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A transcriptomic investigation of handicap models in sexual selection

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Abstract Handicap models link the evolution of secondary sexual ornaments to physiological costs and thus provide a mechanistic explanation for signal honesty in sexual selection. Two commonly invoked models, the immunocompetence handicap hypothesis (ICHH) and the oxidative stress handicap hypothesis (OSHH), propose suppression of immunocompetence or increase of oxidative stress by testosterone, but empirical evidence for both models is controversial and based on morphological and physiological assays. Here, we investigated these two models on the gene transcription level using microarrays to quantify the transcriptomic response of red grouse (Lagopus lagopus scoticus) caecal, spleen and liver tissues to experimental manipulation of testosterone levels. We used a GENEONTOLOGY framework to identify genes related to immune function and response to reactive oxygen species and examined how transcription levels changed under

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F. Mougeot Instituto de Investigaciones en Recursos Cinegéticos (IREC-CSIC), Ronda de Toledo s/n, 13005 Ciudad Real, Spain experimentally increased testosterone levels in birds with parasites present or absent. Contrary to our expectations, testosterone had virtually no effect on gene transcription in spleen and liver. A small number of genes were significantly differentially regulated in caecum, and while their functions and transcription changes are consistent with the ICHH, we found little support for the OSHH. More genes responded to testosterone in the presence rather than absence of parasites, suggesting that handicap mechanisms may be context dependent and more pronounced in the presence of adverse environmental conditions. These findings illustrate the utility of transcriptomics to investigating handicap models, suggest that classic models may not underlie the handicap mechanism, and indicate that novel emerging models involving different mediators and physiological systems should be examined.

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Introduction

One prominent theme in sexual selection research is to elucidate the evolution of extravagant secondary sexual ornaments in males and to elucidate the mechanisms that maintain signal honesty (Andersson 1994). Male ornaments are thought to act as quality signals to females during courtship to obtain the best mate and "good genes" for their offspring (Andersson 1982, 1994). Zahavi's handicap principle suggests that, in order to be reliable indicators of male quality, sexually selected ornaments must be costly to maintain and therefore incur a fitness cost (a "handicap") that cheater males cannot afford to pay (Zahavi 1975). Such a system would be evolutionarily stable (Maynard Smith 1974) and ensure that ornaments are honest quality indicators (Grafen 1990).

Various models have been proposed to explain the link between male quality and honest display of ornaments (Folstad and Karter 1992; Møller and Saino 1994; Olson and Owens 1998; von Schantz et al. 1999; Blount et al. 2003; Alonso-Alvarez et al. 2007; Hill 2011). One classic, still commonly invoked model is the immunocompetence handicap hypothesis (ICHH) by Folstad and Karter (1992). It proposes a trade-off between ornament intensity and immunocompetence (defined as the ability to mount an immune response upon exposure to an antigen) that is mediated by testosterone as both an enhancer of secondary sexual traits and a suppressor of immune function, primarily reflected in increased susceptibility to parasites (Hamilton and Zuk 1982). Therefore, extravagant ornaments advertise a high-quality immune system in males that are able to afford the immunosuppressive effect of testosterone.

Empirical studies have found evidence for a positive correlation between ornament intensity and immunocompetence (e.g. Møller et al. 1998, 1999; Peters et al. 2004; Andersson et al. 2006; Mougeot 2008) and a negative correlation with parasite load (e.g. Møller et al. 1999; Moore and Wilson 2002; Hill and Farmer 2005; Ottová et al. 2005). However, these correlations are not always supported (e.g. Hasselquist et al. 1999; Mougeot and Redpath 2004; Kurtz 2007) and indeed have been shown to be reversed in some cases (Roberts et al. 2004). As a means of testing the predictions of the ICHH, parasite loads and testosterone levels have been manipulated in a broad range of study systems. Reviews and meta-analyses support that experimental parasite challenges decrease testosterone levels (Boonekamp et al. 2008) and ornament intensity (Møller et al. 1999). Various recent studies also support the predicted effects of increased testosterone levels, i.e. enhanced ornaments (e.g. Mougeot et al. 2004; Deviche and Cortez 2005; Kurtz et al. 2007; Roberts et al. 2009b), increased parasite load (e.g. Mougeot et al. 2006; Seivwright et al. 2005; Deviche and Parris 2006; Cox and John-Alder 2007) and decreased immunocompetence (e.g. Mougeot et al. 2004; Oppliger et al. 2004; Deviche and Cortez 2005; Kurtz et al. 2007; Edler et al. 2011; Gil and Culver 2011). However, other studies have failed to find support for the latter (e.g. Buchanan et al. 2003; Mougeot and Redpath 2004; Oppliger et al. 2004; Ros et al. 2006; Alonso-Alvarez et al. 2009; Casagrande and Groothuis 2011; Fuxjager et al. 2011; Ezenwa et al. 2012), and overall support for the ICHH is equivocal at best (Roberts et al. 2004).

