Multiple phenotypic traits predict male mating success in a critically endangered frog

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Abstract
Complex sexual signals spanning multiple sensory modalities may be common in nature, yet few studies have explored how combinations of phenotypic traits influence male attractiveness and mating success. Here, we investigate whether combinations of multiple male phenotypic traits (both within and across sensory modalities) predict male mating and fertilization success in the critically endangered southern corroboree frog, *Pseudophryne corroboree*. We conducted breeding trials in a standardized captive environment where females were given the opportunity to choose between multiple males over the duration of the breeding season. For each male, we measured multiple call traits, aspects of coloration, body size, and age. We found that complex interactions between multiple traits best predicted male mating and fertilization success. In general, males with lower call frequency, lower call rate, and shorter call duration had the highest mating and fertilization success. Fertilization success was additionally linked to male body size and age. These findings suggest that female *P. corroboree* select mates based on a suite of acoustic traits, adding to a growing body of evidence that females use multiple traits to assess male quality. Our results also suggest that females may combine information from multiple signals non-additively. Moreover, our results imply that females gain direct fertility benefits from their mate choice decisions. We argue that understanding female mate choice based on various signals across multiple sensory modalities has important implications for the integration of mate choice into conservation breeding programs and needs to be considered when developing behavior-based captive breeding strategies.

Significance statement
Sexual signals are often highly complex, yet we know little about how multiple signal components both within and across various sensory modalities predict male mating success. We investigated whether combinations of multiple phenotypic traits (within and across sensory modalities) predicted male breeding success in threatened corroboree frogs. We conducted captive breeding trials in a homogeneous environment, where females could choose between multiple males over the duration of a single breeding season. We found that interactions between multiple male traits predicted mating and fertilization success. Males with lower call frequency, call rate, and duration had higher mating success. Fertilization success was also linked to acoustic signals, body size, and age. Understanding male choice for multiple traits further elucidates the complexity of female mate choice. This study is one of the first to consider the conservation implications of multimodal signaling in mate choice.

Keywords Captive breeding · Conservation · Male mating success · Female mate choice · Multiple signals · Multimodal signaling · Amphibian · Conservation

Introduction

Female mate choice is a strong evolutionary factor that can have significant consequences for female fitness (Andersson 1994; Rosenthal 2017). By preferentially mating with certain males, females can significantly improve their reproductive
success, as well as the fitness of their offspring (Petrie 1994; Martin-Wintle et al. 2015; Hartnett et al. 2018; Parrott et al. 2019). In resource-based mating systems, where males provide resources essential for reproduction, females can obtain a combination of both direct and genetic benefits from their mating decisions, as these benefits are not mutually exclusive (Candolin 2003; Bussière et al. 2005). In these systems, females likely use multiple signals to evaluate the quality of potential mates and the probability of securing direct and/or genetic benefits (Candolin 2003). In a diversity of taxa, females discriminate among males based on various condition-dependent secondary sexual traits (such as extravagant courtship displays, ornamentation, or bright coloration) that can act as an honest signal of a male’s capacity to secure resources (Reaney and Backwell 2007), fertility (Robertson 1990), potential to provide paternal care (Montoya and Torres 2015), and/or genetic quality (Montoya and Torres 2015). Such benefits not only stand to elevate female fitness but may also improve the viability of populations (Cally et al. 2019).

Despite the significant fitness benefits that can derive from female mate choice, knowledge of intersexual selection is rarely incorporated into conservation breeding programs (CBPs, encompassing captive breeding and reintroduction) that aim to conserve and manage endangered species (Martin-Wintle et al. 2019). Many CBPs aim to establish viable, self-sustaining populations ex situ, while also providing genetically diverse individuals for population reintroduction in situ. However, for some threatened species, breeding success in captive colonies is unreliable (Wielbickowski 1998). Reproductive failures are often attributed to a lack of knowledge of the reproductive ecology of a target species, including mate choice behavior. Additionally, providing opportunities for mate choice may be disregarded in favor of genetic goals (i.e., individuals are strategically paired to maintain sufficient genetic variation). Yet, a growing body of empirical research has demonstrated that incorporating mate choice into CBPs can significantly bolster reproductive output and offspring fitness (Hartnett et al. 2018; Martin-Wintle et al. 2019), as well as preserve natural reproductive behavior and enhance animal welfare (e.g., by decreasing stress and aggression between mates) (Asa et al. 2011). Considering these benefits, gathering information on mechanisms of mate choice in species targeted for captive breeding has been identified as a crucial step towards integrating knowledge of sexual selection into CBPs (Quader 2005; Asa et al. 2011; Chargé et al. 2014). Specifically, advancing our understanding of factors that determine male attractiveness, and the phenotypic traits used by females to discriminate among potential mates, is likely to facilitate the incorporation of mate choice into captive management, and increase CBP outcomes (Asa et al. 2011; Martin-Wintle et al. 2019). Captive environments provide an ideal opportunity to dissect the mechanisms of female mate choice because mating contexts can be reliably manipulated to uncover the importance of different variables. For instance, breeding environments can be manipulated so that they are relatively homogeneous (compared with natural breeding environments), resulting in less variation in territory/resource quality, and/or climatic conditions (such as temperature and humidity), which can influence variation in male signaling. This enables biologists and managers to more easily examine how variation in male phenotype influences patterns of female mate choice and male mating success in captive breeding colonies.

Among amphibians globally, CBPs are increasingly being established to assist with the recovery of threatened species (Zippel et al. 2011), though there has been little attempt to use knowledge of mate choice to improve the captive breeding and reintroduction of endangered amphibians (but see Settle et al. 2018; Walls and Gabor 2019). This is surprising because for over 50 years, anuran amphibians have been a model group for the study of intersexual selection, and the mechanisms of female mate choice are extremely well understood (Wells 2007). In anuran amphibians, females evaluate potential mates based on multiple secondary sexual characteristics (Gerhardt and Huber 2002; Wells 2007). In particular, there are four main male traits that appear to provide honest signals of male quality: body size, age, call characteristics, and color (Sheldon et al. 2003; Forsman and Hagman 2006). In various anurans, male body size (which is often correlated with age) can predict male mating success, with evidence that females prefer larger and older males (Rausch et al. 2014; Kelleher et al. 2021a). Empirical evidence suggests that larger males may provide direct benefits in the form of higher fertilization success and/or genetic benefits by supplying offspring with good genes (Berven 1987; Robertson 1990; Rausch et al. 2014). Male age can also influence female mate choice in anurans independent of body size (Felton et al. 2006). Females might prefer older males as they have demonstrated their ability to survive, indicating superior genetic quality (Brooks and Kemp 2001). In most anuran species, males advertise to females by calling, and females select males based on a range of call traits, such as call peak frequency, pulse repetition rate, and duration (Gerhardt and Huber 2002). Calls are costly to produce (they are energetically expensive and increase predation and desiccation risk), so call traits can reflect male genetic quality (Gerhardt and Huber 2002). Indeed, there is evidence that by preferentially mating with males that produce longer or more complex calls, females can produce offspring that perform better during larval or juvenile life stages (Welch et al. 1998). Finally, visual traits (such as color) may also indicate male quality. Females have been shown to discriminate between potential mates based on numerous aspects of male coloration, including brightness (luminance), color saturation (chroma), and hue, and this is not necessarily...
restricted to species that exhibit obvious sexual dichromatism (Gomez et al. 2009; Maan and Cummings 2009; Dreher et al. 2017). Assuming that color pigments are costly to acquire and express, variation in color could also provide females with traits that reliably signal male quality (Gomez et al. 2009). Critically, however, sexual signals are often highly complex (Candolin 2003), and a growing number of studies have demonstrated that females use a combination of signal components (both within and across sensory modalities) to assess male quality (Burke and Murphy 2007; Dreher and Pröhl 2014). Based on these findings, mate choice studies should aim to take an integrated, multivariate approach, which considers the influence of multiple traits simultaneously (as opposed to a traditional single trait approach), and tests for interactive effects between various traits (Dreher and Pröhl 2014).

