Genomic conflicts and sexual antagonism in human health: Insights from oxytocin and testosterone

Mökkönen, Mikael; Crespi, Bernard J.

Mökkönen, M., & Crespi, B. J. (2015). Genomic conflicts and sexual antagonism in human health: Insights from oxytocin and testosterone. Evolutionary Applications, 8(4), 307-325. https://doi.org/10.1111/eva.12244

All material supplied via JYX is protected by copyright and other intellectual property rights, and duplication or sale of all or part of any of the repository collections is not permitted, except that material may be duplicated by you for your research use or educational purposes in electronic or print form. You must obtain permission for any other use. Electronic or print copies may not be offered, whether for sale or otherwise to anyone who is not an authorised user.
Genomic conflicts and sexual antagonism in human health: insights from oxytocin and testosterone

Mikael Mokkonen1,2 and Bernard J. Crespi1

1 Department of Biological Sciences, Simon Fraser University, Burnaby, BC, Canada
2 Department of Biological and Environmental Science, University of Jyväskylä, Jyväskylä, Finland

Keywords
genomic imprinting, kinship theory, parental antagonism, parent–offspring conflict, sexual antagonism, sexual conflict.

Abstract
We review the hypothesized and observed effects of two of the major forms of genomic conflicts, genomic imprinting and sexual antagonism, on human health. We focus on phenotypes mediated by peptide and steroid hormones (especially oxytocin and testosterone) because such hormones centrally mediate patterns of physical and behavioral resource allocation that underlie both forms of conflict. In early development, a suite of imprinted genes modulates the human oxytocinergic system as predicted from theory, with paternally inherited gene expression associated with higher oxytocin production, and increased solicitation to mothers by infants. This system is predicted to impact health through the incompatibility of paternal-gene and maternal-gene optima and increased vulnerability of imprinted gene systems to genetic and epigenetic changes. Early alterations to oxytocinergic systems have long-term negative impacts on human psychological health, especially through their effects on attachment and social behavior. In contrast to genomic imprinting, which generates maladaptation along an axis of mother–infant attachment, sexual antagonism is predicted from theory to generate maladaptation along an axis of sexual dimorphism, modulated by steroid and peptide hormones. We describe evidence of sexual antagonism from studies of humans and other animals, demonstrating that sexually antagonistic effects on sex-dimorphic phenotypes, including aspects of immunity, life history, psychology, and behavior, are commonly observed and lead to forms of maladaptation that are demonstrated, or expected, to impact human health. Recent epidemiological and psychiatric studies of schizophrenia in particular indicate that it is mediated, in part, by sexually antagonistic alleles. The primary implication of this review is that data collection focused on (i) effects of imprinted genes that modulate the oxytocin system, and (ii) effects of sexually antagonistic alleles on sex-dimorphic, disease-related phenotypes will lead to novel insights into both human health and the evolutionary dynamics of genomic conflicts.

Introduction
Genomic conflicts are expected to be frequent sources of phenotypic maladaptation, as the optima targeted by different genomic agents may be more or less displaced from the values that maximize survival and reproduction at the level of organisms themselves (Burt and Trivers 2006). For humans, such displacements from fitness maximizing trajectories are expected to frequently manifest as deviations from health, given that alleles associated with greater disease susceptibility are costly for fitness through reduced survival or impaired reproduction. Any such deviations, moreover, should be structured by the nature and directions of genomic conflicts (Frank and Crespi 2011) such that better understanding of these conflicts, and their ongoing natures or potential resolutions, provides important insights into how to understand, prevent, and treat conflict-relevant diseases.

In this article, we provide a review and synthesis of two major forms of genomic conflicts, genomic imprinting, and
sexual antagonism, with regard to their impacts on hormonally mediated, health-related human phenotypes. These two forms of genomic conflict are fundamentally similar in their expected impacts on health and disease, because they both centrally involve different fitness optima for two parties, and they both engender conflicts over optima that can involve the evolution of conflict mechanisms and evolutionary escalation; in both cases, disease-related impacts may ensue due to the deviations from optima and dysregulation of evolved conflict systems (Crespi 2010a; Frank and Crespi 2011). These forms of conflict also encompass two of the major forms of social interactions among sexual organisms, parents with offspring and males with females. We focus mainly on the hormones oxytocin and testosterone due to intense research interest in their physical and psychological effects, their wide-ranging influences on core aspects of development, cognition, and reproduction, and the central roles of oxytocin in mother–offspring interactions and testosterone in male–female differences.

We first discuss how hormones in general, and oxytocin and testosterone in particular, function in the context of evolved systems that mediate phenotypic adaptation and trade-offs. Second, we explain genomic imprinting and sexual antagonism, with regard to salient evolutionary theory, methods for detection and quantification of their effects, examples from the literature from animals and humans, and documented or postulated impacts on human health. In these contexts, we also discuss the similarities and differences between these two forms of conflict. We conclude with suggestions for future studies that integrate the evolutionary theory of genomic conflicts with health applications, especially for hormonally mediated traits. Although we focus predominantly on humans, the considerations below also apply to other sexual organisms, and taxa with genomic imprinting (mainly mammals and flowering plants), and to fitness-related impacts of genomic conflicts other than health and disease.

Functions of oxytocin and testosterone

Oxytocin, a peptide hormone, and testosterone, a steroid hormone, represent two of the best studied endocrine mediators of development, cognition, and behavior in humans and other mammals (McCall and Singer 2012; Auyeung et al. 2013). As such, these hormones exhibit relatively well-understood, as well as pervasive, effects on human adaptive functioning, such that links to maladaptation and disease can be more readily hypothesized and forged.

Oxytocin is produced mainly in the hypothalamus and is released into both the brain, where it modulates neurotransmission, and the peripheral circulation, where it regulates parturition, lactation, and other physiological processes (Gimpl and Fahrenholz 2001; Knobloch and Grinevich 2014). In both settings, oxytocin exerts its effects via binding to oxytocin receptors and stimulation of intracellular signaling. Effects of oxytocin in the brain include modulation of maternal behavior, social bonding, sexual and agonistic behavior, feeding, stress, and anxiety, which are organized via the distributions and densities of oxytocin receptors (Donaldson and Young 2008; Lee et al. 2009; Anacker and Beery 2013; Carter 2014). Neurologically, oxytocin can be considered as a hormonal mediator of social behavior and cognition, through its effects on social attention, social salience (perceived importance of social events), social motivation, social bonding, and social reward (Bethlehem et al. 2014). Early-life behavioral events, especially in infancy, appear to exert ‘organizational’ effects on the oxytocinergic system, mediating sensitivity to its effects (Feldman et al. 2013), although such processes have yet to be clearly elucidated in humans or other mammals. Most generally, increases and decreases in oxytocin, and receptor densities, within different regions of the brain modulate condition-dependent, sex-dependent, and context-dependent behavioral propensities and choices related to social interactions (Hammock and Young 2006). Higher oxytocin activity is thus associated with increased focus on social stimuli and, usually, increased prosocial behavior in the contexts of interactions within one’s own social groups of kin and non-kin, which include parents and offspring, male–female pair bonds, and larger groups defined by kinship or other societal factors (De Dreu 2012; Goodson 2013; Carter 2014). Oxytocin is expected to be of particular importance in humans compared to other animals, given the high levels of complexity to human social interaction and behavior.

Testosterone is produced in the testis, adrenal cortex, and (in females) ovaries, with bodily functions, of course, in the development of male primary and secondary sexual characteristics through activation of androgen receptors. With regard to brain and behavior, prenatal and early postnatal testosterone exerts organizational effects on neurodevelopment and cognitive functions (Auyeung et al. 2013). By comparison, after sexual maturation, such effects center on modulation by testosterone of energetic and time investment trade-offs between mating effort (higher levels) and parental effort (lower levels) in males, as well as trade-offs with investment in other domains such as immune functions (e.g., Klein 2000; Muehlenbein and Bribiescas 2005). In this context, sexual motivation, intrasexual dominance, success in male-typical endeavors, and social status represent primary determinants and arbitrators of success in mating effort, especially for males.

In contrast to oxytocin, testosterone is associated relatively strongly with ‘proself’ and self-oriented, rather than
prosocial and other-oriented, cognition and behavior (Wright et al. 2012). This difference between oxytocin and testosterone is reflected in opposite effects of these two hormones for a remarkable range of phenotypes, including trust (Bos et al. 2010, 2012; Van Ijzendoorn and Bakersmans-Kranenburg 2012), empathy (Domes et al. 2007a,b; van Honk et al. 2011), parenting (Gettler et al. 2011; Feldman et al. 2012; Okabe et al. 2013; Weisman et al. 2014), and amygdala connectivity with social-brain regions such as the orbitofrontal cortex (van Wingen et al. 2010; Volman et al. 2011; Bos et al. 2013; Sripada et al. 2013). Most broadly, these diametric effects frame our consideration of the primary adaptive functions, and maladaptive effects in disease, of these two hormones, as reflecting an axis of social interaction (for oxytocin), compared to an axis of self-orientation, sexuality, and gender (for testosterone). This is a considerable oversimplification, given the interacting roles of other hormones such as estrogens, arginine vasopressin, and cortisol (e.g., van Anders et al. 2011), but it serves as a useful and pragmatic first step in developing the connections of endocrine adaptations with the genetics, and epigenetics, of human risks and manifestations of disease. This conceptualization also structures the contexts and importance of these hormones regarding genomic conflicts, which should thus also be aligned with social, and sexual, conflicts and confluences of interest.

Genomic imprinting

Genomic imprinting refers to silencing of an allele in an offspring according to its parent of origin, either the mother or father (Fig. 1A). Genomic imprinting has evolved under conditions of higher relatedness to maternal kin compared to paternal kin, coupled with increased investment in offspring by females compared to males (Haig 2004, 2013). Usually, such higher relatedness among maternal kin than paternal kin is due to an evolutionary history of higher multiple paternity than multiple maternity within and across broods. By the well-supported kinship theory of imprinting (Ubeda and Haig 2003; Wilkins and Haig 2003; Haig 2013), and for mother–offspring interactions, lower relatedness through paternally inherited than maternally inherited alleles has selected for silencing in fathers (in developing sperm cell lineages) of genes that reduce maternal investment, which leads to higher levels of ‘selfish’ solicitation of resources from the mother, by offspring. In turn, selection in mothers has favored silencing (in developing oocytes) of other genes, specifically those whose expression in offspring leads to increased maternal investment through higher degrees of offspring solicitation of resources or other means. Imprinting effects are predicted in any situations involving relatedness asymmetries between maternal and paternal kin, but such
parent–offspring interactions appear to represent their pri- 
mary selective context throughout mammalian evolution. 

