperm being available at oviposition [25]. In fact, *C. argenteoalba* males insert their pedipalps twice in a single mating bout, using each pedipalp successively (i.e. one pedipalp first, followed by the other). When the insertion successfully ends, the pair separates, and the male repeats its courtship behaviour to make the second insertion using the other pedipalp. The scape typically remains on the epigynum during the first insertion; however, it is absent after the second insertion [8]. This process is logical because if FGM occurred at the first insertion, males could not complete the second insertion, and the second spermatheca would remain empty.

Here, we examine if selection acts on *C. argenteoalba* males to implement FGM after the second insertion in order to maximize paternity success. This question will give us invaluable insights into the evolutionary process and selective mechanisms of FGM. Namely we test two hypotheses. The one-action hypothesis predicts that spiders implement no actions to mutilate the scape during the first insertion, with scape-removal behaviour only being associated with the second insertion. Alternatively, the two-actions hypothesis predicts that a male destroys only one side (right or left) of the scape at the first insertion. Once both insertions (right and left) are complete, the destruction of the scapes on both sides facilitates successful FGM. The rationale for this hypothesis comes from a phenomenon recorded in *L. jeskovi*, in which a claw-like sclerite part of the pedipalp (called tegular apophysis) seems to ‘slash’ the side of the basal part of the scape of females halfway during copulation [9]. Thus, this study aimed to examine which of these two hypotheses is valid in *C. argenteoalba*.

2. Material and methods

I collected both adult and subadult female *C. argenteoalba* from Shimamoto, Osaka and from Nagaoka-Kyo, Japan between 2014 and 2015. All females were found on moulting webs, indicating that they had just completed the final moult (adult) or were ready for the final moult (subadult) [26]. Subadult females were allowed to moult into adults. By following this procedure, I ensure that all adult females were virgins prior to the experiment. Females were released into the observation area (9 m × 2 m) where they built their webs. Individual spiders were identified by variation in the abdominal markings [27] and
the location of their webs. The observation area was not enclosed, but was surrounded by residential buildings. The nearest natural habitat of *C. argenteoalba* was more than 1 km away, with no obvious signs of immigration (e.g. additional webs by immigrating spiders or unknown males residing in the female web) observed during the study. Adult males, for which the mating history was not known, were also collected from the same sites and were maintained in separate vials with wet cotton.

3. Female-exchange experiment

Ten pairs of virgin females and males were subjected to staged mating. When a male successfully inserted one of its pedipalps and completed sperm transfer (I could not determine whether the right or left pedipalp was used, due to the small size of the spiders and short duration of insertion), females were removed from the web. After the male resumed courtship, a different virgin female was introduced to the hub of the experimental web from a nearby web. Soon, the second virgin female was aware of the courting male, and accepted its first, but the male’s second, insertion. The male did not exhibit any further mating behaviour, and left the web. I inspected the second female under the microscope to check for the presence of the scape.

4. Half-eunuch experiment

Adult males were anaesthetized with CO\textsubscript{2}, and the tip of one of their pedipalps was cut off with fine scissors under the microscope. Preliminary observation confirmed that manipulated males could only make one insertion during a single mating event, indicating that the operation successfully disabled the manipulated pedipalp. Which pedipalp was removed (right or left) for each male was randomly determined. Thirty-two experimental females were assigned to four groups. Females from the first group were coupled with a male that had an intact right pedipalp (termed right-hand male, hereafter). After receiving one palpal insertion from the first male, the female was coupled with another male with an intact left pedipalp (hereafter, left-hand male), and received the second insertion. Females from the second group were coupled with a left-hand male first, and then with a right-hand male. The females in the third and fourth groups were coupled with two right-hand and left-hand males, respectively. Owing to issues that arose during the handling of spiders, the sample size was not balanced (*N* = 7, 8, 9 and 8 in groups 1, 2, 3 and 4, respectively). As a result, one right-hand male was used twice as the second male for females in the second and the third groups. Removing the data of this male from the analysis did not affect the qualitative aspect of the results. All other males were used only once. After a female received two insertions, the presence of the scape was inspected under the microscope. The mutilation rate was similar between the first and the second groups, and between the third and fourth groups. As a result, data from the first and second groups were pooled as the contralateral insertion group, and the data from the third and fourth groups as the ipsilateral group. The difference in the frequency of mutilation between the contralateral and ipsilateral insertion groups was examined by Fisher’s exact probability test.

