Functional differences in the hypothalamic-pituitary-gonadal axis are associated with alternative reproductive tactics based on an inversion polymorphism

J.L. Loveland a,*, L.M. Giraldo-Deck a, D.B. Lank c, W. Goymann b, M. Gahr b, C. Küpper a

a Behavioural Genetics and Evolutionary Ecology Research Group, Max Planck Institute for Ornithology, Seewiesen, Germany
b Department of Behavioural Neurobiology, Max Planck Institute for Ornithology, Seewiesen, Germany
c Department of Biological Sciences, Simon Fraser University, Burnaby, Canada

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ABSTRACT

The evolution of social behavior depends on genetic changes, yet, how genomic variation manifests itself in behavioral diversity is still largely unresolved. Chromosomal inversions can play a pivotal role in producing distinct behavioral phenotypes, in particular, when inversion genes are functionally associated with hormone synthesis and signaling. Male ruffs exhibit alternative reproductive tactics (ARTs) with an autosomal inversion determining two alternative morphs with clear behavioral and hormonal differences from the ancestral morph. We investigated hormonal and transcriptomic differences in the pituitary and gonads. Using a GnRH challenge, we found that the ability to synthesize testosterone in inversion carriers is severely constrained, whereas the synthesis of androstenedione, a testosterone precursor, is not. Inversion morphs were able to produce a transient increase in androstenedione following the GnRH injection, supporting the view that pituitary sensitivity to GnRH is comparable to that of the ancestral morph. We then performed gene expression analyses in a second set of untreated birds and found no evidence of alterations to pituitary sensitivity, gonadotropin production or gonad sensitivity to luteinizing hormone or follicle-stimulating hormone across morphs. Inversion morphs also showed reduced progesterone receptor expression in the pituitary. Strikingly, in the gonads, inversion morphs over-expressed STAR, a gene that is located outside of the inversion and responsible for providing the cholesterol substrate required for the synthesis of sex hormones. In conclusion, our results suggest that the gonads determine morph-specific differences in hormonal regulation.

1. Introduction

Genomic rearrangements such as chromosomal inversions frequently provide the genetic variation necessary for the evolution of behavioral diversity associated with mating and reproduction (Chouteau et al., 2017; Gilburn and Day, 1999; Horton et al., 2014b; Küpper et al., 2016; Pearse et al., 2014). At the proximate level, these so-called “supergenes” may contribute to hormonal plasticity by capturing genes required for hormone production and sensitivity [e.g. (Horton et al., 2014a, 2020; Merritt et al., 2020)]. Over time, sequence evolution combined with selection on certain inversion haplotypes may further canalize hormonal profiles into a restricted range that becomes associated with specific behaviors and morphological traits.

Male mating success typically relies on secondary sexual characters that are paired with behaviors, such as territorial aggression and courtship displays. The hypothalamic–pituitary–gonadal (HPG) axis plays a pivotal role in the control of reproduction in vertebrates. At the apex of the HPG axis, gonadotropin-releasing hormone (GnRH) neurons fire synchronously and secrete GnRH to the pituitary in a pulsatile fashion. These pulses stimulate the rhythmic secretion of luteinizing hormone (LH) and follicle-stimulating hormone (FSH) from the pituitary into the blood portal system (Fig. 1) [for details on birds see also (Li et al., 1994; Urbanski, 1984)]. The intermittent stimulation of gonadal tissue by gonadotropins then induces sex steroid synthesis and spermatogenesis in the male gonad.

Sex steroid hormones initiate and maintain social behaviors through genomic and non-genomic routes (Boonyaratankornkit and Edwards, 2007). Among sex hormones, testosterone has been traditionally
invoked to be largely responsible for differences in inter-male aggression (Goodson, 2005; Wingfield et al., 1987) although alternative more integrative mechanisms have been described recently (Goymann et al., 2019; Lipshutz et al., 2019; Wingfield et al., 2019). Testosterone can reach the brain via the bloodstream or by de-novo synthesis in neural tissue (Baulieu, 1997; do Rego and Vaudry, 2016; Tsutsui et al., 1999). Yet, the regulation of social behavior does not only include testosterone but also its androgenic and estrogenic metabolites. The conversion of testosterone into estradiol is a requirement for mediating aggressive and sexual behavior in birds (Ball and Balthazart, 2004; Fusani et al., 2001; Steimer and Hutchison, 1980; Watson and Adkins-Regan, 1989) and estradiol is a known potent activator for sexual behavior in birds (Cornil et al., 2006). Estradiol can also be synthesized from estrone that is made from aromatized androstenedione, a testosterone precursor. Of note, androstenedione is only a weak activator of the androgen receptor, compared to testosterone. Another sex steroid, progesterone, has the ability to mediate aggression in rodent models (e.g. Erpino, 1975; Erpino and Chappelle, 1971; Schneider et al., 2003; Yang et al., 2013), as well as in females of the sex-role reversed black coucals (Centropus grillii) (Goymann et al., 2008). In the rufous hornero (Furnarius rufus), a South American ovenbird, the ratio between testosterone and progesterone has been shown to be a more useful predictor of outcomes of short-term agonistic encounters than testosterone alone (Adreani et al., 2018).

The evolution of diversity in mating repertoires is largely facilitated by testosterone regulation (Ball and Balthazart, 2004; Balthazart et al., 2003; Day et al., 2007; Fusani et al., 2014; Oliveira et al., 2008). Therefore, knowledge of the physiological limits of androgen responsiveness is key to understanding how behavioral diversity can evolve, especially in association with reproductive behaviors. Species that exhibit alternative reproductive tactics (ARTs) are ideal study systems for the proximate regulation of mating behavior because courtship, aggression and secondary sexual traits are in many cases dissociated from male gonadal function (Oliveira et al., 2008). GnRH challenges have proven particularly effective at unmasking when hormonal differences are largely regulated by the social context and interactions therein, and not by an actual physiological limitation to testosterone production (Apfelbeck and Goymann, 2011; Barron et al., 2015; Cain and Pryke, 2017; Goymann and Flores Davila, 2017; Goymann and Wingfield, 2004; Jawor et al., 2006; Spinney et al., 2006). During a GnRH challenge, a standardized amount of GnRH is administered to animals to examine the ability of the pituitary to respond to GnRH and the ability of the gonads to produce and secrete testosterone. In some species with reversible ranks or morphs, the entire HPG axis of subordinate males is suppressed by the social context. For example in the African cichlid Astatotilapia burtoni, the presence and agonistic behavior of dominant males actively keeps other males in a subordinate status (Maruska and Fernald, 2011). Subordinate males experience substantial shrinkage of GnRH neurons, decreased circulating 11-ketotestosterone and reduced spermatogenesis, which leads to smaller testes when the