Imprinted genes are expected to be relatively strongly 
associated with maladaptation and risk of disease, com-
pared to other genes, for several reasons. First, their hap-
loid expression pattern exposes novel, deleterious alleles to 
selection; moreover, under haploidy, dysregulation can rel-
atively easily lead to (approximate) doubling or loss of 
expression, as commonly observed during carcinogenesis 
and some disorders of fetal and childhood growth (Hors-
themeke 2014). Second, imprinting leads to the evolution of 
new, conflict-related molecular mechanisms (such as ligand 
traps and antisense RNA transcripts) that represent novel 
targets for genetic or epigenetic dysregulation. Third, alter-
tations to imprinted genes are expected to generate espe-
cially strong and direct impacts on fitness (and thus 
health), given that they have evolved squarely in the con-
text of strongly fitness-related interactions of, for example, 
mothers with offspring. Fourth, imprinting-related con-
licts can generate situations where both parties are wasting 
important fitness-related resources, but deviations from the 
equilibrium are jointly disfavored. Finally, the conflictual 
nature of imprinted gene systems necessarily produces 
trade-offs in fitness between offspring and mothers; if one 
party ‘wins’, the other is maximally displaced from its opti-

um (Fig. 1A), and any intermediate, more or less stable 
resolution involves the same total displacements from fit-
ness optima distributed across both parties. 

Well-established associations of imprinted genes with 
disorders of placental development, prenatal and postnatal 
growth, risk of type 2 diabetes, cancer, and psychiatric con-
ditions (e.g., Kong et al. 2009; Peters 2014) have been 
reported, although human disease-genetic studies of com-
mon or rare variants have yet to focus on imprinted genes in 
y any systematic or comprehensive manner. Given the 
central roles of oxytocin in mother–offspring interactions 
described above, and human social behavior much more 
generally, do genomic imprinting effects impact this hor-
monal system, and if so how?

Oxytocin, genomic imprinting, and human disease 

Effects of imprinted genes on mammalian phenotypes and 
disease have been measured predominantly by analyzing 
the effects of gene knockouts or knockins in mice, charac-
terizing phenotypes associated with single nucleotide poly-
morphisms in imprinted genes, and by studying the effects 
in humans of imprinting disorders due to deletions or 
duplications of imprinted regions, or epimutations, that 
generate so-called imprinted gene syndromes (Peters 
2014).

A central, key aspect of early postnatal brain and cogni-
tive development is attachment to the mother, which 
mediates both feeding and social-psychological develop-
ment largely through the hormonal effects of oxytocin in 
both the developing child and the mother. As the discovery 
that behavioral and mating system variation in some 
rodents is associated with expression and reactivity to oxy-
tocin and the closely related neuropeptide arginine vasop-
ressin, the roles of oxytocin in human psychology, social 
behavior, and disease have been of increasing interest 
(Bethlehem et al. 2014; Carter 2014). Despite such concen-
trated attention on oxytocin’s effects, no previous studies 
have considered the evidence, and implications for health, 
regarding genomic imprinting and genomic conflicts in the 
oxotocinergic systems of rodents or humans, despite the 
well-established importance of imprinted genes in mother–
offspring interactions (Isles and Holland 2005; Crespi 
2010b, 2011).

Genomic imprinting effects center strongly on conflict 
over solicitation or demands imposed by offspring on the 
mother. Given that oxytocin produced in offspring drives 
bonding and attachment to the mother (Feldman et al. 
2013; Olff et al. 2013), an essential prerequisite to both 
energetic and psychological maternal-resource acquisition, 
a strong prediction from theory is that paternally expressed 
(maternally silenced) imprinted genes should favor 
increased offspring oxytocin production and increased sen-
sitivity via effects on receptor systems. Moreover, higher 
levels of paternally expressed imprinted gene products, and 
higher oxytocin levels in offspring, should be associated 
with increased levels of offspring sucking, and faster post-
natal growth. Conversely, maternal biases to imprinted 
gene expression, as from reductions in paternally expressed 
imprinted gene products, should lead to reduced bonding 
and attachment, less sucking, and slower growth. Such 
effects during infancy are predicted to have especially perva-
sive effects on health given fetal programming of adult 
disease risks and the highly pleiotropic effects of oxytocin 
in physiological and psychological development.

Table 1 summarizes the available evidence from compre-
hensive searches of the relevant literature focusing on 
imprinted genes relating to oxytocin, offspring sucking, 
and postnatal growth. Four imprinted genes, PEG3, NDN, 
MAGEL2, and DLK1, which are all paternally expressed, are 
associated with oxytocin, feeding, and growth as predicted 
from theory, in mice, humans, or both. In all four cases, 
knockouts are known (PEG3, NDN, MAGEL2), or expected 
(DLK1), to exhibit decreases in levels of oxytocin-secreting 
neurons in the hypothalamus, apparently through more or 
less selective reductions in these cell populations. These par-
allel findings fit closely with theory and are especially inter-
esting in the context of the central, highly connected hub 
positions of PEG3 and NDN genes in the imprinted gene 
coexpression network inferred by Varrault et al. (2006) in 
mice. Based on the documented behavioral effects of oxyto-
cin, large-scale reductions in hypothalamic oxytocin-secreting neurons are expected to lead to reduced solicitation of resources by offspring, both energetic (sucking) and behavioral (huddling for warmth and physical contact such as licking and grooming). Reduced solicitation of maternal contact (via activity and crying) and reduced feeding motivation in infancy have also been reported as primary features of the human equivalents of paternal ‘knockouts’ of the NDN and MAGEL2 genes (as well as other genes), Prader–Willi syndrome, which also involves reductions in oxytocinergic neurons in the hypothalamus (Swaab et al. 1995). Reduced expression of DLK1 (most commonly due to maternal uniparental disomy: bearing two maternal copies of chromosome 14) also leads to a set of features, including infant postnatal growth reductions, that overlaps with those of Prader–Willi syndrome (Cox et al. 2004; Crespi 2008), with comparable phenotypes also found in DLK1-knockout mice (Moon et al. 2002).

This convergent set of findings, across genes (PEG3, NDN, MAGEL2, DLK1) and species (humans and mice), is important for human health in at least three ways. First, these data indicate the existence of a set of imprinted genes that impacts directly and strongly on infant feeding and mother–infant bonding-related behavior. The degree to which segregating genetic and epigenetic variation in such genes modulates these phenotypes in humans, however, remains unstudied.

Second, the oxytocinergic system apparently originated in mammals in the context of viviparity, lactation, and mother–offspring bonding, but has since come to be co-opted for bonding between mates and between members of a cooperating social group (Gimpl and Fahrenholz 2001; Anacker and Beery 2013). As a result, alterations to imprinted genes that affect the oxytocin system are expected to have highly pleiotropic effects on human social behavior and psychology, throughout the life span. For example, in mice, knockouts of PEG3 and PEG1/MEST both lead to decreases in maternal behavior (among adult knockout females), including impaired lactation and pup retrieval (Lefebvre et al. 1998; Li et al. 1999), which can most parsimoniously be explained as secondary, long-term, purely maladaptive effects of reduced hypothalamic oxytocin secretion, given that this hormone orchestrates maternal care and feeding of offspring as well as offspring bonding to the mother. In humans, allelic variation in non-imprinted oxytocinergic genes including OXTR modulates the quality of maternal care of infants (e.g., Feldman et al. 2013), but imprinted genes have yet to be studied in this behavioral context. However, Prader–Willi syndrome and related conditions that affect hypothalamic development involve insecure psychological childhood attachment to mothers and represent strong risk factors for psychosis in adulthood (Crespi 2008; Soni et al. 2008). Oxytocin is also considered to be an important hormonal mediator of risk

| Imprinted gene, expression pattern | Evidence regarding oxytocin, infant feeding | References |
|-----------------------------------|--------------------------------------------|------------|
| PEG3 (paternally-expressed gene 3) | Knockout associated with reduced sucking in pups, large reduction in oxytocinergic neurons in adult females, reduced nursing by mothers, in mice | Li et al. (1999), Curley et al. (2004) and Champagne et al. (2009) |
| NDN (ncdin), paternally-expressed | Knockout associated with large reduction in oxytocinergic neurons in hypothalamus, in mice | Muscatelli et al. (2000) |
| GTF2I (General Transcription Factor IIi), paternally-expressed | Knockouts show poor infant sucking, large reduction in oxytocinergic neurons in hypothalamus, impaired oxytocin secretion; knockouts rescued by single postnatal oxytocin injection, in mice | Schaller et al. (2010) and Schaaf et al. (2013) |
| MAGEL2 (MAGE-like 2), paternally-expressed | Knockouts show reduced sucking in mice, deletions involve reduced sucking in humans | Plagge et al. (2004), Geneviève et al. (2005) |
| DLK1 (Delta-like 1 Homolog), paternally-expressed | Highly, selectively expressed in oxytocinergic neurons of hypothalamus after birth, in mice, and affects post-natal growth; locus appears to underlie feeding reductions in human maternal uniparental disomy of chromosome 14 | Buiting et al. (2008), Temple et al. (2009) and Villanueva et al. (2012) |
| GTF2I (General Transcription Factor IIi), maternal bias in expression | Williams syndrome involves deletion of one copy, increased levels of serum oxytocin, increased social behavior; duplications involve separation anxiety in both mice and humans; unknown if gene affects oxytocinergic system | Collette et al. (2009), Dai et al. (2012) and Mervis et al. (2012) |
| PEG1 (MEST) Paternally-expressed gene 1 | Knockout females show reduced, abnormal maternal behavior, comparable to that of PEG3 knockouts; effects on feeding, oxytocinergic system, unstudied | Lefebvre et al. (1998) |

© 2015 The Authors. Evolutionary Applications published by John Wiley & Sons Ltd 8 (2015) 307–325
for other psychiatric conditions such as autism, given its general roles in promoting social attention and bonding (Cochran et al. 2013).

Third, the oxytocin system regulates feeding and satiety, especially for foods with high hedonic value (such as chocolate chip cookies, in one study; Ott et al. 2013). In mammals, feeding is naturally a social behavior motivated by reward (with strong links to the dopaminergic system and nucleus accumbens ‘pleasure center’) (Sabatier et al. 2013). Indeed, in dairy cows, levels of plasma oxytocin in calves are substantially higher if they suckle from their mother than if they drink the same milk from a bucket (Lupoli et al. 2001). These findings, strong links between disturbances to the oxytocin system and obesity (Chaves et al. 2013; Sabatier et al. 2013) (as found, for example, in Prader–Willi syndrome and maternal uniparental disomy 14), and the role of the oxytocin-dopamine system in addiction (Tops et al. 2014), indicate that imprinted genes mediating oxytocin release and responsivity are also expected to show pleiotropic, conflict-associated effects in these important health-related contexts.

The situations described above all involve experimental, mutational, or epimutational reduction of oxytocin levels in imprinted gene systems. In contrast to such reductions, Williams syndrome, which is due to hemizygous deletion of about 20–25 genes on chromosome 7 in humans, provides evidence regarding imprinted gene effects that involved increased levels of oxytocin. This syndrome thus involves ‘hypersocial’ behavior, with overdeveloped social motivation and treatment of strangers as familiar friends (Järvinen et al. 2013), with a hormonal correlate in greatly increased levels of plasma oxytocin (Dai et al. 2012).

