Examining the evidence for major histocompatibility complex-dependent mate selection in humans and nonhuman primates

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Abstract: Compounds of the major histocompatibility complex (MHC) are integral for effective vertebrate adaptive immune response, and are also implicated as cues for sexual selection. The evidence for this is supportive of MHC-based preference for diverse and dissimilar mating partners, in a range of vertebrates from fish to nonhuman primates. However, the evidence for a similar role of these genes and the evolutionary benefits of their diversity in human mate choice has been more controversial. Here, we review the results of 34 primary studies on MHC-mediated mate choice in humans and nonhuman primates in an effort to understand what processes may underscore, or belie, such differences. Both human and nonhuman primate studies show evidence of mate selection for MHC-dissimilarity and diversity, with fewer results in support of mate selection for optimal diversity or for specific “good gene”. In general, stronger support comes from female-choice studies as opposed to male-choice studies (though male preferences and choice are investigated less often). This review suggests that the majority of mate choice results from contemporary human studies are consistent with our evolutionary history, but also reveals that only in humans do we find evidence for preference for mates with similar MHC composition. Overall, we show that contextual nuances, namely, population structure, multiple sensory cues that signal different information, and hormonal influences may explain the conflicting results observed for the role of MHC in human sexual preference, and argue for the exploration of such caveats in less well-studied systems to discern between general patterns and qualities unique to humans.

Keywords: major histocompatibility complex, sexual selection, olfaction, facial attraction, parasite resistance, inbreeding avoidance

Introduction

The role of the major histocompatibility complex (MHC) in mate selection has been investigated for almost 40 years with tantalizing results from both captive and wild populations in a diversity of systems including rodents, fish, birds, reptiles, and primates.1–5 While individual studies show a range of results, from no evidence to strong evidence for MHC-mediated mate choice, overall they proffer a general role for MHC diversity and dissimilarity in mate selection across vertebrates.6 In spite of relatively more effort placed on understanding the importance of MHC for human mating preferences, no such generalities have been distilled from human studies. Instead, excellent reviews have highlighted the equivocal and controversial nature of this topic.7,8

The difficulty in evaluating the role of MHC in mate choice across species is exacerbated by the differences in reproductive strategies between well-researched model species (eg, mice or sticklebacks that procreate early and abundantly) and humans...
(ie, live-slow-die-old strategy). To draw a clearer picture of what may influence patterns of MHC-mediated mate selection, it is worthwhile to re-evaluate the evidence across the primate order, comparing results from humans and nonhuman evolutionary siblings who share more life-history traits and reproductive strategies.

The goal of this review is to synthesize results from classic and contemporary studies of humans and other primates to consider human MHC-mediated mate selection from a primate perspective. We first provide a brief background of sexual selection theory and MHC biological functions, present the evidence for MHC-mediated mate selection in humans, and put those results into context using nonhuman primate studies. In so doing, we look for parallels among our own mating behaviors and those of our most recent ancestors in an effort to gain greater insight into the relationship between MHC diversity and dissimilarity, sexual selection, and offspring fitness. Finally, we discuss conclusions to be drawn from the available evidence and suggest research priorities for future investigations.

**Sexual selection theory and the MHC**

Darwin noticed that plumage, tails, and other exaggerated characters in male animals that should be detrimental to their survival, on the contrary, appeared to flourish with endless variety. He proposed that while these traits did not increase survival, they increased reproductive success by appealing to females. Sexual selection was thus considered an opposing force to natural selection, though more recent theories predict that they can reinforce one another. In a landmark paper, Hamilton and Zuk proposed that females may be selecting bright, showy males because these condition-dependent sexual characters indicate heritable genetic resistance to parasites. Parasites can have major impacts on fitness and can even determine host population dynamics, and animals that can detect and avoid mates likely to be parasitized would gain not only direct benefits but also indirect benefits, or “good genes” for their offspring.