Fig. 1. Overview of the avian hypothalamic-pituitary-gonadal (HPG) axis for testosterone synthesis. Gonadotropin-releasing hormone neurons in the anterior preoptic area of the avian brain send projections to the median eminence where GnRH is released and enters the blood portal vasculature to the anterior pituitary. The inset shows a close-up of the pituitary, which expresses predominantly a specific GnRH receptor (Joseph et al., 2009). The binding of GnRH to its cognate receptor triggers the production of the gonadotropins luteinizing hormone (LH) and follicle-stimulating hormone (FSH) that are released into the bloodstream (Dunn and Millam, 1998). In the testis, binding of FSH to its receptor in Sertoli cells stimulates spermatogenesis and binding of LH to its receptor in Leydig cells initiates a cascade of enzymatic steps that lead to the synthesis of androstenedione, which is then converted to testosterone (Ritchie, 2014). Testosterone begins its circulation through the bloodstream and serves to regulate GnRH, LH and FSH production through a negative feedback loop (Ritchie, 2014). An illustration of a ruff brain was used to represent an avian brain. (Created with BioRender).
HPG axis is under social suppression for weeks (Davis and Fernald, 1990; Francis et al., 1993). However, this can be rapidly reversed if the social environment changes (Burmeister et al., 2005; Maruska et al., 2013). In contrast, in species with genetically determined ARTs, sneaker phenotypes are usually fixed for a lifetime (Oliveira et al., 2008). Such sneaker males tend to have low circulating levels of testosterone but their gonads are of similar size or even larger than those of territorial males, probably because this maximizes their fertilization success with fewer mating opportunities (Fu et al., 2001; Gross, 1996; Taborsky, 1998). This is possible because spermatogenesis actually requires low levels of testosterone to be initiated and maintained (Kustan et al., 2012; Oliveira et al., 2001, 2008). Understanding how genetic changes are able to produce fixed alternative morphs with distinct hormonal profiles is key to building the bigger picture of how morph-specific behaviors with a genetic basis can evolve.

A prominent inversion polymorphism that captures several genes involved in steroid synthesis and metabolism is responsible for three genetically determined male mating morphs - Independents, Satellites, and Faeders - in the ruff, Philomachus pugnax (Küpper et al., 2016; Lamichhaney et al., 2016). During the breeding season, each morph has a distinct appearance and behavior (Hogan-Warburg, 1966; van Rhijn, 1973). Independent males have dark colored plumage and are overtly aggressive, whereas Satellites and Faeders, the two other ARTs, seldom exhibit aggressive behavior. Satellite males, the smallest of the three morphs, lack ornament plumage and do not perform male-typical courtship behaviors but rather adhere to a pure sneaking strategy and have testes that are up to 2.5 times larger than those of Independent males (Jukema and Piersma, 2006). Faeder males have been documented to mount females, as well as other males in the wild, and mate successfully in captivity (Jukema and Piersma, 2006; Küpper et al., 2016; Lank et al., 2013). These morphological and behavioral differences are linked to striking hormonal differences in testosterone among the morphs: Independent males have high levels of testosterone compared to Satellites and Faeders (Küpper et al., 2016). Conversely, Satellites and Faeders have high levels of androstenedione, a testosterone precursor, compared to Independent males. Androstenedione, on its own, may act as a weak activator of androgen receptors and additionally, may be converted to testosterone or estrogen in target tissues including the brain (Soma et al., 2003). Indeed, experimentally elevating androstenedione in Independents results in increased aggression but is ineffective at eliciting any aggressive behavior in Satellites (albeit it increases courtship behavior in Satellites, Morgan, 2009).

Remarkably, the determination of morph type is controlled by a 4.5 Mb autosomal inversion.Independents have two copies of the ancestral non-inverted haplotype, whereas Satellites and Faeders are both heterozygous for the inversion. Satellites and Faeders each have different inversion haplotypes: the Faeder inversion haplotype arose first and the Satellite haplotype is the result of subsequent recombination between the Faeder inversion and one or several ancestral Independent alleles (Küpper et al., 2016; Lamichhaney et al., 2016). The presence of several genes in the inversion region (i.e., HSD17B2, SDR42E1, CYSB5S) with known involvement in sex steroid synthesis and metabolism suggests an inversion-based direct effect on hormone production and regulation. In view of the relatively high levels of circulating androstenedione in inversion morph males compared to Independent males, the HSD17B2 gene, which is located within the inversion region and encodes the enzyme that converts the testosterone back to its precursor androstenedione, has been suggested as a candidate that could provide a direct explanation for this hormonal difference among morphs (Küpper et al., 2016; Lamichhaney et al., 2016). Yet, pilot data did not provide clear evidence of a key role for testicular expression differences in HSD17B2 among morphs (Loveland et al., in preparation). This suggests that perhaps a more fruitful avenue to understanding how the inversion disrupts sex steroid synthesis requires examining the entire HPG axis, and in particular its ability to mount maximal androgenic hormonal responses, rather than just gene expression differences between morphs in testicular tissue. Because the ability to produce testosterone may be a crucial trait directly controlled by genes within the inversion, identifying where within the HPG axis this disruption occurs is a necessary first step towards deciphering the underlying molecular mechanisms that generate morphological, neurobiological and behavioral differences across morphs.

In this study, we aimed to investigate the physiology and molecular mechanisms that maintain hormonal differences in adult male ruffs during the breeding season. First, we performed a GnRH challenge to characterize the physiological range of testosterone that each morph can produce. We tested whether inversion morphs were capable of mounting a typical testosterone surge following a GnRH injection. We measured testosterone, androstenedione and corticosterone to examine the response range of each hormone during the GnRH treatment. Second, in a separate set of untreated birds, we measured the expression of key genes in the pituitary and gonads to assess hormonal sensitivity and production within the HPG axis. We measured the expression of genes required for pituitary sensitivity to sex steroids, gonadotropin-releasing hormone sensitivity (GnRH receptors), gonadotropin production (FSH) and gonadal sensitivity to gonadotropins (LH and FSH receptors), as well as the expression of the gene encoding the steroidogenic acute regulatory protein (StAR) which is essential for making cholesterol available for the synthesis of steroid hormones. We hypothesized that morph differences in the expression of any of these genes would indicate which tissue types and processes (i.e. hormone production or sensitivity) along the HPG axis would be responsible for the low circulating testosterone levels observed in inversion morphs.

