

Evidence of Amino Acid Diversity–Enhancing Selection within Humans and among Primates at the Candidate Sperm-Receptor Gene *PKDREJ*

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Sperm-egg interaction is a crucial step in fertilization, yet the identity of most interacting sperm-egg proteins that mediate this process remains elusive. Rapid evolution of some fertilization proteins has been observed in a number of species, including evidence of positive selection in the evolution of components of the mammalian egg coat. The rapid evolution of the egg-coat proteins could strongly select for changes on the sperm receptor, to maintain the interaction. Here, we present evidence that positive selection has driven the evolution of *PKDREJ*, a candidate sperm receptor of mammalian egg-coat proteins. We sequenced *PKDREJ* from a panel of 14 primates, including humans, and conducted a comparative maximum-likelihood analysis of nucleotide changes and found evidence of positive selection. An additional panel of 48 humans was surveyed for nucleotide polymorphisms at the *PKDREJ* locus. The regions predicted to have been subject to adaptive evolution among primates show several amino acid polymorphisms within humans. The distribution of polymorphisms suggests that balancing selection may maintain diverse *PKDREJ* alleles in some populations. It remains unknown whether there are functional differences associated with these diverse alleles, but their existence could have consequences for human fertility.

Sperm-egg recognition is crucial for successful fertilization and therefore is key to reproduction.¹ A priori, it could be expected that fertilization and the proteins that mediate it would be highly conserved. However, rapid evolution and striking divergence of numerous reproductive proteins between closely related species have been found in diverse taxa.² Both sperm and egg gamete-recognition molecules exhibit adaptive evolution, suggesting some form of coevolutionary chase. This rapid, adaptive divergence could be the reason the process of sperm-egg recognition often exhibits species specificity.

The mammalian egg coat comprises at least three glycoproteins with zona pellucida (ZP) domains: ZP1, ZP2, and ZP3.^{3,4} The ZP3 protein was thought to be the primary inducer of the sperm acrosome reaction in mouse,^{5,6} although recent studies suggest that sperm recognize a three-dimensional structure that may require all three ZP proteins.^{3,7} Positive selection has been shown to promote the rapid divergence of ZP2 and ZP3.^{8,9} Two clusters of amino acids reported to be involved in the species-specific induction of the acrosomal reaction in mouse^{5,6} have evolved by positive selection, indicating that the selective pressure may be related to sperm-egg interaction.^{8,9} The identity of the sperm receptor that interacts with mammalian ZP proteins remains unknown and controversial; a minimum of 10 candidates have been proposed.^{10,11} If ZP3 and ZP2 egg proteins have undergone rapid evolution, sperm receptor or receptors for these proteins might also have experienced significant selective pressure to maintain the func-

tional relationship with their cognate protein or protein complex. Such coevolution has been demonstrated for interacting sperm-egg recognition proteins in abalone.¹² Hence, evidence of strong, positive selection on a candidate sperm surface provides an additional characteristic that might suggest its further study for a potential receptor role in mammalian fertilization.

We examined one of the sperm-receptor candidate proteins, *PKDREJ*, for evidence of rapid adaptive evolution among primates. This large, intronless gene encodes an ~8-kb transcript in humans.¹³ Its sequence reveals a significant region of homology with members of the PKD gene family. Members of this gene family code for a number of membrane-bound proteins that form calcium ion channels and play important roles in cell-to-cell and cell-to-extracellular matrix interactions.¹⁴ The family includes the gene responsible for human polycystic kidney disease, polycystin-1,¹⁵ and for *suREJ*, a sea urchin protein important in fertilization.¹⁶ *suREJ* localizes to the plasma membrane over the acrosomal vesicle. Experimental evidence suggests that *suREJ* binds to the fucose sulfate polymers of the sea urchin egg jelly and initiates the acrosomal reaction by activating gated calcium channels. This is likely done through interactions among the two C-type lectins, the REJ domain, and the two forms of the polymer.¹⁷ All PKD family genes appear to contain a subunit of a non-specific cation channel,¹³ the REJ domain followed by a GPS unit, a transmembrane region, the LHD/PLAT domain, and a variable number of transmembrane domains.