The MHC, an ancient chromosomal region found in virtually all vertebrates, is the perfect emblem of “good genes” as it is associated with immune defense and mating preferences in a wide array of species, including humans, mice, and nonmodel species. The MHC represents a set of loci that plays a role in presenting self and foreign particles to T lymphocytes, enabling nonself-recognition for immune activation. MHC genes exist in a multigene family with two main subclasses, Class I and Class II, which are responsible for presenting antigenic peptides of intracellular and extracellular parasites, respectively. Genes within these classes are among the most polymorphic in the genome, and there is great variability in allelic diversity among loci (Figure 1). MHC alleles differ in their binding affinities for specific pathogen proteins (eg, binding supertypes) so that the degree of polymorphism at MHC loci is related to the variety of parasites recognized by the immune system. Among human populations, spatial patterns in MHC polymorphism have been explained in relation to both massive epidemics associated with human colonization and to regional parasite burdens, indicative of a strong role for parasites in shaping the microevolution of MHC. Individual MHC variability generates a selective advantage through enhanced clearance of and resistance to multiple infections. In addition, specific human MHC genotypes confer adaptive resistance to individual parasites, including malaria, hepatitis B, and HIV.

Humans are often considered exempt from some of the processes acting on “natural” animal communities due to our ability to modify our environment and control the impact of potential selection events (eg, immunization, sanitation, quarantine, etc). Nevertheless, humans have interacted with parasites throughout our evolutionary history and are currently known to be infected by over 1,400 parasitic species. On a global scale, parasites are currently responsible for nearly 50 percent of mortality in humans under the age of 45. Given the importance of parasites as selective pressure in natural systems, it follows that ubiquitous coevolving parasites should have played an equally powerful role in human biological evolution.

![Figure 1: Simplified map of the human MHC showing the range of allelic diversity.](http://www.ebi.ac.uk/imgt/hla/stats.html)

**Abbreviation:** MHC, major histocompatibility complex.
Role of MHC heterozygosity in mate selection and maintenance of fitness

Due to its role in immune response, MHC polymorphism is attributed to antagonistic coevolution with parasites. Although it is clear that allelic diversity at MHC loci is associated with pathogens, other studies have pointed to disassortative mating and kin recognition, either for inbreeding avoidance or to increase offspring genetic diversity, in maintaining heterozygosity and allelic variation across the MHC. Simulation studies show that MHC-mediated sexual selection can effectively maintain polymorphism over a wide range of demographic scenarios, and a comparative study across 117 mammal species indicated that a larger relative testes size (as a proxy for sexual selection) is universally associated with increased sequence diversity at a highly polymorphic Class II locus (DRB locus). In other words, there is both theoretical and cross-species support for the role of sexual selection in maintaining MHC diversity. Further, a number of studies have shown support for a direct mechanism allowing sexual selection on the MHC, including those focused on mice, fish, and humans that show animals can discern individuals with similar or dissimilar MHC genotypes based on olfactory cues. Allele-specific volatile byproducts of MHC-peptide binding are thought to be secreted in sweat and urine and these combinations reveal the allelic composition of the MHC genotype. Because of the possibility that some allele combinations are more beneficial than others under a given selection pressure, a condition-dependent selection mechanism could result in fitness advantage. However, it is not well understood how MHC-mediated cues for mate suitability impact fitness across changing selective landscapes.

Utility of MHC in mating preferences

MHC-dependent mating preferences can fall under four nonmutually exclusive categories:

1. Preference for dissimilar (or complementary) genotypes – to increase offspring heterozygosity and thus increase their immunocompetence and reduce the probability of immunopathology from deleterious recessive alleles. In this case, highly polymorphic MHC loci may serve as indicators of genome-wide diversity and a preference for MHC dissimilarity could also assist with inbreeding avoidance in low-diversity populations.

2. Preference for intermediate dissimilarity – to obtain optimal rather than maximal diversity for offspring, providing enough protection from pathogens while also avoiding the cost of having too many alleles that present self-peptides and deplete autoreactive T-cell diversity. Here, as in (1), the preferred mate depends on the genotype of the chooser; in this instance, females with a low number of MHC alleles choose mates with more (dissimilar) alleles, and more diverse females choose males with fewer alleles.

3. Preference for mates that possess a particular fitness-enhancing allele or supertype – to acquire direct benefits associated with high-quality mates and indirect benefits associated with heritable “good genes” for offspring. “Good genes” are predicted to be relatively rare in the population, stemming from the coevolutionary “arms race” with parasites. Specifically, parasite evolution is thought to drive the diversity of host MHC alleles in a negative-frequency dependent manner, leading to a time lag between a newly arisen MHC allele and widespread resistance. Hence, the most common alleles are predicted to be the most immunologically ineffective.