2. Materials and methods

2.1. Birds and housing

We used a total of 69 adult ruff males from a captive breeding flock at Simon Fraser University, with a mean age ± standard error (SE) of 5.44 ± 0.62 yrs. This captive population was originally established from eggs collected near Oulu, Finland, in 1985, 1989 and 1990 (Lank et al., 2013). We conducted all experiments and sample collection during three breeding seasons (see specific dates in relevant sections below). Males were housed in an outdoor aviary in a same-sex pen with visual access to females in adjacent pens and had unrestricted access to food and water at all times prior to being either tested in the GnRH challenge experiment or killed for tissue collection. Males devoted to tissue sampling were part of other behavioral experiments; they were performing morph-typical social behaviors immediately before being killed and no pharmacological manipulations of any kind were performed prior to collection. All males had spent at least 3 weeks in these same-sex pens before testing or sample collection. An exception was that four inversion morph birds in the GnRH experiment had been housed with females and had had opportunities to mate, but we do not have information as to whether or when they mated on the days we conducted the experiment. Other than those four birds, as far as we know all birds could have had experience in mating within the colony during previous years. All housing and procedures (permit #1232B-17) were approved by the Animal Care Committee of Simon Fraser University operating under guidelines from the Canadian Council on Animal Care.

2.2. GnRH challenge

We performed the GnRH challenge on 34 unique male birds (2018: 8 Independents, 2019: 11 Independents, 7 Satellites, 9 Faeders). We performed these experiments on the following dates: in 2018 on June 11th and 12th, and in 2019 on May 30th and 31st, June 2nd and June 11th. In 2018, we sought to test the efficacy of the dosage and the sampling time points, and we did not test all morphs. Specifically, in 2018, a subset of Faeders (n = 6) were given GnRH to establish the appropriate dose.
These individuals were GnRH injected again in 2019 and only the 2019 data were used for subsequent analyses. In addition, one Independent male had insufficient blood sampled in 2018, but was re-sampled in 2019 and we only used the 2019 data for subsequent analyses. In both years, among Independents, we randomly assigned males to either the GnRH or the saline group. Because of sample size considerations and because we were interested in morph differences in the response to GnRH, we assigned all Satellites and Faeders to the GnRH group, similar to other studies (Cain and Pryke, 2017; Goymann and Wingfield, 2004). When possible, we injected the same proportions across treatment/morph groups daily to control for day effects. For each bird, the workflow was as follows. We captured the bird by net and brought it indoors, where we weighed and bled it from the left wing vein. The time from capture to first bleed was 4.17 ± 0.26 min (mean ± SE); this constituted the pre-challenge baseline sample. Immediately after blood sampling, we gave the bird an intramuscular GnRH injection (Bachem, H-3106) as performed in other studies with similar dose per body weight [e.g. (Apfelbeck and Goymann, 2011; Bergeon Burns et al., 2014; Goymann and Flores Davila, 2017; Jawor et al., 2006; Susan DeVries et al., 2011)] aimed at the pectoral muscle with a dose of 12.5 μg per 150 g body weight or a corresponding volume of saline, and placed it in a box. The time points for sampling were based on the above-mentioned studies, and we confirmed these were effective for the ruff in our first year of testing (2018). Three boxes were used for keeping the birds between blood sampling intervals (LxWxH dimensions: 80 × 50 × 45 cm). The inside of each box was kept dark and the boxes were kept in the same room where the experimenters were handling birds. As injections were performed staggered, a box held from 1 to 3 birds at any time. We took two more blood samples at 30 and 90 min post-injection from the right wing and either right or left leg, respectively. We drew a maximum of 300 μl of blood each time, kept it on ice for the duration of testing time, and then centrifuged it at 3000 rpm for 15 min. We collected plasma and stored it at −20 °C for up to four weeks before transporting it on dry ice to the lab in Germany and subsequent storage at −80 °C until analysis.

2.3. Hormone analysis

Plasma testosterone, androstenedione, progesterone and corticosterone concentrations were determined by radioimmunoassay (RIA) following previously published protocols for extraction and measurement (Goymann et al., 2001, 2006, 2008). For testosterone, androstenedione and corticosterone, plasma samples were extracted with dichloromethane (DCM) after overnight equilibration (4 °C) of the plasma with 1500 dpn of tritiated testosterone, androstenedione, or corticosterone (Perkin Elmer, Rodgau, Germany). The organic phase was then separated from the aqueous phase by plunging the extraction tubes into a methanol-dry ice bath and decanting the dichloromethane phase into a new vial. This extraction step was repeated twice to increase extraction efficiency. Then, the DCM phase was dried under a stream of C, dried samples resuspended in phosphate buffered saline with 1% gelatine (PBSG) and left overnight at 4 °C to equilibrate. Progesterone was extracted with ethyl ether after overnight equilibration (4 °C) of the plasma with 1500 dpn of tritiated progesterone (Perkin Elmer, Rodgau, Germany). Separation of the organic and aqueous phase was conducted similar to the procedure described for the other hormones. This kind of extraction effectively removes substances that could interfere with the assay. For all hormone extracts, an aliquot of the redissolved samples was transferred to scintillation vials, mixed with 4 ml scintillation fluid (Packard Ultima Gold) and counted to an accuracy of 2.5. For the RIA, a standard curve was set up in duplicates by serial dilution of stock standard testosterone and androstenedione ranging from 0.39–200 pg, and cortisol and progesterone from 1.95–1000 pg. Testosterone, androstenedione, corticosterone or progesterone antisera (all from Esoterix Endocrinology, Calabasas, CA, USA) were added to the respective standard curve, the controls and to duplicates of each sample (100 μl). After 30 min the respective testosterone, androstenedione, corticosterone or progesterone labels were added and the assays incubated for 20 h at 4 °C. Then, bound and free fractions were separated at 4 °C by adding 0.5 ml dextran-coated charcoal in PBSG assay buffer. After 14 min incubation with charcoal, samples were spun (3600 g, 10 min, 4 °C) and supernatants decanted into scintillation vials at 4 °C. After adding 4 ml scintillation liquid (Packard Ultima Gold) vials were counted. Standard curves and sample concentrations were calculated with Immunofit 3.0 (Beckman Inc. Fullerton, CA), using a four parameter logistic curve fit. Data were analyzed in three assays for testosterone, two assays for androstenedione, and one assay for corticosterone and progesterone, respectively. The lower detection limits of the standard curves were determined as the first value outside the 95% confidence intervals for the zero standard (Bmax) and were 3.6–4.9 pg/ml for testosterone, 6.7–8.1 pg/ml for androstenedione, 57.0 pg/ml for corticosterone, and 27.6 pg/ml for progesterone. The intra-assay coefficient of variation as determined by extracted chicken plasma pools was 1.5%, 3.0% and 10.4% for the testosterone assays, twice 14.8% for the androstenedione assays, 13.2% for the corticosterone assay and 24.6% for the progesterone assay. The inter-assay coefficient of variation as determined by extracted chicken plasma pools for the three testosterone assays was 5.9% and for the two androstenedione assays was 1.3%. Because the testosterone antibody used shows significant cross-reactions (44% according to the specification sheet provided by the manufacturer) with 5α-dihydrotestosterone our testosterone measurement may include a fraction of 5α-dihydrotestosterone.