5. Results

In the female exchange experiment, none of the second females lost their scapes. In the half-eunuch experiment, 10 of the 15 females from the contralateral insertion group and four of the 17 females from the ipsilateral insertion group lost their scapes. This difference was statistically significant (*p* = 0.031).

6. Discussion

The results of the female-exchange experiments showed that when females received only one palpal insertion, no FGM occurred, even when it was the second insertion for males. In comparison, the half-eunuch experiments showed that the incidence of FGM was significantly higher in females that received two contralateral insertions than in females that received ipsilateral insertions. Thus, results supported the two-actions hypothesis over the one-action hypothesis. The results of the present study suggest that, during palpal insertion, *C. argenteoalba* males slash the side of the scape, similarly to that recorded in *L. jeskovi* [9]; thus, slashing both right and left sides of the scape facilitates FGM.

In the half-eunuch experiment, some females from the ipsilateral insertion group lost their scapes. A previous study showed that *C. argenteoalba* females lost their scapes only after receiving one palpal
insertion, although at a low rate (only two in 44 mutilations observed) [8]. Thus, cutting both sides of the scape might not always be required for FGM. As argued in the study on *L. jeskovi* [9], additional action, such as twisting off the scape, might be involved in successful mutilation. In the typical mating ritual of *C. argenteoalba*, a copulating male and female hang from a mating thread by the third and the fourth legs, and their bodies often rotate along the vertical line, passing through the point of genital conjunction immediately after the timing of insertion. This body rotation might generate a twisting force, which, if strong enough, might cause genital mutilation, even when only one side of the scape is cut. The necessity of additional action is logical because if the scape was removed at the exact timing of the second slashing, which is considered to occur when the pedipalp clasps the scape, subsequent palpal insertion would fail.

FGM seems to require two insertions. Two-actions FGM might be essential for the efficiency of securing paternity because when a male has made only a single palpal insertion, failing to insert the palp a second time, the scape remains attached to the female. This event might be expected when a female cannibulates a male after the first palpal insertion or when the mating sequence was interrupted after the first insertion by a disturbance, such as sudden change in weather, the destruction of the web or when two males simultaneously court the same female. If males only mutilated the scape during the second insertion, as expected by the one-action hypothesis, the scape of a female that had received only one insertion would remain undamaged. Consequently, a second male could make two successful palpal insertions. Assuming that two males inject a similar amount of sperm per insertion, the sperm of the first male would represent one-third of the sperm mixture. By contrast, if the male cut the scape halfway during the first insertion, as expected by the two-actions hypothesis, the opportunity for a second male to make two palpal insertions would be lowered. Assuming that males exhibit no preference in the use of their pedipalps, the second male might insert its pedipalp into a contralateral genital opening. Then, the scape would detach, preventing the second insertion. Without cryptic female choice, in this instance, the paternity share of the first male would be raised to half. Thus, cutting the scape halfway is considered beneficial to secure paternity, even if this action alone does not lead to mutilation.

FGM appears to be related to sexual conflict. Under sexual conflict, males often enforce costs on their mating-partner females and females exhibit counter-adaptations [28,29], and theoretical study revealed that FGM may evolve even when females suffer fecundity costs [30]. In some insects, males damage the internal genitalia of females in copula and females resist behaviourally to this harmful copulation [31,32]. In *C. argenteoalba*, females do not seemingly exhibit any behaviour to resist mutilation [8]. Nevertheless, an arms race on the efficiency of mutilation might occur between males and females, in the form of antagonistic genital coevolution [33]. In *Cyclosa*, some species other than *C. argenteoalba* do not exhibit FGM (personal observation). Comparison of the morphology of genitalia, especially the basal part of the scape in females and tegular apophysis in males, among species that do and do not exhibit FGM might provide information about the evolution of genitalia, including details on the mechanism of mutilation.

