Sexual conflict over mating in red-sided garter snakes (*Thamnophis sirtalis*) as indicated by experimental manipulation of genitalia

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Sexual conflict over mating can result in sex-specific morphologies and behaviours that allow each sex to exert control over the outcome of reproduction. Genital traits, in particular, are often directly involved in conflict interactions. Via genital manipulation, we experimentally investigated whether genital traits in red-sided garter snakes influence copulation duration and formation of a copulatory plug. The hemipenes of male red-sided garter snakes have a large basal spine that inserts into the female cloaca during mating. We ablated the spine and found that males were still capable of copulation but copulation duration was much shorter and copulatory plugs were smaller than those produced by intact males. We also anaesthetized the female cloacal region and found that anaesthetized females copulated longer than control females, suggesting that female cloacal and vaginal contractions play a role in controlling copulation duration. Both results, combined with known aspects of the breeding biology of red-sided garter snakes, strongly support the idea that sexual conflict is involved in mating interactions in this species. Our results demonstrate the complex interactions among male and female traits generated by coevolutionary processes in a wild population. Such complexity highlights the importance of simultaneous examination of male and female traits.

1. Introduction

Sexual conflict results when the evolutionary interests of males and females are not optimized simultaneously during mating, fertilization and/or parental care [1–3]. During mating, conflict often occurs when the ideal copulation duration is different for males and females [1,4–6]. For females, copulation times beyond those required for optimal sperm transfer may be detrimental owing to decreased ability to choose other mates as fathers of her offspring [7–9], increased risk of injury and predation, and/or decreased feeding rates [10]. Therefore, females are expected to favour shorter copulations especially if sperm transfer occurs quickly [11]. For males, however, longer copulations can serve not only to transfer more sperm, but also to mate-guard the female, thus males are expected to typically favour longer copulations than females. When males transfer other substances during copulation, for example a copulatory plug, conflict is likely to be more intense if these secretions negatively affect female fitness but positively affect male fitness [5,12,13]. Male genital morphology has been shown to be associated with sexual conflict during copulation in many taxa [14–20].

Sexual conflict over mating is a prominent feature of the mating system of red-sided garter snakes (*Thamnophis sirtalis parietalis*) [21,22]. In this species, the sex...
ratio at spring emergence is strongly male-biased. Several
dozens of males will congregate around a newly emerged
damselforming ‘mating balls’ in which males court her and
attempt copulation [23, 24]. When the female is ready to
mate, she gapes her cloaca so copulation can occur. Mating is
effectively random with respect to male size in the large aggre-
gations of males (6–24) a female encounters as she emerges
[25, 26]. Females, therefore, do not seem able to choose larger,
fitter males prior to copulation (garter snakes exhibit indeter-
minate growth, thus large males are successful at foraging
and evading predators [27, 28]). It also seems unlikely that
females use features of the hemipene to assess mates during
copulation because small males with small hemipenes are
just as successful as large males with large hemipenes [29].
Females may also be forced to copulate if males can elicit a
cloacal gaping response owing to oxygen deprivation in the
female using caudocephalic waves during courtship [30].

Sexual conflict may be further intensified by the gelatinous
copulatory plug that a male deposits within the female’s
cloaca. The plug delays female remating [31], prevents sperm
ejection from unwanted males, prevents sperm leak-
age and acts as a spermatophore from which sperm are
liberated over the course of 2 days [32]. These features of the
red-sided garter snake mating system suggest that sexual con-
flict over copulation duration, rather than cryptic female choice
(CFC), is an important selective pressure operating during the
female’s first mating in the den [22]. As the female moves
away from the den, the plug from her first mating dissolves and she
may mate again in the smaller, less dense aggregations in small
aspen groves surrounding the den where larger males are more
prevalent and successful [23–25]. Thus, female choice may be
important when females move away from the den into the
aspen groves.