The southern corroboree frog, *Pseudophryne corroboree*, is a terrestrial breeding anuran and is one of Australia’s most critically endangered vertebrate species. Since the appearance of the deadly pathogen *Batrachochytrium dendrobatidis* (Bd, the amphibian chytrid fungus) in Australia in the 1970s, *P. corroboree* has experienced extreme population declines (Hunter et al. 2010), with less than 50 mature individuals currently remaining in the wild. The species is now the subject of a large-scale, multi-institutional CBP established in 2003 (McFadden et al. 2013). While protocols have been established that allow *P. corroboree* to be successfully bred in captivity, very little is currently known about factors influencing male mating success and patterns of mate choice. Female mate choice is apparent in captive breeding enclosures, where there is a distinct reproductive skew (approximately 30–50% of males receive successful matings) (McFadden et al. 2013). This skew indicates that some males are more attractive to females than others. Additionally, a proportion of females fail to breed annually if they do not encounter a suitable male (McFadden et al. 2013). These observations point towards female mate choice being an important component of the *P. corroboree* mating system. Male sexual coercion or male-male competition is not expected to strongly influence patterns of male mating success in *P. corroboree* because females move between the nests of multiple males before mating, and during these interactions males show no signs of attempting to force copulation (MSM, personal observation). Moreover, there is a distinct absence of male physical aggression, or other traits typically associated with intense intrasexual selection in anurans (e.g., nuptial spines, enlarged forearms, heads, or body size). In a recent field study in northern corroboree frogs (*Pseudophryne pengilleyi*), parentage analysis also provided limited evidence for alternative mating tactics such as satellite or sneaking behavior and the incidence of nest takeover was very low (Kelleher et al. 2021a). While *P. corroboree* males do have a distinct territorial call, these appear to function in establishing spacing between males at the time when choruses are first established, with no evidence that resident males suppress advertisement calling by their neighbors (suggestive of a dear enemy effect). A recent, preliminary investigation into female mate preferences in *P. corroboree* suggested that male dominant call frequency may be important in mate evaluation (Kelleher et al. 2021b); however, females may use a suite of different traits to assess potential mates. Advancing our understanding of factors that influence mate choice in *P. corroboree* may provide conservation managers with knowledge needed to develop behavior-based captive breeding strategies, which may maximize colony reproductive output and population viability.

The aim of the present study was to investigate patterns of male mating success in *P. corroboree* to inform the CBP for this species. Specifically, we investigated whether variation in multiple male phenotypic traits (call characteristics, coloration, age, body size) predicted: (1) male mating success and (2) male fertilization success. These aims were addressed by conducting breeding trials in a captive colony where male nest quality was standardized and females were able to choose between multiple potential mates.

**Methods**

**Study species**

The southern corroboree frog (*P. corroboree*) is a small (23–30 mm snout-vent length) terrestrial anuran from the family Myobatrachidae. The species is characterized by its striking bright yellow and black longitudinal stripes, located on its dorsal surface (Fig. 1). *P. corroboree* is endemic to Kosciuszko National Park in south-eastern Australia and its distribution is restricted to areas above 1300 m in elevation. *P. corroboree* females reach sexual maturity between 3 and 4 years old in captivity, and 4 and 5 years old in the wild, while males reach sexual maturity between 2 and 3 years old in captivity and 3 and 4 years old in the wild (Hunter 2000). *P. corroboree* breeds annually, with the breeding season commencing in mid to late summer (Osborne 1991). Male *P. corroboree* construct small, terrestrial nests (depressions) in sphagnum bogs or wet tussock grasslands, adjacent to ephemeral pools, or watercourses (Osborne 1991). Males call to attract females, with peak calling activity typically occurring from January to February (Pengilley 1971). Females oviposit within the nest chamber and then leave, while males remain with the eggs for several weeks (Osborne, 1991). Female clutch size ranges from 16 to 40 eggs (mean = 26.4) (Osborne 1991). Observations in nature and in captivity indicate that females can be monandrous and lay all their eggs in one nest (Pengilley 1973;
However, there is also evidence that some females may exhibit sequential polyandry and partition their clutch between the nests of multiple males (Pengilley 1973; McFadden et al. 2013). Of note, sequential polyandry is common in other closely related *Pseudophryne* species, such as *Pseudophryne bibronii* (Byrne and Keogh 2009). Fertilized eggs undergo intracapsular development inside the nest chamber, until they enter diapause, where development is suspended at Gosner stage 27 (Osborne 1991). Hatching is triggered by heavy autumn rain, which floods nest sites. Tadpoles are flushed into adjacent ephemeral pools or water bodies, where they continue to develop until they metamorphose in spring.

**Study animals and husbandry**

All frogs used in the present study were initially collected as eggs from extant wild populations between 2003 and 2012. After eggs were collected, they were transported to Taronga Zoo, NSW, Australia, where they were hatched and reared to adulthood in a biosecurity facility. Outside of the breeding season, all frogs were housed in same-sex groups (2–8 frogs) in clear, plastic enclosures (28 cm $L \times 17$ cm $W \times 18$ cm $H$). Each enclosure contained a base layer of fine aquarium gravel, covered by a layer of sphagnum moss (Brunnings, Australia). Frogs were fed *Acheta domestica* crickets (6–10 days old) twice weekly. Crickets were dusted with calcium powder before every feed (calcium with vit. D$_3$, Rep-Cal Research Labs, USA). Enclosures were flushed with reverse-osmosis (R.O.) water twice weekly to prevent the accumulation of detritus and nitrogenous waste. Inside the facility, frogs were exposed to natural ambient light from a nearby window and provided with UV-enriched light supplied by a single Reptisun lamp (36” 10.0 UV-B T8, Reptisun) suspended above the enclosures, approximately 36 cm from the frogs. UV lights were controlled by an external light sensitive switch (HPM, NSW, Australia) which detected changes in outside lighting conditions and simulated the natural local photoperiod. The facility was temperature-controlled, and ambient air temperature inside the facility was cycled annually, to reflect natural seasonal changes, which included a 6-week hibernation period. Temperature throughout the year ranged from 5 °C during the winter hibernation period, to a maximum of 20 °C during the breeding season. During the winter hibernation period, feeding ceased and resumed once the temperature was above 15 °C.

