Mate-copying in Drosophila: a matter of taste or disgust?

Guillaume Lespagnol

May 18, 2019

Contents

1	Intr	roduction	2
2	Material and Methods		5
	2.1	Fly maintenance	5
	2.2	Fly stains and crossings	5
3	General experimental procedures		6
	3.1	Acceptance/Rejection experiment	7
	3.2	Neuronal blockade experiment	8
	3.3	Mate-copying score	8
	3.4	Statistical analyses	9
	3.5	Ethical statements	10
4	Results		10
	4.1	A/R Experiment	10
	(Dai	nchin et al. 2018, p. 118). Or several: Danchin et al. (e.g. 2018, p. 118) a	nd
Da	anchir	n et al. (but see 2018, p. 1789)	

1 Introduction

In the animal kingdom, you must win to mates. In some species, winning is a matter of weight, strength or weapons (Clutton-Brock 1982; Anderson and Fedak 1985). In others it's a matter of charms in a much more peaceful contest, and the breeders are chosen by the other sex. The mate-choice (i.e intersexual selection) can lead to captivating and highly complex traits, such as courtships or songs in birds (Danchin et al. 2005). Nevertheless, many species do not exhibit such traits and the choice is based on much more ambiguous and discreet signs. "The most difficult and important act of choice is the choice of a mate" (Fisher 1915), in context of mate-choice making a mistake can be very expensive since it impact individual's offspring. To avoid mistakes, many species have acquired the capacity to learn from the observation of others and can therefore use social learning in numerous decision-making processes.

Social learning can take different form, as teaching or copying. The latter is simpler, and even exists in non-social invertebrates (Coolen et al. 2005; Laidre Mark E. 2010). Many differents behaviours can be copied (Thornton Alex et al. 2010; Leeuwen et al. 2014). It's particularly beneficial when individual learning is costly (time consuming or dangerous; also see (Webster M.M and Laland K.N 2008)) as in mate-choice. Therefore, copying the mate-choice of potentially older conspecifics can be a reliable choice for naive individuals to avoid unnecessary mistakes.

Mate-copying is a form of social learning in which the observation of a sexual interaction between conspecifics biases the subsequent mate-choice decision of the observer (Brown et al. 2011). It has been first demonstrated in fishes (Dugatkin Lee Alan and Godin Jean-guy J. 1992), followed by observations in many vertebrates (Galef and White 1998; Yorzinski and Platt 2010) and recently in invertebrates (Mery et al. 2009; Fowler-Finn et al. 2015). Its benefits are double-sided, it allows naive individuals to avoid mistakes and make sure that their descendant will be preferred by conspecifics. Interestingly,

in population with genetic preferences, mate-copying can override them (Dugatkin 1996; Witte and Ryan 1998). At larger scale, mate-copying can even shape preferences of entire populations and potentially lead to formation of local traditions. Such traditions transmitted vertically and horizontally, for a trait-based preference and possibly for a long time can even lead to the appearance of animal culture (Brooks 1998; Danchin et al. 2018).

The existence of culture in non-human species has long been disputed (Laland and Hoppitt 2003) but is increasingly accepted among scientists (Aplin et al. 2015; Whitehead 2017). The list of animals exhibiting a form of cultural tradition is growing constantly (Schaik et al. 2003; Thornton Alex et al. 2010; Whiten 2017), and one of the most recent may surprise many, Drosophila melanogaster (Danchin et al. 2018). Very few, if none, species have been studied as deeply and for whom knowledge in every scientific field (genetics, development, neuroscience...) is as extensive than Drosophila Melanogaster. Thus existence of mate-copying in this species represents a wondeful opportunity to understand the obscure neuronal root of an evolutive behavior broadly shared in the animal kingdom.

In drosophila, depending on if the stimulus is aversive or appetitive, different groups of neurons are involved (Busto et al. 2010; Vogt et al. 2014). Therefore, our first step was to test if mate-copying implies aversive or appetitive memory, by isolating different kinds of stimulus contained in copulation of conspecifics. We considered that a rejection represent an aversive stimulus and acceptance of copulation an appetitive stimulus for an observer female. So, we created two treatments by presenting a male rejected by a female ("Rejection" treatment) or a male accepted by a female ("Acceptance" treatment), for which we measured the inclination to copy.