4. Preference for diverse (heterozygous) mates – to receive direct benefits associated with mates with better body-condition (e.g., higher intrasexual competitive ability, reduced risk of contracting infection, greater provisioning ability), or indirect benefits associated with “good-genes-as heterozygosity.” Theoretically, it has been shown that offspring of heterozygous males have higher relative fitness in fluctuating environments, and that female choice favoring heterozygous males is selected for under most scenarios. The preference for more heterozygous males may also be confounded with a preference for specific “good genes” or for similar genotypes, since more heterozygous individuals, on average, will harbor more rare alleles and will share more alleles.

Mate selection for MHC or inbreeding avoidance?

There are significant fitness consequences of inbreeding, and examples in wildlife include lower birth weight, survival, reproduction, and resistance to disease. Mouse studies show mice shun mates with parental genotypes to avoid incestuous mating, suggesting that mice use MHC-associated cues primarily for inbreeding avoidance, rather than increasing immunogenetic diversity, per se. On the other hand, experimentally controlled natural matings of red junglefowl (Gallus gallus) showed that MHC, not background relatedness,
drives female sperm use bias for MHC-dissimilar males. In humans and other primates, a greater risk of pregnancy failure is associated with high MHC similarity among mates. In addition, offspring conceived by MHC-similar spouses may be at greater risk for autoimmune disease associated with homozygosity.

In addition to the consequences of inbreeding connected to MHC allele sharing between mates, partners sharing specific combinations of MHC alleles linked to killer immunoglobulin-like receptor (KIR) genes may be at greater risk for pregnancy disorders. KIRs are receptors expressed by natural killer cells and T cells that bind to MHC Class I glycoproteins expressed on all nucleated cells; these interactions activate or inhibit immune response to varying degrees, influencing susceptibility to and protection against infection and autoimmunity. Specific MHC-KIR combinations can result in excessive inhibition or activation of natural killer cells and influence pregnancy outcomes; in particular, KIR A haplotypes lack genes that encode receptors for MHC-C2 ligands, which would normally help activate natural killer cells. This is important because the activation of natural killer cells helps form the placenta and enlarge maternal blood vessels, providing nourishment to the growing fetus. Mothers homozygous for KIR A have increased risk for preeclampsia, a condition during pregnancy characterized by endovascular abnormality that can develop into deadly seizures (eclampsia) and the main cause of pregnancy-related mortality worldwide. Suggestive evidence of selection against inhibitory combinations of MHC-KIR haplotypes comes from inverse frequencies of KIR A haplotypes and MHC-C2 in a study conducted across 58 human populations. In effect, strong selection against hazardous maternal-fetal genotypes is driving the availability of those genotypes in space. Other evidence of coevolution between MHC and KIR genes is related to their role in disease risk and the survival benefit conferred by these immune receptors may preserve their allelic diversity in spite of their reproductive disadvantage. Nonhuman primates and humans both have greater diversity at KIR genes than mice, suggesting a shared reliance on these genes for immune functioning that occurred relatively recently. MHC-KIR combinations as well as epistatic interactions with other immunity-related genes may be underlying the associations detected in primate MHC mate selection studies.

High costs of pregnancy make female mammals generally choosier about potential mates; in the case of humans, women tend to be more sexually cautious than men, but the goal of choosiness is still unclear. This is because mate selection for MHC diversity may be simply a by-product of preference for genome-wide diversity. In this case, MHC genes may be acting as a marker for assessing relatedness between individuals, and recent genomic studies have shown that Nigerian Yoruban couples with high MHC similarity also have high genome-wide relatedness. This lack of independence between genomic regions can affect interpretations of the strength of MHC-mediated mate choice, and has only recently been examined. Nonhuman primate studies in the wild, obligated to rely on neutral markers for parentage assignment, have usefully incorporated both neutral and MHC relatedness to reveal significant relative importance of the MHC in mate selection.