Given this mixed support, alternative models have been suggested. One common alternative is the oxidative stress handicap hypothesis (OSHH), which extends the ICHH beyond the immune system to include the antioxidant machinery. Specifically, it posits that the trade-off that ensures honest signalling predominantly involves resistance to oxidative stress rather than solely immunocompetence (von Schantz et al. 1999; Alonso-Alvarez et al. 2007). As in the ICHH, the underlying mechanism is testosterone dependent: Testosterone increases the metabolic rate (Buchanan et al. 2001; Oppliger et al. 2004; Alonso-Alvarez et al. 2007) and may consequently increase the production of reactive oxygen species (ROS) that may cause tissue damage or compromise an immune system response (Alonso-Alvarez et al. 2007). However, it may also simultaneously enhance the availability of antioxidant carotenoids, which may be either allocated to compensate for an excess in ROS production or deposited into carotenoid-dependent ornaments (Blas et al. 2006; McGraw et al. 2006). Therefore, only males with high-quality antioxidant machinery can afford the increased ROS production induced by high testosterone levels and still divert available antioxidant carotenoids into ornaments.

Correlational research on the OSHH has focussed on the relationship between ornament intensity and oxidative balance, rather than covariance with testosterone levels. Ornament intensity has been shown to predict oxidative DNA damage (Freeman-Gallant et al. 2011) and resistance to oxidative stress during an immune challenge (Pérez-Rodríguez et al. 2010). Experimental tests of the OSHH have largely supported the underlying theory: Increased testosterone levels increase oxidative stress (e.g. Alonso-Alvarez et al. 2007; Kurtz et al. 2007; Mougeot et al. 2009) but also antioxidant defences (e.g. Blas et al. 2006; McGraw et al. 2006; Mougeot et al. 2009) including carotenoids (e.g. Alonso-Alvarez et al. 2008; Martínez-Padilla et al. 2010). Manipulated parasite loads influence carotenoid and oxidative balance (e.g. Martínez-Padilla et al. 2007; Mougeot et al. 2007, 2009, 2010a), but there are inconsistent interactions between testosterone and parasites on ornament intensity across populations, suggesting a role of environmental context (Martínez-Padilla et al. 2010; Vergara et al. 2012a). Crucially, the relevancy of the OSHH for carotenoid-based ornaments depends on the assumptions of a testosterone-induced increase in the bioavailability of carotenoids (Blas et al. 2006) as well as a possible antioxidant function of carotenoids, though the latter is controversial (Costantini and Møller 2008; Martínez et al. 2009; Vinkler and Albrecht 2010).

All previous studies that have tested handicap models have used manipulations of hormone levels or parasite loads to capture subsequent morphological or physiological phenotype changes (Møller et al. 1999; Roberts et al. 2004). Examining morphology or physiology alone disregards the complex interactions of gene products and physiological systems in different tissues that underpin phenotype expression (Casagrande and Groothuis 2011; Ezenwa et al. 2012). Here, we provide an alternative, more fundamental approach based on the transcriptomic phenotype rather than the physiological phenotype. Gene transcription is the initial response to environmental stimuli and can therefore be argued to represent a crucial stage in the generation of phenotypic variation. In spite of confounding regulation of gene expression downstream of gene transcription (Brockmann et al. 2007), transcriptome profiling has been shown to be a useful approach to characterising the physiological response to experimental treatments and to identifying candidate genes (e.g. Pemberton et al. 2011; Thompson et al. 2011; Debes et al. 2012; Matzkin 2012). Here, we use microarrays to quantify the effect of experimentally increased testosterone levels on transcription levels of genes involved in immune response and ROS defence. The resulting patterns of up- and down-regulation of these genes in different tissues can then be used to assess whether the transcriptomic response conforms to the expectations of the ICHH or OSHH.

Our focal study system is the red grouse (Lagopus lagopus scoticus Latham) and its main gastro-intestinal nematode parasite Trichostrongylus tenuis Mehlis. This parasite exhibits a direct life cycle. Infective larvae are ingested with heather shoots and reside in the caecal mucosa where adult worms cause marked damage, resulting in weight loss, poor overall condition and potentially compromised survival and lowered fecundity (Watson et al. 1987; Hudson et al. 1992; Delahay et al. 1995). Prevalence of infection in grouse populations is greater than 90 % (Wilson 1983), and although grouse mount an immune response, they cannot acquire immunity and thus bear chronic worm burdens for life (Shaw and Moss 1989). The grouse-nematode system has previously been used to test the ICHH and OSHH using morphological and physiological assays. Folstad and Karter (2005) and Mougeot et al. (2006) found that male grouse implanted with testosterone retained a higher parasite load after a standardised parasite challenge than control grouse. The implanted grouse also developed larger ornaments than control grouse and therefore supported both assumptions of the ICHH (Mougeot et al. 2006). More recently, a factorial field experiment involving experimentally manipulated testosterone levels and parasite loads was carried out to examine the interactions of testosterone and parasites on ornament intensity and oxidative balance (Mougeot et al. 2009, 2010a; Martínez-Padilla et al. 2010). Increased testosterone levels enhanced ornaments and increased oxidative damage as well as circulating antioxidants (Mougeot et al. 2009). Removing parasites also enhanced ornaments, and increased parasite loads and testosterone levels together increased oxidative damage, suggesting an oxidative stress handicap in agreement with the OSHH (Mougeot et al. 2009, 2010a).

Based on the same factorial field experiment presented in Mougeot et al. (2009), Webster et al. (2011b) recently conducted a microarray experiment to characterise the interactive effects of parasite loads and testosterone levels on the caecum transcriptome. By comparing transcription levels of genes across treatment contrasts with different parasite loads and testosterone levels, they identified a set of genes that were up-regulated when parasites were present and subsequently down-regulated when testosterone levels were increased. Here, we use the same raw microarray data that were generated by Webster et al. (2011b) but employ a conceptually different analytical approach on a different set of data. We a priori target genes related to immune system function and response to ROS production, and then examine transcription changes of these genes in caecal, spleen and liver tissues of testosterone-treated grouse compared to control grouse across three parasite load treatments. These three tissues represent both local and global physiological states, as they include the site of parasitic infection and organs centrally important for homeostasis (Mougeot and Redpath 2004; Bankoti and Stäger 2012).