Then we investigated deeper neuronal mechanisms of mate-copying by searching which group of neurons is required for mate-copying. Indeed, the neuronal mechanisms of non-social learning is very well known in Drosophila (reviewed in Cognigni et al. (2018)). Regarding their roles in the latter, two brain structures are particularly prone to be involved in mate-copying, the central complex and the mushroom body. The central complex, localized in the center of the insect brain plays a major role in decoding visual information.

It receives visual inputs from the rest of the brain and controls vision-related behaviors, memory and learning (Guo et al. 2017). The mushroom body is an integrative center involved in learning, memory, decision-making and visual associative memory. Notably, specific groups of dopaminergic neurons localized in mushroom bodies are involved in the acquisition of aversive and appetitive visual memory (Liu et al. 2012; Vogt et al. 2014). On the contrary, despite a rich repertoire of well-studied social processes (Pasquaretta et al. 2016; Teseo et al. 2016; Dawson et al. 2018), neuronal mechanisms of social learning are still poorly understood. However, a recent study have found that dopamine is required in mate-copying (Monier et al. 2018). Dopamine is a neurotransmitter, that drive a variety of brain function among which the formation of appetitive and aversive memory (Riemensperger et al. 2005; Sitaraman et al. 2008; Alekseyenko et al. 2010; Berry et al. 2012; Yamamoto and Seto 2014). Dopamine is produced in dopaminergic neurons but we do not know which are involved in mate-copying. However, we do know some of the neurons required for non-social visual memory. Vogt et al. (2014) showed that neurons labeled by TH-GAL4 and Ddc-GAL4 transgene are essential in in visual learning. Given that mate-copying involves visual learning, we expect these neurons to be required. Thanks to Liu et al., we know that UAS-GAL4 technology can be used to block specific sets of neurons (Liu et al. 2012). Precisely, by coupling UAS-GAL4 with the Shibire protein, it is possible to temporary silence specific groups of neurons and investigate in what processes they are essential. Our goal was to use a similar technique but concerning mate-copying. We used mutant flies expressing GAL4 in TH-labeled or Ddc-labeled neurons, that we have silenced to see if those neurons are required in mate-copying (for more details, see "Fly stains and crossings" section). We thus created two treatments, depending of the group of the neurons silenced, for wich we calculated mate-copying scores. If one group is involved in mate-copying, the corresponding treatment will exhibit a mate-copying score similar to random choice, due to the incapacity of mutants to learn.

2 Material and Methods

2.1 Fly maintenance

We used the common Canton-S strain of D.melanogaster (wild-type, and UAS / Gal4 lines described above). Flies were raised and kept in 30 ml tubes containing standard corn flour-yeast-agar medium at $25^{\circ} \pm 1^{\circ}$ C and 56 ± 4 % humidity with a 12:12H light:dark cycle. Humidity and temperature were controlled and adjusted continuously with two independents automatic humidifiers and one manual heater. Medium was cooked every 3 weeks and stored at 4°C until use. Flies were manipulated with a hand-made mouth aspirator made of a glass pipette, tubing and gauze.

Every morning, adult flies were removed from the breeding vials so that the newly emerged flies collected within the 6-8 hours were virgin. For Canton-S strain, 120 males and 120 females were used daily for breeding (20 tubes with 6 males and 6 females in each) and all other adults were euthanized in a freezer. For mutant strains, all adults were used for breeding.

Virgins were sexed without anesthesia, by gentle aspiration and then kept in unisex groups of 7 females or 14 males until experiments. Both demonstrator and observer flies were 3 or 4 days old. Males and females were used only once as females are reluctant to re-mate (Chapman et al. 2003) and reject males they just saw copulating (Loyau et al. 2012). After experiments, all flies were put in a food vial and cold-euthanized at the end of the day.

2.2 Fly stains and crossings

For the second experiment, we used two mutant genotypes, Ddc-GAL4/w+;;UAS-Shits/+ and w+/w-;;UAS-Shits/TH-GAL4, obtained by crossing homozygous lines.

UAS-Shits is a transgene that contains an GAL4-specific enhancer, UAS (Upstream Activating Sequence) driving the production of Shibire protein in cells where GAL4 is present. Shibire is a thermosensitive protein that inhibit neuronal activity at restrictive temperature (30°C) by preventing vesicle recycling (Kitamoto 2001). Ddc-GAL4 and

TH-GAL4 drive production of a transcriptional activator (GAL4) only in specific subsets of dopaminergic neurons. GAL4 activates the expression of genes downstream to UAS. Ddc-GAL4 labels neurons involved in appetitive olfactory memory:the blockade of these neurons by Shibire protein has been shown to impair the acquisition of such memory at restrictive temperature.TH-GAL4 labels neurons involved in aversive olfactory memory (Liu et al. 2012).