Indeed, a wide range of the behavioral, cognitive, and neurological phenotypes of Williams syndrome closely parallel the effects of oxytocin administration to healthy individuals, for such traits as increased social approach (Kemp and Guastella 2011; Järvinen et al. 2013), trust (Zhong et al. 2012; Godbee and Porter 2013), increased gaze toward faces and eyes (Guastella et al. 2008; Porter et al. 2010), and increased empathy (Fidler et al. 2007; Hurlemann et al. 2010).

In Williams syndrome, behavioral effects have been linked with deletion of the GTF2I gene (Sakurai et al. 2011), and GTF2I, although it has not been demonstrated to be imprinted, demonstrates a strong maternal-chromosome bias in expression (Collette et al. 2009), such that reduced gene dosage (as in Williams syndrome, compared to control individuals) creates a paternal bias in expression and behavior. This syndrome thus appears to represent, in part, the converse of situations where oxytocin is decreased due to loss of paternal expression (as for PEG3 and NDN), and it implicates oxytocinergic effects of imprinted genes even more broadly in human social behavior and cognition.

Despite the considerations discussed above, the degree to which genomic imprinting conflicts impacting the oxytocin system mediate human health and disease remains incompletely understood. Increasing evidence, however, indicates that early development of the oxytocin system in humans exerts lifelong effects on social behavior, stress, subjective well-being, and risk for a broad swath of psychiatric conditions (Cochran et al. 2013; Bethlehem et al. 2014). Analyses of this system that take account of genomic conflicts should yield more comprehensive and more clearly explicable results, as well as elucidating mechanisms that provide new opportunities for health promotion and development of novel therapies. Indeed, in MAGEL2-knockout mice, a single postnatal injection of oxytocin is sufficient to rescue infant feeding deficits (Schaller et al. 2010), and in humans, failure to establish effective breastfeeding is the main cause of clinic visits by mothers and infants (Wells 2003).

Sexual antagonism

Conflict of interests that involve genomic imprinting are sometimes referred to as parental antagonism, in that the bulk of such conflicts center on the divergent interests of paternally inherited and maternally inherited genes, within offspring or other individuals subject to asymmetric relatedness of paternal and maternal genes to social interactants (Haig 2013; Haig et al. 2014). In contrast to parental antagonism, evolutionary conflict of interests between the sexes are commonly referred to as sexual antagonism, or intralocus sexual conflict (van Doorn 2009; Pennell and Morrow 2013) (Fig. 1B,C).

Sexual antagonism can be defined as a situation whereby females and males express a trait with the same or highly overlapping genetic basis but with different fitness optima (Bonduriansky and Chenoweth 2009) that are not jointly achieved by both sexes. In such circumstances, alleles are sexually antagonistic to the extent that they increase the fitness of one sex but decrease the fitness of the other (Fig. 2). Sexual antagonism has been predicted from simple models to represent a pervasive impediment to phenotypic optimization by selection (e.g., Connallon and Clark 2014a), yet its expected and apparent impacts on human health and disease have seldom been investigated (Morrow and Connallon 2013).

Sexual antagonism differs from parental antagonism (genomic imprinting) in several important ways. First, parental antagonism involves genomic conflict between two parties (types of genes) within the same individual, which can be considered as a single genomic ‘niche’ for interacting genes (Patten et al. 2013). As a result, optimal trait values cannot be jointly achieved for maternal genes and paternal genes at the same time, so the conflicts cannot be resolved in any evolutionary sense unless one side...
irrevocably wins. By contrast, sexual antagonism involves two different 'niches' for genes, males and females, such that the evolution of fully sex-specific trait expression, where each sex reaches its optimum, is feasible in principle and could resolve the conflict, at least in the short term (Pennell and Morrow 2013; Haig et al. 2014). Second, whereas parental antagonism generates organism-level maladaptations along an axis of resource demands imposed by offspring on mothers during development (affecting physical and psychological resource acquisition), sexual antagonism is predicted to generate sex-specific forms of maladaptation along an axis of male–female differences, such that male trait values will be displaced toward (or away from) those of females, and vice versa, each to some degree. As for genomic imprinting effects, deviations from optimal phenotypes are expected to manifest primarily as deviations in health, such that particular alleles increase the health of one sex but decrease health of the other, or decrease the health of both. Third, sexually antagonistic alleles and effects are expected to be concentrated on sex chromosomes (the X and Y in mammals), because of the different inheritance patterns of the sex chromosomes and autosomes. Recent theoretical work has demonstrated that X-linked genes should favor females, while autosomal genes should favor male fitness optima (Frank and Crespi 2011; Connallon and Clark 2011; but see also Patten and Haig 2009 for a model predicting male-biased alleles on the X chromosome). Furthermore, the transmission of sex chromosomes also leads to different levels of relatedness along lineages when comparing males and females relative to their ancestral sex chromosomes (Rice et al. 2010). Empirical work from Drosophila indicates that the X chromosome is a hotspot for sexually antagonistic genes (Gibson et al. 2002; Innocenti and Morrow 2010), although it is currently unclear whether this finding applies to species, such as mammals, that possess a higher ratio of autosomal to sex chromosomal DNA and also differ in the nature of dosage compensation. Most generally, to the extent that sex chromosomes represent foci for sexually antagonistic effects, they may also differentially mediate the health effects of such conflicts, unless antagonisms are also more readily resolved to sex-specific optima for sex chromosomal genes than for autosomal genes.

In this review, we focus on the classic scenario of sexual antagonism, in which opposing directional selection acts on a trait to create an evolutionary tug-of-war for shared alleles between the sexes. We note, however, that there are several potential relationships between genotype, phenotype, and fitness that can reflect the operation of sexual antagonism; for example, alleles with sexually dimorphic or opposing phenotypic effects can result in one sex achieving and the other sex overshooting its fitness optimum (Turelli and Barton 2004; Connallon and Clark 2014b) even in the absence of sex-differential selection. Information on the specific contexts and forms of selection on phenotypes, and genetic basis of phenotypic variation in each sex, are thus ultimately required for a thorough understanding of how sexual antagonism impacts human health and disease.

**Detection and measurement of sexual antagonism**

How common and important is sexual antagonism in nature, and how do its consequences affect risk and forms of human disease? Demonstration of sexually antagonistic selection for a given trait requires that several criteria be met. First, the two sexes must differ in fitness optima for some phenotype, with a shared genetic basis between males and females. With regard to hormonally mediated traits, Table 2 provides a suite of evidence, for humans and other mammals, that the two sexes not only differ but show opposite phenotypes in response to some hormonally mediated stimulus. To the extent that either males or females deviate from their sex-specific fitness-maximal responses in such circumstances, such opposite effects may represent especially likely conditions for sexual antagonism, because, unless male and female responses are uncoupled, a favorable allele that augments a response in one direction, for one sex, should necessarily be unfavorable in the other sex.

Second, as noted above, selection must be opposite in direction between the sexes for a sexually antagonistic trait, such that an adaptive allelic effect in one sex is associated with a maladaptive effect in the other sex. Assessment of such effects is commonly carried out through measurements of phenotypic selection (such as selection gradient analyses) that examine a phenotypic value against a measure of fitness, partitioned by sex (Fig. 2). Various fitness components have been used to quantify this pattern, with lifetime reproductive success providing the most...
evolutionarily salient information. Counting numbers of offspring provides a simple and straightforward proxy of lifetime reproductive success that also allows for cross-species comparisons (Jones 2009) and yields an indication, given genetic information on the bases for the phenotype, of how genotypic frequencies might be expected to shift in a population across generations.

Several specific approaches have commonly been used to detect and quantify effects of sexual antagonism. First, within sibships, segregating sexually antagonistic alleles should tend to increase the fitness of one sex but decrease the fitness of the other sex. Such effects are expected to manifest both in terms of components of fitness, such as numbers of offspring, and in terms of risks and severity of sex-associated diseases that are observed or predicted to decrease fitness. Such family-based data can also be used to compare father–daughter and mother–son fitness measures using regression methods (Falconer and MacKay 1996; Foerster et al. 2007), with sexual antagonism indicated by lower fitness of cross-sex, compared to same-sex, parent–offspring pairs. These approaches are notably amenable for use with large epidemiological databases that include pedigrees as well as health information, as described in more detail below for the case of schizophrenia.

Second, quantitative genetic methods have been used to infer the operation of sexual conflict, given that this process is expected to generate negative genetic correlations between the fitness of males compared to females (i.e., genetic trade-offs; Robinson et al. 2006; Foerster et al. 2007; Schroderus et al. 2010). This approach, which is amenable for use with multiple-generation pedigree data, represents a generalization of the simple family-based methods described above and can be used to explore the impacts of sexually antagonistic gene expression over more than one generation. Such genetically based life-history perspectives can be powerful tools for quantifying genetic conflict and other traits affecting health and disease in humans (Stearns and Koella 2008; Gluckman et al. 2009), because they are designed to elucidate the sources, mechanisms, and effects of genetically based trade-offs.

Third, artificial selection experiments that involve directional selection on one sex for specific sex-associated traits, and tracking of phenotypes and measures of fitness in offspring of both sexes, have been used to test predictions regarding sexually antagonistic effects (e.g., Mills et al. 2012; using bank voles). Such selection experiments may also focus directly on the sex-differential fitness consequences of putative sexually antagonistic alleles, by breeding or selecting individuals for specific candidate genotypes and determining the sex-specific effects on associated phenotypes and components of fitness, preferably in relatively natural environments (Mokkonen et al. 2011). These methods generate experimental models of how sexual antagonism may be operating in the field, whose results
can be compared with patterns in natural populations. Research on \textit{Drosophila} genetics has provided the most extensive experimental evidence for sexually antagonistic genes through such selective breeding, in the laboratory environment (e.g., Rice 1992; Chippindale et al. 2001), as well as via studies of drift in small populations (Heskeht et al. 2013). A simple extension of this general approach, which has yet to be employed, involves testing for sex-specific positive selection in natural populations and determining the phenotypic and fitness effects of the derived and ancestral alleles in males and females. This method would be especially useful, and straightforward, for human populations and for connecting sexually antagonistic alleles with consequences for health.

Fourth, sex-biased patterns of gene expression have been used to test for effects of sexual antagonism and associations of sexual antagonism with the relative strengths of selection on males compared to females (Connallon and Knowles 2005; Innocenti and Morrow 2010; Griffin et al. 2013). A simple extension of this general approach, which has yet to be employed, involves testing for sex-specific positive selection in natural populations and determining the phenotypic and fitness effects of the derived and ancestral alleles in males and females. This method would be especially useful, and straightforward, for human populations and for connecting sexually antagonistic alleles with consequences for health.

Finally, theoretical predictions regarding expected concentrations and effects of sexually antagonistic alleles on sex chromosomes can be leveraged to test for sexual conflict (e.g., Manik and Ellegren 2009; Allen et al. 2013), given, for example, that alleles of benefit to females should be found differentially on the X chromosome in mammals and that sexually antagonistic drive should generate genomic conflicts, and harmful fitness-reduced effects, among opposite-sex siblings (Rice et al. 2008, 2010).