Under the ICHH, increased testosterone levels are expected to compromise the immune system, identifiable from changes in the underlying transcriptomic response. This can be mediated either directly through downregulation of immune genes or up-regulation of genes that negatively regulate components of the immune system or compensate for down-regulated components elsewhere (Schmid-Hempel 2003). We therefore hypothesise that immune genes become significantly up- or down-regulated in grouse with experimentally increased testosterone levels compared to grouse with natural levels. Similarly, central to the OSHH, increased testosterone levels are expected to increase ROS production and a concomitant defence response, i.e. antioxidant mobilisation, to combat oxidative stress. In consequence, genes involved in ROS response, particularly in mobilising antioxidants, should become upregulated. Alternatively, genes that negatively regulate antioxidant mobilisation may become down-regulated. We therefore hypothesise that ROS response genes become treated grouse and control grouse in relevant genes would be inconsistent with the respective handicap models on the transcriptomic level.

Materials and methods

Field experiment and microarray assays

Details of the field experiment and microarray assays are described in Mougeot et al. (2009) and Webster et al. (2011b). Briefly, a factorial field experiment involving manipulation of testosterone levels and parasite loads in 40 male grouse was carried out in Edinglassie (Scotland) and Catterick (England) moors in autumn 2006. The birds were implanted with silastic tubes containing either testosterone proprionate (treatment T) or saline (treatment I) and were either administered with the anthelmintic levamisole hydrochloride (treatment A), approximately 5,000 L3 *T. tenuis* larvae following anthelmintic treatment (treatment P) or not manipulated for parasite load at all (treatment N). The treatments were shown to be effective for the duration of the experiment (Mougeot et al. 2009; Webster et al. 2011b).

Caecum, spleen and liver samples were taken three to four weeks after experimental treatment, and RNA was extracted for gene transcription analysis (Webster et al. 2011b). A bespoke grouse microarray was constructed from 9,600 clones, representing 5,925 unique sequence transcripts (contigs), obtained by standard cDNA libraries and suppressive subtractive hybridisation (SSH) from grouse with high versus low parasite loads (Webster et al. 2011a). All possible combinations (hereafter referred to as contrasts) of the six treatment groups were assayed on microarrays in a loop design using RNA extracts of sufficient quality, i.e. with an RNA integrity number (RIN) greater than 8.5 following Agilent 2100 Bioanalyzer analysis (Webster et al. 2011b). High-quality RNA extracts were available for all 40 birds from spleen and liver tissue, but only for 27 birds from caecal tissue due to quality control failures (RIN<8.5) (Table 1).

Data analysis

In order to examine the assumptions of the ICHH and OSHH, we first identified those contigs on the microarray that were involved in immune system function or ROS response before we analysed microarray data. We achieved this by interrogating all contig clone sequences against protein databases to identify gene products and then retrieving GENEONTOLOGY (GO) annotations (The Gene Ontology **Table 1** Sample sizes (RNA extracts) in microarray experiments for three tissues and six treatment groups, comprising male grouse with either increased (T) or natural (I) testosterone levels, and either anthelmintic-treated (A), natural (N) or increased parasite loads (P)

	Caecum	Spleen	Liver
TA	4	9	9
IA	3	9	9
TN	4	4	4
IN	3	4	4
TP	7	7	7
IP	6	7	7
Total	27	40	40

Consortium 2000) to characterise each gene product in terms of biological process, molecular function and cellular component. We performed all database interrogations in December 2011 using the BLAST2GO framework (Conesa et al. 2005; Conesa and Götz 2008). Contrary to the original annotation strategy by Webster et al. (2011b), we employed a hierarchical search strategy. Firstly, all contigs were queried against the SWISSPROT database, using the NCBI Basic Local Alignment Search Tool (BLAST; http://www.ncbi.nlm. nih.gov/blast/Blast.cgi) with an expectation value cutoff of 10^{-10} . Then, those contigs without a BLAST result were queried against the CHICKEN GENBANK PROTEIN database (gp/ 9031.10804//gga genbank prot), and subsequent failures were finally queried against the ZEBRAFINCH GENBANK PROTEIN database (gp/59729.12898//tg genbank prot). Each identified gene product was mapped and annotated $(10^{-10} \text{ expecta-})$ tion cut-off) for GO identifiers, and GO annotations were augmented by querying the ANNEX database (Myhre et al. 2006). GO annotations were then screened for GO:0002376 immune system process (Go level 2), GO:0006979 response to oxidative stress (Go level 4) and GO:0016209 antioxidant activity (GO level 2), including parent terms in all cases. The GO term response to oxidative stress covers gene products involved in all physiological processes that respond to exposure to ROS, whereas antioxidant activity covers antioxidant enzymes, e.g. superoxide dismutase, catalase and peroxidase.