As white recessive mutation w- impairs fly vision (Götz 1964), we used mutant females with one wild-type copy of the white gene for the experiments. To do so, we crossed w+;;Shits males with females from each Gal4 line. To obtain w+;;Shits strain from a w-;;Shits strain, we crossed males w-;;Shits with females w+;;TM2/TM6b over two generations and selected TM2 non TM6B flies only, with CO2 anesthesia, and we then isolated homozygous w+;;Shits progeny.

TH-GAL4 and Ddc-GAL4 lines were provided by Guillaume Isabel in the same Canton-S background as the Wild-type strain.

3 General experimental procedures

Artificial male phenotypes were created by dusting virgin males with pink or green powder (Mery et al. 2009). Each vial of males was randomly assigned to a color. Before the experiment started, males were placed in a clean vial to remove the excess of dust for at least 20-30 min. Experiments took place in the same tube set-up and a similar but slightly modified speed-learning protocol than described in Dagaeff et al. (2016) (Figure 1, see also specific experiment section).

Demonstrator and observer flies were placed in two compartments of double plastic tubes, separated by a thin glass partition and closed by cotton plugs. All replicates were run in blocks of six trials with cardboard barriers between experimental set-ups, to prevent information exchange between the flies and disturbance from the surroundings.

During the demonstration, we always showed two different male phenotypes (color) to the observer females, with one favorite that copulated with the demonstrator female.

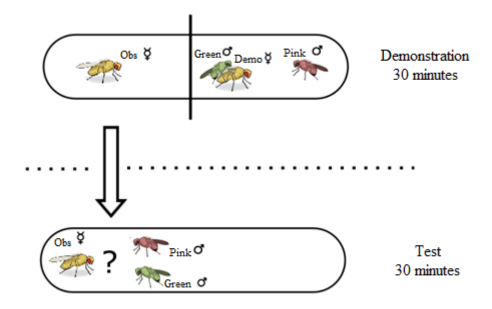


Figure 1: Légende ici.

If demonstrator females refused to copulate, the trial was discarded. Specifics of demonstrations for each experiment are described above (see specific experiment section).

Once the demonstrations were over, we started the mate-copying test by introducing a couple of colored virgin males (one of each color) in front of the observer female and we removed the partition, allowing the female to freely choose between males for 30 min. The partition was put back in place when all three flies were in the same side of the tubes, to promote proximity between flies. During that time, we recorded the time of first courtship for each male, the time when copulation started and the color of the chosen male. The onset of the courtship was defined as the first wing-extension of a male (Figure 2).

3.1 Acceptance/Rejection experiment

During this first experiment, we tested whether mate-copying is achieved through aversive or appetitive memory. To do so, we split the negative and positive information given by the usual mate-copying demonstration. A classical demonstration, in which a demonstrator female chooses between two males, contains a rejection (negative information) of a male and an acceptance (positive information) of the other one. We thus created three

demonstration treatments: (1) a control where a demonstrator female freely chooses between two males, (2) an "acceptance" treatment with one accepted male copulating with a demonstrator female, and (3) a rejection treatment with one male actively rejected by a female (Figure 3).

3.2 Neuronal blockade experiment

This second experiment aimed at exploring the mechanisms underlying the results of the Acceptance/Rejection experiment by discovering one group of dopaminergic neurons involved in mate-copying.

The demonstration was similar to classical protocol (Figure 1), but we used Ddc-GAL4/w+;;UAS-Shits/+ (treatment "Ddc") and w+/w-;;UAS-Shits/TH-GAL4 mutants (treatment "Gal4") as observer females. The use of these mutants allowed us to have a temporal control on specific sets of neurons presumably involved in appetitive (Ddc) or aversive (TH) memory, thanks to the thermosensitive activation of Shibire. During the demonstration and 30 minutes before, observer mutant females were heated to a restrictive temperature of 33°C, thanks to a heating mat under the tube of these females. At 33°C, Shibire protein blocks the neurons in which it is expressed, and thus the acquisition of appetitive memory should be blocked in observer females of "Ddc" treatment, and the acquisition of aversive memory in "TH" females.

After all copulations ended, demonstrator males and females were removed. Observer females were then stored individually in clean tubes at 25°C for 3-4 hours to ensure that labelled neurons are no more blocked, then we proceeded to a classical test at 25°C.