Related to the problem of linkage between MHC and genome-wide diversity within individuals is the complication of population structure. MHC allele frequencies vary across ethnic groups and populations, and self-preference of subpopulations is characterized by distinctive frequencies of alleles. If the study subjects express racial or ethnic mate preferences that are correlated with MHC allele frequencies, then substructuring of alleles within a larger population may erroneously show selection for MHC similarity or dissimilarity. Most studies have acknowledged this caveat and have tried to control it by restricting their study population to individuals of similar ancestry. For example, homogeneous origins have been demarcated based on race (ie, skin color), and inferred from national or ethnic surnames. However, many human populations have substantial, yet not apparent, admixture, making it difficult to standardize samples. Further, preference for dissimilar or similar individuals may just as well be a real effect moderated by the diversity of the potential mate pool. The majority of studies finding MHC-dissimilarity preferences in humans and other primates that were statistically significant have used relatively homogeneous and isolated populations. For example, the Hutterite population showing evidence for disassortative MHC mate choice in Ober et al’s classic study descended from approximately 68 founders from the mid-1700s Russia. In contrast, studies finding preferences for MHC similarity have been conducted on more genetically variable populations or populations with strong sociocultural substructuring.
populations, they have a preference for mates with a greater number of shared MHC alleles to balance the competing costs of inbreeding and outbreeding.\textsuperscript{54}

**Studies of mate preference cues in humans**

One of the most controversial issues regarding MHC-mediated preferences in humans is the mechanism of detecting genetic diversity in others. The specificity of MHC composition may be ascertained through the volatile peptides from urine, sweat, and axillary (armpit) glands, and as mentioned previously, there is increasing support from studies representing a variety of taxa that animals can identify MHC genotypes based on olfactory cues.\textsuperscript{42,46–49} The MHC-regulated differences in bacterial flora composition between individuals could allow for indirect association between MHC and body odors,\textsuperscript{104} perhaps where commensal microflora metabolize secreted MHC-peptide fragments and excrete them as volatile odors.\textsuperscript{41} Exciting recent evidence suggests that genetically determined microbial communities can influence scent marks of mice,\textsuperscript{105} and that more individual MHC Class II diversity is associated with less diverse gut microbiota in wild sticklebacks.\textsuperscript{106} This also highlights another potential reason to maintain optimal rather than maximal diversity – to preserve symbiotic microbiome diversity.

It is reasonable to ask if modern human practices such as wearing perfumes and hygienic routines make any historic MHC influence obsolete.\textsuperscript{97} Using 18 natural ingredients that have been in use as fragrances for over 5,000 years (eg, cinnamon, sandalwood, jasmine, and rose), Milinski and Wedekind experimentally tested the prediction that perfume preference for one’s self is specific to MHC genotype.\textsuperscript{107} They found significant correlations between MHC genotype and perfumes selected “for self”, but not “for partner”, in line with the hypothesis that instead of disguising or masking natural body odor, perfumes may function to amplify MHC-related individual body odors. Interestingly, in a recent study, Milinski et al used artificially synthesized MHC-peptide ligands and a functional magnetic resonance imaging (fMRI) machine to test if humans possess the sensory facility to recognize MHC-associated olfactory cues. They found that female students preferred the scent of “self” to “nonself” peptides applied to their arms, and showed these preferences in brain activation scans, suggesting the presence of an internal reference for MHC genotype detection.\textsuperscript{108} This may help explain how humans can identify MHC-dissimilar mates.

In the first direct test of MHC-mediated odor preference in humans, Wedekind et al found MHC-dissimilar preferences in female Bern University students who were asked to rate the pleasantness of T-shirts worn two consecutive nights by male coeds.\textsuperscript{99} Not all odor-preference studies corroborate another (Figure 2), and it has been suggested that variation in methodologies may explain the differences in their findings.\textsuperscript{7} For example, the source of odor stimulus (T-shirts in contact with the majority axillary region vs cotton sachets in contact with the chest), treatment of the stimulus (same-day presentation to rats vs freezing and presentation weeks later), rating protocol and phrasing of survey questions, and number and identity of the MHC loci typed all may have nontrivial effects on the strength, if not direction, of the results.\textsuperscript{7}