Ethics. The experiments were performed in accordance with the guidelines for ethological studies from the Japan Ethological Society.

Data accessibility. All data are available in the text.

Authors’ contribution. I am the sole author of this study and conducted all the work by myself.

Competing interests. I declare I have no competing interests.

Funding. This work was partly supported by a JSPS Grant-in-Aid for Scientific Research (C) (nos. 26440251 and 17K07576).

Acknowledgements. I thank Dr Keizo Takasuka and Dr Atushi Ushimaru for providing suggestions and assistance in conducting the half-eunuch experiment.

References

1. Barbosa M, Maguran A. 2006 Female mating decisions: maximizing fitness? *J. Fish Biol.* **68**, 1656–1661. (doi:10.1111/j.0022-1370.2006.01033.x)

2. Jennions MD, Petrie M. 2000 Why do females mate multiply? A review of the genetic benefits. *Biol. Rev.* **75**, 21–64. (doi:10.1017/S0006323999005432)

3. Slatyer RA, Mazur BS, Backwell PR, Jennions MD. 2012 Estimating genetic benefits of polyandry from experimental studies: a meta-analysis. *Biol. Rev.* **87**, 1–33. (doi:10.1111/j.1469-185x.2011.00182.x)

4. South A, Lewis SM. 2011 The influence of male ejaculate quantity on female fitness: a meta-analysis. *Biol. Rev.* **86**, 299–309. (doi:10.1111/j.1469-185x.2010.00194.x)

5. Sherman PW. 1989 Mate guarding as paternity insurance in Idaho ground squirrels. *Nature* **338**, 418–420. (doi:10.1038/338418a0)

6. Wolfner M. 2002 The gifts that keep on giving: physiological functions and evolutionary dynamics of male seminal proteins in *Drosophila*. *Hereditas* **88**, 85. (doi:10.1038/sj.hdy.680007)

7. Dixon AF, Anderson MJ. 2002 Sexual selection, seminal coagulation and copulatory plug formation in primates. *Folia Primatol.* **73**, 63–69. (doi:10.1159/000064784)

8. Nakata K. 2016 Female genital mutilation and monandry in an orb-web spider. *Biol. Lett.* **12**, 20150912. (doi:10.1098/rsbl.2015.0912)