3.3 Mate-copying score

As in previous studies (Danchin et al. 2018; Monier et al. 2018; Nöbel et al. 2018), a mate-copying score evaluated female's tendency to copy the choice of the demonstrator. A mate-copying score of 1 was assigned to females that copulated with the color preferred by demonstrator females and a score of 0 in the opposite case. For each treatment, a

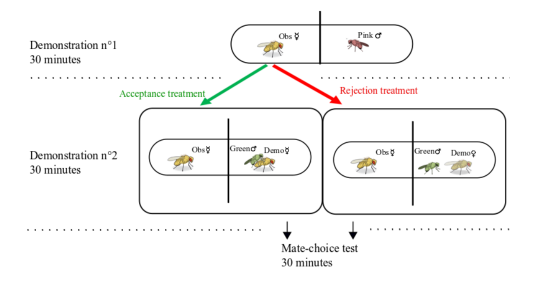


Figure 2: Légende ici.

mate-copying index was calculated as the mean of mate-copying scores per treatment, a random choice indicated by a value of 0.5. All replicates where only one male courted the female before copulation were discarded because in these situations the female was not unambiguously in a position to make a choice between the two colors.

3.4 Statistical analyses

All statistical analyses were performed with the R software version 3.5.1 (R Core Team, 2018). For each treatment, the difference from a random choice was tested with a binomial test. Mate-copying scores were then analyzed in a generalized linear mixed model (GLMM, package lme4 (Bates et al. 2015)). Starting models contained the following fixed effects: treatment, normalized air pressure (air pressure in Toulouse-Blagnac weather station, at the time of the beginning of the experiment, minus mean air pressure), normalized air pressure variation within the six preceding hours and all interaction between these three variables, experimenter effect and its interaction with treatment. A random "block" effect was also introduced in the models to account for the non-independence of observer flies from the same block of 6 tubes-set up trained and tested in parallel. The significance of fixed effects was tested using Wald chi-square tests included in ANOVA function (car

package, Fox and Weisberg (2018)). Model simplification was achieved by successive withdrawal of the non-significant terms in a backward selection approach, using P-values and starting with the highest-order interaction. The final model was chosen as the one with the lowest Akaike Information Criteria (AIC, Akaike, 1969). Comparisons between treatments were done using post-hoc X² tests.

3.5 Ethical statements

Behavioral observations of D. melanogaster required no ethical approval and complied with French laws regarding animal welfare. We kept the number of flies used in this study as small as possible. We handled flies by gentle aspiration without anesthesia to minimize damage and discomfort. After the experiments, individuals were euthanized in a freezer at -20°C.

4 Results

4.1 A/R Experiment

In total, we tested 850 females among which 530 copulated, including 192 with double-courtship, 64 for each treatment. First we tested for female's color preference with binomial test but neither in the demonstration (N = 850, 426 females copulated with green males and 424 copulated with pink males; binomial test: P = 0.973) nor the test (N = 192, 90 copulated with pink and 102 with green males; binomial test: P = 0.427) was there any significant difference between the two colors.

For each treatment, the difference from random choice was tested with a binomial test, acceptance (where the observer female sees a copulation, N = 64, P < 0.001) and control (N = 64, P = 0.03) were both significantly different from random, but rejection treatment (where the observer female sees a male rejected without copulation, N = 64, P = 0.382) was not (Figure 4).

To test for the significance of mate copying among treatments, we built a global

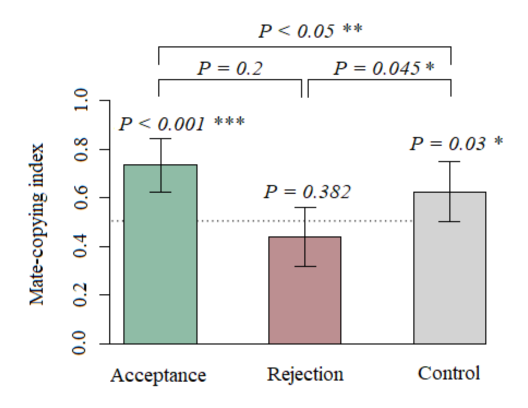


Figure 3: Légende ici.