2.4. Tissue collection

We sampled male birds in the morning between 8:00–12:15, except for two birds in 2019 that were sampled at 13:00 and 15:00, respectively. The dates of collection were: 11th–17th June (2017), 5th–16th June (2018), and 8th–13th June (2019). We captured each bird with a standard net and placed it in a bag for transport indoors. Once indoors, the bird was weighed and killed by rapid cervical dislocation followed by immediate decapitation to minimize stress (average time from capture to sacrifice, mean (range) = 5.35 (1–14) minutes). We collected blood by vein puncture prior to killing and/or directly from the decapitation site. We dissected tissue samples including gonads, which we weighed, rinsed briefly in phosphate-buffered saline (PBS) and dried with a tissue before putting one gonad in RNAlater (Ambion, cat#AM7021) and the other to freeze in Neg50 (Thermoscientific) medium or in aluminum foil on dry ice; the former were used for RNA extractions. Pituitaries were removed from the skull and placed immediately in RNAlater. These samples stored in RNAlater were kept at 4 °C for 1–3 days per manufacturer’s instructions and then stored frozen at −20 °C for approximately one month until transporting to the Max Planck Institute for Ornithology in Seewiesen on dry ice and then stored at −80 °C until RNA extraction.

2.5. RNA extraction and cDNA synthesis

We extracted RNA from 35 pituitaries and 35 gonads (17 Independents, 9 Satellites, 9 Faeders) using the RNasy MiniKit (Qiagen) according to the manufacturer’s protocol. One modification for gonads was that 200 μl of the lysed homogenate was mixed with 300 μl lysis buffer, passed through a QIAShredder column to remove excess genomic DNA before starting the RNA extraction protocol. We measured RNA concentration with a Nanodrop and assessed RNA quality with the Bioanalyzer RNA nanochip (Agilent) in a subset of pituitary samples (N
to further analysis due to low yield or RNA integrity = 25) and all gonad samples. We excluded two pituitary and four gonad samples from further analysis due to low yield or an RNA integrity number (RIN) below 5.4. For the remaining samples, the average RIN for pituitaries and gonads was 9.8 and 7.4, respectively. We synthesized a total of 500 ng (pituitary) and 1 μg (gonad) of RNA into cDNA using the iScript cDNA synthesis kit (Bio-Rad) in 20 μl reactions according to manufacturer’s instructions. We diluted cDNA ten-fold before use as template in qPCR assays.

2.6. Primer design for target genes

To measure expression of genes essential to HPG axis function and steroid synthesis, we designed primers for ten genes and tested expression in pituitary and testis as follows: FSHB, GNRH-R1, GNRH-R3, AR, PGR, ESR1, ESR2 in pituitary only; FSHR, LHR, STAR in testis only. To assess pituitary sensitivity to GnRH we examined gene expression of gonadotropin-releasing hormone receptors (GNRHRs) I and III (hereinafter GNRH-R1 and GNRH-R3). Avian GNRH receptors have varying names in the literature (e.g. (Bédécarrats et al., 2006)); based on sequence identity conservation between ruff and chicken sequences we followed the nomenclature of Joseph et al. (2009). We included the follicle-stimulating hormone beta subunit (FSHB) to gauge pituitary production of FSH and follicle-stimulating hormone receptor (FSHR) to assess gonadal sensitivity to FSH. Likewise, we designed primers for the gene encoding the luteinizing hormone receptor (LHR) to assess gonadal sensitivity to its corresponding gonadotropin. We selected the gene encoding the steroidogenic acute regulatory protein (StAR) because it was possible to detect differences in cholesterol metabolism, which is essential for steroid synthesis. To test for sensitivity to sex steroids, we designed primers for genes encoding the estrogen receptors alpha and beta (ESR1 and ESR2), progesterone receptor (PGR) and androgen receptor (AR). For genes that we did not detect by name in the current ruff genome release (NCBI Calidris pugnax Annotation Release 100, software version 6.5) we searched for homologs using the respective chicken, turkey or quail sequence (NCBI) as query against the ruff genome with BLAST (Altschul et al., 1990). We designed primers with NCBI Primer-BLAST (Ye et al., 2012) and ensured that each amplicon spanned at least one exon-exon boundary and when possible we designed at least one primer within a given pair to anneal across an exon-exon junction. Genes encoding GnRH receptors were confirmed after cross-validation with alignments to other bird species because their names in the ruff genome were different loci for ‘gonadotropin-releasing hormone II receptor-like’. All primers and results from standard curves are listed in Table S1. Despite our best efforts, we were unable to locate the gene encoding the beta subunit of the luteinizing hormone with these methods, possibly because of the high GC content of this gene in avian species (chicken 72.12%, turkey 67.94%, quail 62.5%). Both FSH and LH use a common alpha subunit, but because their respective beta subunits are hormone-specific (Proudman et al., 1999) we could not use the alpha subunit as a proxy for LH production. We reasoned that even if we were able to bioinformatically retrieve this sequence in the ruff, designing useful qPCR primers would have been difficult due to the numerous stretches of Gs and Cs.

2.7. Primer design for reference genes

We tested a panel of 11 genes in gonad- and pituitary-derived cDNA for their suitability as reference genes for use in qPCR (Table S1, based on (Loveland et al., 2014; St-Pierre et al., 2017; Zinowi-Kramer et al., 2014)). The best two genes for each tissue type that did not show differences between morphs were chosen after analysis with GeNorm and NormFinder.