This strategy identified 326 contigs in total. In order to investigate the transcriptomic response of these contigs to testosterone, we retrieved microarray data from the treatment contrasts TA vs. IA, TN vs. IN and TP vs. IP, representing the transcriptomic response to increased testosterone levels T (versus natural levels I) under anthelmintic treatment A, natural chronic parasite infection N and parasite challenge P, respectively. For each contig and contrast, we retrieved the relative difference in transcriptomic response between treatments (fold change) and the associated p value for the null hypothesis of no differential response (see Webster et al. (2011b) for technical details). As a means of

correcting for false rejections of the null hypothesis among these 326 contigs, we adjusted the p value of each contig to account for the false discovery rate (FDR) (Benjamini and Hochberg 1995) using the R package fdrtool (Strimmer 2008). This package calculates FDR adjusted p values (=q values) using a method that keeps the false nondiscovery rate minimal. We performed FDR correction for each contrast separately and used the p value distribution of the 326 identified contigs only rather than all contigs on the microarray as done previously (Webster et al. 2011b). Contigs were identified as significantly differentially regulated when q < 0.05. We estimated the statistical power of the microarray data to detect significantly differentially regulated contigs by calculating the expected false discovery rate (Efdr) from the p value distributions of all 5,925 contigs using the R package locfdr (Efron 2007). An Efdr below 0.4 suggests high power, i.e. a high probability of detecting truly significant p values in the dataset.

Results

Of the 5,925 unique DNA contigs on the microarray, 1,864 (31.5 %) yielded a BLAST hit after cumulative interrogation of three protein databases, and of these 1,781 were successfully GO annotated (Table 2). Of these, 282 contigs were annotated with GO:0002376 immune system process (hereafter: immune genes) and 65 with either GO:0006979 response to oxidative stress (64 contigs) or GO:0016209 antioxidant activity (13 contigs) (hereafter: ROS response genes). A number of contigs were annotated with both immune system and ROS response categories, resulting in 326 identified contigs in total. The top BLAST hit descriptions for all identified contigs are presented in Tables S1 and S2. The set of immune genes covered various prominent constituents of the immune system (e.g. complement system, T-cell surface complexes, immunoglobulins and ubiquitin/proteasome-related complexes), whereas the set of ROS response genes covered a range of prominent antioxidant enzymes (e.g. catalase, superoxide dismutase, glutathione S-transferase, peroxiredoxins and glutaredoxins).

The effect of testosterone treatment on transcription regulation of these focal genes was weak overall, particularly so in liver and spleen tissue (Table 3). There was no significant effect at all in liver tissue, as the distribution of p values was near uniform in all contrasts, resulting in FDR-corrected p values of q >> 0.05 for all contigs. This was also the case in spleen tissue, apart from the TN vs. IN contrast, in which three immune genes were significantly differentially regulated. Accordingly, statistical power was estimated as considerably low in these tissues (Table 3). In comparison, numbers of significantly differentially regulated genes and estimated power were considerably higher in caecal tissue. The changes in caecal gene transcription in each contrast are illustrated as volcano plots that plot log₂ fold changes (measure of biological significance) against $-\log_{10} q$ (measure of statistical significance) (Fig. 1).

Across the three tissues, six immune genes were significantly down-regulated and one was significantly up-regulated in grouse with natural chronic parasite infection (TN vs. IN), whereas 17 were significantly down-regulated and three were significantly up-regulated in grouse with a parasite challenge (TP vs. IP) (Table 4). These contigs represent not only some prominent immune-system components, such as immunoglobulin chains, cell-surface proteins and complement factors, but also components involved in cell proliferation and motility, e.g. myosin, gelsolin and coronin (Table S3). A single ROS response gene (actin) was significantly up-regulated in the TN vs. IN contrast, whereas four ROS response genes were significantly down-regulated in the TP vs. IP contrast, none of which were prominent antioxidant enzymes (Table S4). No significantly differentially regulated immune genes and ROS response genes were found in grouse treated with an anthelmintic (TA vs. IA contrast).

The overall transcriptomic response to testosterone in caecum, extended beyond the focal immune and ROS response genes, covered a wide range of physiological processes in the three treatment contrasts under examination. A number of

 Table 2
 Gene product identification results after hierarchical BLAST interrogation of 5,925 contigs against three databases (SWISSPROT, then CHICKEN GENBANK PROTEIN, then ZEBRAFINCH GENBANK PROTEIN) followed by GENEONTOLOGY (GO) annotation

	SWISSPROT	CHICKEN	ZEBRAFINCH	Total	%
Protein database hits					
BLAST hits	1,801	60	3	1,864	31.5
GO hits	1,746	33	2	1,781	30.1
GO term hits					
GO:0002376 immune system process	280	2	0	282	4.8
G0:0006979 response to oxidative stress+G0:0016209 antioxidant activity	65	0	0	65	1.1

The total numbers of hits are also given as the percentage of the total dataset (5,925 contigs)

Table 3 Significantly differentially regulated contigs (q < 0.05) in response to testosterone (increased testosterone levels T compared to natural levels I) in three tissues, three parasite treatment groups (anthelmintic A, natural chronic infection N and parasite challenge P) and selected GO terms

	TA vs. IA			TN vs. IN			TP vs. IP		
	Caecum	Spleen	Liver	Caecum	Spleen	Liver	Caecum	Spleen	Liver
All go terms	18	0	0	18	3	1	103	0	0
GO:0002376 immune system process	0	0	0	4	3	0	20	0	0
GO:0006979 response to oxidative stress+ GO:0016209 antioxidant activity	0	0	0	1	0	0	4	0	0
Power estimate (Efdr)	0.42	0.58	0.69	0.49	0.60	0.62	0.28	0.58	0.63

Efron's expected false discovery rate (Efdr) is given for each tissue/contrast combination as an estimate of power (Efdr ≤ 0.4 indicates high power) (Efron 2007)

genes were significantly up- or down-regulated in either contrast (Fig. 1, Table 3), and the most frequent level-2 and level-3 GO terms in the biological process ontology are presented in Fig. 2.