Copulation duration and plug deposition can be affected
by female behavioural strategies. In a related species, the
checkered garter snake (Thamnophis marcianus), copulatory
plug formation is prevented when females perform axial
rotations (body rolls) that terminate copulations early [33].
This body-roll behaviour also leads to significantly shorter
copulation durations in the plains garter snake (Thamnophis
radix) [34]. Thus, copulation duration may be important for
plug formation and female body-roll behaviour may indicate
sexual conflict over mating [34].

In addition to body rolls, female genital musculature
may also influence plug formation and copulation duration.
The cranial pouch of the cloacal urodaeum of females (hence-
forth vaginal pouch) is highly muscularized [35, 36]. The ease
with which a male intromits his hemipene and the period he
maintains intromission could depend on the muscular contrac-
tions of the vaginal pouch [32]. Therefore, copulation duration
and plug deposition are likely both mediated by female behav-
ior and by interactions between female and male genitalia
during copulation.

Male genital morphology has been shown to be associated
with sexual conflict during copulation in many taxa [14–18].
In North American natricine snakes, taxa with bilobed hemi-
penes have longer copulation durations than those with
cylindrical hemipenes [34]. Male hemipenes in T. sirtalis are
relatively simple cylindrical structures [34], with a ring of ker-
atinized spines and a large sharp basal spine located on the
lateral aspect of the everted hemipenes that may also be associ-
ated with conflict over copulation duration (figure 1). The
basal spine is the largest hemipenial spine and the first to
make contact with the female cloaca at intromission, and has
been hypothesized to act as a grappling hook that allows the
male to anchor and evert his hemipene into the female
cloaca [37]. However, the adaptive significance of the basal
spine has yet to be tested experimentally.

One way to test adaptive hypotheses of sexual conflict
during mating is to manipulate both females and males such
that the putative adaptations for control over conflict are ren-
dered ineffective [1, 38–41]. Here, we conducted experiments
in red-sided garter snakes during the female’s first mating at
the den where sexual conflict is most likely to be important.
We hypothesized that the basal spine allows the male’s hemi-
penes to maintain position during copulation despite female
body rolls or vaginal contractions used to terminate copu-
lation. To test this hypothesis, we ablated the basal spine of
the male hemipenes to determine whether males could initiate
and maintain intromission. We predicted that, if sexual conflict
over copulation duration and plug size occurs in this system,
spine-ablated males would be able to copulate, but for shorter
periods and produce smaller copulatory plugs. Alternatively, if
the basal spine is required for normal hemipene eversion as
hypothesized by Pisani [37], then spine-ablated males should
not be able to copulate at all. Female garter snakes perform
body rolls to terminate copulations, but we also hypothesized
that they use muscles of the vaginal pouch to mediate copu-
lation duration and plug deposition. To test this hypothesis,
we held females at intromission to prevent body rolls that
may terminate copulation immediately. These females were
either injected with saline or a local anaesthetic in the cloaca
and vaginal pouch region to reduce female control over the
course of copulation. We predicted that males would be able
to deposit larger plugs in anaesthetized females and these
females would be subjected to longer copulations even
though they could perform body rolls after intromission
because they would not be able to use their cloacal and vaginal
structures to terminate copulations.

2. Material and methods
(a) Study species

Thamnophis sirtalis parietalis in Manitoba exhibit a condensed
period of mating (four to six weeks) during the spring when
they emerge from limestone hibernacula. Males emerge earlier than females and remain at the den sites longer, resulting in a strongly male-biased sex ratio [42].

All experiments were carried out on individuals collected from Inwood, Manitoba, Canada (50°31.58′ N, 97°29.71′ W). All individuals used in the mating trials were identified with a letter written in permanent marker (Sharpie) on a 0.25 × 0.25 cm piece of 3M, Nexcare adhesive tape affixed to their head. All animals were housed in nylon arenas (1 × 1 × 1 M) and were provided water ad libitum until mating trials. For the trials, newly emerged, unmated females (typically still cold and slow) were collected as they emerged from the ground.