2.8. Quantitative PCR (qPCR)

We conducted and report all qPCR assays according to the Minimum Information for Publication of Quantitative Real-Time PCR Experiments (MIQE) guidelines (Bustin et al., 2009). We ran assays on a LightCycler 480 II (Roche) machine using the SooAdvanced Universal SYBR Green Super mix (Bio-Rad) in 384-well plates (Roche) and each reaction was run in duplicate. We performed qPCR assays to generate the full data set across eight plates (each plate always balanced for morph) and standard curves on four plates. Each well consisted of a 10 μl reaction containing 1× SooAdvanced Universal SYBR Green Super mix, 340 nM of each primer and either 3.75 ng (pituitary) or 7 ng (gonad) of cDNA (i.e. 1.5 μl of the ten-fold cDNA dilution assuming a one-to-one correspondence between input RNA and synthesized cDNA; this practice of synthesizing the same amount of RNA into cDNA for each sample is standard and recommended practice (Bustin et al., 2009; Ståhlberg et al., 2004)). The cycling conditions were: pre-incubation step (95 °C for 30 s), 45 cycles (95 °C for 10 s, annealing and extension at 60 °C for 30 s) with acquisition at the end of each cycle, followed by a melt curve (95 °C for 5 s with 5 acquisitions per °C from 65 °C to 97 °C with a 0.11 °C ramp rate). We performed calculations from the raw amplification data in the LightCycler 480 Software (version 1.5.1.62). On every plate we confirmed that each primer pair produced a single melt curve peak in the presence of cDNA template that was different from any primer dimer that may form when only water is used as template. Negative controls showed either no amplification or the presence of an unambiguous primer dimer peak. To test for any possible genomic carryover from RNA extraction to cDNA synthesis, we ran the qPCR with cDNA synthesis negative controls (i.e. no reverse transcriptase) for a subset of samples; as expected, none showed any amplification.

We ran a standard curve with serially diluted cDNA (1:10 to 1:100000) from one sample of an Independent male to calculate the amplification efficiency of each primer pair, and efficiency was calculated separately for each tissue. This dilution range for ESR2 primers was too wide, so for these primers we used serially diluted cDNA in a lower range (1:10 to 1:40). Efficiency was calculated with the Absolute Quantification tool and 2nd Derivative Maximum method which uses the formula [Efficiency = 10−1/slope] based on the quantification cycle (Cq, termed crossing point (Cp) in the software) and log concentration of template in each well. Efficiencies are listed in Table S1, all were above 1.92. We confirmed that the Cq values for the selected reference genes did not vary between Independents (n = 5) and inversion morphs (n = 6, 3 Satellites and 3 Faeders) (pituitary HPRT1 Mann-Whitney p = 0.662, RPL30 Mann-Whitney p = 0.429; gonads HPRT1 t-test p = 0.380, RPL32 t-test p = 0.5). We calculated the relative mRNA levels of target genes with the best reference genes for each tissue (HPRT1 and RPL30 for pituitaries, HPRT1 and RPL32 for gonads) with the Advanced Relative Quantification analysis, which is based on the ∆∆CT method by (Livak and Schmittgen, 2001) using the actual primer efficiencies from the standard curve instead of the preassigned value of 2.

2.9. Statistical analyses

We used log-transformed hormone data for statistical analyses, but plotted non-transformed values in the figures. We calculated fold-change at the 30 min time point relative to baseline using non-transformed values. We used only Independent males to test for the effect of treatment (control vs. GnRH). We analyzed samples for testosterone in 2018 and 2019 and for androstenedione and corticosterone in 2019. In an initial model, we used a two-way ANOVA on the difference between baseline and the 30 min time point in Independent males only to test for the main effect of year or interaction between year and treatment in testosterone levels. As neither the interaction between year and treatment nor year showed a significant effect (year x treatment in testosterone levels. As neither the interaction between year and treatment nor year showed a significant effect (year x treatment = 0.396, p = 0.593; year F(1,14) = 1.921 p = 0.187; treatment F(1,14) = 15.07, p = 0.002) we pooled the individuals by treatment group across both years for the final model.

To compare the hormonal changes between saline and GnRH-injected Independent males we prepared a mixed model with time, treatment and their interaction as fixed factors and ID as random factor.
For posthoc tests, we tested whether the 30 and 90-min time points were different from baseline for each treatment group, and report Dunnett’s tests. To assess whether treatment groups differed clearly at each time point we used Sidak’s tests.

To assess hormonal variation across different morphs in response to the GnRH challenge, we compared GnRH-injected individuals only. We used a mixed model with morph, time and their interaction as fixed factors and ID as random factor. For posthoc tests, we tested whether the 30 and 90-min time points were different from baseline for each morph, and report Dunnett’s test. To test for morph differences at each time point we performed the Tukey’s test.

For comparisons among morphs in fold-change for testosterone and androstenedione, we analyzed morph differences as the relative change at 30 min from baseline (i.e. fold-change) with Kruskal-Wallis (KW) tests because sample sizes were too small to test for normality. We performed posthoc tests when the KW test was significant and used Dunnett’s test to assess which morphs differed from one another.

For qPCR data on nine genes (six tested in pituitary and three in gonads) we analyzed variation in expression between morphs at each time point we performed the Tukey’s test to assess which morphs differed from one another.

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3. Results

3.1. GnRH challenge in Independent males

The GnRH challenge resulted in different circulating testosterone concentrations in saline and GnRH-injected Independent males (Table S2, Fig. 2A). We found a strong interaction between time and treatment \((F(2,22) = 14.06, p < 0.001)\). Saline-injected birds had testosterone levels lower in post-injection samples relative to baseline, whereas, as predicted, GnRH-injected birds had higher levels of testosterone after 30-min post-injection relative to baseline (Dunnett’s \(p = 0.066\) (Fig. 2A). We also found a treatment effect for circulating androstenedione concentrations \((F(1,9) = 8.129, p = 0.019, \text{Table S2, Fig. 2B})\). At the 30-min time point, GnRH-injected Independent males had higher androstenedione levels compared to baseline (Dunnett’s \(p = 0.014)\), whereas saline-injected males did not differ between baseline and the 30-min sample (Dunnett’s \(p = 0.971)\). At the 90-min time point there was no clear difference between treatment groups nor to the baseline measurements in androstenedione concentrations.

![Graph A](image1.png)  
**Fig. 2.** Hormone response in Independent ruff males after a GnRH challenge. 
Time-course of steroid hormone levels in Independent males before and after a GnRH or control injection are shown for (A) testosterone, (B) androstenedione and (C) corticosterone 0, 30 and 90 mins post-injection. Individual data points are superimposed on box plots. The “0” timepoint indicates pre-injection levels.
were instead caused by functional differences in Leydig cells, in processes and pathways that occur after (i.e. downstream from) LHR activation (Fig. 5A). Consistent with this, morphs differed in STAR expression (ANOVA F(2, 28) = 6.971, p = 0.003, FDR-adjusted p = 0.016) (Fig. 5B). Yet, the observed STAR expression differences were not in the predicted direction. Faeders had the highest expression levels, exhibiting statistically clear differences compared to Independents (Tukey’s p = 0.002), but not to Satellites (Tukey’s p = 0.087). There was no clear difference between Independents and Satellites (Tukey’s p = 0.481). In agreement with previous reports, among the untreated birds (i.e. those not part of the GnRH experiment) morphs differed in circulating testosterone levels (Kruskal-Wallis p = 0.005) (Fig. 5C): inversion morphs had similarly low levels (Satellite vs. Faeder, Dunn’s p > 0.99), with Faeders (Dunn’s p = 0.008), but not Satellites (Dunn’s p = 0.064) exhibiting clearly lower levels than Independents.