To evaluate the impacts of sexual conflict on human health, the approaches described above must be dovetailed with information on sex differences in genotype-phenotype associations that are relevant to risks and manifestations of disease (Ober et al. 2008; Gilks et al. 2014). Such dimorphisms are expected to be driven by sexual differences in gene expression (and other allelic effects) that trace, ultimately, to the causes of human sexual differentiation. Steroid and peptide hormones represent among the most important such causes, and as such, they should be the primary mediators of sexual conflict in its influences on human health and disease. How, then, might sexual antagonism be mediated by genetically based effects of hormonal variation?

\textbf{Sexual antagonism and genetically based hormone systems}

In changing social and abiotic environments, hormones provide effective mechanisms for regulating behavioral responses appropriate to a given sex, situation, and stimulus. Selection should thus favor different regulatory phenotypes specific to each sex (Clutton-Brock 2007; Kokko and Jennions 2008), yet responses to selection may be constrained due to the largely shared genome.

As described above, hormones provide primary mechanisms for development, physiology, and behaviors to operate in context-specific and sex-specific manners. For example, plasma testosterone is found in much higher concentrations in males, while females demonstrate higher concentrations of oxytocin, at least in most studies (Altomus et al. 1999; Carter 2007; Pierrebomber et al. 2010; Weisman et al. 2013). Currently, testosterone is the best studied hormone with regard to evidence for sexual antagonism in humans and other animals, and the developmental and physiological mechanisms that underlie such effects are relatively well understood (Ketterson et al. 2005; Stearns and Koella 2008; Hau and Wingfield 2011). The role of testosterone in mediating sexual antagonism is further bolstered by the fact that the androgen receptor gene, which is important for the functional response to testosterone, is located on the X chromosome in humans.

We next describe evidence for sexual antagonism, especially as mediated by effects of testosterone (and related steroid hormones), in non-human animals, and we briefly relate this evidence to human health and disease. We then focus directly on evidence regarding sexual antagonism in humans, in the context of the methods discussed above, and discuss the relevance of such evidence to disease. We concentrate specifically on several areas of central importance to human health: life-history phenotypes, behavior, immune function, and psychology.

\textbf{Sexual antagonism, testosterone, and life-history traits}

The organizational and activational effects of testosterone on physiological and behavioral traits make it an important mediator of vertebrate life-history traits and provide a core context for effects of sexual antagonism. The influence of this hormone can thus mediate the timing of reproduction, the growth and development of offspring, and, most importantly, sex-specific traits that affect components of survival and reproduction.

A series of studies of bank voles (\textit{Myodes glareolus}) has demonstrated that the physiological and behavioral effects of testosterone confer advantages in male--male competition: Males with higher circulating plasma testosterone levels achieve greater reproductive success (Mills et al. 2007, 2009). However, there is also direct sexual antagonism over testosterone levels, whereby responses to artificial selection increase the fitness of males at the expense of females through antagonistic effects on mating behavior (Mills
et al. 2012; Mokkonen et al. 2012). As testosterone is known to affect sexual motivation and dominance behavior in mammals, males will, to a point, achieve greater mating success with greater testosterone production. The sexual antagonism is realized because female propensity to mate has been shown to decrease with selection for testosterone; brothers and sisters thus cannot both benefit from selection for testosterone. At the population level, the benefit that testosterone confers on males in intrasexual competition is highly dependent on the social environment of individuals, as too many dominant males in the population will result in lower average reproductive success (Mokkonen et al. 2011); this frequency dependence can maintain genetic polymorphisms when there are competing sex-specific optima in a population. Such patterns may also be influenced by age: A quantitative genetic study on bank voles (M. Mokkonen, E. Koskela, T. Mappes and E. Schroderus, unpublished data) has shown that sexual antagonism over fertility increases with the age of females, presumably due to the sex-differential declines in fertility that are mediated by sex hormones. Potential parallels with human life history are of interest because the reproductive life span of females appears to be increasing over time (Stearns et al. 2010); older females are predicted to express sexually antagonistic alleles more strongly, which may influence risks of reproductive disorders in particular.

Observations of other mammalian species have provided notable evidence of sexual antagonism in natural environments, with parallels to the well-studied bank voles described above. Thus, for example, red deer males that had higher fitness in terms of lifetime reproductive success sired daughters that generally had lower fitness (Foerster et al. 2007). Likewise, Soay sheep experience sexually antagonistic selection over horn phenotype (Robinson et al. 2006). Body mass in mountain goats has also been found to be consistent with sexual antagonism, given that paternal body mass is negatively correlated with daughter’s body mass, but positively correlated with son’s body mass (Mainguy et al. 2009). Finally, data on bighorn sheep show that longer-horned males sire sons with higher viability, but daughters with lower viability (Martin et al. 2014); such effects appear to be mediated in part by effects of male testosterone levels on social rank (Martin et al. 2013).

What are the expected consequences of such findings for human health and well-being, given that such sexually antagonistic effects on physiological, behavioral, and life-history traits are commonly found across mammals? The most general predicted phenotypic pattern would be that male-beneficial effects of higher testosterone would be associated with lower health and fitness among females; conversely, female-beneficial effects of hormonal phenotypes that differentially involve benefits on females (such as those involving higher estradiol, and oxytocin) would be expected to have negative impacts upon males. As described in more details below, patterns consistent with this expectation have been reported for some human phenotypes, although only a few of them have been directly analyzed for association with physical or psychological health. Additionally, there is considerable opportunity for including models and predictions of sexual antagonism to understanding pathology of fertility-related diseases. Given that sexual antagonism results in reduced reproductive success in one sex, these fitness costs may differentially manifest in disease-related aspects of reproductive physiology, among both males and females.

Sexual antagonism, testosterone, and immune function

One of the key features of testosterone that has drawn the interest of evolutionary biologists is the pleiotropic nature of this hormone in mediating several important fitness-related functions and trade-offs (Hau and Wingfield 2011). Immunosuppressive effects of testosterone production and response sensitivity have been a central focus of research in this area (Folstad and Karter 1992; Mills et al. 2010; Hau and Wingfield 2011), in that they apparently represent a fundamental life-history trade-off that may render some individuals more susceptible to disease. To the extent that males benefit more than females from testosterone production, and if such production is subject to effects of sexually antagonistic alleles, then females may be hypothesized to differentially suffer immune system costs from relatively high testosterone.

Evidence that testosterone and immune function are subject to effects of sexual antagonism comes from a series of studies of bank voles. Thus, Mills et al. (2010) showed that male voles subject to artificial selection on immune function showed correlated responses to selection in levels of testosterone. Experimental administration of testosterone to males also led to reduced immune function (in the highest testosterone group), as well as increased social status and reproductive success (Mills et al. 2010). Most importantly, Schroderus et al. (2010) demonstrated, using quantitative genetic methods, that immune function and testosterone levels were subject to negative genetic correlations in both males and females, such that selection for higher testosterone in males will result in reduced immune function among females.

Among humans, activity and vigor of the immune system is notably higher among females than males (Bouman et al. 2005; Oertelt-Prigione 2012), and negative associations have been demonstrated between testosterone levels and immune system activity across many studies (Muehlenbein and Bribiescas 2005). The medical relevance of such associations is demonstrated, for example, by lower anti-
Sexual antagonism in human populations

Studies focusing on human sexual antagonism are limited but increasing. Stulp et al. (2012) recently used long-term cohort data to demonstrate intralocus sexual conflict on human height, showing that sisters had higher reproductive success (numbers of children) than their brothers among shorter sib-pairs, but brothers had higher success than sisters among average height pairs. Given evidence of a negative association between height and reproductive success among females, but a stabilizing-selection pattern among males favoring average height, these data indicate that neither sex is expected to achieve its sex-specific optimum for this trait.

A study by Bolund et al. (2013) using pedigree data from a pre-industrial Finnish population highlighted the usefulness of combining phenotypic selection gradient analysis with genetic data to assess sexual conflict in human populations. In this population, the phenotypic selection gradients were opposite in direction between the sexes for first and last reproduction, and reproductive life span, but these traits also showed positive, rather than negative, genetic correlations between the sexes, with regard to fitness (Bolund et al. 2013). Sexual antagonism was therefore not realized in this case, despite selection operating antagonistically between the sexes. By contrast, Stearns et al. (2012), using a combination of phenotypic selection data and information on genetic correlations, determined that sexually antagonistic selection was acting on several important human health measures such as height, weight, blood pressure, glucose levels, cholesterol levels, and age at first birth, and constrained joint evolutionary responses to selection in a manner expected to exert impacts on sex-specific human health and disease.

Additional evidence has emerged, for various nonmedical traits, that hormones are implicated in human sexual antagonism. Garver-Apgar et al. (2011) demonstrated that masculine males and females reported higher ‘mate values’ for brothers relative to sisters, suggesting that the influence of higher testosterone increased perceived mate value in males, but resulted in lower perceived mate value for ‘androgenized’ females. Studies of facial attractiveness scores between siblings added support to the hypothesis that sexual antagonism operates in such situations, apparently mediated by effects of testosterone (Manning et al. 2000; Mitchem et al. 2014). These studies would represent effects of sexual antagonism to the extent that perceived mate values were associated with differential fitness by sex in sibships, and especially if such facial-feature and mate-choice effects were linked with hormone-associated genetic variation.

Hormonal effects on human phenotypes have also commonly been indexed by finger 2D:4D ratio, whereby a lower ratio reflects greater prenatal testosterone exposure relative to estradiol, and thus, greater ‘masculinization’ (Lutchmaya et al. 2004). Manning et al. (2000) demonstrated that measures of male reproductive success were higher for individuals with lower (more ‘masculine’) 2D:4D ratios, while the opposite was true for females. This finding was replicated in a larger study (Manning and Fink 2008), but not in a sample of women from Finland (Helle 2010). Although further data are needed, especially regarding the mechanisms linking 2D:4D with measures of fitness, the former two studies are consistent with sexual antagonism effects mediated by prenatal ratios of testosterone to estradiol. Such evidence for antagonism effects is important for health given the evidence for associations of 2D:4D ratios with risks for diseases such as cancer (e.g., Muller 2013) and coronary artery disease (e.g., Wu et al. 2013). In principle, sexual antagonism theory thus predicts opposite effects on males and females in such disease risks versus protection, mediated by loci that influence production and sensitivity to steroid hormones especially during early development.

Finally, human handedness shows one of the primary signatures of sexual antagonism: Female mixed-handers report themselves as more masculine, and male mixed-handers report themselves as more feminine, compared to more lateralized individuals (Tran et al. 2014). Such effects appear to be mediated in part by genetic variation in the androgen receptor, in that among females, left handedness is associated with a larger number of microsatellite repeats, but in males, it is associated with fewer repeats (Medland et al. 2005; see also Hampson and Sankar 2012).

Sexual antagonism and human behavior, psychology, and psychiatric conditions

A primary signature of sexual antagonism in human behavior-related phenotypes, from evidence available to date, is...
increased fitness of one sex, combined with decreased fitness of related individuals of the other sex, in the context of a genetically based, sexually dimorphic trait whose expression is mediated by hormones. As described in detail below, this pattern has been reported and substantially replicated in the context of two human phenotypes: sexual orientation and schizophrenia. The former phenotype is not, of course, directly associated with health, although it may be indicative of the potential for sexually antagonistic effects in behavioral and psychological traits linked with risk of disease. By contrast, schizophrenia and the related psychiatric conditions, such as bipolar disorder, major depression, and borderline personality disorder, represent among the most important human health conditions with regard to personal, social, and economic impacts. New perspectives on the causes of these conditions from the evolutionary theory of sexual antagonism should offer novel insights into understanding them and driving strategies for the collection of new data.