Discussion

We here provide a novel perspective to investigating handicap models in sexual selection by studying the transcriptomic rather than the physiological phenotype. Specifically, we studied the transcriptomic response of red grouse to testosterone in three key tissues in order to investigate the main assumptions of the immunocompetence handicap hypothesis (ICHH) and oxidative stress handicap hypothesis (OSHH). We hypothesised that immune genes (ICHH) or genes involved in response to ROS (OSHH) become significantly up- or down-regulated in grouse treated with testosterone implants compared to control grouse. Only a small proportion of identified genes were significantly differentially regulated, with inconsistencies across tissues and parasite treatment groups. The OSHH results in particular are somewhat at odds with the physiological response to testosterone of the same birds, which was previously reported to support the OSHH (Mougeot et al. 2009). We discuss observed discrepancies between the transcriptomic and physiological phenotype and highlight potential pitfalls in the transcriptomic approach.

Transcriptomic evidence for the ICHH and OSHH

The main assumption of the ICHH is that testosterone simultaneously acts as an enhancer of male secondary traits, such as epigamic ornaments, and a suppressor of immunocompetence (Folstad and Karter 1992). In red grouse, experimentally increased testosterone levels have been shown to enhance ornaments (Mougeot et al. 2004, 2009), increase parasite load (Seivwright et al. 2005; Mougeot et al. 2006) and decrease T-cell-mediated immunity and overall condition (Mougeot et al. 2004). Given these findings, some transcriptomic evidence of a compromised immune system would be expected in testosterone-treated grouse, either directly through down-regulation of immune genes or upregulation of genes that negatively regulate components of the immune system or compensate for down-regulated components (Schmid-Hempel 2003). The volcano plots for immune genes in caecal tissue were overall asymmetrical with some bias towards (non-significant) down-regulation. Most of the significantly differentially regulated immune genes were indeed down-regulated, but the heavy-chain component of ferritin (also up-regulated in spleen) and the coxsackievirus and adenovirus receptor were up-regulated by testosterone in grouse with parasites present. Both of these compounds may have immunosuppressive properties (Carson and Chapman 2001; Gray et al. 2001), so up-regulation by testosterone is fully compatible with the expectations. As such, the transcriptomic response in caecal tissue of grouse with a parasite challenge may provide transcriptomic evidence for an immunosuppressive function of testosterone according to the ICHH. However, the absence of a transcriptomic response in liver and (in most cases) spleen tissue, the weak response in grouse with natural parasite loads and the lack of response in grouse treated with an anthelmintic weaken the evidence and are grounds for caution.

We further examined the main assumption of the OSHH that increased testosterone levels increase ROS production and consequently enhance the physiological response to ROS (Alonso-Alvarez et al. 2007). ROS production is expected to result in mobilisation of innate antioxidant defences in an attempt to regain a positive oxidative balance (Finkel and Holbrook 2000), even if this cannot be attained due to a low-quality oxidation defence system. Testosterone-treated grouse have been shown to experience an increased total antioxidant status and a concomitant increase in oxidative stress handicap might be operating (Mougeot et al. 2009). Therefore, an up-regulation of ROS response genes (i.e. antioxidants) or a down-regulation of genes that negatively



Fig. 1 Transcriptomic response of caecal tissue to increased (T) contrasted with natural testosterone levels (I) in three parasite treatment groups (anthelmintic A, natural chronic infection N and parasite challenge P). Volcano plots relate biological significance (fold change) to statistical significance (FDR adjusted q value) of gene transcription

changes. Each *dot* represents one contig, GO-annotated with GO:0002376 *immune system process* (*row A*), GO:0006979 *response* to oxidative stress or GO:0016209 antioxidant activity (*row B*), or any GO term (*row C*). Black dots are statistically significant (q<0.05; *dashed grey line* indicates cut-off)

regulate antioxidant mobilisation would be expected in testosterone-treated grouse. However, none of the identified antioxidant genes were significantly differentially regulated in any contrast. Instead, a component of the cytoskeleton (actin) was significantly up-regulated in grouse with natural parasite levels, and four genes that do not possess anti-oxidant properties were significantly down-regulated in grouse with a parasite challenge. The up-regulation of actin may indicate a compensatory response to oxidative damage to the cytoskeleton (Farah et al. 2011), but these results provide no evidence for the OSHH under our initial hypothesis that anti-oxidants become up-regulated.

A possible explanation for the observed lack of enhanced ROS response may be that there was no increased ROS production and therefore no concomitant ROS response, which would be inconsistent with the main assumption of