**Captive breeding design**

To determine whether male phenotypic traits predict breeding success in *P. corroboree*, captive breeding trials were conducted in eight glass breeding tanks (tank A, B, C, D, E, F, G, or H; 135 cm $L \times 55$ cm $H$ and 55 cm $W$, Supplementary Fig. 1) that were purposefully built for the captive breeding of *P. corroboree* at Taronga Zoo (McFadden et al. 2013). Each breeding tank was open at the top and contained a base layer of white aquarium gravel and a thick plant layer of live sphagnum moss (Supplementary Fig. 1). In nature, male corroboree frogs form choruses and cluster together in small groups (often around the edge of ephemeral pools or around small pieces of vegetation), so the size of the breeding area and proximity of male nest sites within the breeding tanks reflects natural conditions. Of note, the breeding tanks are suitable for assessing mate choice because *P. corroboree* call and breed deep within the live sphagnum moss, which limits the potential for sound to reverberate. Lighting to the breeding tanks was provided by one UV-enhanced lamp (36” 10.0 UV-B T8, Reptisun) and one plant grow light bulb (F30W Gro-Lux T8, Sylvania), which were both controlled by an external light sensitive switch (HPM, NSW, Australia) which simulated the natural local photoperiod. During captive breeding trials (19 December 2018–15 March 2019), frogs were maintained on a 20 °C/17 °C day/night temperature cycle. The breeding tanks were hydrated by an internal automated misting system. Reverse-osmosis filtered water was delivered once per day for one minute through three spray nozzles situated above each tank (below the tank lid). This watering system ensured that the live sphagnum moss was 100% saturated at all times (confirmed via measurements with a moisture probe meter; MPM-160-B, 12-bit resolution, ICT International Pty Ltd, Armidale, Australia).
This design ensured that the nesting environments were homogeneous across all breeding tanks.

In each captive breeding trial (N = 8, one trial per breeding tank), females were given a choice between multiple males simultaneously for the duration of the breeding season (i.e., live males were present for the entirety of the breeding season). For the purpose of this study, to allow for considerable variation among males (and potential for female discrimination), eight males and eight females were placed in each breeding tank. This resulted in each female being able to choose between a maximum of eight different males in a breeding chorus, which simulated a natural breeding chorus, where males aggregate in small groups and acoustically interact. Males and females grouped together remained in their designated breeding tanks for the duration of the breeding season. While the operational sex ratio is not known for wild *P. corroboree*, observations indicate that males remain in the chorus for an extended period assessing males before mating (D. Hunter, personal communication). Additionally, female arrival to breeding sites in *Pseudophryne* species is controlled by a combination of climatic and social conditions and reasonably synchronized (O’Brien et al. 2021), so there are typically groups of females assessing groups of males at the same time. Moreover, this design follows the current *P. corroboree* captive breeding protocol where groups of multiple females (N = 2–8) and multiple males (N = 2–8) are assigned to breeding tanks based on a maximum avoidance of inbreeding genetic breeding scheme (OEH 2012). For this study instead of sorting individuals based on genetics, males were sorted into breeding tanks based on their pre-breeding body weight (weight as of the 19th of November, range = 1.46–2.65 g, mean = 2.03 g), as male mating and fertilization success is related to body size in various anuran species, and is often linked to male call characteristics (Wells 2007). Males were weighed and then ordered from largest to smallest. The eight largest males were then randomized into each of the eight tanks (tank A, B, C, D, E, F, G, or H). Then, the eight smallest males were randomized into each of the eight tanks based on their pre-breeding body weight (weight as of the 19th of November, range = 1.46–2.65 g, mean = 2.03 g), as male mating and fertilization success is related to body size in various anuran species, and is often linked to male call characteristics (Wells 2007). Males were weighed and then ordered from largest to smallest. The eight largest males were then randomized into each of the eight tanks (tank A, B, C, D, E, F, G, or H). Then, the eight smallest males were randomized into each of the eight tanks. This process continued until all 64 males had been assigned to breeding tanks and ensured that small, medium, and large size males were present in every tank. The mean body size of males did not differ between breeding tanks (ANOVA: \( F_{7,56} = 0.0919, p = 0.9986 \), Supplementary Fig. 2), nor did variance in body weight (Levene’s test: \( F_{7,56} = 0.1749, p = 0.9894 \), Supplementary Fig. 2). Visibly gravid females were also assigned randomly to tanks via pre-breeding body weight (weight as of the 19th of November 2018; range = 2.74–4.44 g, mean = 3.37 g). The mean body size of females did not differ between breeding tanks (ANOVA: \( F_{7,56} = 0.1106, p = 0.9974 \), Supplementary Fig. 3) nor did variance (Levene’s test: \( F_{7,56} = 0.0594, p = 0.9997 \), Supplementary Fig. 3). At the time of the study, males ranged in age from 6 to 15 years old post metamorphosis, and females ranged in age from 6 to 14 years old post metamorphosis (known from husbandry records based on when eggs were collected from the wild).

**Data collection**

Captive breeding trials commenced at the beginning of the breeding season, 19 December 2018, when males started to advertise to females by calling. Males (N = 64) were removed from same-sex housing and transferred into their designated breeding tanks (tanks A–H). Males were placed into the breeding tanks 6 weeks earlier than females. This was done to allow males to establish individual nest sites (as they would do in the wild) within the tanks before introducing females and is consistent with the current *P. corroboree* captive breeding protocol (McFadden et al. 2013). Immediately prior to being placed into the tank, each male was weighed using a portable balance, and snout-vent length (SVL) was measured with calipers. Next, each male was digitally photographed to enable quantification of their yellow dorsal coloration at a later date (see the “Male color analysis” section). Frogs were photographed inside a custom-built, portable photography light arena. The arena consisted of two opaque cylindrical containers stacked on top of each other, with integrated white LED strip lights attached around the inside walls of the arena. This prevented shadowing and created a standardized lighting environment for all photos. Photographs were taken with a Canon EOS 7D camera with a standard lens (Canon EFS 18–55 mm lens) that was positioned on top of the arena, with the camera lens placed through a viewing hole at the top of the arena. The distance between the camera lens and the frog was 28 cm. Photographs were taken in raw format with the following settings: ISO = 400, f = 11, shutter speed = 1/200. All color photographs included an X-rite ColorChecker Passport (X-rite, USA), which consists of 24 colored and grayscale squares against which the colors in each photo are standardized.

On the 4th of January 2019, approximately 2 weeks after males started actively calling, occupied male nest sites were identified and flagged on the surface of the moss (male nests are below the surface of the moss) using a small, white plastic disc marked with an individual nest ID number. Nest sites were flagged again on the 14th, 17th, and 23rd of January 2019, and call recordings started on the 29th of January 2019, once the majority of nests had been identified and peak calling activity was observed. Call recordings were taken three times a week during periods of high calling activity (07:00 h and 14:00 h) for the duration of the study. All males that were consistently calling during the recording periods were recorded using a Marantz Professional handheld solid-state recorder (Model No. PMD661MKII, D&M Holdings Inc. Tokyo, Japan) connected to a directional microphone.
(NTG2, Rode Microphones, Sydney, Australia) which was held approximately 30 cm above the focal male. Calls were recorded in .wav format at a sampling rate of 44.1 kHz with a 16-bit resolution.