(b) Experiment 1: treated males mating in arenas with untreated females

We collected 42 actively courting males directly from the den and placed them in circular nylon mating arenas (45 cm diameter × 75 cm tall). We standardized male experience by allowing all males to mate once in a larger arena (1 × 1 × 1 M) on the same day prior to assigning them to experimental groups. During these initial mating trials, we introduced two unmated females to the arena until copulation commenced. The mating pair was then gently moved to a smaller enclosure where copulation could be closely monitored to record copulation duration (±10 s). We also recorded which hemipene was used during copulation as it may correlate with plug mass [29].

After mating, we measured and weighed, and assigned randomly to one of two groups: ablated or control. All males were lightly anaesthetized with 0.0015 ml of 0.5% methohexital sodium per gram body mass administered subcutaneously at the juncture between the dorsal and ventral scales approximately 4 cm from the head of the snake [43]. Ten minutes after anaesthesia, each male received two 0.1 µl bilateral, subcutaneous injections of the local anaesthetic 2% Lidocaine HCl. The injection sites were both between the first and second dorsal scale rows, four subcaudal scales caudal to the cloaca. Five to 10 min after Lidocaine injections, both hemipenes were everted manually exposing the basal spines, which were then cut close to its base with comeoscleral scissors from both hemipenes of the experimental males. Clipping the spine resulted in very little or no bleeding. Control males were treated in the same way except that the basal spine was touched but not clipped with the scissors. Snakes were then placed in a 20 gal (75.7 l) aquarium situated on a heating pad (30°C) and monitored every 10 min until righting reflex returned [43]. Males of this species regain courtship behaviour quickly after invasive surgeries and show no deficit in courtship ability [44], and all the males engaged in courtship during the experimental matings.

Males were housed indoors in two small circular arenas (45 cm diameter × 75 cm tall) for 4 days, given water ad libitum and then were taken back to the den site and placed together in a nylon arena (1 × 1 × 1 M). Two newly emerged unmated females were introduced into the arena and observed until copulation occurred. If a male achieved intromission, we timed copulation duration in the same way as in the first mating. We recorded copulation duration even in situations where a female rolled her body to dislodge a male prior to depositing a copulatory plug. If the pair remained in copula for more than a few seconds, we gently removed the pair to an empty arena. In order to keep the number of courting males constant, we introduced untreated males with tape placed over their cloaca to prevent copulation while still allowing courtship. We continued introducing newly emerged unmated females and untreated males until the end of the day. Soon after the termination of copulation, we removed the copulatory plug using a blunt probe [32,45] and measured plug mass (±0.01 g). We also visually inspected females for the presence of semen inside the cloaca. All trials were conducted in a single day and 25 s matings were recorded (14 ablated versus 11 control males).

(c) Statistical analysis

Copulation duration was analysed using two-way repeated measures ANOVA in ScaMPlot v. 11.0 with treatment and male mating number as factors. Logistic regression could not be used to test for the effect of copulation duration on plug deposition as there was complete separation in the data (plugs were produced in all copulations over, and none under, 400 s). Average copulation duration ranges from 480 to 1200 s in this species (reviewed in [34]). We set a cut-off (300 s) to obtain a test of independence of the effect of copulation duration on plug deposition using $\chi^2$ with Yates continuity correction. We then used the same test to determine whether treatments differed significantly in the number of copulations above and below the 300 s threshold.