Morphs differed in the expression of the progesterone receptor in the pituitary (ANOVA F(2,30) = 11.32, p < 0.001, FDR-adjusted p = 0.002) (Fig. 6A). Faeders had lower PGR expression than the other two morphs (Independents vs. Faeders, Tukey’s p < 0.001; Satellites vs. Faeders, Tukey’s p = 0.030). Satellites showed intermediate expression values, but the expression was not clearly different from that of Independents (Tukey’s p = 0.281). None of the other sex hormone receptors ESR1, ESR2 or AR showed clear differences in pituitary expression among morphs (all, FDR-adjusted p > 0.8). There was no relationship between PGR expression and circulating progesterone levels when all males were pooled (Spearman r = 0.005, p = 0.980). However, within morphs we detected a positive relationship in Faeders (Spearman r = 0.810, p = 0.022), but not in the other two morphs (Satellites Spearman r = −0.028, p > 0.99; Independents Spearman r = 0.05, p = 0.88) (Fig. 6B). We tested for morph-independent correlations between gene expression (nine genes: pituitary FSHB, GNRH-R3, AR, ESR1, ESR2, PGR, and gonadal LHR, FSHR, STAR) and androgens (testosterone, androstenedione and progesterone). We did not have all three hormone measurements for all birds due to insufficient plasma in some samples (testosterone, N = 31; androstenedione, N = 21; progesterone, N = 23). Only the association between GNRH-R3 expression in the pituitary and testosterone was statistically clear after the FDR adjustment (Pearson r = 0.682, p < 0.001; FDR-adjusted p = 0.003), however, we note that this was driven by the values of a single Independent male with the highest testosterone levels.

4. Discussion

Breeding ruffs show striking differences in behavior, appearance and androgen physiology across three morphs that only differ at the genomic level by a small autosomal inversion. Exactly how the inversion causes these hormonal and behavioral differences is unknown. In one set of birds, we tested the physiological range of androgenic production across the three morphs and in another set of untreated birds, we analyzed gene expression along the HPG axis to discern which tissue types and processes are likely to give rise to androgenic hormonal differences in this species. We found that the two inversion morphs, Satellites and Faeders, had generally a similar physiological response to the GnRH challenge, both inversion morphs were incapable of synthesizing high levels of testosterone, but notably, were able to produce high levels of androstenedione.

Because of the presence of a large inversion that contains approximately 1000 genes (Thomas et al., 2008; Thorneycroft, 1975). In this species, the white morph harbors the inversion and both males and females have brighter plumage, higher aggression and lower parental care during the breeding season compared to the tan morph, which is the presumed ancestral morph
because it is homozygous for the non-inverted allele (Falls and Kopachena, 1993; Ficken et al., 1978; Horton et al., 2013). As in ruffs, there are morph differences in testosterone levels: white morph males have higher testosterone levels than tan males. However, this difference is only seen in wild-caught birds, not in captive ones, perhaps because of differences in the stability of the social environment or induced stress, which could potentially suppress testosterone production via increased corticosterone (Spinney et al., 2006). In contrast, in ruffs testosterone differences among morphs in the breeding season are highly pronounced even in captivity (Küpper et al., 2016, this study), suggesting that in the wild this difference may be even more pronounced and/or the effect of the inversion on circulating testosterone levels is less labile to the influence of the social environment. In white-throated sparrows, white-striped inversion morph males show a more robust increase in testosterone after a GnRH challenge compared to tan-striped males, even though the LH surge caused by the injection is not different between the morphs (Spinney et al., 2006). In that study, the authors suggested that testosterone differences might stem from morph differences in the expression of LHR and/or steroidogenic enzymes in the gonads. In contrast, in our study species, inversion morphs are the less aggressive morphs and have lower testosterone compared to the ancestral morph (Independents); accordingly, their testosterone response to GnRH is markedly subdued. Further, we confirmed that there are no morph differences in gonadal LHR expression among morphs despite the presence of a subdued testosterone response of the ruff inversion morphs compared to that of Independents. In addition, males of the inversion morph in the white-throated sparrow have higher FSHR expression in gonads compared to non-inverted tan males (Zinzow-Kramer et al., 2014); yet, we did not find any morph differences in FSHR expression among ruff morphs.

We predicted that if the cause of low testosterone in inversion morphs were due to a direct constraint from the inversion itself, testosterone levels after GnRH administration would show either a subdued increase or no increase at all in Satellites and Faeders. Alternatively, if low testosterone in inversion morphs were regulated by the social context, we expected that testosterone levels in inversion morphs would increase after the GnRH injection. Our results provide strong support for the former, that is, a direct effect of the inversion on the capacity to produce testosterone. This suggests that the inversion irreversibly disrupts testosterone synthesis in Faeders and Satellites. Our results from the expression of key genes along the HPG axis in untreated males point towards the gonads as the most likely site of physiological

![Figure 3](image-url)

**Fig. 3.** Ruff morphs differ in their androgenic response after a GnRH challenge. Hormone levels in Independent (blue), Satellite (red) and Faeder (yellow) males before and after a GnRH injection are shown for testosterone and androstenedione in top and bottom panels, respectively. Both Satellites and Faeders are heterozygous for a dominant autosomal inversion. Individual data points are superimposed on box plots. The time course from pre- to post injection is shown in (A and C) and fold-change at 30 min relative to pre-injection levels is shown in (B and D), the dashed line at ‘1’ fold-change indicates ‘no change’. The “0” time point indicates pre-injection levels. Box plots extend to the 25th and 75th percentiles, the line is drawn at the median and the whiskers extend to the minimum and maximum values. To provide informative detail to the illustration of testosterone levels for all morphs, note the introduction of y-axis breaks in A and B. See Table 1 for statistics on A, B. Asterisk denotes $p < 0.05$. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
differentiation that leads to the observed low circulating testosterone levels in inversion morphs. First, we found that expression in the pituitary is broadly consistent with no differentiation in pituitary function between morphs, though Satellites did show a trend for higher FSHB expression. However, we cannot rule out the possibility that there are morph differences in the amount of GnRH that is naturally produced, as we have not studied properties of GnRH neurons or endogenous GnRH production. In other species with fixed ARTs, differences in the number and size of GnRH neurons between morphs exist, suggesting that there might also be differences in how much GnRH is synthesized and released. For example, in the white-throated sparrow, females of the inversion morph have fewer, but larger, GnRH-immunoreactive neurons in the septo-preoptic area (Lake et al., 2008). Likewise, in theplainfin midshipman, Porichthys notatus, territorial courting males have larger GnRH neurons than type II non-territorial sneakers (Grober et al., 1994). However, we did not find differences in GNRHR3 expression between morphs, which would have been indicative of morph differences in GnRH receptor expression and/or endogenous GnRH production.