Females (N = 64) were placed into their designated breeding tanks on January 31, 2019. To determine male mating success, all male nest sites were checked once a week on a Friday for six consecutive weeks (the entire breeding season). The first nest checks occurred 1 week after females first entered the breeding tanks (8 February 2019), with subsequent checks made on the following dates: February 15, February 22, March 1, March 8, and March 15, 2019. During each nest check, we identified the resident male (i.e., the male calling from the flagged nest), using their unique dorsal and ventral patterns, and recorded whether eggs were present or absent, and counted total egg number. When eggs were present, they were removed from the male’s nest and transferred into plastic containers on sphagnum moss. Male mating success was recorded as mated (yes/no) and the total number of eggs received over the entire breeding season (6-week period). We used the total number of eggs a male received as a measure of mating success because it was not possible to ascertain the total number of matings (females may partition their clutch among multiple nests). However, a higher egg number can be assumed to indicate a greater number of matings. This approach follows that of a recent field study in P. corroboree sister species, the northern corroboree frog, Pseudophryne pengilleyi (Kelleher et al. 2021a). After the breeding season was complete, on the 20th of March 2019, all eggs were assessed to determine whether they were fertile, based on whether there was evidence of embryo development. We then calculated the proportion of fertilized eggs for each male by dividing the number of fertilized eggs by the total number of eggs received. In regard to parentage, a recent field study in P. corroboree sister species, P. pengilleyi, found that a resident male is typically the genetic sire of all the eggs within their nest (Kelleher et al. 2021a), so we have assumed that is the case in this study. Of note, it was not possible to record data blind because this study required knowledge of individual male identity.

**Male call analysis**

While advertising to females, P. corroboree males produce two main calls: a two-part call and a one-part call (Pengilley 1971; Osborne 1991). Oscillograms and spectrograms of both call types are presented in Fig. 2. The two-part call is comprised of two distinct components: a longer initial component, typically followed by a clear interphase, and then a shorter, pulsatile second component (Fig. 2; Pengilley 1971). Call recordings were analyzed using the sound analysis software Raven Pro v 1.5 (Cornell Lab of Ornithology, Bioacoustics Research Program, 2014). Temporal call traits (call duration, pulse repetition rate, and call rate) were obtained from oscillograms. Frequency information was obtained using Fast Fourier transformation, with a Blackman window function (window size = 1024 samples, 3-db bandwidth = 70.7 Hz, DFT = 2048 samples, grid spacing = 21.5 Hz, with 50% overlap) (following Kelleher et al. 2021a). For each male, we measured call traits (duration, frequency, and pulse repetition rate) for each component of the two-part call (1st and 2nd component, Fig. 2) and the one-part call (Fig. 2). For both the two-part and one-part call, multiple individual calls were analyzed (85% of males had between 5 and 12 calls analyzed for each call type), over 1 and 3 call recordings taken throughout the breeding season. This level of sampling is comparable to other studies investigating the effect of male call traits on mating success in frogs (e.g., Pröhl 2003; Smith and Roberts 2003b; Felton et al. 2006; Kelleher et al. 2021a). Average call traits for each call component were calculated as follows: average call duration (length from the beginning of the first pulse to the end of the last pulse of the call, seconds), average pulse repetition rate (number of pulses in the call divided by the call duration, pulses/sec), and average call peak frequency (frequency occurring at the highest amplitude, Hz). Call rate was calculated as the total number of calls (combined total number of two-part and one-part calls, ranging from 5 to 32 calls) given per minute from a male’s longest, continuous call recording (ranging from 30 s to 4 min). This resulted in ten call variables measured for each male: average duration of the 1st and 2nd components of the two-part call, and the one-part call, average frequency of the 1st and 2nd components of the two-part call, and one-part call, average pulse rate of the 1st and 2nd components of the two-part call, and one-part call and call rate. Of note, a small number of males (N = 7) had their average call duration, frequency, and pulse rate calculated from less than 5 calls for one call type (i.e., for either the two-part or one-part call). Although this level of sampling is comparable to other studies (Smith and Roberts 2003b; Kelleher et al. 2021a), the statistical analysis (outlined below in the “Statistical analysis” section) was run with these males excluded to ensure they were not driving patterns observed. As this was not the case, these males were retained to increase power of the analysis.

**Male color analysis**

Male P. corroboree color analysis followed methods described in Kelleher et al. (2021a Appendix). In brief, we obtained average red (R), blue (B), and green (G) pixel values from the yellow stripes on the entire dorsal surface (head, body, and legs) of each individual male using a custom Matlab script (Mathworks Inc., Natick, MA, USA) written by JAE. After standardized RGB values were calculated from the color standards in each digital photo, a second
custom Matlab script written by JAE was used to determine a color space based upon calculations of hue, chroma, and luminance (detailed descriptions of hue, chroma, and luminance calculations are described in Cadena et al. 2017, 2018). Hue represents the color, measured as the angle relative to the axis (Endler 1990; Cadena et al. 2018), chroma is the “saturation” of the coloration and is measured as the distance from the origin (Endler 1990; Cadena et al. 2018), and luminance represents the brightness of the coloration and was calculated as the sum of the standardized RGB values (Endler 1990; Cadena et al. 2018). We did not use an eye model, because data on cone absorption spectra are not available for this species and family. In any case, our question here is simply whether these crude measures of color predict mate choice.

**Statistical analysis**

To investigate the influence of male phenotypic traits on male mating and fertilization success in *P. corroboree*, we measured 16 explanatory variables; (1) body weight, (2) age, (3) snout-vent length (SVL), (4) luminance, (5) hue, (6) chroma, (7) call rate, (8) average duration of two-part call, 1st component, (9) average duration of two-part call, 2nd component, (10) average duration of one-part call, (11) average frequency of two-part call, 1st component, (12) average frequency of two-part call, 2nd component, (13) average frequency of one-part call, (14) average pulse rate of two-part call 1st component, (15) average pulse rate of two-part call, 2nd component, and (16) average pulse rate of one-part call. Of the 64 males included in this study, 47 males were included in the statistical analysis. Twelve males were excluded as they failed to advertise to females, and an additional five males (four from tank A and one from tank B) were excluded during call analysis due to overlap on call recordings with neighboring males (we were unable to determine which call belonged to a focal male). Of note, four females died of unknown causes before laying eggs (two from tank D and two from tank H).

Prior to analysis, we checked for collinearity among all explanatory variables using a Pearson’s correlations matrix (see Supplementary Fig. 4). Variables with a significant correlation coefficient $\geq 0.5$ were considered to be highly correlated (Zuur et al. 2009) so one of the two affected

![Fig. 2 a Oscillogram and b spectrogram of a *P. corroboree* male advertisement call, consisting of two call types—a two-part call (with a 1st and 2nd component) and one-part call](image-url)
variables was removed from further analysis. Male age and hue were negatively correlated ($r = -0.50$, $p = 0.0004$) and chroma and luminance were positively correlated ($r = 0.79$, $p = < 0.0001$), so hue and chroma were removed from further analysis. The average frequency of all calls were highly correlated (average frequency 1st component–2nd component: $r = 0.85$, $p = < 0.001$; average frequency 1st component–one-part call: $r = 0.87$, $p = < 0.001$; and average frequency 2nd component–one-part call: $r = 0.87$, $p = < 0.001$), so one measure of frequency (average frequency one-part call) was taken forward in the analysis. Finally, the average pulse rate of the 1st component and average pulse rate of the one-part call were positively correlated ($r = 0.77$, $p = < 0.001$) so the average pulse rate of the 1st component was removed from the analysis. Independent variables taken forward in the analysis were selected based on important predictor variables in a recent study investigating predictors of male mating success in *P. corroboree* sister species, *P. pengilleyi* (Kelleher et al. 2021a).

We used linear models to examine the potential effect of male body weight, SVL, and age on male call characteristics: call rate, call duration (all call components), call frequency (one-part call), and pulse rate of the 2nd component and of the one-part call. Before fitting models, all explanatory variables were first standardized to have a mean of zero and a standard deviation of 1 as variables were measured on vastly different scales (Zuur et al. 2009). This improves model convergence and calculates coefficient estimates on the same scale (Zuur et al. 2009). Linear models were fit for each call variable separately, with standardized body weight, SVL, and age entered as continuous explanatory variables. Call rate, average pulse rate, and average frequency of the one-part call were all log-transformed to improve normality.