A series of ten studies, across multiple ethnicities, has provided evidence supporting a hypothesis of increased fecundity in maternal (but not paternal) relatives of homosexual men (review in Ciani and Pellizzari 2012). Concomitant reduced offspring production of homosexual men, and evidence that some degree of X chromosome linkage for underlying alleles is most compatible with the available genetic association and pedigree data, has led Ciani and colleagues (Ciani et al. 2008) to infer that a hypothesis of sexually antagonistic alleles provides the best explanation for this otherwise-paradoxic trait (Iemmola and Ciani 2009; Ciani and Pellizzari 2012; Ciani et al. 2012). The genetic and environmental mechanisms that mediate such effects remain unclear and complex, although hormones (Balthazar 2011) or X-linked epigenetic factors (Rice et al. 2012) appear to play some roles.

A notable health implication of this pattern of results follows from the finding that female relatives (mothers and maternal aunts) of male homosexuals exhibited significantly reduced complications in pregnancy and fewer gynecological disorders, than did female relatives of heterosexual men (Ciani et al. 2012). Determining the mechanisms that mediate such positive effects should have direct implications for female reproductive health and for health-related pleiotropic effects of the relevant causal factors in homosexual men. More generally, these findings represent clear evidence for sexual antagonism in human development, cognition, behavior, and reproduction and indicate that its effects are likely to also promote the maintenance of variation in more directly health-related traits.

A series of studies on the epidemiology of schizophrenia has focused on the hypothesis that fitness-related benefits in the relatives of schizophrenics may help to explain the high prevalence of this disorder, and its substantial heritability, in the context of its strongly negative impacts on reproduction by patients (van Dongen and Boomsma 2013). None of these studies have reported fitness benefits to relatives sufficiently strong (under realistic models) to maintain schizophrenia risk alleles, but a notable number of them (Bassett et al. 1996; McGrath et al. 1999; Haukka et al. 2003; Weiser et al. 2009; meta-analysis in Bundy et al. 2011) have found evidence that female relatives of individuals with schizophrenia exhibit significantly higher fertility than female control individuals, even though male relatives of schizophrenics show greatly reduced fertility.

Most recently, a large study using a Swedish database of over two million individuals (Power et al. 2013) demonstrated strong effects supporting these smaller, more heterogeneous studies: Sisters of schizophrenic individuals had significantly increased numbers of children compared to controls (whereas brothers had significantly fewer); moreover, sisters of individuals with bipolar disorder also showed significantly increased fecundity (brothers showing no difference), and both sisters and brothers of individuals with major depression demonstrated such increases. This study and virtually all previous reports have also shown that schizophrenia itself has substantially more adverse effects on male than female reproduction; this disorder is also more deleterious in its effects on cognitive abilities among males than females (e.g., Abu-Akel and Bo 2013).

Such sex differences in schizophrenia prevalence and severity have usually been attributed to protective effects of estrogen in females, although influences from other hormones appear also to be involved (reviews in Mendrek and Stip 2011; Hayes et al. 2012). Additional evidence consistent with effects of sexual antagonism in schizophrenia risk comes from functional-imaging studies that demonstrate ‘masculinization’ of female schizophrenia patients and ‘feminization’ of male patients (Mendrek 2007; Mendrek et al. 2007), mediated in part by levels of testosterone (Mendrek et al. 2011), which suggests dysregulation directly along a male–female axis.

Considered together, these data indicate that schizophrenia demonstrates clear evidence of risk mediation, in part, by effects of sexual antagonism. As such, some notable proportion of schizophrenia risk alleles should exhibit beneficial effects among females (with regard, presumably, to functions ultimately linked with reproduction, and associated cognitive–emotional processes) in addition to their deleterious effects among males (with regard to schizophrenia risk, and its associated, underlying phenotypes). A primary implication of these results is that studies of schizophrenia that do not account for sex differences in the genetic and gene by environment bases of risk will yield results that are at best incomplete, and at worst incorrect. The findings also suggest that the X chromosome, as a nexus for sex-
ually antagonistic effects, may exhibit an especially important influence on schizophrenia risk, as a suite of previous studies has also suggested (e.g., Crespi 2008; Crow 2008; Goldstein et al. 2011, 2013).

The degree to which the patterns of findings described above, for sexual orientation and schizophrenia, generalize to other health-related conditions remains unknown, but relevant theory predicts that any human disease showing sex differences in incidence, severity, or manifestations should be a candidate for genetic risks mediated in part by sexual antagonism. Among psychiatric conditions, autism demonstrates among the strongest evidence for such sex differences and potential effects of sexual antagonism, with a striking male bias in prevalence, clear hormonal effects on risk, greater severity among females, effects from X-linked loci, and disorder-associated enhancements in some performance traits related to perception, visual-spatial skills, and mechanistic cognition, which are concentrated among males (Marco and Skuse 2006; Mottron et al. 2006; Treffert 2009; Baron-Cohen et al. 2011; Robinson et al. 2013; Teatero and Netley 2013). As such, genetic risk of autism may engender hormone-associated alleles that tend, overall, to benefit males, but also tend to increase risk of autism and other testosterone-linked diseases among females. This hypothesis is supported by evidence for elevated incidence of testosterone-related disorders in the female relatives of autistic individuals (Ingudomnukul et al. 2007) by high rates of testosterone-related ‘steroidopathies’ among women with autism spectrum disorders (Pohl et al. 2014) and by higher levels of serum testosterone in females (but not males) with autism, compared to controls (Schwarz et al. 2011; Bejerot et al. 2012). A general prediction of the hypothesis is that some ‘autism risk’ alleles should also confer benefits to males in healthy populations (e.g., in visual-spatial abilities), but also be associated with costs to females.

Conclusions

One of the primary uses of evolutionary biology in the study of human health and disease is indicating what novel data would be especially useful to collect. Genomic conflicts provide a paradigmatic example of such evolutionary insights, because their dynamics and predictions are well beyond the purview of standard medical frameworks for studying and understanding disease. We have analyzed two of the primary forms of genomic conflicts, genomic imprinting (parental antagonism) and sexual antagonism, because they both predict conflict-driven, health-related deviations from optimal phenotypes for one or both of the parties involved, in the core fitness-related domains of parent–offspring and male–female interactions, respectively. With regard to genomic imprinting effects, we have described evidence, from theory and empirical work, that imprinted genes impact upon development of the human oxytocinergic system, a system that mediates lifelong psychological health and well-being. Thus far, genetic studies of this system have focused almost exclusively on the non-imprinted genes OXTR, OXT, and CD38; our results indicate that genetic, epigenetic, and functional studies of the imprinted genes PEG3, NDN, MAGEL2, DLK1, and GTF2I should lead to important new insights into the roles of oxytocin in human health, especially with regard to infant resource solicitation, mother–infant interactions, and their downstream psychological effects. With regard to sexual antagonism, our review suggests that some notable proportion of human sex-differential risks of disease incidence and severity may be attributable to such conflicts, especially in the context of positive versus negative effects of phenotypes mediated by testosterone, estrogen, and other hormones in males compared to females. As regards analyzing the genetic bases of human disease risks, such considerations should compel data collection and analysis that explicitly tests hypotheses of protection from disease in one sex, but increased risks of disease in the other, attributable to effects from the same locus. In addition to helping uncover new sources of disease risk, the study of sexual antagonism should thus also demonstrate beneficial sex-specific phenotypes, such as increased reproductive health among females or males, due to favorable sex-specific alleles. Studies of human sexual antagonism should dovetail directly with ongoing genetic association, large-scale epidemiological, and selection–measurement analyses, such that testing for sex-antagonistic effects mainly requires increased awareness of its expected prevalence, dynamics, and impacts on risks and phenotypes of sex-differential disease.

Acknowledgements

The authors would like to thank D. Haig for useful discussions. This research was supported by NSERC and the Academy of Finland (Grant #257729 to MM).

Literature cited

Abu-Akel, A., and S. Bo 2013. Superior mentalizing abilities of female patients with schizophrenia. Psychiatry Research 210:794–799.

Albers, H. E. 2012. The regulation of social recognition, social communication and aggression: vasopressin in the social behavior neural network. Hormones and Behavior 61:283–292.

Allen, S. L., R. Bonduriansky, and S. F. Chenoweth 2013. The genomic distribution of sex-biased genes in Drosophila serrata: X chromosome demasculinization, feminization, and hyperexpression in both sexes. Genome Biology and Evolution 5:1986–1994.

Altemus, M., K. R. Jacobson, M. Debellis, M. Kling, T. Pigott, D. L. Murphy, and P. W. Gold 1999. Normal CSF oxytocin and NPY levels in OCD. Biological Psychiatry 45:931–933.
Bos, P. A., J. Panksepp, R. M. Bluth Carter, C. S. 2014. Oxytocin pathways and the evolution of human behavior. Frontiers in Behavioral Neuroscience 7:185. Review.

van Anders, S. M., K. L. Goldey, and P. X. Kuo 2011. The steroid/peptide theory of social bonds: integrating testosterone and peptide responses for classifying social behavioral contexts. Psychoneuroendocrinology 36:1265–1275.

Auyeung, B., M. V. Lombardo, and S. Baron-Cohen 2013. Prenatal and postnatal hormone effects on the human brain and cognition. Pflügers Archiv: European Journal of Physiology 465:537–571.

Balthazart, J. 2011. Minireview: hormones and human sexual orientation. Endocrinology 152:2937–2947.

Baron-Cohen, S., M. V. Lombardo, B. Auyeung, E. Ashwin, B. Chakrabarti, and R. Knickmeyer 2011. Why is autism spectrum conditions more prevalent in males? PLoS Biology 9:e1001081.

Bassett, A. S., A. Bury, K. A. Hodgkinson, and W. G. Honer 1996. Reproductive fitness in familial schizophrenia. Schizophrenia Research 21:151–160.

Bejerot, S., J. M. Eriksson, S. Bonde, K. Carlström, M. B. Humble, and E. Eriksson 2012. The extreme male brain revisited: gender coherence in adults with autism spectrum disorder. British Journal of Psychiatry 201:116–123.

Bethlehem, R. A., S. Baron-Cohen, J. van Honk, B. Auyeung, and P. A. Bos 2014. The oxytocin paradox. Frontiers in Behavioral Neuroscience 8:48.

Bolund, E., S. Bouwhuis, J. E. Pettay, and V. Lummaa 2013. Divergent selection on, but no genetic conflict over, female and male timing and rate of reproduction in a human population. Proceedings of the Royal Society B: Biological Sciences 280:20132002.

Bonduriansky, R., and S. F. Chenoweth 2009. Intralocus sexual conflict. Trends in Ecology and Evolution 24:280–288.

Bos, P. A., D. Terburg, and I. van Honk 2010. Testosterone decreases trust in socially naive humans. Proceedings of the National Academy of Sciences USA 107:9991–9995.