While odor is an important cue for female sexual preference, it is just one of multiple cues used to evaluate potential mates.\textsuperscript{109} An alternative or complementary mechanism for the relationship between MHC and mate choice in humans comes from female perception of male facial attractiveness. In a study of female mate choice, males who were heterozygous at three key MHC loci were judged as “more attractive” than males homozygous at single or multiple loci.\textsuperscript{96} Also, the skin of these males alone (in the absence of facial context) was judged as “healthier” than that of homozygotes.\textsuperscript{96} On a population level, the association between attractiveness and perceived immune status is manifested through an increased value on physical attractiveness in cultures exposed to high rates of parasitism.\textsuperscript{109} Data from 29 cultures show that areas with relatively higher pathogen prevalence value a mate’s physical attractiveness more than people in areas with little pathogen incidence.\textsuperscript{109} The avoidance hypothesis supposes that individuals avoid mating with persons capable of spreading infection,\textsuperscript{110} and predicts that physical attractiveness may be a signal of current health and lack of transmissible parasite load.\textsuperscript{109} Alternatively, relative beauty may be an indication of mate quality and good genes, and sexual selection may discriminate individuals with regard to their pathogen resistance.\textsuperscript{111} For example, two features that are cross-culturally associated with male attractiveness are prominent cheekbones and chins.\textsuperscript{112} As these traits develop during and after puberty in response to testosterone production\textsuperscript{113} and testosterone is an immunosuppressant,\textsuperscript{114} individuals who cannot afford the cost of a high level of testosterone production would display this honest advertisement of their gene quality in their facial features. Thus, facial attractiveness may be an honest indicator of “good genes” and preferences for good-gene indicator traits have been shown
to be relatively robust and consistent across cultures.115
Results so far have been inconclusive about the relative importance of sensory information on mating preferences, and multiple cues from different sensory modalities may be integrated for determining an optimal mate (Figure 3). For example, assortative preferences for facial similarity and disassortative preferences for odor may result in optimal degree of outbreeding.95,116

One explanation for the inconsistency in the strength and direction of odor and facial preference studies is variance in the context under which experiments were performed. Female primates (including humans) undergo periodic ovulation cycles with increased sexual behavior at mid-cycle, presumably in rhythm with ovulation.117 In naturally cycling women, changes occur in the concentration levels of several hormones, and these fluctuations are related to changes in women's attractiveness to men, as well as to women's sexual desire and motivation for sexual attention.118

It has been argued that females' estrous mate preferences

Figure 2 Summary of results for MHC-associated mate selection in humans.
Notes: Significant tests for preferences for (A) dissimilarity and (B) diversity and/or heterozygosity vary by the "chooser" investigated. Significance also varies within sex, and (C) shows the number of significant results (for both dissimilar and diversity preferences) according to women taking contraceptive pills and those normally cycling. The solid line in (A and B) denotes nonsignificant results. Trend refers to results with P-values between 0.05 and 0.1. *Jacob et al's work54 supports a preference for an intermediate number of shared alleles. Data from the recent review by Havlíček and Roberts,7 and additional studies were identified up from 2008 to January 2015 via Web of Science using the topic "HLA" or "MHC" or "major histocompatibility complex" and "mate choice" or "mate selection" or "mate preference" and searching within results for "human". Abbreviation: MHC, major histocompatibility complex.

Figure 3 Information from multiple phenotypic cues (eg, visual and odor) may be integrated to inform mate preference.
weigh cues and signals of intrinsic genetic benefits more heavily than outside the fertile phase. Interestingly (and perhaps disturbingly), Wedekind et al’s study on olfactory preference among Bern University students found a reversed preference, for odors of MHC-similar men, in a subset of women who were taking contraceptive pills. Subsequent studies have found similar contrasting preferences between pill- and nonpill-using females, whereby in pill-users the predicted dissimilarity preference was either nonsignificant, or reversed (Figure 2C). Contraceptive pills (and other hormone-supplementation methods) developed to avoid pregnancy work in various biochemical ways to prevent either ovulation, fertilization, or implantation of an embryo; however, both the mechanism through which they act and the subsequent effects on female mood and sexuality is surprisingly poorly understood. These reviews also point out that there are intrinsic difficulties in the design of empirical studies on hormone effects on mate choice in humans, such as the fact that cross-sectional studies do not account for the under-representation of women who discontinue pill use after experiencing ill effects, rendering it difficult to confirm the pill’s direct effects on female behavior. Some authors go so far as to argue that after confounding factors and appropriate response variables have been controlled, there remains no convincing evidence that the menstrual phase has any effect on the detection of mate genetic quality at all. Therefore, intra-individual preference consistency should be confirmed, and moderating factors should be investigated.