model including the effects of experimenter, treatment, normalized air pressure (actual air pressure minus global mean of air pressure), normalized air pressure variation (for the last six hours) and the interaction between air pressure and variation of air pressure. Only the treatment had a significant effect on mate-copying (GLMM, χ^2 : N=192, $\chi^2=10.447$, p=0.005), air pressure (GLMM, χ^2 : N=192, $\chi^2=1.831$, P=0.176) were non-significant. The interaction between air pressure and variation of air pressure was close to be significant (GLMM χ^2 : N=192, $\chi^2=2.946$, P=0.086), as we could have expected in regard of the results of Dagaeff et al. (2016). No significant difference has been found between control and acceptance treatments ($\chi^2=1.76$, P=0.18; Fig4), but both are different from rejection treatment (acceptance - rejection: $\chi^2=11.62$, P<0.005; control - rejection: $\chi^2=4.52$, P=0.033; Fig4).

Voir Figure (??).

References

- Alekseyenko, Olga V., Carol Lee, and Edward A. Kravitz (May 2010). "Targeted Manipulation of Serotonergic Neurotransmission Affects the Escalation of Aggression in Adult Male Drosophila melanogaster". In: 5.5, e10806.
- Anderson, Sheila S. and Michael A. Fedak (Aug. 1985). "Grey seal males: energetic and behavioural links between size and sexual success". In: 33.3, pp. 829–838.
- Aplin, Lucy M. et al. (Feb. 2015). "Experimentally induced innovations lead to persistent culture via conformity in wild birds". In: 518.7540, pp. 538–541.
- Bates, Douglas et al. (Oct. 2015). "Fitting Linear Mixed-Effects Models Using lme4". In: 67.1, pp. 1–48.
- Berry, Jacob A. et al. (May 2012). "Dopamine Is Required for Learning and Forgetting in Drosophila". In: 74.3, pp. 530–542.
- Brooks, R. (Feb. 1998). "The importance of mate copying and cultural inheritance of mating preferences". In: 13.2, pp. 45–46.
- Brown, Culum, Kevin Laland, and Jens Krause (June 2011). Fish Cognition and Behavior. John Wiley & Sons.
- Busto, Germain U., Isaac Cervantes-Sandoval, and Ronald L. Davis (Dec. 2010). "Olfactory learning in Drosophila". In: 25.6, pp. 338–346.
- Chapman, Tracey et al. (Aug. 2003). "The sex peptide of Drosophila melanogaster: female post-mating responses analyzed by using RNA interference". In: 100.17, pp. 9923–9928.
- Clutton-Brock, T. H. (Jan. 1982). "The Functions of Antlers". In: 79.2, pp. 108–124.
- Cognigni, Paola, Johannes Felsenberg, and Scott Waddell (2018). "Do the right thing: neural network mechanisms of memory formation, expression and update in Drosophila". In: 49, pp. 51–58.
- Coolen, Isabelle, Olivier Dangles, and Jérôme Casas (Nov. 2005). "Social Learning in Noncolonial Insects?" In: 15.21, pp. 1931–1935.
- Dagaeff, Anne-Cecile et al. (Nov. 2016). "Drosophila mate copying correlates with atmospheric pressure in a speed learning situation". In: 121, pp. 163–174.

- Danchin, Étienne, Luc-Alain Giraldeau, and Frank Cézilly (Mar. 2005). Écologie comportementale: Cours et questions de réflexion. Dunod.
- Danchin, Etienne et al. (Nov. 2018). "Cultural flies: Conformist social learning in fruitflies predicts long-lasting mate-choice traditions". In: 362.6418, pp. 1025–1030.
- Dawson, Erika et al. (Sept. 2018). "Social environment mediates cancer progression in Drosophila". In: 9, p. 3574.
- Dugatkin Lee Alan and Godin Jean-guy J. (Aug. 1992). "Reversal of female mate choice by copying in the guppy (Poecilia reticulata)". In: 249.1325, pp. 179–184.
- Dugatkin, L A (Apr. 1996). "Interface between culturally based preferences and genetic preferences: female mate choice in Poecilia reticulata." In: 93.7, pp. 2770–2773.
- Fisher, R. A. (Oct. 1915). "The evolution of sexual preference". In: 7.3, pp. 184–192.
- Fowler-Finn, Kasey D. et al. (Dec. 2015). "The complexities of female mate choice and male polymorphisms: Elucidating the role of genetics, age, and mate-choice copying". In: 61.6, pp. 1015–1035.
- Fox, John and Sanford Weisberg (Sept. 2018). An R Companion to Applied Regression. SAGE Publications.
- Galef BENNETT G., JR and DAVID J. White (Mar. 1998). "Mate-choice copying in Japanese quail, Coturnix coturnix japonica". In: 55.3, pp. 545–552.
- Götz, Karl Geokg (June 1964). "Optomotorische Untersuchung des visuellen systems einiger Augenmutanten der Fruchtfliege Drosophila". In: 2.2, pp. 77–92.
- Guo, Aike et al. (Dec. 2017). "Vision, Memory, and Cognition in Drosophila". In: Reference Module in Neuroscience and Biobehavioral Psychology.
- Kitamoto, T. (May 2001). "Conditional modification of behavior in Drosophila by targeted expression of a temperature-sensitive shibire allele in defined neurons". In: 47.2, pp. 81–92.
- Laidre Mark E. (May 2010). "How rugged individualists enable one another to find food and shelter: field experiments with tropical hermit crabs". In: 277.1686, pp. 1361–1369.
- Laland, Kevin N. and William Hoppitt (Jan. 2003). "Do animals have culture?" In: *Evolutionary Anthropology: Issues, News, and Reviews* 12.3, pp. 150–159.