In those matings that produced a plug, the plug mass data were normally distributed with equal variance, so we fit these models using generalized linear models (Gaussian distribution; identity link function) in R v. 2.15.3, [46]. Previous studies have demonstrated the effects of male and female size and hemipene (left or right) on plug mass [29]. We employed multi-model selection using the Akaike’s information criterion corrected for small sample sizes (AICc) on regression models (R: MuMIn package [47]). AICc allowed us to identify a set of model parameters that best fit our data while adding penalties for extra parameters [48–50]. Our initial candidate model included the following parameters: treatment, male and female size (snout-to-vent length, svl), hemipene, copulation duration as main effects, and male size by female size and treatment by copulation duration interactions. We hoped to use the best model (lowest AICc value) to describe our observed data; however, the top-ranked model included only the intercept. Instead, we identified the only significant model-averaged parameter, which was the interaction between copulation duration and treatment ($p = 0.0405$: electronic supplementary material, table S1). Thus, we constructed a model that included the main effects of treatment and copulation duration and their interaction. The analysis (ANCOVA) and graphing of this model was conducted in XSTAT v. 2012.6.02.

(d) Experiment 2: treated females mating in the natural den with untreated males

Twenty-four unmated females were collected 1 day before mating trials. Females were housed overnight in a nylon arena (1 × 1 × 1 M) and were provided water ad libitum. The next day, females were returned to Inwood quarry for assisted mating trials. Females randomly assigned to the local anaesthesia treatment (N = 12) received two 30 µl bilateral injections of 0.5% Marcaine (Bupivacaine HCl) directly lateral to the cloaca between the first and second dorsal scale rows approximately 30 min prior to mating trials [51,52]. Control females (N = 12) received injections of saline instead of Marcaine in the same locations. Immediately after the injections, females were placed in small arenas (45 cm diameter × 75 cm tall) and after 30 min females were placed in a natural aggregation of males within the center of the den. We had determined in experiment no. 1 that female body rolls could dislodge a male and prevent the hemipene from anchoring to the cloaca at intromission. Because we were interested in the female’s ability to affect copulation duration via cloacal and/or vaginal control, with the thumb and index finger we held each female approximately 1 cm caudal of the cloaca while males courted her, and then used a blunt probe to lift the ventral scale that covers the opening to her cloaca as soon as a male aligned with the female [53]. Females
from both treatments were handled the same. These assisted matings resulted in males evert ing a hemipene into the female’s cloaca after only 1–10 s, so the females were restrained for a very short interval. We released the female once intromission occurred, and the pair then copulated naturally. Therefore, females could roll during copulation after intromission and we noted that some females from each treatment rolled during copulation. When intromission occurred, a stopwatch was started immediately. After 1 min of copulation, the pair was gently moved from the large natural aggregation to a small circular arena (45 cm diameter × 75 cm tall) where they were constantly observed until copulation terminated and the duration was recorded (± 10 s). Once copulation was completed, the copulatory plug was removed and plug mass was measured (± 0.01 g).

(e) Statistical analysis
The copulation duration and plug mass (matings that produced a plug) satisfied normality and equal variance assumptions so we fit these models using generalized linear models (Gaussian distribution; identity link function) in R. We employed multi-model AICc selection as described in experiment 1. Our initial candidate model included the following parameters: treatment, male and female size (svl), hemipene as main effects variables and male by female size interaction. The top-ranked model of plug mass included treatment, female size (svl) and copulation duration as main effects, and the interaction of copulation duration with treatment. We plotted the plug mass model using the standardized residuals of plug mass given female size to better illustrate the effect of the interaction between treatment and copulation duration on plug mass (figure 5). The analysis and graphing of the best models were conducted in XLSTAT v. 2012.6.02 and/or SIGMAPLOT v. 11.

3. Results

(a) Experiment 1: spine-ablated males

(i) Copulation duration
As predicted, copulation duration was significantly shorter in spine-ablated males than control males, but not different in the first copulation pre-treatment (figure 2). Spine-ablated males (n = 14): copulation duration at first mating (pre-ablation; $\bar{X} \pm$ s.e.m., 855 ± 57 s; at second mating (post-ablation) 249 ± 73 s. Control males (n = 11): copulation duration at first mating $\bar{X} \pm$ s.e.m., 819 ± 68 s; at second mating 643 ± 90 s. Eight of the 14 ablated males copulated for less than 1 min and failed to deposit a copulatory plug. Visual inspection of the female cloaca revealed that five of these males did not inseminate the female, but three males did. When spine-ablated males achieved copulation, their hemipene shafts were much further outside the female cloaca compared with matings by intact males, and the hemipene was held in place by the crown of smaller spines in the middle of the hemipene (figure 1).