To investigate predictors of male mating success we used generalized linear mixed-effects models (GLMM) with a zero altered negative binomial (ZANB) distribution and log link function. The ZANB GLMM models the data in two parts: (1) a binomial logistic regression with male mating success (yes or no) as the response variable and (2) a negative binomial GLMM of the count data, with the total number of eggs as the response variable (excluding zeros) (Zuur et al. 2009). As the data were over dispersed and zero-inflated, a ZANB distribution was used in place of a Poisson or standard negative binomial distribution. Our sample size ($N = 47$) limited the number of explanatory variables that could be fitted in the same model, so we used a two-step model selection process (following Kelleher et al. 2021a). First, we constructed 11 separate ZANB models each with the number of eggs as the response variable and one single explanatory variable as a predictor (see Supplementary Table 3). All standardized explanatory variables were entered into models as continuous predictors, and breeding tanks (A–H) were included in every model (in both the count and binomial part of the model) as a random effect. We then used an information-theoretic model selection process to rank the 11 single variable ZANB models based on their AIC$_c$ value with a correction for a small sample size (AIC$_{c}$). We compared these 11 separate models based on their AIC$_c$ value and examined the model coefficients to determine whether the variable in question was important in the binomial or count part (or both) of the ZANB model. Variables that were within delta AIC$_c$ < 2 of the top-ranked single variable model were retained for further analysis, these were as follows: age, call rate, luminance, all call durations, and average call frequency (see Supplementary Table 3). Next, we constructed 41 candidate ZANB models (including an intercept-only model) using these variables and ranked them based on their AIC$_c$ value (see Supplementary Table 4 for a full list of candidate models). Due to our sample size, only interactions with call rate were considered, as call rate is often an important predictor of male mating success in anurans (Pröhl 2003; Wells 2007) and no model was constructed with complexity exceeding the limits of the sample size. The model with the lowest AIC$_c$ value was considered to provide the best fit but models within delta AIC$_c$ < 2 were considered to be plausible alternatives (Zuur et al. 2009; Symonds and Moussalli 2011). When multiple candidate models were within delta AIC$_c$ < 2 of the top-ranked model, estimated coefficients and $p$-values were derived from model averaging (Zuur et al. 2009).

To determine predictors of male fertilization success, we used GLMMs with a binomial distribution and logit link function (following methods outlined in Zuur et al. 2009). For males that achieved mating success ($N = 31$), the proportion of fertilized eggs was entered as a response variable, with the total number of eggs overall entered as a model weight (Zuur et al. 2009) and breeding tanks (A–H) entered as a random effect. The model selection followed the same process described above. After modeling each explanatory variable separately, we retained the top six variables (based on their AIC$_c$ value), plus one measure of body size (SVL) for further investigation (see Supplementary Table 5). These variables were as follows: age, call rate, average pulse rate (2nd component), average duration (1st and 2nd component), SVL, and average frequency. As for mating success, we then constructed 41 candidate models (including an intercept-only model, see Supplementary Table 6) and ranked them according to their AIC$_c$ value. The model with the lowest AIC$_c$ value was considered to provide the best fit, but models within delta AIC$_c$ < 2 were also considered as alternative models (Zuur et al. 2009). Predictions were calculated for statistically significant predictors from the averaged model. We investigated model fit by visually examining diagnostic plots of residual versus fitted values for each predictor variable in the top-ranked model(s). As a supplementary analysis, we also investigated predictors of the total number of fertilized eggs.

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of fertilized eggs using a GLMM with a negative binomial distribution and a log link function. We followed the same model selection process as described above and the analysis showed similar results to the proportion analysis, both for the top-ranked predictor variables and candidate models (see Supplementary Tables 7 and 8). All statistical analyses were completed in R version 3.5.1 (R Development Core Team 2018). GLMM analyses were conducted using the glmmTMB package (Brooks et al. 2017), while AICc and model averaging were conducted using the MuMIn package (Bartoń 2019).

Results

Variation in aspects of male phenotype

Of the 64 males included in the captive breeding trials, 81% (52/64) established individual nest sites and actively advertised to females by calling. The remaining 19% of males (12/64) failed to establish nest sites and were excluded from further analysis. Five advertising males were also excluded from further analysis due to call overlap with neighboring males (see Male call analysis and Statistical analysis sections). Thus, a total of 47 males were included in the statistical analysis.

Of the males included in the analysis (N = 47), body weight ranged from 1.61 to 2.80 g, with a mean (± SEM) of 2.23 g (± 0.04). Male SVL ranged from 23.5 to 27.8 mm, with a mean (± SEM) of 26.34 mm (± 0.14). There was a significant, positive correlation between male body weight and SVL (r = 0.38, p = 0.008, Supplementary Fig. 4). Male age ranged from 6 to 14 years, with a mean (± SEM) of 9.70 years (± 0.38). There was no significant correlation between male body weight or SVL and male age (Supplementary Fig. 4). Variation in male call and color traits are summarized in Supplementary Table 1. There was a significant positive relationship between SVL and male call rate; larger males had higher call rates (Supplementary Table 2). Additionally, there was a significant positive relationship between male body weight and average pulse rate of the one-part call; larger males had higher pulse repetition rates (Supplementary Table 2). Body weight, age, and SVL did not influence any of the other male call characteristics (see Supplementary Table 2).

Variation in male mating and fertilization success

Of the 64 males included in the captive breeding trials, 53% (34/64) of males gained matings (number of eggs > 0). Of the males included in the analyses (N = 47), 65% (31/47) of males received matings (number of eggs > 0). The total number of eggs in a male’s nest over the entire breeding season ranged from 3 to 95 eggs (Fig. 3), with an average of 28.6 eggs per nest. Of the males that achieved mating success (N = 31), fertilization rates ranged from 0 to 91.3% with an average fertilization rate of 48.6%.

Predictors of male mating success

Mating probability

There were 12 competing models (AICc < 2) to explain the probability of a male mating (Supplementary Table 4). These models contained different additive combinations of male age, luminance, and average call duration (of the 1st and 2nd component) as predictor variables (Supplementary Table 4). Male age, luminance, and average duration (1st component) appeared to have a positive effect on mating probability (Table 1) and average duration (2nd component) appeared to have a negative effect on mating probability (Table 1). However, there was no significant effect of any of these predictors on mating probability (Table 1).

Mating success (total number of eggs)

The best model to explain variation in the total number of eggs in a male’s nest contained the interaction between call rate and average call frequency of the one-part call (Supplementary Table 4). There was one competing model within 2 AICc points of this model, which contained the interaction between call rate and average call duration (Supplementary Table 4). There was a significant effect of both interactions on the number of eggs in a male’s nest (Table 1).

The effect of call duration (one-part call) on male mating success depended on call rate. Males with short calls and a low call rate had the highest numbers of eggs (Fig. 4a). When call rate was high, males with longer calls had higher egg numbers than males with shorter calls (Fig. 4a).
The effect of call frequency on male mating success also depended on call rate. Overall, males with lower call frequencies had higher numbers of eggs than males with higher call frequencies (Fig. 4b). When call frequency was low, call rate did not significantly influence mating success (Fig. 4b). But, when call frequency was high, males with lower call rates had higher mating success than males with higher call rates.