Tissue	Fold	Fold change		Gene product (BLAST description)	GO terms			Accessions	
	TA	TN	ТР		С	F	Р	Protein	Contig
GO:000237	76 immı	ine system	process						
Caecum	-	-	-1.79	Ubiquitin-40S ribosomal protein S27a	5	2	35	P79781	GW704812
Caecum	-	-	-1.63	Ig lambda chain V-1 region	3	1	2	P04210	GW704581
Caecum	-	-1.80	-1.40	Ig lambda chain V-1 region	2	2	2	P04210	GW700881
Caecum	-	—	-1.16	Caspase recruitment domain-containing protein 11	5	4	22	Q9BXL7	GW704745
Caecum	_	-	-1.14	HLA class II histocompatibility antigen gamma chain	5	2	12	P04233	GW704842
Caecum	-	-	-1.02	T-cell surface glycoprotein CD3 delta chain	1	2	6	Q764N2	GW705191
Caecum	-	_	-0.90	Myosin-11	24	9	41	P10587	GW706306
Caecum	_	-	-0.83	Dual specificity protein phosphatase 1	5	4	33	P28562	GW706824
Caecum	_	_	-0.82	Myosin-11	24	9	41	P10587	GW703178
Caecum	_	_	-0.77	Complement factor I	5	1	4	Q61129	GW703417
Caecum	_	_	-0.77	Kininogen-1	2	2	12	P01044	GW704416
Caecum	_	-1.16	-0.69	Gelsolin	12	6	27	O93510	GW704663
Caecum	_	_	-0.65	Ras-related C3 botulinum toxin substrate 2	16	15	62	P15153	GW705819
Caecum	_	_	-0.64	Immunoglobulin J chain	2	1	1	P01592	GW699715
Caecum	_	_	-0.59	YTH domain family protein 2	0	0	1	Q4R5D9	GW705307
Caecum	_	_	-0.58	Lymphocyte antigen 6E	3	0	3	Q90986	GW704464
Caecum	-	_	-0.40	72 kDa type IV collagenase	8	8	46	Q90611	GW705867
Caecum	_	—	0.41	Coxsackievirus and adenovirus receptor	4	1	8	P78310	GW706459
Caecum	_	—	0.48	Coxsackievirus and adenovirus receptor homolog	7	2	11	Q9R066	GW700528
Caecum	-	_	0.56	Ferritin heavy chain	3	3	12	P08267	GW700124
Caecum	-	-1.30	_	Myosin-9	24	9	41	P14105	GW704927
Caecum	_	-0.72	_	Coronin-1C	6	4	10	Q9ULV4	GW705584
Spleen	-	-1.42	_	Elongator complex protein 1	4	6	5	O95163	GW701060
Spleen	_	-0.94	—	Plastin-2	10	5	13	P13796	GW702767
Spleen	_	1.01	—	Ferritin heavy chain	3	3	12	P08267	GW699682
GO:000697	79 respo	onse to oxi	dative stre	ess+G0:0016209 antioxidant activity					
Caecum	_	_	-0.83	Dual specificity protein phosphatase 1	5	4	33	P28562	GW706824
Caecum	_	_	-0.76	Serine/threonine-protein kinase Sgk1	7	8	25	Q6U1I9	GW705373
Caecum	-	_	-0.61	Bile salt export pump	7	7	20	095342	GW701767
Caecum	-	_	-0.40	72 kDa type IV collagenase	8	8	46	Q90611	GW705867
Caecum	_	1.41	_	Actin	6	2	15	P60009	GW699904

Table 4 Characterisation of significantly differentially regulated contigs (q < 0.05) in response to testosterone

Each row represents one contig and provides statistically significant fold changes in three parasite treatment groups (anthelmintic TA, natural chronic infection TN and parasite challenge TP). The gene product description is given alongside the number of GO annotations in cellular component (C), molecular function (F) and biological process (P) ontologies, and database accessions for the protein SWISSPROT hit and the original contig sequence (GENBANK EST)

the OSHH and contradict a presumed key metabolic effect of testosterone (Buchanan et al. 2001; Oppliger et al. 2004; Muehlenbein and Bribiescas 2005; Alonso-Alvarez et al. 2007). This is unlikely to be the case because oxidative damage was increased in testosterone-treated grouse, suggesting that an excess of ROS was present (Mougeot et al. 2009). Alternatively, an enhanced ROS response might not have been necessary because a sufficient testosteroneinduced increase in carotenoid availability (Blas et al. 2006; McGraw et al. 2006; Alonso-Alvarez et al. 2008) fully compensated for excess ROS (but see Costantini and Møller 2008; Martínez et al. 2009; Vinkler and Albrecht 2010). This may be possible, given that the GO terms we used to select genes may only cover innate ROS defenses rather than dietary carotenoids. Indeed, testosterone enhanced comb redness and increased circulating carotenoid levels in one of the two study populations (Martínez-Padilla et al. 2010). However, the birds still experienced oxidative damage (Mougeot et al. 2009), suggesting that ROS could not be fully compensated for. Therefore, an enhanced



Fig. 2 Most frequent level-2 and level-3 GO annotations in the biological process ontology of all contigs significantly differentially regulated in response to testosterone

mobilisation of innate ROS defences would be expected and is indeed evidenced by an increase in the birds' total antioxidant status (Mougeot et al. 2009), even when carotenoid levels were not increased (Martínez-Padilla et al. 2010). An increase in innate ROS defences, however, is not reflected in the transcriptomic data, suggesting that caecum, spleen and liver may not be the ideal tissues for investigating oxidative stress in red grouse. Nevertheless, four genes that are involved in response to oxidative damage, rather than antioxidant defence, were significantly down-regulated by testosterone in the presence of a parasite challenge, suggesting that testosterone-induced suppression of oxidative damage responses might be an alternative mechanism.