(ii) Copulatory plug mass
Only six of 14 spine-ablated males deposited a plug, compared with eight of 11 control males. The likelihood of plug deposition was significantly greater for copulations lasting more than or equal to 5 min than those less than 5 min (14/16 copulations more than 5 min produced a plug, versus 0/9 copulations less than 5 min; $\chi^2$ test of independence with Yates’ continuity correction: $\chi^2 = 14.52, p = 0.00014$). Copulations of spine-ablated males were more likely to be below the 5 min threshold (8/14 versus 1/10 for the control group; $\chi^2 = 3.94, p = 0.026$).

Copulatory plug mass was affected by both copulation duration and treatment. Plug mass increased with copulation duration in spine-ablated males but not in control males; this relationship was primarily driven by the small plugs made by males that copulated for very short periods (figure 3). Because the top-ranked model using the AICc model selection procedure was the intercept only model, we built a model based on the only significant model-averaged parameter, which was the interaction between copulation duration and treatment. We chose the model that included the main effects and the interaction between them (AICc = −56.8, $\Delta$i = 4.15; $R^2 = 0.445$). In this model, both copulation duration and treatment are significant predictors of plug mass (see electronic supplementary material, table S1).
Group/C22

Conflict interactions over copulation duration and copulatory

We found evidence that genital features play a role in sexual

4. Discussion

We found evidence that genital features play a role in sexual conflict interactions over copulation duration and copulatory

Figure 4. Copulation duration as a function of female treatment in locally anaesthetized females (Marcaine treated) versus saline-injected (control) females. Copulation was much longer in Marcaine-treated females. Bars in the graph represent the s.e.

(b) Experiment 2: anaesthetized females

(i) Copulation duration

Females whose cloaca and vagina were anaesthetized had longer copulation durations than control females. The top-ranked model for the copulation duration data only included ‘Treatment’ as a factor (AICc = 348.4, Δi = 0). When female body rolls were prevented at intromission, and their cloacal and vaginal tissues were anaesthetized (Marcaine treated), copulation duration was significantly longer in anaesthetized (M) than saline-injected (S) females (figure 4; t-test: t_{22} = 2.374, p = 0.027. Group, \( \bar{X} \) ± s.e.m.: M, 908 ± 96 s; S, 609 ± 82 s). Five females rolled during copulation (two M-treated and three saline-injected), rolling did not shorten copulation duration in this experiment (ANOVA with rolls nested within treatments, F_{3,20} = 2.246, p = 0.114), but we may not have the statistical power to detect the effect of rolls on copulation duration.

(ii) Copulatory plug mass

The best fit model for plug mass included treatment, female size (svl) and copulation duration as main effects and the interaction of copulation duration with treatment (AICc = -97.0, Δi = 0, R^2 = 0.434), all of which were highly significant (see electronic supplementary material, table S2). Overall, larger females received larger plugs, and the treatments had opposite effects on plug mass. After controlling for the effect of female body size (svl), Marcaine-treated females received smaller plugs as copulation duration increased. In the control females, after controlling for the effect of female body size (svl), longer copulation durations yielded larger plugs (figure 5). The longest copulation duration for the Marcaine-treated females is within the observed range for copulation durations in this species [29,31,34,45,53].