Birds have two GnRH isoforms (GnRH1 and GnRH2) and two genes for GnRH receptors (GNRHR1 and GNRHR3). Originally, we had intended to measure expression of both GnRH receptors to determine whether both were expressed in the pituitary and whether one was more...

**Fig. 4.** Ruff morphs do not differ in the expression of key genes in the pituitary and gonads that regulate gonadotropin sensitivity and production. Illustration of the end products for key genes responsible for pituitary sensitivity to GnRH and gonadotropin production and gonadal gonadotropin sensitivity (left panel). Box plots of pituitary expression of GNRHR3 and FSHB, and testis expression of FSHR and LHR in Independents (blue), Satellites (red) and Faeders (yellow) (right panel). Gene expression is plotted as abundance relative to two reference genes, see Methods for details. Box plots extend to the 25th and 75th percentiles, the line is drawn at the median and the whiskers extend to the minimum and maximum values. Individual data points are superimposed on box plots. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

**Fig. 5.** Gonadal STAR expression is atypical in ruff inversion morphs during the breeding season. A) Schematic representation of regulatory interactions between STAR and testosterone in gonadal Leydig cells. A series of enzymatic reactions convert cholesterol into androstenedione denoted for simplicity by the dashed arrow. Androstenedione to testosterone conversion is reversible through HSD17B2 activity (bold), encoded by a gene inside the inversion shared by Satellites and Faeders. B) Male morphs differed in levels of STAR expression in the testis and C) in testosterone measured from circulating blood plasma. In both cases, only Faeders and Independents showed a clear statistical difference (asterisks, adjusted p < 0.01 from posthoc tests). According to assumed feedback mechanisms between testosterone and STAR protein expression (depicted in A), inversion morphs, in particular Faeders, appear to have such low levels of circulating testosterone that they are unable to down-regulate STAR expression in a typical manner. Box plots extend to the 25th and 75th percentiles, the line is drawn at the median and the whiskers extend to the minimum and maximum values. Individual data points are superimposed on box plots; panel C the y-axis break was introduced to provide informative detail on the full range of testosterone levels. (Created with BioRender).
abundant than the other. In chicken (Gallus gallus), the GnRH-R3 receptor is thought to be the major receptor for regulating pituitary gonadotroph function, in part because its levels in the pituitary are 1400-fold higher than GnRH-R1 (Joseph et al., 2009). The primers we had designed for GNRRH-R1 did not pass quality control criteria, most likely because this gene has low expression (see Table S1). Therefore, we were unable to examine GNRRH-R1 expression in the pituitary. However, in line with the study by Joseph et al. (2009) we observed GNRRH-R3 expression levels similar to the expression seen in chicken, suggesting that GNRRH-R3 is the major functional receptor for GnRH1 in the pituitary.

Second, we found that, among untreated males, both inversion morphs had an apparent chronic upregulation of STAR expression in the gonads. Genes early in a biosynthetic pathway can be major regulators of the synthesis of particular end products, in part due to negative feedback mechanisms that ensure prompt and tight regulation (Manna and Stocco, 2005). The StAR protein is essential to making cholesterol available for the synthesis of sex steroids (Lin et al., 1995) (Fig. 5A). To achieve this, the StAR protein transports cholesterol from the outer to inner mitochondrial membrane and then from that point on, through a series of enzymatic steps cholesterol can be converted into any of several steroid hormones (Stocco, 2001). In vertebrates, the STAR gene is typically regulated by several hormones, including testosterone, and its function has been suggested to be a rate-limiting step for testosterone synthesis, despite not being a ‘strict’ enzymatic protein (Houk et al., 2004; Manna and Stocco, 2005). In the dark-eyed junco (Junco hyemalis) STAR expression levels predict individual variation in testosterone among males, with higher expression in males with relatively high circulating testosterone levels (Rosvall et al., 2016). Accordingly, we had initially hypothesized that inversion morphs would have reduced STAR expression compared to Independent males. However, the relatively high levels of androstenedione in inversion morphs added complexity to this prediction. Our STAR expression result suggests that inversion morphs have the substrate needed (i.e. cholesterol) to mount a steroidal response (also evidenced by GnRH-treated inversion morphs that were able to elevate androstenedione levels). Yet still, elevated STAR levels even in untreated inversion morphs are indicative of a failed attempt to elevate testosterone levels. Moreover, in dark-eyed juncos testis mass predicts testosterone responsiveness and subspecies differ in the expression of luteinizing hormone receptor (Bergeon Burns et al., 2014). The underlying genetic mechanism for this variation is not known. In ruffs, the inversion determined morphs also differ in relative testis mass, but this variation is decoupled from testosterone responsiveness and we did not find morph differences in the expression of luteinizing hormone receptor in testes. These species-specific responses in passerines and shorebirds point to naturally occurring plasticity in different parts of the HPG axis, which deserves further study.

One reason for the low circulating levels of testosterone could be an increased back conversion from testosterone into androstenedione in both inversion morphs (Fig. 5A) as has been suggested previously (Küpper et al., 2016; Lamichhaney et al., 2016). Inversion morphs have high levels of androstenedione during the breeding season (Küpper et al., 2016, this study). The gene encoding HSD17B2, an enzyme that converts testosterone to androstenedione, is located within the inversion region and its immediate surrounding sequence contains several deletions in the inversion haplotypes (Küpper et al., 2016; Lamichhaney et al., 2016). Interestingly, we observed a moderate increase of androstenedione levels in both inversion morphs 30 mins post-injection of GnRH. This could indicate that in the inversion morphs testosterone is converted back to androstenedione rather quickly, such that at our sampled time points the short-lived transient increase in testosterone in inversion morphs had already passed. However, we think that such a scenario is unlikely as gonadal HSD17B2 levels are similar across morphs (Loveland et al., in preparation). Rather we suggest that the lack of a robust increase in testosterone in inversion morphs after a GnRH injection results from an existing impairment in androstenedione to testosterone conversion leading to a lack of testosterone availability in the first place.