Bos, P. A., J. Panksepp, R. M. Bluthé, and J. van Honk 2012. Acute effects of steroid hormones and neuropeptides on human social-emotional behavior: a review of single administration studies. Frontiers in Neuroendocrinology 33:17–35.

Bos, P. A., J. van Honk, N. F. Ramsey, D. J. Stein, and E. J. Hermans 2013. Testosterone administration in women increases amygdala responses to fearful and happy faces. Psychoneuroendocrinology 38:808–817.

Bowman, A., M. J. Heineman, and M. M. Faas 2005. Sex hormones and the immune response in humans. Human Reproduction Update 11:411–423.

Buining, K., D. Kanber, J. I. Martín-Suárez, W. Lieb, P. Terhal, B. Albrecht, S. Purmann et al. 2008. Clinical features of maternal uniparental disomy 14 in patients with an epimutation and a deletion of the imprinted DLK1/GTL2 gene cluster. Human Mutation 29:1141–1146.

Bundy, H., D. Stahl, and J. H. MacCabe 2011. A systematic review and meta-analysis of the fertility of patients with schizophrenia and their unaffected relatives. Acta Psychiatria Scandinavica 123:98–106.

Burt, A., and R. Trivers 2006. Genes in Conflict: The Biology of Selfish Genetic Elements. Belknap, Cambridge, MA.

Carter, C. S. 2007. Sex differences in oxytocin and vasopressin: implications for autism spectrum disorders? Behavioural Brain Research 176:170–186.

Carter, C. S. 2014. Oxytocin pathways and the evolution of human behavior. Annual Review of Psychology 65:17–39.
gene. Proceedings of the Royal Society B: Biological Sciences 271:1303–1309.
Dai, L., C. S. Carter, J. Ying, U. Bellugi, H. Pourmajafi-Nazarloo, and J. R. Korenberg 2012. Oxytocin and vasopressin are dysregulated in Williams Syndrome, a genetic disorder affecting social behavior. PLoS One 7:e38513.
De Dreu, C. K. 2012. Oxytocin modulates cooperation within and competition between groups: an integrative review and research agenda. Hormones and Behavior 61:419–428.
Ditzen, B., U. M. Nater, M. Scharer, R. La Marca, G. Bodenmann, U. Ehlert, and M. Heinrichs 2013. Sex-specific effects of intranasal oxytocin on autonomic nervous system and emotional responses to couple conflict. Social Cognitive and Affective Neuroscience 8:897–902.
Domes, G., M. Heinrichs, A. Michel, C. Berger, and S. C. Herpertz 2007a. Oxytocin improves “mind-reading” in humans. Biological Psychiatry 61:731–733.
Domes, G., M. Heinrichs, J. Gläscher, C. Büchel, D. F. Braus, and S. C. Herpertz 2007b. Oxytocin attenuates amygdala responses to emotional faces regardless of valence. Biological Psychiatry 62:1187–1190.
Domes, G., A. Lischke, C. Berger, A. Grossmann, K. Hauenstein, M. Heinrichs and S. C. Herpertz 2010. Effects of intranasal oxytocin on emotional face processing in women. Psychoneuroendocrinology 35:83–93.
Donaldson, Z. R., and L. J. Young 2008. Oxytocin, vasopressin, and the neurogenetics of sociality. Science 322:900–904.
van Dongen, J., and D. I. Boomsma 2013. The evolutionary paradox and the missing heritability of schizophrenia. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics 162B:122–136.
van Doorn, G. S. 2009. Intralocus sexual conflict. Annals of the New York Academy of Sciences 1168:52–71.
Falconer, D. S., and T. F. C. MacKay 1996. Introduction to Quantitative Genetics. 4th edn. Pearson Prentice Hall, London, UK.
Feldman, R., O. Zagoory-Sharon, O. Weisman, I. Schneiderman, I. Gordon, R. Maoz, I. Shalev et al. 2012. Sensitive parenting is associated with plasma oxytocin and polymorphisms in the OXTR and CD38 genes. Biological Psychiatry 72:175–181.
Feldman, R., I. Gordon, M. Influs, T. Gutbir, and R. P. Elstein 2013. Parental oxytocin and early caregiving jointly shape children’s oxytocin response and social reciprocity. Neuropsychopharmacology 38:1154–1162.
Fidler, D. J., S. L. Hephburn, D. E. Most, A. Philofsky, S. J. Rogers, and W. E. MacLean Jr 2007. Emotional responsivity in young children with Williams syndrome. American Journal on Mental Retardation 112:194–206.
Foerster, K., T. Coulson, B. C. Sheldon, J. M. Pemberton, T. H. Clutton-Brock, and L. E. Kruuk 2007. Sexually antagonistic genetic variation for fitness in red deer. Nature 447:1107–1110.
Folstad, I., and A. J. Karter 1992. Parasites, bright males, and the immuno-competence handicap. The American Naturalist 139:603–622.
Frank, S. A., and B. J. Crespi 2011. Pathology from evolutionary conflict, with a theory of X chromosome versus autosome conflict over sexually antagonistic traits. Proceedings of the National Academy of Sciences USA 108(Suppl. 2):10886–10893.
Furman, D., B. P. Hejblum, N. Simon, V. Jovic, C. L. Dekker, R. Thiebaut, R. J. Tibshirani et al. 2014. Systems analysis of sex differences reveals an immunosuppressive role for testosterone in the response to influenza vaccination. Proceedings of the National Academy of Sciences USA 111:869–874.
Garver-Apgar, C. E., M. A. Eaton, J. M. Tybur, and M. E. Thompson 2011. Evidence of intralocus sexual conflict: physically and hormonally masculine individuals have more attractive brothers relative to sisters. Evolution and Human Behavior 32:423–432.
Geneviève, D., D. Sanlaville, L. Faire, M. L. Kottler, M. Jambou, P. Gosset, D. Boustani-Samara et al. 2005. Paternal deletion of the GNAS imprinted locus (including Gnasxl) in two girls presenting with severe pre- and post-natal growth retardation and intractable feeding difficulties. European Journal of Human Genetics 13:1033–1039.
Gettler, L. T., T. W. McDade, A. B. Feranil, and C. W. Kuzawa 2011. Longitudinal evidence that fatherhood decreases testosterone in human males. Proceedings of the National Academy of Sciences USA 108:16194–16199.
Gibson, J. R., A. K. Chippindale, and W. R. Rice 2002. The X chromosome is a hot spot for sexually antagonistic fitness variation. Proceedings of the Royal Society B: Biological Sciences 269:499–505.
Gilks, W. P., J. K. Abbott, and E. H. Morrow 2014. Sex differences in disease genetics: evidence, evolution and detection. Trends in Genetics 30:453–463.
Gimpel, G., and F. Fahrenholz 2001. The oxytocin receptor system: structure, function, and regulation. Physiological Reviews 81:629–683.
Gluckman, P. D., A. Beedle, and M. Hanson 2009. Principles of Evolutionary Medicine. Oxford University Press, Oxford.
Godbee, K., and M. A. Porter 2013. Attribution of negative intention in Williams syndrome. Research in Developmental Disabilities 34:1602–1612.
Goldstein, J. M., S. Cherkzerzian, L. J. Seidman, T. L. Petryshen, G. Fitzmaurice, M. T. Tsuang, and S. L. Buka 2011. Sex-specific rates of transmission of psychosis in the New England high-risk family study. Schizophrenia Research 128:150–155.
Goldstein, J. M., S. Cherkzerzian, M. T. Tsuang, and T. L. Petryshen 2013. Sex differences in the genetic risk for schizophrenia: history of the evidence for sex-specific and sex-dependent effects. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics 162B:698–710.
Goodson, J. L. 2013. Deconstructing sociality, social evolution and relevant nonapeptide functions. Psychoneuroendocrinology 38:465–478.
Griffin, R. M., R. Dean, J. L. Grace, P. Rydén, and U. Friberg 2013. The shared genome is a pervasive constraint on the evolution of sex-biased gene expression. Molecular Biology and Evolution 30:2168–2176.
Guastella, A. J., P. B. Mitchell, and M. R. Dadds 2008. Oxytocin increases gaze to the eye region of human faces. Biological Psychiatry 63:3–5.
Haig, D. 2004. Genomic imprinting and kinship: how good is the evidence? Annual Review of Genetics 38:553–585.
Haig, D. 2013. Coadaptation and conflict, misconception and muddle, in the evolution of genomic imprinting. Heredity, 113:96–103.
Haig, D., F. Øbeda, and M. M. Patten. 2014. Specialists and generalists: the sexual ecology of the genome. In S. Gavrilets, and W. R. Rice, eds. Sexual Conflict, pp. 37–48. Cold Spring Harbor Perspectives, Cold Spring Harbor, NY.
Hammock, E. A., and L. J. Young 2006. Oxytocin, vasopressin and pair bonding: implications for autism. Philosophical Transactions of the Royal Society of London Series B, Biological Sciences 361:2187–2198.
Hampton, E., and J. S. Sankar 2012. Hand preference in humans is associated with testosterone levels and androgen receptor gene polymorphism. Neuropsychologia 50:2018–2025.
Hau, M., and J. C. Wingfield 2011. Hormonally-regulated trade-offs: evolutionary variability and phenotypic plasticity in testosterone signalling pathways. In: T. Flatt, and A. Heyland, eds. Mechanisms of Life History Evolution, pp. 349–364. Oxford University Press, Oxford.

Genomic imprinting, sexual antagonism and hormones
Genomic imprinting, sexual antagonism and hormones

Haukka, J., J. Suvisaari, and J. Lönnqvist 2003. Fertility of patients with schizophrenia, their siblings, and the general population: a cohort study from 1950 to 1959 in Finland. The American Journal of Psychiatry 160:460–463.

Hayes, E., E. Gavrilidou, and J. Kulkarni 2012. The role of oestrogen and other hormones in the pathophysiology and treatment of schizophrenia. Schizophrenia Research and Treatment 2012:540273.

Helle, S. 2010. Digit length ratio (2D:4D) and variation in key life-history traits and fitness in contemporary Finnish women. Behavioral Ecology 21:1061–1066.

Hesketh, J., K. Fowler, and M. Reuter 2013. Genetic drift in antagonistic genes leads to divergence in sex-specific fitness between experimental populations of Drosophila melanogaster. Evolution 67:1303–1510.

Hoge, E. A., E. Anderson, E. A. Lawson, E. Bui, L. E. Fischer, S. D. Khandage, L. F. Barrett et al. 2014. Gender moderates the effect of oxytocin on social judgments. Human Psychopharmacology 29:299–304.

Hollis, B., D. Houle, Z. Yan, T. J. Kawecki, and L. Keller 2014. Evolution under monogygous feminizes gene expression in Drosophila melanogaster. Nature Communications 5:3482.

van Honk, J., D. J. Schutter, P. A. Bos, A.-W. Kruijt, E. G. Lentjes, and S. Baron-Cohen 2011. Testosterone administration impairs cognitive empathy in women depending on second-to-fourth digit ratio. Proceedings of the National Academy of Sciences USA 108:3448–3452.