Figure 5. Standardized residuals of plug mass given female snout-to-vent length plotted as a function of copulation duration. This analysis only includes those matings that produced a plug. There are separate linear regressions for each treatment to illustrate the significant interaction between treatment and copulation duration (Marcaine: open triangles, dashed regression line, R^2 = 0.641, \( p = 0.002 \); saline control: filled circles, solid regression line, R^2 = 0.239, \( p = 0.152 \)).

(a) Spine-ablation

As predicted, copulation duration decreased when the hemipenial basal spine was ablated. The decrease was mostly driven by matings of less than 1 min that were typically terminated by female body rolls. These results support our initial hypothesis that the basal spine allows males to gain and maintain intromission despite female behaviours that shorten copulation. Overall, short copulations were less likely to produce copulatory plugs. As predicted, spine-ablated males produced smaller copulatory plugs, and this relationship was primarily owing to very small plugs produced during short copulations.

Although the basal spine helps males maintain intromission despite female body rolls, the spine is not required for males to evert their hemipenes inside the female as suggested by Pisani [37]. This is unsurprising given that the basal spine is absent in non-natricine snakes, which have functional hemipenes [54–58]. However, the spine allowed a better fit between male and female genitalia, because spine-ablated hemipenes [54–58]. However, the spine allowed a better fit between male and female genitalia, because spine-ablated males often appeared to be farther outside the female cloaca (CR Friesen 2009, 2011, & PLR Brennan, 2011, personal observation). Full eversion and inflation of the hemipene takes place a few seconds after copulation begins (CR Friesen 2009, 2011, PLR Brennan, 2011 & RT Mason 1999–2011, personal observation), but males begin inseminating females almost immediately upon eversion and prior to plug formation [11,31]. Males who were unable to make a plug, may therefore still gain some fertilizations, but with the shortened copulation duration in this species versus other snakes [31,59], the plug became necessary to prevent sperm leakage [32] and a potential source of sexual conflict.

Traits involved in sexual conflict may have originally evolved as a result of male–male competition [60]. Male–
male competition in red-sided garter snakes is manifested in ‘tail wrestling’ rather than body-combat, and males constantly push their tails in between a female and any other courting male [61]. Many natrixine snakes, all of which have basal hemipene spines, mate in aggregations, and have no overt male combat [62]. Having a basal spine may help to quickly initiate copulation during tail wrestling, but also to prevent competitors from interrupting intromission when multiple males’ tails are proximate to the female’s cloaca. The evolution of the basal spine allows males to gain more control over copulation duration, forcing females to evolve some counter trait to regain some control, leading to sexually antagonistic coevolution.

(b) Anaesthetized females

Anaesthetization of the female cloaca and vaginal pouch increased copulation duration as we had predicted. In this experiment, we restrained all females (treated and control) during the initiation of copulation to minimize female body rolls during intromission, but females could roll after the hemipene was anchored in place, but this did not affect copulation duration in this study. Thus the increased copulation duration in anaesthetized females was then likely due to relaxation of the musculature of the vaginal pouch and cloaca. This suggests that these structures, in addition to body rolls, are involved in female control of copulation duration. Interestingly, average copulation duration has been reported to be about 840–1176 s [29,34], and in our first experiment, with unassisted mating, copulation duration was 840 ± 43 s. However, in our second experiment, when we prevented females from rolling, copulation duration of control females was much shorter (609 ± 82 s). It is possible that control females used their vaginal and cloacal tissues to make copulations much shorter than if we had allowed normal unassisted copulations to occur because they were prevented from rejecting unwanted males during intromission.