We report a novel link between progesterone receptor expression in the pituitary and circulating progesterone in the Faeder morph, which has the more ancient inversion. This result while somewhat unexpected, offers an interesting avenue for future exploration. PGR transcript abundance was highest in Independents followed by Satellites and then Faeders; and only in Faeders, PGR transcript abundance was positively correlated to circulating progesterone levels. Assuming the same binding properties between progesterone and its receptor among morphs, a reduction in PGR would require a greater amount of progesterone in Faeders for the same receptor occupancy [i.e. “law of mass action” (Nussey and Whitehead, 2001)]. In black coucals (Centropus grillii), a sex-role reversed species, progesterone inhibits aggression in females (Goymann et al., 2008) and in the rufous hornero, the ratio between testosterone and progesterone can predict outcomes of short-term agonistic encounters better than testosterone alone (Adreani et al., 2018). In the tree lizard (Urosaurus ornatus), fixed color morphs result from organizational effects of both progesterone and testosterone levels and most striking, is that experimental manipulation of progesterone

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**Fig. 6.** Morphs differ in progesterone receptor expression in the pituitary. A) PGR expression in Independents (blue), Satellites (red) and Faeders (yellow). Gene expression is plotted as abundance relative to two reference genes, see Methods for details. Box plots extend to the 25th and 75th percentiles, the line is drawn at the median and the whiskers extend to the minimum and maximum values. Individual data points are superimposed on box plots. B) Relationship between progesterone and PGR expression in each morph (colors as in A), a significant correlation is only present in Faeders. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
levels with a single injection on the day of hatching is sufficient to induce males to become the orange-blue morph (Moore et al., 1998). In another study, progesterone implants in castrated males triggered aggressive behavior, albeit to a lesser degree than testosterone implants (Weiss and Moore, 2004). While these evidences implicate progesterone in aggression regulation in lizards and birds, whether and how pituitary PGR expression and Faeder behavior and physiology could be linked is less clear. Perhaps, because Faeder males have been subjected to selection that favors a non-aggressive phenotype, the progesterone receptor may have lost any functional role in relation to male aggression altogether in this morph and as such, its expression is reduced.

Given that the inversion is the main location of genetic differentiation among morphs, we expect that constraints on testosterone production are life-long. Accordingly, inversion morph males will experience significant differences in exposure to androgens early in development compared to Independent males. In birds, how the hormonal environment in early development affects the sexual differentiation of the brain and behavior is complex and species-specific (Adkins-Regan, 1987; Balthazart and Ball, 1995; Gahr, 2004). To date, this has not been investigated in ruffs. Nonetheless, we expect brain development to be influenced by an interaction between the hormonal environment, sex chromosomes and inversion genotype, ultimately leading to adult neural circuits for morph-specific behaviors. The earliest detected differences between ruff morphs are between chicks. At hatching, chicks of all morphs are similar in size but quickly develop different growth trajectories (Giraldo-Deck et al., 2020). Still, behavioral differences across male morphs may be influenced by organizational effects of testosterone. Consistent with a hypothesis for strong organizational effects, wherein de-masculinization of the brain occurs due to estrogen levels in females, adult ruff females that are given testosterone implants grow male-like plumage and exhibit male-typical behaviors according to their genotype (Lank et al., 1999). These results demonstrate that genotype-specific neural circuits for male behaviors are present in both sexes and can be activated by testosterone. In addition, this suggests that since the inversion is present in all cells, the same constraints that lead to low circulating testosterone levels in inversion morphs could also be present in the de-novo production of testosterone in neural tissue, which could further accentuate neurobiological differences among morphs. Sneaker males may have evolved to have high levels of glucocorticoids as an adaptive response that enables the maintenance of a stable mating phenotype (Arterbery et al., 2010; Knapp and Neff, 2007). Furthermore, it is possible that acute increases in androgens are also accompanied by increased corticosterone. We therefore investigated whether morphs differ in the rate at which they can increase and decrease levels of this stress hormone. However, we did not find a morph difference in the degree to which GnRH affected corticosterone, rather corticosterone increased in both saline and GnRH injected groups. This suggests that ruff morphs experienced handling stress in a similar way.

In the white-throated sparrow, recent studies have elucidated how two genes from within the inversion, estrogen receptor alpha (ESR1) and vasoactive intestinal peptide (VIP), are responsible for morph behavioral differences in aggression, song frequency and parental care in specific regions of the brain (Horton et al., 2014a, 2020; Merritt et al., 2020; Sun et al., 2018). Notably, because in the white-throated sparrow breeding season testosterone differences between morphs disappear under captive social settings, the picture that has emerged is that these key genes within the inversion affect neural gene expression that is associated with estrogen sensitivity and production of the VIP hormone in specific brain areas. Yet, importantly, sex steroid hormone synthesis is not as strongly influenced by the inversion than as we report here for the ruff. From these two exemplar fixed ART species, a picture of complex gene regulation is emerging showing that variation in gene expression may be caused by sequence divergence in regulatory regions that are the consequence of suppressed recombination (e.g. cis-effects in ESR1 and VIP in white-throated sparrows) as well as trans-effects, where the presence of the inversion affects the expression of genes that are outside of the inversion and functionally associated with androgen production.

5. Conclusions

We investigated how inversion morphs respond hormonally to exogenous GnRH in male ruffs. We found that the ability to synthesize testosterone is severely disrupted in the two inversion morphs, Faeders and Satellites. There was no evidence for widespread reduced functioning or throughput in early stages of the testosterone synthesis pathway, as inversion morphs show over-expression of STAR, which provides substrate for synthesis of sex hormones. Given the absence of clear differences across morphs in the expression of key genes in the pituitary for HPG axis function (Fig. 4), we argue that morph differences in the capacity to produce testosterone originate within the gonads, and notably, are not due to morph differences in gonadotropin production nor sensitivity. In support, we found that inversion morphs were able to produce a transient increase in androstenedione, a testosterone precursor, following the GnRH injection, supporting the view that their sensitivity to GnRH is comparable to that in Independents. Taken together, our study shows that the chromosomal inversion may have profound effects on the expression of genes involved in steroid regulation that are located outside of the inversion region.

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CRediT authorship contribution statement

JLL conceived the GnRH-related and gene expression elements of the project; JLL, CK and DL designed experiment conditions and sample collection; JLL, LG and DL performed the GnRH challenge experiment; WG supervised hormone measurements; MG provided lab space and reagents, JLL collected and processed all samples and analyzed the data. JLL wrote the initial draft with input from CK, all authors edited and approved the final manuscript.

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