In conjunction, there is little transcriptomic evidence for the two focal handicap models in red grouse. However, caution may be warranted because the weak transcriptomic response may potentially reflect limitations in our study design and data analysis rather than evidence to accept the null hypothesis. The experimental treatments of the underlying field study were certainly effective, given that testosterone implants caused a two- to threefold increase in testosterone levels, and worm counts of parasite purged and parasite challenged birds differed by a factor of six (detailed in Mougeot et al. 2009; Webster et al. 2011b). Moreover, the microarray data for caecum have been validated by quantitative PCR, using independent RNA extracts, and showed congruence even for contrasts with low sample sizes such as TA vs. IA (Webster et al. 2011b). Nevertheless, statistical power to detect significantly differentially regulated genes might have been compromised by relatively low sample sizes in some treatment groups (particularly in caecum). We note, however, that in spite of larger sample sizes, spleen and liver yielded virtually no significantly differentially regulated genes, suggesting that this was not due to low power. Crucially, in spite of potential sample size limitations, some significant transcriptomic response to testosterone treatment was detected in two out of three tissues. The response was strongest in caecal tissue, but nearly absent in spleen and entirely absent in liver tissue, in spite of their immunological and homeostatic importance (Mougeot and Redpath 2004; Bankoti and Stäger 2012). This inconsistency highlights that tissue choice is critical in addressing questions that involve the transcriptome. Although studying various tissues allows for identifying tissues involved in response to a treatment, restriction to only a few tissues may miss those tissues most relevant to the response of interest. While the caecum is the site of parasite infection in grouse and should therefore be the best site for investigating immune system processes (Watson et al. 1987; Shaw and Moss 1989; Webster et al. 2011b), oxidative stress processes may have been better represented in tissues more sensitive to oxidative stress, such as brain or heart (Floyd 1999; Bayeva and Ardehali 2010).

Moreover, we aimed to minimise the occurrence of false positives throughout the analysis process as we were interested in testing specific predictions of gene transcription rather than merely characterising a treatment effect. We therefore employed moderately stringent criteria during BLAST searches and GO annotation. As a consequence, only 30.1 % of all original contigs were GO annotated and only few contigs were annotated with GO terms of interest. This sample may have missed potentially relevant genes that were not captured by the microarrays or were not annotated. Nevertheless, the sample included a wide range of prominent components of the immune system and the antioxidant machinery. Assuming that these complex physiological systems are underpinned by a large number of genes, particularly if they are assumed to be a major component of condition as signalled by ornaments (Rowe and Houle 1996), even a limited gene sample that may miss some relevant genes can be assumed to be appropriate for investigating handicap mechanisms.

We used FDR correction of p values to correct for multiple significance testing and chose a stringent threshold of

5 % FDR to ensure a low rate of false positives, as is common in microarray experiments (e.g. Thompson et al. 2011; Debes et al. 2012; Matzkin 2012). In order to ascertain to what extent the numbers of identified discoveries depended on the set of p values chosen to calculate q values, we re-calculated q values for the 326 contigs of interest from the p values of all 5,925 contigs, thus providing a more conservative approach to FDR. This approach had no effect on results in spleen and liver, and in caecum it removed only one ROS response contig each from the TN vs. IN and TP vs. IP contrasts, and one and two immune system contigs from the TN vs. IN and TP vs. IP contrasts, respectively. Given that both approaches produced very similar results, we are confident that our statistical approach is robust. We note that using a higher FDR threshold, or even uncorrected p values rather than FDR, to identify significantly differentially regulated contigs naturally would have resulted in more identified contigs in all cases, but also in an unacceptably large number of false positives.

Apart from these statistical caveats, there are two potential limitations associated with a transcriptomic approach and selection of genes through GENEONTOLOGY annotations. Firstly, the transcriptome does not necessarily predict the proteome at the final stage of gene expression. Gene transcription is only the initial step of gene expression, and various posttranscriptional regulatory mechanisms are recognised (Day and Tuite 1998; Brockmann et al. 2007). These mechanisms confound the assumed simple relationship between mRNA and final protein, so that only as little as 20-40 % of variation in protein concentration may be accounted for by the variation in mRNA levels (Brockmann et al. 2007). It is possible that the previously published physiological patterns in field studies on grouse are achieved predominantly downstream of gene transcription. However, this is somewhat difficult to reconcile with studies that have demonstrated the utility of transcriptomic profiling to characterising gene expression and identifying candidate genes (e.g. Pemberton et al. 2011; Thompson et al. 2011; Debes et al. 2012; Matzkin 2012). Secondly, an assumption of our study design is that all contigs selected for either immune system process or response to oxidative stress respond to testosterone treatment as a result of these very processes. However, given that a gene is rarely involved in a single biological process only, an entirely separate process may be responsible for a change in gene expression in response to testosterone. GENEONTOLOGY annotation reflects this complexity by allowing multiple annotation of a single gene product (The Gene Ontology Consortium 2000). In this study, only a small proportion of significantly differentially regulated genes were annotated for a single biological process alone. Nevertheless, it is worth noting that, for example, not all immune genes with a single GO annotation for biological process responded significantly to testosterone treatment, contrary to expectations based on the ICHH.

Wider implications for handicap models

In summary, our results indicate that the immunosuppressive effect of testosterone may at least partially be effected by gene-transcription changes in caecum and (to a much smaller extent) spleen under parasite load. This may be consistent with the ICHH and the results of previous grouse field studies. Contrarily, there is no transcriptomic evidence for an increase in antioxidant activity in caecum, spleen and liver, which is inconsistent with the OSHH and previous field study results. However, we have shown that testosteroneinduced suppression of a response to oxidative damage, rather than an increase in ROS defences, might confer the oxidative stress handicap, and we encourage to consider this alternative mechanism in future studies.