Horsthemke, B. 2014. In brief: genomic imprinting and imprinting diseases. The Journal of Pathology 232:483–487.

Hurler, P. W., A. Lin, O. A. Onur, M. X. Cohen, T. Baumgartner, S. Metzler, I. Drizbok et al. 2010. Oxytocin enhances amygdala-dependent, socially reinforced learning and emotional empathy in humans. Journal of Neuroscience 30:4999–5007.

Iemmola, F., and A. C. Ciani 2009. New evidence of genetic factors influencing social and mating system affect sex and species differences in immune function among vertebrates. Behavioural Processes 81:49–56.

Kemp, A. H., and A. J. Guastella 2011. The role of oxytocin in human affect: a novel hypothesis. Current Directions in Psychological Science 20:222–231.

Ketterson, E. D., V. Nolan Jr, and M. Sandell 2005. Testosterone in females: mediator of adaptive traits, constraint on sexual dimorphism, or both? The American Naturalist 166:585–589.

Klein, S. L. 2000. Hormones and mating system affect sex and species differences in immune function among vertebrates. Behavioural Processes 51:149–166.

Knobloch, H. S., and S. Grinevich 2014. Evolution of oxytocin pathways in the brain of vertebrates. Frontiers in Behavioral Neuroscience 8:31.

Kokko, H., and M. D. Jennions 2008. Parental investment, sexual selection and sex ratios. Journal of Evolutionary Biology 21:919–948.

Kong, A., V. Steinthorsdottir, G. Masson, G. Thorleifsson, P. Sulem, S. Besenbacher, A. Jonasdottir et al. 2009. Parental origin of sequence variants associated with complex diseases. Nature 462:868–874.

Kubiszanski, L. D., W. B. Mendes, A. A. Appleton, J. Block, and G. K. Adler 2012. A heartfelt response: oxytocin effects on response to social stress in men and women. Biological Psychology 90:1–9.

Lee, H. J., A. H. Macbeth, J. H. Pagani, and W. S. Young 2009. Oxytocin: the great facilitator of life. Progress in Neurobiology 88:127–151.

Lefebvre, L., S. Viville, S. C. Barton, F. Ishino, E. B. Keverne, and M. A. Surani 1998. Abnormal maternal behaviour and growth retardation associated with loss of the imprinted gene Mest. Nature Genetics 20:163–169.

Li, L., E. B. Keverne, S. A. Aparicio, F. Ishino, S. C. Barton, and M. A. Surani 1999. Regulation of maternal behavior and offspring growth by paternally expressed Peg3. Science 284:330–333.

Lupoli, B., B. Johansson, K. Unnas-Moberg, and K. Svennersten-Sjuanja 2001. Effect of suckling on the release of oxytocin, prolactin, cortisol, gastrin, cholecystokinin, somatostatin and insulin in dairy cows and their calves. The Journal of Dairy Research 68:175–187.

Lutchmaya, S., S. Baron-Cohen, P. Raggatt, R. Knickmeyer, and J. T. Manning 2004. 2nd to 4th digit ratios, fetal testosterone and estradiol. Early Human Development 77:23–28.

Mainguy, J., S. D. Côté, M. Festa-Bianchet, and D. W. Coltman 2009. Father-offspring phenotypic correlations suggest intralocus sexual conflict for a fitness-linked trait in a wild sexually dimorphic mammal. Proceedings of the Royal Society B: Biological Sciences 276:4067–4075.

Manck, E. H., and H. Ellegren 2009. Sex-linkage of sexually antagonistic genes is predicted by female, but not male, effects in birds. Evolution 63:1464–1472.

Manning, J. T., and B. Fink 2008. Digit ratio (2D:4D), dominance, reproductive success, asymmetry, and socioeconomicity in the BBC Internet Study. American Journal of Human Biology 20:451–461.

Manning, J. T., L. Barley, J. Walton, D. J. Lewis-Jones, R. L. Trivers, D. Singh, R. Thornhill et al. 2000. The 2nd-4th digit ratio, sexual dimorphism, population differences, and reproductive success: evidence for sexually antagonistic genes? Evolution and Human Behavior 21:163–183.

Marco, E. J., and D. H. Skuse 2006. Autism-lessons from the X chromosome. Social Cognitive and Affective Neuroscience 1:183–193.

Martin, A. M., H. Presseau-Gauvin, M. Festa-Bianchet, and F. Pelletier 2013. Male mating competitiveness and age-dependent relationship between testosterone and social rank in bighorn sheep. Behavioral Ecology and Sociobiology 67:919–928.

Martin, A. M., M. Festa-Bianchet, D. W. Coltman, and F. Pelletier 2014. Sexually antagonistic association between paternal phenotype and offspring viability reinforces total selection on a sexually selected trait. Biology Letters 10:20140043.

McCall, C., and T. Singer 2012. The animal and human neuroendocrinology of social cognition, motivation and behavior. Nature Neuroscience 15:681–688. Review.

McGrath, J. J., J. Hearle, L. Jenner, K. Plant, A. Drummond, and J. M. Barkla 1999. The fertility and fecundity of patients with psychoses. Acta Psychiatrica Scandinavica 99:441–446.

Medland, S. E., D. L. Duffy, A. B. Spurdle, M. J. Wright, G. M. Geffen, G. W. Montgomery, and N. G. Martin 2005. Opposite effects of androgen receptor CAG repeat length on increased risk of left-handedness in males and females. Behavior Genetics 35:735–744.
Genomic imprinting, sexual antagonism and hormones

Mendrek, A. 2007. Reversal of normal cerebral sexual dimorphism in schizophrenia: evidence and speculations. Medical Hypotheses 69:896–902.

Mendrek, A., and E. Stip 2011. Sexual dimorphism in schizophrenia: is there a need for gender-based protocols? Expert Review of Neurotherapeutics 11:951–959.

Mendrek, A., A. Mancini-Marie, C. Fahim, and E. Stip 2007. Sex differences in the cerebral function associated with processing of aversive stimuli by schizophrenia patients. The Australian and New Zealand Journal of Psychiatry 41:136–141.

Mendrek, A., N. Lakis, and J. Jiménez 2011. Associations of sex steroid hormones with cerebral activations during mental rotation in men and women with schizophrenia. Psychoneuroendocrinology 6:1422–1426.

Mervis, C. B., J. Dida, E. Lam, N. A. Crawford-Zelli, E. J. Young, D. R. Henderson, T. Onay et al. 2012. Duplication of GTF2I results in separation anxiety in mice and humans. American Journal of Human Genetics 90:1064–1070.

Mills, S. C., A. Grapputo, E. Koskela, and T. Mappes 2007. Quantitative measure of sexual selection with respect to the operational sex ratio: a comparison of selection indices. Proceedings of the Royal Society B: Biological Sciences 274:143–150.

Mills, S. C., A. Grapputo, I. Jokinen, E. Koskela, T. Mappes, T. A. Oksanen, and T. Poikonen 2009. Testosterone-mediated effects on fitness-related phenotypic traits and fitness. The American Naturalist 173:475–487.

Mills, S. C., A. Grapputo, I. Jokinen, E. Koskela, T. Mappes, and T. Poikonen 2010. Fitness trade-offs mediated by immunosuppression costs in a small mammal. Evolution 64:166–179.

Mills, S. C., E. Koskela, and T. Mappes 2012. Intralocus sexual conflict for fitness: sexually antagonistic alleles for testosterone. Proceedings of the Royal Society B: Biological Sciences 279:1889–1895.

Mitchem, D. G., A. M. Purkey, N. M. Grebe, G. Carey, C. E. Garver-Apgar, T. C. Bates, R. Arden et al. 2014. Estimating the sex-specific effects of genes on facial attractiveness and sexual dimorphism. Behavior Genetics 44:270–281.

Mokkonen, M., H. Kokko, E. Koskela, J. Lehtonen, T. Mappes, H. Martiskainen, and S. C. Mills 2011. Negative frequency-dependent selection of sexually antagonistic alleles in Myodes glareolus. Science 334:972–974.

Mokkonen, M., E. Koskela, T. Mappes, and S. C. Mills 2012. Sexual antagonism for testosterone maintains multiple mating behaviour. Journal of Animal Ecology 81:277–283.

Moon, Y. S., C. M. Smas, K. Lee, J. A. Villena, K. H. Kim, E. J. Yun, and H. S. Sul 2002. Mice lacking paternally expressed Pref-1/Dlk1 display growth retardation and accelerated adiposity. Molecular and Cellular Biology 22:5585–5592.

Morrow, E. H., and T. Connallon 2013. Implications of sex-specific selection for the genetic basis of disease. Evolutionary Applications 6:1208–1217.

Mottron, L., M. Dawson, I. Soulières, B. Hubert, and J. Burack 2006. Enhanced perceptual functioning in autism: an update, and eight principles of autistic perception. Journal of Autism and Developmental Disorders 36:27–43.

Muehlenbein, M. P., and R. G. Bribiescas 2005. Testosterone-mediated immune functions and male life histories. American Journal of Human Biology 17:527–558.

Muller, D. C. 2013. Second to fourth digit ratio: a predictor of disease in later life? Maturitas 75:1–2.

Muscatelli, F., D. N. Abrous, A. Massacrier, I. Boccaccio, M. Le Moal, P. Cau, and H. Cremer 2000. Disruption of the mouse Necdin gene results in hypothalamic and behavioral alterations reminiscent of the human Prader–Willi syndrome. Human Molecular Genetics 9:3101–3110.

Ober, C., D. A. Loisel, and Y. Gilad 2008. Sex-specific genetic architecture of human disease. Nature Reviews Genetics 9:911–922.

Oertelt-Prigione, S. 2012. The influence of sex and gender on the immune response. Autoimmunity Reviews 11:A479–A485.

Okabe, S., K. Kitano, M. Nagasawa, K. Mogi, and T. Kikusui 2013. Testosterone inhibits facilitating effects of parenting experience on parental behavior and the oxytocin neural system in mice. Physiology and Behavior 118:159–164.

Olff, M., J. L. Frijling, L. D. Kubzansky, B. Bradley, M. A. Ellenbogen, C. Cardoso, J. A. Bartz et al. 2013. The role of oxytocin in social bonding, stress regulation and mental health: an update on the moderating effects of context and interindividual differences. Psychoneuroendocrinology 38:1883–1894.

Ott, V., G. Finlayson, H. Lehnert, B. Heitmann, M. Heinrichs, J. Born, and M. Hallschmidt. 2013. Oxytocin reduces reward-driven food intake in humans. Diabetes 62:8418–8425.

Patten, M. M., and D. Haig 2009. Maintenance or loss of genetic variation under sexual and parental antagonism at a sex-linked locus. Evolution 63:2888–2895.

Patten, M. M., F. Ubeda, and D. Haig 2013. Sexual and parental antagonism shape genomic architecture. Proceedings of the Royal Society B: Biological Sciences 280:20131795.

Pennell, T. M., and E. H. Morrow 2013. Two sexes, one genome: the evolution of sexual and parental antagonism. Science 339:1310.