Human vs nonhuman primate populations

In common with many other primates, human females are thought to be sensitive to potential cues of indirect benefits. Strong selection for indirect benefits of mate choice typically occurs in polygynous and polyandrous mating systems, and there is evidence of a polygamous human heritage from comparative studies of sexual dimorphism and Y-chromosome diversity, reviewed by Roberts and Havlíček. Indeed, nonhuman primate studies are necessary to fully understand the evolution of MHC-mediated mate selection in humans because they allow us to track the evolutionary trajectory of the preference and to study the selective pressures favoring the traits’ emergence and maintenance. Complementing similar behavioral traits, nonhuman primates share a number of genetic traits, including orthologous MHC genes, with humans. Further, identical MHC allele motifs are associated with resistance to related pathogens, for example, to simian immunodeficiency virus (SIV), the predecessor of HIV. One difficulty with the study of human MHC-mediated mate choice is that current humans have ways of manipulating our microbiota, odor, and physical attributes, modifying signals of condition. Our relatively recent environmental modifications have dramatically increased human lifespan and health while at the same time increasing the difficulty for detecting evolutionary associations between mating preferences and fitness benefits. Our ancestral condition may be most closely reflected in remote tribes, like those of the Amazon that are chronically infected with marginally debilitating macroparasites like helminthes, transiently infected with bacteria (staphylococci, streptococci, diarrheal bacteria), and occasionally infected with endemic zoonotic diseases. However, the very thing that makes these populations so important to study, their relative isolation, is exactly what makes these population studies so challenging, namely, their inaccessibility. In lieu of observing our prehistoric ancestors, nonhuman primates can serve as a window to the past. Nonhuman primates are thought to represent human origins more closely than current human populations, and have potentially fewer confounding cultural, socioeconomic, and technological factors (eg, antibiotics, perfume, deodorants). Perhaps the most glaring example of our differences is the recent breakdown in limits to dispersal, facilitated by modern modes of travel. Whereas many prior criticisms have emphasized interracial/social influences on human assortative mating behavior, it is likely that the population structure of most wild or semi-wild primates more closely resembles that of our recent ancestors who only rarely dispersed over long distances. On the other hand, detecting selection for optimal MHC diversity may be difficult if the population is already at its equilibrium diversity optimum. Studying communities with a high proportion of disparate migrants (eg, college student studies) may therefore offer some advantage along with the challenge of contemporary signal distortion.

Results

Support for the specific functions of MHC-mediated mate choice

We report here that the vast majority of the 34 primary studies on MHC-mediated mate choice in primates have focused on humans (n=27), with just seven studies yet published on various other species including lemurs (Microcebus murinus and Cheirogaleus medius), rhesus macaques (Macaca mulatta), mandrill (Mandrillus sphinx), baboons (Papio ursinus), and...
the Sichuan snub-nosed monkey (Rhinopithecus roxellana) (Table S1). Nevertheless, several important patterns emerge from comparing results across these studies (Figures 2 and 4). One is that nearly half of all tests in both humans and nonhuman primates show a significant association between MHC genotype and mate selection. Another is that evidence for MHC-mediated mate preference is more often significant when tested in females than when tested in males or inferred from mating pairs. However, studies on female preference were twice as numerous as studies on male preference (human: 20 F, 9 M; nonhuman primates: 10 F, 5 M) (Table S1). This may reflect an expectation bias determining the allocation of study resources when preferences for indirect benefits, if observed at all, are predicted to be stronger for females and so easier to detect. Lastly, when evidence for each of the main hypotheses concerning MHC-involvement in sexual selection (listed below) has been lacking or conflicting, similar contextual factors present reasonable alternative hypotheses.