### Predictors of fertilization success

The best model to explain variation in male fertilization success (proportion of fertilized eggs) contained male age and the interaction between call rate and average call frequency as predictor variables (Supplementary Table 6). There was one competing model within 2 AICc points of this model, which contained average call frequency and the interaction between call rate and SVL as predictor variables (Supplementary Table 6).

There was a significant effect of male age on male fertilization success; older males had a higher proportion of fertilized eggs (Table 2 and Fig. 5). There was a significant decrease in the proportion of fertilized eggs with increasing call frequency, but the strength of this relationship depended on a male’s call rate (Fig. 6a). For males with lower call frequencies, fertilization success did not vary with call rate (Fig. 6a). But, for males with higher call frequencies, fertilization success varied with call rate; males with lower call rates had higher fertilization success than males with higher call rates.

The effect of body size (SVL) on fertilization success also depended on a male’s call rate. Small males had a high proportion of fertilized eggs irrespective of call rate (Fig. 6b). Large males with low call rates had higher fertilization success than large males with higher call rates (Fig. 6b).

### Discussion

Understanding factors that influence female mate choice can potentially aid in the captive breeding of threatened species. Here, we investigated whether variation in male body size, age, calling behavior, and coloration predicted male reproductive outcomes in critically endangered *P. corroboree*. Our results revealed that both male mating and fertilization success were predicted by a complex interplay between multiple phenotypic traits. In particular, variation in male mating success (total egg number received) was strongly explained by a combination of interactions between male call rate, call duration, and call frequency. Fertilization success was also associated with call rate and call frequency, in addition to male body size and age. These findings suggest that females are using multiple traits (call traits and physical traits) to assess male quality and fertility and using them in different combinations. More broadly, these results add to a large body of empirical evidence that female mate choice decisions are based on information gained from multiple male traits (Candolin 2003; Starnberger et al. 2014; Mitoyen et al. 2019). Moreover, these results highlight that studies that only consider the influence of a single trait may fail to capture to complexity of mate choice (Taylor et al. 2007; Gould and Augustine 2020).

### Effect of calling behavior

Males with low-frequency calls had higher mating and fertilization success than males with high-frequency calls (irrespective of call rate), a pattern that is relatively common in anurans (Wells 2007; Lesbarrères et al. 2008). This outcome suggests that *P. corroboree* males with low-frequency calls are highly attractive and is in line with the results of our recent manipulative phonotaxis study, which found that *P. corroboree* females significantly preferred low-frequency

| Table 1 | Model-averaged coefficient estimates of variables from the best fitting ZANB generalized linear mixed models of male mating success in *P. corroboree* (N=47). Significant effects are indicated in bold |
|---|---|
| **Binomial logistic probability model** | **β** | **SE** | **z** | **p** |
| Intercept | −0.764 | 0.341 | 2.165 | 0.030 |
| Age | 0.599 | 0.363 | 1.600 | 0.110 |
| Luminance | 0.560 | 0.359 | 1.510 | 0.131 |
| Average duration (1st component) | 0.629 | 0.349 | 1.743 | 0.081 |
| Average duration (2nd component) | −0.653 | 0.376 | 1.683 | 0.092 |
| **Negative binomial count model** | **β** | **SE** | **z** | **p** |
| Intercept | 3.685 | 0.188 | 18.865 | <0.001 |
| Call rate | −0.324 | 0.169 | 1.860 | 0.063 |
| Average frequency | −0.243 | 0.109 | 2.168 | 0.030 |
| Average duration (one-part call) | −0.079 | 0.110 | 0.699 | 0.485 |
| Call rate * average duration (one-part call) | 0.416 | 0.149 | 2.695 | 0.007 |
| Call rate * average frequency | −0.325 | 0.111 | 2.841 | 0.005 |
male advertisement calls (Kelleher et al. 2021b). Our findings strongly suggest that call frequency is an honest signal of male fertility and fertilization potential, as males with low call frequencies also had high fertilization success. This finding implies that female *P. corroboree* may preferentially select males with low call frequencies to receive direct fertility benefits. This finding is particularly noteworthy, as a direct relationship between male call frequency and fertilization success has rarely been reported in anurans. One example comes from a study on a wild population of the smooth toadlet (*Uperoleia laevigata*), where female mate choice based on male call frequency (which is related to male body size) also resulted in enhanced fertilization success (Robertson 1990). Interestingly, very few studies have investigated relationships between anuran male call traits and fertility, but there is evidence for such associations in birds and insects (Wagner and Harper 2003; Simmons et al. 2010). Therefore, the potential for females to assess male fertility based on acoustic signals may be more widespread than currently appreciated and requires further examination. Evidence to date has illustrated that fertility is generally related to visual signals (e.g., Lifjeld et al. 2011), but it is possible that acoustic signals may also provide strong and reliable indicators of male fertilization potential.

Calling behavior was also related to male mating and fertilization success in three other key and interactive ways: (1) males with less attractive, higher frequency calls had higher breeding success if they called at a lower rate, (2) males with short calls had higher mating success than males with longer calls, and males with short calls had the highest mating success if they called at a lower rate, and (3) call rate had little effect on mating success if calls were longer. These relationships were surprising as they contrast with past studies in anurans, which have reported that (1) females typically prefer males that invest more in calling (e.g., have longer calls or call at a higher rate) (Gerhardt and Huber 2002; Wells 2007) and (2) males with unattractive calls (e.g., high call frequency or shorter calls) can increase their relative attractiveness by increasing their calling effort (e.g., by calling at a higher rate or over more nights) (Morris and Yoon 1989; Smith and Roberts 2003b). In regard to call rate, we found that there was a positive association between call rate and body size (larger males had higher call rates), and large males with high call rates had very low fertilization success (see the “Effect of age and body size” section). This may explain why males with high call rates generally had lower success (and a high call rate did not improve the attractiveness of high frequency or short calls), as females may have been avoiding larger males due to the costs associated with low fertility (discussed below). Interestingly, however, call rate had little effect on mating success when call duration increased. There is no obvious explanation for this association. However, evidence from studies in other anurans, as well as birds and insects, suggest that females are often particularly sensitive to calls that contain a greater amount of energy (i.e., longer, faster rate, louder), possibly due to a pre-existing bias in the sensory system (Ryan and Keddy-Hector 1992). For example, longer calls have been shown to elicit a heightened stimulatory response from females (Ryan and Keddy-Hector 1992), and this stimulation can supersede the effect of other call traits (such as call rate). Moreover, there is also evidence that this increased stimulation can modify or reverse preferences for other traits (Klump and Gerhardt 1987; Ryan and Keddy-Hector 1992). Additionally, these relationships might exhibit a high level of complexity because anuran mating behavior is often plastic. For instance, there is often considerable among- and within-individual variation in male calling behavior, and males can use different calling strategies depending on social context (Byrne 2008) such as the strategies being used by competitors (Lesbarrères and Lodé 2002). Additionally, males can also adopt different calling strategies depending on body size or age (Smith and Roberts 2003b). Female anurans can also exhibit reversible phenotypic plasticity in mate preferences, and mate choice can be influenced by a multitude of factors such as female age, reproductive state, prior social experience, and social interactions between chorusing males (such as decoy effects) (Jennions and Petrie 1997; Ryan et al. 2019).