- Leeuwen, Edwin J. C. van, Katherine A. Cronin, and Daniel B. M. Haun (Nov. 2014). "A group-specific arbitrary tradition in chimpanzees (Pan troglodytes)". In: 17.6, pp. 1421–1425.
- Liu, Chang et al. (Aug. 2012). "A subset of dopamine neurons signals reward for odour memory in Drosophila". In: 488.7412, pp. 512–516.
- Loyau, Adeline et al. (Oct. 2012). "When not to copy: Female fruit flies use sophisticated public information to avoid mated males". In: 2, p. 768.
- Mery, Frédéric et al. (May 2009). "Public versus personal information for mate copying in an invertebrate". In: 19.9, pp. 730–734.
- Monier, Magdalena et al. (2018). "Dopamine and Serotonin Are Both Required for Mate-Copying in Drosophila melanogaster". In: 12, p. 334.
- Nöbel, Sabine, Etienne Danchin, and Guillaume Isabel (Sept. 2018). "Mate-copying for a costly variant in Drosophila melanogaster females". In: 29.5, pp. 1150–1156.
- Pasquaretta, Cristian et al. (Mar. 2016). "How social network structure affects decision-making in Drosophila melanogaster". In: 283.
- Riemensperger, Thomas et al. (Nov. 2005). "Punishment Prediction by Dopaminergic Neurons in Drosophila". In: 15.21, pp. 1953–1960.
- Schaik, Carel P. van et al. (Jan. 2003). "Orangutan cultures and the evolution of material culture". In: 299.5603, pp. 102–105.
- Sitaraman, Divya et al. (Apr. 2008). "Serotonin is necessary for place memory in Drosophila". In: 105.14, pp. 5579–5584.
- Teseo, Serafino, Liisa Veerus, and Frédéric Mery (June 2016). "Fighting experience affects fruit fly behavior in a mating context". In: 103.
- Thornton Alex, Samson Jamie, and Clutton-Brock Tim (Dec. 2010). "Multi-generational persistence of traditions in neighbouring meerkat groups". In: 277.1700, pp. 3623–3629.
- Vogt, Katrin et al. (Aug. 2014). "Shared mushroom body circuits underlie visual and olfactory memories in Drosophila". In: 3. Ed. by Mani Ramaswami, e02395.

- Webster M.M and Laland K.N (Dec. 2008). "Social learning strategies and predation risk: minnows copy only when using private information would be costly". In: 275.1653, pp. 2869–2876.
- Whitehead, Hal (July 2017). "Gene-culture coevolution in whales and dolphins". In: 114.30, pp. 7814–7821.
- Whiten, Andrew (July 2017). "Culture extends the scope of evolutionary biology in the great apes". In: 114.30, pp. 7790–7797.
- Witte, Klaudia and Michael J. Ryan (Jan. 1998). "Male body length influences mate-choice copying in the sailfin molly Poecilia latipinna". In: 9.5, pp. 534–539.
- Yamamoto, Shinya and Elaine S. Seto (2014). "Dopamine Dynamics and Signaling in Drosophila: An Overview of Genes, Drugs and Behavioral Paradigms". In: 63.2, pp. 107–119.
- Yorzinski, Jessica L. and Michael L. Platt (Feb. 2010). "Same-Sex Gaze Attraction Influences Mate-Choice Copying in Humans". In: 5.2.