The effect of the interaction between treatment and copulation duration on copulatory plug mass was complex. We predicted that longer copulations would result in increased plug mass and, therefore, that Marcaine-treated females would have larger plugs deposited within them. However, plug mass was smaller than expected in Marcaine-treated females after accounting for female body size, and this deficit increased with copulation duration. This could mean that the cloacal and vaginal tissues provide a surface against which males can pack their copulatory plug material and that anaesthesia relaxes the tissues so that this packing of material is less efficient such that it takes longer to deposit the plug. The size and density of the copulatory plug may depend on those same muscular contractions of the vaginal pouch owing to a change in volume of the pouch and/or time in copulo. Anecdotally, in experiment 1, we observed that most of the plugs produced by spine-ablated males were less formed, less dense and extruded around the hemipene. Those plugs produced during control mating were more solid and compact, perhaps because the tip of the hemipene was further from the vaginal pouch walls, but we did not quantify this variable. Further anecdotal evidence that the vaginal pouch musculature limits plug size comes from a field observation in which a male deposited a plug within the rectum of a female (CR Friesen, PLR Brennan & RT Mason, 2011, personal observation). The plug deposited in the rectum was very large and not densely packed, possibly because the female could not squeeze the rectum to limit space.

It can be very difficult to distinguish between sexual conflict and CFC hypotheses of genital evolution [41], and here we hypothesized that sexual conflict rather than CFC explains the role of the basal spine in increasing copulation duration. Under the CFC hypothesis, the basal spine could be a stimulatory trait to the female. If this were the case, then spine ablation would result in shorter copulation duration because females rejected males that failed to properly stimulate them rather than because males could not remain attached when females were rolling or actively using their vaginal muscles to eject the male as proposed by sexual conflict. According to CFC, when females were anaesthetized (i.e. experimentally ‘blind’ [41]), copulation duration should have been shorter because females could not sense the males’ stimulation and would be more likely to reject them [63], contrary to our findings. The basal spine may cause physical harm to the female genitalia as females often bleed during and after copulation, likely as a result of the basal spine penetrating the cloacal mucosa (CR Friesen 2006–2011 & PLR Brennan, 2011, personal observation). Damage to the female by the basal spine during mating suggests conflict during mating rather than a stimulatory function hypothesized by CFC. Furthermore, the epithelial tissue of the female cloaca in which the basal spine embeds is thickly stratified and keratinized [64], which may be a female counter-adaptation to minimize damage owing to sexual conflict as has evolved in bedbugs with traumatic insemination [65]. Even if males with bigger, more damaging spines achieved longer copulations, this would not represent evidence of a process of female choice, but rather an expected outcome of conflict interactions [60]. We cannot, however, rule out that males may expect resistance and interpreted the anaesthetized females as weakened and unfit leading them to reduce allocation of their ejaculate accordingly, if males mate multiply during the breeding season [66]. Male mate choice is often under-appreciated [67]. Male garter snakes have been demonstrated to preferentially court larger, more fecund females [62,68–72]. There is evidence that males allocate more plug material but not more sperm to larger females [45] and males are significantly less likely to deposit a plug after two matings [32]. Ejaculates can be costly to produce [73,74], and plug production appears to be energetically expensive for male red-sided garter snakes [75]. Given these facts, male ejaculate adjustment with respect to female size and condition deserves further investigation in this species. In this study, we measured copulatory plug mass and copulation duration, but other variables related to plug formation may be important in conflict interactions. For example, the optimal rate at which the plug dissolves would be expected to be higher for females than males but the optimal rate of dissolution may vary with male quality. Females might benefit from the evolution of female enzymes that breakdown the plug matrix to coevolve with the protein composition of the plug, which would allow them to mate sooner after mating within the den, where mating is random with respect to male size. In conclusion, we have documented that manipulation of genital traits changes copulation duration and plug size in a manner mostly consistent with predictions of sexual conflict over mating using a new and tractable model species. These are wild populations of animals so although they are manipulated, they exhibit typical mating behaviours which are difficult to observe in other species of vertebrates.
This provides a novel vertebrate model system that is especially suited for manipulation and controlled experiments. Procedures performed on animals were approved by Yale (IACUC 2009-11290) and Oregon State University (IACUC 2009-11 ACUP-3738), and the research was conducted under permit from Manitoba Conservation (WBI1240).

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