In spite of some weak transcriptomic support in the red grouse system, the broader literature suggests that the ICHH and OSHH are probably not always biologically realistic. The simplistic model of the original ICHH has been considerably questioned since a meta-analysis (Roberts et al. 2004) found no effect of testosterone on immunocompetence when controlled for multiple studies on the same species. Since 2004, various further studies have tested the ICHH, but positive results (e.g. Deviche and Cortez 2005; Deviche and Parris 2006; Cox and John-Alder 2007; Kurtz et al. 2007; Casagrande and Groothuis 2011; Edler et al. 2011; Gil and Culver 2011) are contrasted with negative results (e.g. Oppliger et al. 2004; Ros et al. 2006; Alonso-Alvarez et al. 2009; Casagrande and Groothuis 2011; Fuxjager et al. 2011; Ezenwa et al. 2012). Similarly, although the main mechanism of the OSHH is often empirically supported (e.g. Alonso-Alvarez et al. 2007, 2008; Mougeot et al. 2007, 2009, 2010a; Pérez-Rodríguez et al. 2010; Freeman-Gallant et al. 2011), environmental context may play a major role (Martínez-Padilla et al. 2010; Vergara et al. 2012a; Vergara and Martínez-Padilla 2012). Therefore, alternative models may be required to explain Zahavian handicaps.

A prominently discussed alternative mechanism is the physiological stress hypothesis (Møller and Saino 1994), in which the trade-off between carotenoid ornaments and body condition is not mediated by testosterone but is dependent on stress hormone levels, such as corticosterone, a glucocorticoid stress hormone involved in the mobilisation of energy sources and induction of characteristic stress behaviour (Buchanan 2000; Sapolsky et al. 2000). Environmental stressors may cause physiological stress that becomes manifested by increased corticosterone levels (e.g. Dahl et al. 2012). Chronically elevated corticosterone levels may be immunosuppressive (e.g. Berger et al. 2005) and corticosterone

may correlate with testosterone levels (Besedovsky et al. 1986; Evans et al. 2000; Casto et al. 2001; Buchanan et al. 2003; Owen-Ashley et al. 2004; Mateos 2005). Testosterone and corticosterone may interact to, for example, have an immuno-enhancing effect, which has been shown in *Taeniopygia guttata* (zebra finch), although this incurred substantial physiological costs (Roberts et al. 2007, 2009a). Recent correlational evidence for a stress-mediated handicap mechanism comes from research on human males, whose facial characteristics correlate with cortisol (human equivalent of corticosterone) levels and immunocompetence, which itself is affected by an interaction between testosterone and cortisol (Moore et al. 2011; Rantala et al. 2012).

In red grouse, variation in corticosterone levels has been shown to explain variation between individuals in parasite load and testosterone-induced ornament enhancement (Bortolotti et al. 2009). Similarly, more recent work suggests that handicap mechanisms are context-dependent and ornaments signal condition reliably only in adverse environmental conditions, i.e. under stress (Vergara et al. 2012a, b; Vergara and Martínez-Padilla 2012). In our study, more immune genes and ROS response genes were significantly differentially regulated in the presence of parasites (particularly of a parasite challenge) than in absence. This effect may be partially caused by higher power caused by larger sample size in the parasite challenge group. However, even between contrasts with identical sample sizes (TA vs. IA and TN vs. IN), more genes responded under natural parasite infection than under anthelmintic treatment. This could be speculated to indicate a stressmediated handicap mechanism in grouse as a testosteronedependent handicap mechanism may be more pronounced in the presence of a stressor (i.e. parasites) (Bortolotti et al. 2009; Vergara et al. 2012a, b). Future work could test this more explicitly using transcriptomic profiling in response to manipulation of testosterone and corticosterone levels to disentangle the interaction between these hormones.

As an alternative perspective, Hill (2011) put forward a fundamentally different view on condition-dependent characters and signals. He defines condition as the ability to maintain vital cellular processes during environmental challenges, determined by a combination of genotypic, epigenetic and somatic states, and proposes to decouple honest signalling from fitness costs and rather link it with the functionality of cellular processes. This view is intuitively elegant, but its examination is considerably constrained by our understanding and ability to disentangle biochemical processes. Currently, the consensus view on handicaps from empirical studies is that the main mediator is probably corticosterone or an interaction of several hormones and systems (e.g. Costantini et al. 2008; Bortolotti et al. 2009; Casagrande and Groothuis 2011; Moore et al. 2011; Ezenwa et al. 2012; Rantala et al. 2012). Driven by the recognition of physiological stress as a major factor in trade-offs (Bortolotti et al. 2009; Mougeot et al. 2010b), and given that a simple universal relationship between physiological systems and life history may not exist (Versteegh et al. 2012), evolutionary ecology should focus on the individual rather than species or higher taxa because each individual will resolve trade-offs in its own way, given genetic makeup, life history, energetic balance and environmental context (Bortolotti et al. 2009; Martínez-Padilla et al. 2010; Vergara et al. 2012a; Vergara and Martínez-Padilla 2012; Versteegh et al. 2012).

This work has provided a novel perspective on investigating the assumptions of handicap models and our case study on grouse transcriptomic data points to alternative handicap models and ideas currently discussed in the literature. Moreover, it has highlighted the limitations in experimental design and analysis that need to be overcome if transcriptomic approaches are to be used to disentangle the mechanisms behind Zahavian handicaps.

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Conflicts of interest The authors declare that they have no conflicts of interest.

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