Selection for similarity/dissimilarity among mates

Tests of preference for more MHC-similar or dissimilar mates revealed approximately half of all studies with significant results showing strong evidence of selection for dissimilarity across humans and nonhuman primates (Figures 2A and 4A). In humans, men showed significant facial and odor preferences only for MHC-dissimilar women.39,114 Interestingly, several human studies also showed evidence of preference for MHC partner similarity, potentially reflecting assortative mating amongst widely mixed populations. Preferences for similarity included women’s facial and odor preferences for MHC-similar men,95,96,98 and couples displaying more MHC-similarity than randomly assigned partners.54,85–88,93,103 However, excessive MHC-similarity may be undesirable and was shown to predict deviation from fidelity in both humans and other primates. Female fat-tailed dwarf lemurs (Cheirogaleus medius) with extra-pair mates tended to have higher supertype overlap with their social mates than did females who did not have extra-pair copulations.5 Similarly for humans, women’s stronger desire for extra-pair partners and sexual dissatisfaction with current mates was associated with more MHC allele overlap with those partners.100

Selection for intermediate dissimilarity and diversity

Evidence for optimal MHC-dissimilarity/diversity in mates is weak, though the majority of human and other primate studies have not explicitly tested for it. Two nonhuman primate studies tested for mate choice for optimal diversity and found nonsignificant results.89,135 and two human studies found suggestive evidence of selection for optimal offspring diversity. Derti et al compared genome-wide single nucleotide polymorphisms (SNPs) between European–American mates and found some evidence that partners avoid extreme similarity at MHC genes89 (though another study using overlapping datasets have interpreted these results to support MHC-dissimilarity136); Jacob et al found that women preferred odors of men with more alleles matching those inherited paternally, but not maternally,44 suggesting that women employ an odor preference that provides them optimal allele matching.

Selection for mates with greater MHC diversity

Where an effect was detected, both humans and nonhuman primates universally showed preferences for MHC-indicating cues (in humans) or choice of actual mates (humans and other primates), signaling more diverse MHC haplotypes or supertypes than their own (or than other individuals in the population) (Figures 2B and 4B).85,89,96,99,134,137 Additionally, this relationship has been detected solely as female preference for males. Similar findings were detected by Kamiya et al.

**Figure 4** Summary of results for MHC-associated mate selection in nonhuman primates.

**Notes:** Significant tests for preferences for (A) dissimilarity and (B) diversity and/or heterozygosity vary by the “chooser” investigated. The solid line in (A) and (B) denotes nonsignificant results. Data from the recent review by Setchell and Huchard,43 and additional studies were identified up from 2009 to January 2015 via Web of Science using the topic “MHC” or “major histocompatibility complex” and “mate choice” or “mate selection” or “mate preference” and searching within results for “primate”.

**Abbreviation:** MHC, major histocompatibility complex.
who recently conducted a meta-analysis across vertebrates and found evidence for diversity (and dissimilarity), but only when multiple loci within individuals were considered. This may indicate the presence of hotspots of selection on specific components of the MHC, or that using more loci increases power to detect significant associations.

Selection for “good genes”

To date, there is only limited support for mate selection directed by “good genes” in humans and other primates. In a Dutch study, specific combinations of alleles had significantly increased frequencies in spouses than the background population, though preferential associations of disparate MHC alleles could not be distinguished from the linkage of alleles within haplotypes. Another study indicated selection for “good genes”, although in this case, common alleles, rather than rare ones, predicted the attractiveness of women’s scents to men. Rare alleles are expected to be advantageous in parasite defense, but as they are projected to increase in frequency until their commonness hinders their utility, allele frequency at any given time is a poor predictor of their current quality. In fat-tailed dwarf lemurs, specific MHC supertypes were more common in chosen social mates than random males, but three studies in other nonhuman primates found no evidence of selection for specific MHC supertypes. Additional investigations are needed to help resolve the generality of “good genes” preferences.