Collectively, our findings suggest that female *P. corroboree* combine information from multiple acoustic traits non-additively and that the strength of preference for one trait has the potential to be altered by the strength of other traits (Candolin 2003; Smith and Roberts 2003a). Similar results have been reported for other anurans (Smith and Roberts 2003a; Burke and Murphy 2007; Castellano and Rosso 2007). For example, phonotaxis trials in gray treefrogs (*Hyla versicolor*) have shown that females appear to weigh call duration more heavily than call rate and that they discriminate between males in favor of long calls at a slow rate over short calls at a higher rate (Gerhardt 2001). In the barking tree frog (*Hyla gratiosa*), females also combine signal components non-additively, and preferences for call duration and call rate vary depending on the percentage difference between the two stimuli (Burke and Murphy 2007). It is possible that female *P. corroboree* may weigh call rate more heavily when assessing males with higher call frequency or shorter calls, as opposed to when call frequency is low (which is highly attractive and an indicator of male quality) or when call duration is long (as this may act as a hyper stimulus). It is important to consider that patterns of female mate choice observed here could also reflect among-individual variation in female preferences for particular traits (e.g., some females prefer high call frequency while others prefer low frequency) (Jennions et al. 1995); however, our recent manipulative phonotaxis study in *P. corroboree* did
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**Effect of age and body size**

We found no relationship between male body size and age in our study population, yet our results revealed that both these traits independently predicted male fertilization success. Older *P. corroboree* males had significantly higher fertilization success than younger males. There are several possible explanations for this finding. One possibility is that older males have greater fertilization potential (functional fertility) because they have higher sperm production or superior quality sperm (Sheldon 1994; Brooks and Kemp 2001). Although the relationship between functional fertility and age is yet to be explicitly examined in anurans, studies in birds, fish, and invertebrates have shown that older males often possess a greater fertilization capacity, and that by selecting older males as mates females can insure against functional infertility (Schmoll et al. 2007). An alternative explanation is that older males have higher fertilization success because they are of higher genetic quality (Brooks and Kemp 2001). Viability indicator models of sexual selection predict that male age may reliably indicate intrinsic genetic quality and that older males signal their viability via their proven ability to survive (Brooks and Kemp 2001). Critically, intrinsic genetic quality may also be associated with superior sperm performance (Locatello et al. 2006), with evidence in various taxa that males of higher genetic quality have increased sperm viability, motility, and longevity (Locatello et al. 2006; Evans et al. 2007); these are sperm traits known to influence fertilization success. Obtaining direct fertility benefits from mate choice is likely to be particularly important to females in polygynous mating systems, where males mate consecutively with multiple females, and highly successful males risk becoming sperm depleted (Scarponi and Godin 2018). In the *P. corroboree* mating system males are polygynous (Kelleher et al. 2021a), and in the present study the most successful males would have mated with at least three females (estimated based on female average clutch size and the total number of eggs they received throughout the entire season). Additionally, *P. corroboree* males have a small relative testes size compared with other myobatrachid frogs (Byrne et al. 2002), so highly successful males may be at a particularly high risk of becoming sperm depleted. Accordingly, there may be strong selection on *P.
Corroboree females to assess and choose males based on fertility (Scarponi and Godin 2018).

The relationship between body size and fertilization success was more complex. We found that the effect of body size on fertilization success depended on a male’s call rate, and, additionally, call rate was positively associated with male body size (larger males called at higher rates). Smaller males generally had higher fertilization success than larger males, regardless of call rate. Large males with high call rates had very low fertilization success. This finding was unexpected because larger males (that often invest more in sexual displays as they have greater energy reserves) are often reported to have high fertilization success, and females typically prefer large males as they provide direct fertility benefits (Pujolar et al. 2012). Conversely, our findings are in line with numerous studies demonstrating that smaller males may also have enhanced fertilization success (Klaus et al. 2011; Young et al. 2013). Smaller males may have higher fertilization success because they preferentially invest in sperm production, at the expense of investing in growth or elaborate sexual signals (Stearns 1989; Simmons et al. 2017; Lüpold et al. 2019). Such life-history tradeoffs have been reported in various taxa, with strong negative relationships reported between male body size, sexual signaling intensity, and fertility (Evans 2010; Sherman et al. 2010). For instance, in the quacking frog (Crinia georgiana), there is a negative relationship between male body size and sperm viability (proportion of live sperm), suggesting that smaller males allocate more resources to sperm production (Dziminski et al. 2010). Similarly, in Chinook salmon (Oncorhynchus tshawytscha), smaller subordinate males have higher sperm velocity and higher fertilization success than larger, dominant, ornamented males (Flannery et al. 2013; Young et al. 2013). It is possible that P. corroboree males also preferentially and strategically invest in certain fitness-determining traits over their lifetime. Corroboree frogs are long-lived; individuals have been reported to live for up to 9 years post metamorphosis in the wild (Hunter 2000), and 20 years post metamorphosis in captivity (MSM personal observation). P. corroboree are also characterized by indeterminate growth (as are most reptiles and amphibians) and thus likely have ample time and opportunity to strategically allocate resources to sperm production at the expense of growth and/or other sexually selected traits. Such strategic allocation could account for the intriguing lack of correlation between male age and body size in this study population (older males were not necessarily larger). An alternative explanation is that larger males may be too heavy to support during amplexus, making it difficult for females to oviposit optimally, resulting in lower fertilization success. This has been reported in the smooth toadlet (U. laevigata) where heavy males (>70% of female body weight) impede...

### Table 2
Model-averaged coefficient estimates of variables from the best fitting generalized linear mixed models of male fertilization success (proportion of fertilized eggs) in *P. corroboree* (*N* = 31). Significant effects are indicated in bold.

| Model terms                | β     | SE    | z     | p     |
|----------------------------|-------|-------|-------|-------|
| Intercept                  | −0.055| 0.482 | 0.109 | 0.913 |
| Call rate                  | −1.262| 0.130 | 9.296 | <0.001|
| SVL                        | −0.513| 0.134 | 3.658 | <0.001|
| Average frequency          | −0.537| 0.190 | 2.767 | 0.006 |
| Age                        | 0.312 | 0.078 | 3.825 | <0.001|
| Call rate * average frequency | −0.441| 0.109 | 3.829 | <0.001|
| Call rate * SVL            | −0.802| 0.146 | 5.228 | <0.001|

### Fig. 5
The effect of male age (years) on male *P. corroboree* fertilization success (proportion of fertilized eggs). Line is model prediction with 95% confidence intervals.
successful fertilization, due to the inability of females to carry them during prolonged and arduous oviposition (Robertson 1990). In *P. corroboree*, oviposition is presumed to occur over a protracted period of at least several hours (possibly longer) and may even include a period of preovipositional clasping (which may also last several hours), as reported in three other closely related, terrestrial breeding species, *Pseudophryne bibronii, Pseudophryne dendyi,* and *Pseudophryne semimarmorata* (Woodruff 1976). In this study, the average *P. corroboree* male was approximately 66% of the average female’s body weight. Thus, it seems plausible that *P. corroboree* males above a certain size threshold may be too large for the majority of females to support during this period of lengthy amplexus, reducing fertilization success. Overall, regardless of the mechanism, our results show that multiple traits may reflect a male’s fertilization capacity and suggest that female mate choice decisions may be based on male fertility.