9. Mougenot P, Prügel J, Thom U, Steinhef Philipp OM, Kupryanov I, Uni G. 2015 Securing paternity by mutilating female genitalia in spiders. *Curr. Biol.* **25**, 2980–2984. (doi:10.1016/j.cub.2015.09.074)
10. Foelix R. 2010 Biology of spiders. Oxford, UK: Oxford University Press.
11. Uhl G, Nessler SH, Schneider J. 2007 Copulatory mechanism in a sexually cannibalistic spider with genital mutilation (Araneae: Araneidae: Argiope bruennichi). Zoology 110, 398–408. (doi:10.1016/j.zool.2007.07.003)
12. Sparks TC, Keogh DP, Pary RA. 1996 Energetic costs of mate guarding behavior in male stream-dwelling isopods. Oecologia 106, 166–171. (doi:10.1007/BF00328595)
13. Kuntner M, Kralj-Fišer S, Schneider JM, Li D. 2009 Mate plugging via genital mutilation in nephilid spiders: an evolutionary hypothesis. J. Zool. 277, 257–266. (doi:10.1111/j.1469-7998.2008.00533.x)
14. Uhl G, Nessler S, Schneider J. 2010 Securing paternity in spiders? A review on occurrence and effects of mating plugs and male genital mutilation. Genetica 138, 75–104. (doi:10.1007/s10709-009-9388-5)
15. Kralj-Fišer S, Gregorič M, Zhang S, Li D, Kuntner M. 2011 Eunuchs are better fighters. Anim. Behav. 81, 933–939. (doi:10.1016/j.anbehav.2011.02.018)
16. Li D, Oh J, Kralj-Fišer S, Kuntner M. 2012 Remote copulation: male adaptation to female cannibalism. Biol. Lett. 8, 512–515. (doi:10.1098/rsbl.2011.1200)
17. Lee QQ, Oh J, Kralj-Fišer S, Kuntner M, Li D. 2012 Emasculation: gloves-off strategy enhances eunuch spider endurance. Biol. Lett. 8, 733–735. (doi:10.1098/rsbl.2012.0285)
18. Andrade MCB. 1996 Sexual selection for male sacrifice in the Australian redback spider. Science 271, 70–72. (doi:10.1126/science.271.5245.70)
19. Shine R, Olsson MM, Mason RT. 2000 Chastity belts in garter snakes: the functional significance of mating plugs. Biol. J. Linn. Soc. 70, 377–390. (doi:10.1111/j.1095-8312.2000.tb01229.x)
20. Avila FW, Siot LK, LaFlamme BA, Rubinstein CD, Wollnfer MF. 2011 Insect seminal fluid proteins: identification and function. Annu. Rev. Entomol. 56, 21–40. (doi:10.1146/annurev-ento-120709-144823)
21. Foellmer MW. 2008 Broken genitals function as mating plugs and affect sex ratios in the orb-web spider Argiope aurantia. Evol. Ecol. Res. 10, 449–462.
22. Koprowski JL. 1992 Removal of copulatory plugs by female tree squirrels. J. Mammal. 73, 572–576. (doi:10.2307/1382026)
23. Taylor ML, Price TAR, Wedell N. 2014 Polyandry in nature: a global analysis. Trends Ecol. Evol. 29, 376–383. (doi:10.1016/j.tree.2014.04.005)
24. Tanikawa A. 1992 A revisional study of the Japanese spiders of the genus Glosso MENG (Araneae: Araneidae). Acta Arachnol. 41, 11–85. (doi:10.2476/asjaa.41.11)
25. Foellmer MW, Fairbairn DJ. 2004 Males under attack: sexual cannibalism and its consequences for male morphology and behaviour in an orb-weaving spider. Evol. Ecol. Res. 6, 163–181.
26. Takasuka K, Yusa T, Ishigami T, Nakata K, Matsuura R, Ikeda K, Maeto K. 2015 Host manipulation by an ichneumonid spider ectoparasitoid that takes advantage of preprogrammed web-building behaviour for its cocoon protection. J. Exp. Biol. 218, 2226–2232. (doi:10.1242/jeb.122739)
27. Nakata K, Shigemiy Y. 2015 Body-colour variation in an orb-web spider and its effect on predation success. Biol. J. Linn. Soc. 116, 954–963. (doi:10.1111/bij.12400)
28. Stockley P. 1997 Sexual conflict resulting from adaptations to sperm competition. Trends Ecol. Evol. 12, 154–159. (doi:10.1016/S0169-5347(97)01000-8)
29. Marlow EH, Arnqvist G. 2003 Costly traumatic insemination and a female counter-adaptation in bed bugs. Proc. R. Soc. Lond. B 270, 2377–2381. (doi:10.1098/rspb.2003.2514)
30. Mouginit P, Uhl G, Fromhage, L. 2017 Evolution of external female genital mutilation: why do males harm their mates? R. Soc. open Sci. 4, 171195. (doi:10.1098/rsos.171195)
31. Blaundenhorn Wu, Hosken DJ, Martin OF, Reim C, Teuschl Y, Ward PL. 2002 The costs of copulating in the dung fly Sepsis cynipsea. Behav. Ecol. 13, 353–358. (doi:10.1093/beheco/13.3.353)
32. Crudgington HJ, Siva-Jothy MT. 2000 Genital damage, kicking and early death. Nature 407, 855–856. (doi:10.1038/35038154)
33. Brennan PL, Prum RO. 2015 Mechanisms and evidence of genital coevolution: the roles of natural selection, mate choice, and sexual conflict. Cold Spring Harb. Perspectives Biol. 7, a017749. (doi:10.1101/cshperspect.a017749)