Peters, J. 2014. The role of genomic imprinting in biology and disease: an expanding view. Nature Reviews Genetics, 15:517–530.

Pierrehumbert, B., R. Torrisi, D. Laufer, O. Halfon, F. Ansermet, and M. Popovic 2010. Oxytocin response to an experimental psychosocial challenge in adults exposed to traumatic experiences during childhood or adolescence. Neuroscience 166:168–177.

Plagge, A., E. Gordon, W. Dean, R. Boiani, S. Cinti, J. Peters, and G. Kelsoy 2004. The imprinted signaling protein XL alpha s is required for postnatal adaptation to feeding. Nature Genetics 36:818–826.

Polli, A., S. Cassidy, B. Auyeung, and S. Baron-Cohen 2014. Uncovering stereotypy in women with autism: a latent class analysis. Molecular Autism 5:27.

Pointer, M. A., P. W. Harrison, A. E. Wright, and J. E. Mank 2013. Masculinization of gene expression is associated with exaggeration of male sexual dimorphism. PLoS Genetics 9:e1003697.

Porter, M. A., T. A. Shaw, and P. J. Marsh 2010. An unusual attraction to the eyes in Williams–Beuren syndrome: a manipulation of facial affect while measuring face scanpaths. Cognitive Neuropsychiatry 15:505–530.

Power, R. A., S. Kyaga, R. Uher, J. H. MacCabe, N. L. Kendler, P. McGuffin et al. 2013. Fecundity of patients with schizophrenia: evidence and speculations. Medical Hypotheses 82:1426–1439.

Rice, W. R., S. Gavrilets, and U. Friberg 2008. Sexually antagonistic “zygotic drive” of the sex chromosomes. PLoS Genetics 4:e1000313.
Rice, W. R., S. Gavrilets, and U. Fribег 2010. The evolution of sex-specific grandparental harm. Proceedings of the Royal Society B: Biological Sciences 277:2727–2735.

Rice, W. R., U. Fribег, and S. Gavrilets 2012. Homosexuality as a consequence of epigenetically canalized sexual development. The Quarterly Review of Biology 87:343–368.

Rilling, J. K., A. C. Demarco, P. D. Hackett, X. Chen, P. Gautam, S. Stair, E. Haroon et al. 2014. Sex differences in the neural and behavioral response to intranasal oxytocin and vasopressin during human social interaction. Psychoneuroendocrinology 39:237–248.

Robinson, M. R., J. G. Pilkington, T. H. Clutton-Brock, J. M. Pemberton, and L. E. B. Kruuk 2006. Live fast, die young: trade-offs between fitness components and sexually antagonistic selection on weaponry in Soay sheep. Evolution 60:2168–2181.

Robinson, E. B., P. Lichtenstein, H. Anchensarer, F. Happé, and A. Ronald 2013. Examining and interpreting the female protective effect against autistic behavior. Proceedings of the National Academy of Sciences USA 110:5258–5262.

Sabatier, N., G. Leng, and J. Menzies 2013. Oxytocin, feeding, and satiety. Frontiers in Endocrinology 4:35.

Sakurai, T., N. P. Dorr, N. Takahashi, L. A. McInnes, G. A. Elder, and J. D. Buxbaum 2011. Haploinsufficiency of G6f2α, a gene deleted in Williams Syndrome, leads to increases in social interactions. Autism Research 4:28–39.

Schaaf, C. P., M. L. Gonzalez-Garay, F. Xia, L. Potocki, K. W. Gripp, B. Zhang, B. A. Peters et al. 2013. Truncating mutations of MAGEL2 cause Prader–Willi phenotypes and autism. Nature Genetics 45:1405–1408.

Schaller, F., F. Watrin, R. Sturny, A. Massacrier, P. Szepetowski, and F. Muscatteri 2010. A single postnatal injection of oxytocin rescues the lethal feeding behaviour in mouse newborns deficient for the imprinted Magel2 gene. Human Molecular Genetics 19:4895–4905.

Schroederus, E., I. Jokinen, M. Koivula, E. Koskela, T. Mappes, S. C. Mills, T. A. Oksanen et al. 2010. Inter- and intra-sexual trade-offs between testosterone and immune system: implications for sexual and sexually antagonistic selection. The American Naturalist 176:E90–E97.

Schwarz, E., P. C. Guest, H. Rahmoune, L. Wang, Y. Levin, E. Ingudomnukul, L. Ruta et al. 2011. Sex-specific serum biomarker patterns for the imprinted Magel2 gene. Human Molecular Genetics 20:3922–3926.

Soni, S., J. Whittington, A. J. Holland, T. Webb, E. N. Maina, H. Boer, and D. Clarke 2008. The phenomenology and diagnosis of psychiatric illness in people with Prader–Willi syndrome. Psychological Medicine 38:1505–1514.

Sripada, C. S., K. L. Phan, I. Labuschgane, R. Welsh, P. J. Nathan, and A. G. Wood 2013. Oxytocin enhances resting-state connectivity between amygdala and medial frontal cortex. The International Journal of Neuropsychopharmacology 16:255–260.

Stearns, S. C., and J. C. Koella 2008. Evolution in Health and Disease. 2nd edn. Oxford University Press, Oxford.

Stearns, S. C., S. G. Byars, D. R. Govindaraju, and D. Ewbank 2010. Measuring selection in contemporary human populations. Nature Reviews Genetics 11:611–622.

Stearns, S. C., D. R. Govindaraju, D. Ewbank, and S. G. Byars. 2012. Constraints on the coevolution of contemporary human males and females. Proceedings of the Royal Society of London. Series B: Biological Sciences 279:4836–4844.

Stulp, G., B. Kuijper, A. P. Buunk, T. V. Pollet, and S. Verhulst 2012. Intralocus sexual conflict over human height. Biology Letters 8:976–978.

Swaab, D. F., J. S. Purba, and M. A. Hofman 1995. Alterations in the hypothalamic paraventricular nucleus and its oxytocin neurons (putative satiety cells) in Prader–Willi syndrome: a study of five cases. The Journal of Clinical Endocrinology and Metabolism 80:573–579.

Teatero, M. L., and C. Netley 2013. A critical review of the research on the extreme male brain theory and digit ratio (2D:4D). Journal of Autism and Developmental Disorders 43:2664–2676.

Temple, I. K., V. Shrubb, M. Lever, H. Bullman, and D. J. Mackay. 2009. Isolated imprinting mutation of the DLK1/GTL2 locus associated with a clinical presentation of maternal uniparental disomy of chromosome 14. BMJ Case Reports, 2009:pii:bcr06.2009.1997. DOI:10.1136/bcr.06.2009.1997.

Thompson, R. K., George, J. C. Walton, S. P. Orr, and J. Benson 2006. Sex-specific influences of vasopressin on human social communication. Proceedings of the National Academy of Sciences USA 103:7889–7894.

Tops, M., S. L. Koole, H. IJzerman, and F. T. Buismen-Pijlman. 2014. Why social attachment and oxytocin protect against addiction and stress: insights from the dynamics between ventral and dorsal corticostriatal systems. Pharmacology, Biochemistry, and Behavior 119:39–48.

Tran, U. S., S. Stieger, and M. Voracek 2014. Handedness and sex roles: mixed-handers are less sex-congruent stereotyped. Personality and Individual Differences 66:10–13.

Treffert, D. A. 2009. The savant syndrome: an extraordinary condition. A synopsis past, present, future. Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences 364:1351–1357.

Turelli, M., and N. H. Barton 2004. Polygenic variation maintained by balancing selection: pleiotropy, sex-dependent allelic effects and G × E interactions. Genetics 166:1053–1079.

Ubeda, F., and D. Haig 2003. Dividing the child. Genetica 117:103–110.

Uzefovsky, F., I. Shavev, S. Israel, A. Knafo, and R. P. Ebstein 2012. Vasopressin selectively impairs emotion recognition in men. Psychoneuroendocrinology 37:576–580.

Van Ijzendoorn, M. H., and M. J. Bakermans-Kranenburg, 2012. A sniff of trust: meta-analysis of the effects of intranasal oxytocin administration on face recognition, trust to in-group, and trust to out-group. Psychoneuroendocrinology 37:438–443.

Varrault, A., C. Guyed, A. Delalbre, A. Bellmann, S. Housami, C. Aknin, D. Severac et al. 2006. Zac1 regulates an imprinted gene network critically involved in the control of embryonic growth. Developmental Cell 11:711–722.

Veennema, A. H., R. Breedwold, and G. J. De Vries 2013. Sex-specific modulation of juvenile social play by vasopressin. Psychoneuroendocrinology 38:2554–2561.

Villanueva, C., S. Jacquier, and N. de Roux 2012. DLK1 is a somato-dendritic protein expressed in hypothalamic arginine-vasopressin and oxytocin neurons. PLoS One 7:e36134.

Volman, I., I. Toni, L. Verhagen, and K. Roelofs 2011. Endogenous testosterone modulates prefrontal-amygdala connectivity during social emotional behavior. Cerebral Cortex 21:2282–2290.

Weiser, M., A. Reichenberg, N. Werbeloff, H. Halperin, E. Kavitz, R. Yaffe, and M. Davidson 2009. Increased number of offspring in first degree relatives of psychotic individuals: a partial explanation for the persistence of psychotic illnesses. Acta Psychiatrica Scandinavica 119:466–471.

Weisman, O., O. Zagoory-Sharon, I. Schneiderman, I. Gordon, and R. Feldman 2013. Plasma oxytocin distributions in a large cohort of women and men and their gender-specific associations with anxiety. Psychoneuroendocrinology 38:694–701.
Weisman, O., O. Zagoory-Sharon, and R. Feldman 2014. Oxytocin administration, salivary testosterone, and father-infant social behavior. Progress in Neuro-Psychopharmacology and Biological Psychiatry 49:47–52.

Wells, J. C. 2003. Parent-offspring conflict theory, signaling of need, and weight gain in early life. The Quarterly Review of Biology 78:169–202.

Wilkins, J. F., and D. Haig 2003. What good is genomic imprinting: the function of parent-specific gene expression. Nature Reviews Genetics 4:359–368.

van Wingen, G., C. Mattern, R. J. Verkes, J. Buitelaar, and G. Fernández 2010. Testosterone reduces amygdala–orbitofrontal cortex coupling. Psychoneuroendocrinology 35:105–113.

Wright, N. D., B. Bahrami, E. Johnson, G. Di Malta, G. Rees, C. D. Frith, and R. J. Dolan. 2012. Testosterone disrupts human collaboration by increasing egocentric choices. Proceedings of the Royal Society of London. Series B: Biological Sciences 279:2275–2280.

Wu, X. L., D. Y. Yang, W. H. Chai, M. L. Jin, X. C. Zhou, L. Peng, and Y. S. Zhao 2013. The ratio of second to fourth digit length (2D:4D) and coronary artery disease in a Han Chinese population. International Journal of Medical Sciences 10:1584–1588.

Zhong, S., M. Monakhov, H. P. Mok, T. Tong, P. S. Lai, S. H. Chew, and R. P. Ebstein 2012. U-shaped relation between plasma oxytocin levels and behavior in the trust game. PLoS One 7:e51095.