While our findings shed light on patterns of mate choice in *P. corroboree*, there are some limitations that should be mentioned. First, it is possible that routine checks of male nests may have resulted in an increase in male movement and that some level of nest swapping may have occurred in between checks, adding noise to the data. Second, it should be recognized that we did not examine maternal effects. Although our design minimized the potential for such effects by randomizing the allocation of females to breeding tanks (and ensured that each tank received females across a range of body sizes), we were unable to explicitly test whether different female phenotypes had an influence on fertilization success or embryo development. There is also the potential for fertilization success to have been influenced by male–female interactive effects, with certain phenotypic combinations having more pronounced impacts on reproductive outcomes (Byrne et al. 2021). Additionally, we were unable to assess whether there were any male-male effects or female-female effects on mating success. In regard to male-male effects, intra-sexual competition could impact female access to preferred males and the allocation of eggs to certain males, but not others. Male *P. corroboree* rarely engage in the types of physically aggressive behavior frequently observed in other anurans that are subject to intense male-male competition (e.g., wrestling, grappling, ramming) (Byrne and Roberts 2004; Wells 2007; de Sa et al. 2020). Although it is possible that male *P. corroboree* may interfere with each other’s calling behavior and courtship by altering their call type (e.g., by producing more territorial calls), or by adjusting specific call characteristics (e.g., by strategically elevating their call rate in the presence of females) (Schwartz et al. 2001; Byrne 2008). Such dynamic vocal interactions between neighboring males could potentially influence the relative attractiveness of particular males within tanks and modify female mate choices (Schwartz et al. 2001). In regard to female-female interactions, aggression has never been observed in female *P. corroboree*, though interference could take more subtle forms, such as females monopolizing a male by remaining in amplexus for extended periods. However, the fact that there is a mating bias towards a subset of males indicates that there is unlikely to be any major constraint on females mating with preferred males. This bias, however, does raise the possibility that females copy mate choices of other females, as reported for various taxa (reviewed in Witte et al. 2015). Moving forward, controlling for maternal effects, and potential effects of courtship interference, should be factored into future studies investigating mate choice in *P. corroboree*.

**Broad implications**

Taken together, our results reveal that under captive conditions, female *P. corroboree* use multiple male traits within a sensory modality (multicomponent signal) to select mates and that fertilization success is predicted by multiple traits (both acoustic and physical traits). We expect that this is also likely to be the case in nature, as our recent field study in the closely related *P. pengilleyi* revealed that male mating success was predicted by multiple male traits (pulse repetition rate and age) (Kelleher et al. 2021a). Moreover, our results show that multiple male traits predict fertilization success, suggesting that *P. corroboree* females may gain direct fertility benefits from their mate choice decisions. This knowledge significantly advances our understanding of sexual selection processes in *P. corroboree*. Our findings also advance our understanding of the direct benefits gained from mate choice in anurans, as it is one of the first studies to demonstrate a link between male call traits (such as frequency and call rate) and fertilization success in a frog. More broadly, our findings contribute to a rapidly growing body of evidence in mammals, birds, anurans, fish, and invertebrates, showing that female mate choice is often based on suites of traits (Candolin 2003; Stange et al. 2017; Berson and Simmons 2018; Gould and Augustine 2020). Our study also draws attention to the complexity of female mate choice in highly dynamic acoustic environments (such as anuran breeding choruses) (Richardson and Lengagne 2010), highlighting that a single trait approach to understanding mate choice is unlikely to accurately reflect sexual selection processes (Starnberger et al. 2014).

**Conservation implications**

Our findings show that multiple male phenotypic traits predict successful breeding in a captive environment. We propose that this information may be used to develop behavior-based captive breeding protocols in *P. corroboree*. Currently, the *P. corroboree* captive breeding protocol involves
Fig. 6 The effect of the interaction between a male call rate and average call frequency (Hz) and b male call rate and SVL (mm) on male *P. corroboree* fertilization success (proportion of fertilized eggs). Lines are model predictions with 95% confidence intervals. In line with Fig. 4, call rate has been categorized (calculated using the 25th, 50th, and 75th percentiles) in order to visualize the interaction with frequency and SVL.

offering females a choice between 2 and 8 males based on a maximum avoidance of inbreeding (MAI) genetic breeding scheme (OEH 2012). Based on our results, we suggest that conservation managers consider manipulating the composition of males available to females in order to bolster breeding success. For instance, it may be beneficial to increase the relative number of older males, or smaller males presented to females, to increase the fertilization success of clutches. Another possibility is that conservation managers may be able to manipulate male attractiveness by using knowledge of acoustic signals. By understanding which call traits are attractive, conservation managers may be able to predict which individual males are likely to be preferred by females. Males could then be strategically arranged in breeding choruses to either increase or decrease the relative attractiveness of individual males, and subsequently their probability of obtaining mating success. It may also be possible to construct “artificial” choruses using playback speakers where call traits of “neighboring” males are manipulated to increase the relative attractiveness of unmated males (e.g., using an approach similar to Lea and Ryan 2015). It is important to note that although *P. corroboree* can be bred successfully in captivity, integrating knowledge of mate choice has the potential to enhance total reproductive output by encouraging a greater proportion of females to mate, partitioning mating success among a larger pool of males, and increasing the proportion of viable, fertilized eggs. Perhaps more importantly, these types of behavioral manipulations may assist with the genetic management of *P. corroboree*, by facilitating the breeding of “unattractive” but genetically valuable males. This approach may increase the total number and genetic diversity of offspring available for reintroduction into the wild. Though it should be kept in mind that selective pressures in captivity may differ from those experienced in the wild, necessitating some consideration of genetic and phenotypic predictors of post-release performance and survival. Manipulation of signals used in mate choice to improve captive breeding output has only been achieved in a few mammal species (Roberts and Gosling 2004; Parrott et al. 2019), so its application to amphibian captive breeding programs remains novel and promising.

Our findings also have implications for the application of female mate choice to conservation breeding programs globally. Our results highlight the need for empiricists to consider how multiple signal components, across various sensory modalities, may interact to influence mate choice and subsequent reproductive outcomes within CBPs. For example, if conservation managers attempt to pair individuals for breeding based on knowledge of one signal component that influences mate choice, yet females actually require multiple signals to accurately assess potential mates, breeding pairs may still be incompatible, leading to reduced reproductive success (Noer et al. 2017). Moreover, empiricists may also need to take into account how specific captive breeding environments may influence which traits are important. For instance, in homogeneous captive breeding environments, females may place greater emphasis on male phenotypic traits as opposed to variation in resource quality (such as nest sites) (Lifjeld and Slagsvold 1988). Overall, we recommend that future research aiming to incorporate mate choice into CBPs takes a multivariate approach and identifies how multiple traits interact to influence male attractiveness before integrating knowledge of female mate choice into captive management.

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Author contribution PGB, AJS, and SRK conceived the study. SRK and MSM collected the data. JAE developed color quantification methods and designed and wrote the color analysis program. SRK analyzed the call and color data. SRK and MGS ran the statistical analyses. SRK wrote the manuscript with input from all authors.

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Data availability Data is available as supplementary material.

Code availability Not applicable.

Declarations

Ethics approval All work conducted during this study was approved by the Taronga Conservation Society Australia Animal Ethics Committee (AEC protocol number—3a/10/16). The nest checks conducted...
during this study are in line with the current corroboree frog breeding protocol, and all viable eggs obtained from this study were reintroduced into the wild as part of the corroboree frog recovery plan.

**Consent for publication**  All authors have given their consent for publication.

**Conflict of interest**  The authors declare no competing interests.

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