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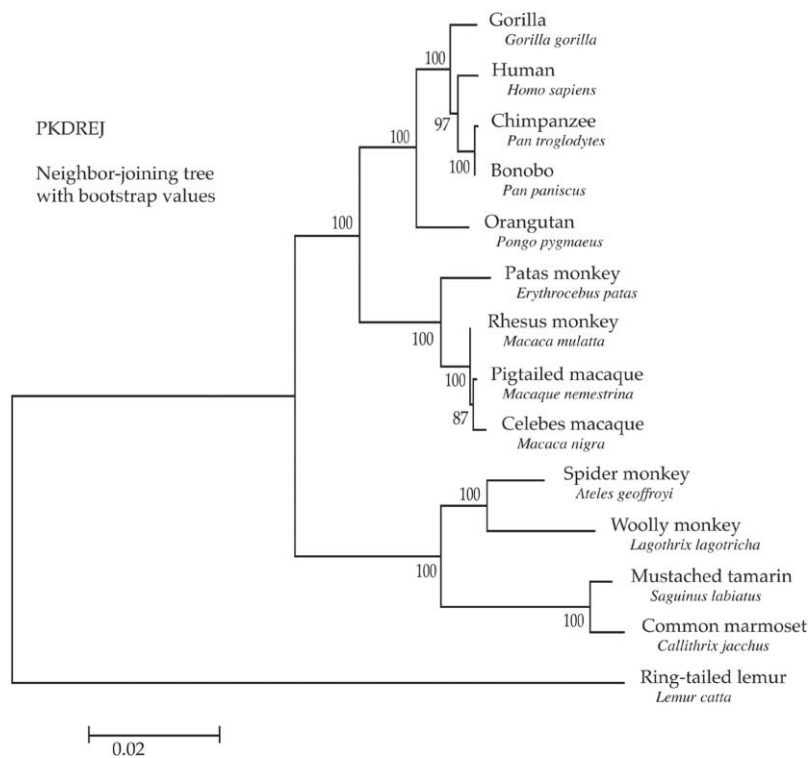


Figure 1. Neighbor-joining tree of the *PKDREJ* locus from the 14 primate species studied. Scale bar represents Kimura 2-parameter distances.

Four lines of indirect evidence support the idea that *PKDREJ* might function as a sperm surface receptor for the egg ZP. First, *PKDREJ* is a homologue to sea urchin REJ molecules demonstrated to be involved in sperm-egg interaction. In a phylogenetic analysis, *PKDREJ* groups with *suREJ* compared to the other mammalian PKD genes (W. J. Swanson and J. Gatesy, unpublished material). Second, the gene encoding *PKDREJ* shows testis-specific expression in both mouse¹⁸ and human,¹³ consistent with the expectation for a sperm protein involved in fertilization. Third, the PKDREJ protein localizes to the extracellular portion of the acrosomal region of spermatozoa.¹⁸ This is the region where sperm receptors for the ZP are expected to be located.¹⁸ Fourth, functional characterization of PKDREJ modulation of G-protein signaling is consistent with its potential role in the ZP-induced acrosome reaction.¹⁹ In the current study, we tested whether *PKDREJ* shows signs of rapid adaptive evolution, as has been observed in mammalian egg-coat proteins^{8,9} and other sperm and egg proteins.²⁰

We determined the *PKDREJ* sequence from 13 nonhuman primates and 48 humans, to conduct statistical tests of inter- and intraspecific sequence variation. We found evidence that the divergence of *PKDREJ* has been promoted by adaptive evolution. Analysis of variation in the d_N/d_S ratio among sites pinpointed particular codons that show signs of positive selection, indicating that they may be important for the function of PKDREJ. By examination

of variation both within and between species, we have documented intra- and interspecies diversifying selection. Our results suggest that PKDREJ shows evolutionary characteristics expected for a sperm receptor for the mammalian egg-coat ZP glycoproteins.^{13,18}

Material and Methods

DNA Samples

A panel of genomic DNAs from 13 nonhuman primate species was used. Animals were chosen to be sufficiently diverged to test for the presence of positive selection but close enough to allow ease of sequence alignment. Identifiers in parentheses indicate the sample numbers from Coriell Cell Repositories' NIA Aging Cell Repository DNA Panel-Primate Panel: Phylogenetic PRP00001. Samples included chimpanzee (*Pan troglodytes*; NG06939), bonobo (*Pan paniscus*; NG05253), gorilla (*Gorilla gorilla*; NG05251), Sumatran orangutan (*Pongo pygmaeus*; NG12256), patas monkey (*Erythrocebus patas*; NG06116), Celebes crested macaque (*Macaca nigra*; NG07101), pigtailed macaque (*M. nemestrina*; NG08452), rhesus monkey (*M. mulatta*; NG07109), woolly monkey (*Lagothrix lagotricha*; NG05356), black-handed spider monkey (*Ateles geoffroyi*; NG05352), red-chested mustached tamarin (*Saguinus labiatus*; NG05308), common marmoset (*Callithrix jacchus*; NA07404), and ring-tailed lemur (*Lemur catta*; NG07099). To survey human polymorphisms for PKDREJ, we used DNAs from 48 individuals who comprised an African American panel of 25 individuals (NA17101–NA17140) and a CEPH European panel of 23 individuals (NA06990, NA07019, NA07348, NA07349, NA10830–