Conclusion, caveats, and recommendations

MHC-mediated mate choice in humans is a topic of great interest to biological, psychological, and social sciences, but is also highly controversial. In fact, of the 27 studies on human MHC-associated mate selection reviewed here, one-third (nine studies) have provoked direct criticism and published correspondence between authors and critics (Table S1). These discussions conclude that the most critical factor to consider is context, including within and among individuals and within and among populations. We briefly mention two caveats for the results of MHC-dependent mate selection studies. The first is that nonhuman primate studies have mostly focused on mate choice from pairs with one or more offspring and looked for differences between the pair components and the larger population; in contrast, many of the human studies were based on the mate preference of individual subjects for visual or olfactory cues. The eventual choice for a mate may be modified from the original preference by costs of choosiness (eg, from intersexual conflict or intrasexual competition).

The second caveat is that mate choice inferred from paternity in studies may potentially be biased in favor of genetic dissimilarity if differential embryo mortality caused by MHC similarity and/or inbreeding restricts childless pairs from study. Only half of the human studies based on paired couples included any information on the presence of children, and all nonhuman primate pairs, which only found evidence for MHC-dissimilarity, were defined by the presence of at least one offspring. Yet a degree of preference for MHC similarity in human couples may reflect ancestry-related mate selection and not necessarily MHC-mediated selection. Studies that use many individuals and control for genome-wide effects to isolate MHC-mediated mate selection will become more frequent as new methodologies (including large-scale MHC genotyping of nonmodel primate species and genome-wide SNP data) become more feasible.

The intense scrutiny of the evidence concerning MHC-mediated mate selection in humans can help serve as a guide for the design and interpretation of studies in other, particularly nonmodel, organisms. While many human behaviors can be argued as entirely “unnatural”, they also highlight the importance of complicating factors (eg, hormone cycles and migration rates) that researchers of nonmodel species should be aware of within their own systems (eg, variance in social structure, population substructure, or population density), because they may influence the strength and direction of MHC-mediated mate selection, real or apparent. Perhaps the most salient take-home message from human studies is that MHC-associated mate selection differs between populations. Even setting aside the complications from sociocultural and economic conditions, human populations do show evidence for MHC-mediated mate selection, but the mode and direction of that selection are not consistent across populations. The variance in results can more adequately be delineated in a meta-analysis, and a clearer pattern of the drivers of differences in preference and choice await a quantitative comparison across studies.

In conclusion, we find that human studies are controversial with much debate over discordant results, but a thorough examination of findings from primate studies shows that this inconsistency is entirely consistent. The commonalities among the majority of human and nonhuman primate studies appear to be that the MHC is likely to be involved in mate-choice decisions for many populations, evidence for MHC-mediated mate selection is stronger for females than males, and the variation in results reflects issues surrounding...
multiple cues, population structure, and social influences. More generally, lack of evidence for mate preference or choice may reflect a greater cost than gain for expressing a preference, and search costs, opportunity costs, and trade-offs with other preferences are dependent on context and demography. Greater exploration across nonhuman primate species with varying population and social structures, in addition to testing for preferences based on individual cues, will undoubtedly feed back into helping understand the relationship between MHC composition and sexual selection in humans.

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Disclosure

The authors report no conflicts of interest in this work.

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**Supplementary material**

Table S1 Compilation of 34 primary literature reports on MHC and mate choice in primates and humans; for each primary reference, taxon, support for specific mechanisms, mate choice cue, and whether offspring were produced for mates is indicated, as well as whether a direct response or commentary was published.

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Females prefer dissimilar male scents, and males prefer dissimilar female scents. Human MHC class II heterozygous males are more attractive than homozygous males and heterozygous females. In a semi-free-ranging primate population, human mate choice depended on MHC similarity. MHC-dependent mate choice was observed in the HapMap European American dataset, supporting the hypothesis of MHC-dependent mate choice. Coetzee et al. found that common HLA alleles associated with health, but not facial attractiveness. Laurent et al. observed that MHC-dependent mate choice in humans: why genomic patterns from the HapMap European American dataset support the hypothesis. Derti et al. responded to “MHC-dependent mate choice in humans: why genomic patterns from the HapMap European American dataset support the hypothesis” by Derti and Roth. Garver-Appgar et al. found that major histocompatibility complex alleles, sexual responsivity, and unfaithfulness in romantic couples.

References


40. Stewart-Williams S, Thomas AG. The ape that thought it was a peacock: does evolutionary psychology exaggerate human sex differences? *Psychol Inq.* 2013;24(3):137–168.