Table 1. Maximum-Likelihood Estimates of Selection

Model and Parameters	l	Positively Selected Sites
M0 (one ratio): $d_N/d_S = .3914$	-14831.677	None
M1 (neutral): $p_0 = .632$ and $\omega_0 = .071$ $p_1 = .368$ and $\omega_1 = 1.000$	-14743.041	Not allowed
M2 (selection): $p_0 = .656$ and $\omega_0 = .092$ $p_1 = .326$ and $\omega_1 = 1.000$ $p_2 = .018$ and $\omega_2 = 3.170$	-14739.552	314, 351, 1129, 1147, 1367, 1480, 1522, 345, 351, 387, 436, 528, and 547
M3 (discrete): $p_0 = .448$ and $\omega_0 = .000$ $p_1 = .497$ and $\omega_1 = .623$ $p_2 = .054$ and $\omega_2 = 2.398$	-14738.992	286, 312, 314, 345, 351, 387, 436, 528, 547, 629, 797, 870, 891, 953, 972, 1010, 1052, 1074, 1120, 1129, 1136, 1147, 1305, 1367, 1422, 1480, 1492, 1497, 1522, 1534, 1553, 1564, 1610, 1663, 1666, 1673, 1731, 2034, 2106, and 2209
M7 (β): $p = .103$ and $q = .145$	-14744.192	Not allowed
M8 (β and ω): $p_0 = .967$, $p = .229$, and $q = .402$ $p_1 = .033$ and $\omega = 2.703$	-14739.187	286, 312, 314, 345, 351, 547, 797, 870, 972, 1129, 1147, 1305, 1367, 1422, 1480, 1497, 1522, 1553, and 1666
M8a (β and fixed ω): $p_0 = .646$, $p = 7.695$, and $q = 99.0$ $p_1 = .355$ and $\omega = 1.000$	-14743.051	Not allowed

NOTE.— p is the proportion of sites in each class, ω is the d_N/d_S ratio, and l is the log likelihood.

NA10861, NA12547, NA12548, and NA12560) used by Seattle-SNPs for nucleotide-variation discovery.

PCR and Sequencing

PKDREJ is a gene corresponding to the region 45030225–45037883 on chromosome 22 (GenBank accession number NM_006071; March 2006 assembly). Primers were designed using PRIMER3 version 0.2²¹ and were based on the known human sequence. The primers and conditions for PCR and sequencing are available on request. Species-specific primers were designed from the sequence obtained from the closest known relative to the desired species. PCR products were diluted fivefold with deionized water, were cycle sequenced using BigDye version 3.1, were ethanol precipitated, and were analyzed on an ABI 3100 automated sequencer (Applied Biosystems).

Data Analysis

Primate *PKDREJ* sequences were imported into Sequencher 4.2 (Gene Codes) for manual assembly. A consensus sequence was created from multiple overlapping reads and was aligned with the human reference sequence from the UCSC Genome Browser. The 5' 600-bp region of the *PKDREJ* coding region (~300 aa) is a GC-rich region that could not be successfully amplified or sequenced for the majority of species or for the human panel. The exported consensus sequences were aligned by eye in Se-AL version 2.0 (Oxford Evolutionary Biology). A neighbor-joining tree was constructed using Kimura 2-parameter distances in MEGA3.²² A maximum-likelihood tree was produced from the 14 primate species sequences with use of DNAML in the Phylip version 3.5 package²³ for use in PAML. We used an HKY+G model with empirical base frequencies, a transition:transversion ratio of 3.6, and a gamma-distribution-shape parameter of 0.1715. The model was chosen using hierarchical likelihood-ratio tests as implemented

in ModelTest.²⁴ Maximum likelihood-based methods²⁵ were used to detect the presence of adaptive evolution on the amino acid sequence of PKDREJ. These tests were implemented using CODEML in the PAML package (v. 3.14). CODEML allows the use of models with d_N/d_S ratios that vary among sites.²⁶

A likelihood-ratio test was used to examine the data for codons with d_N/d_S ratios significantly >1. This was done by comparing the likelihood of a null model, without selection, against the likelihood of the same model that includes an additional class of sites whose d_N/d_S ratio was free to vary. The neutral models included a model with a single d_N/d_S ratio averaged across all sites (M0), a model with a d_N/d_S class between 0 and 1 and a class with $d_N/d_S = 1$ (M1), and a model with the d_N/d_S ratio that assumes a beta distribution limited to the interval 0,1 (M7). The selection models (M2 and M8) add one additional class of sites with d_N/d_S estimated from the data. To test for variation in the d_N/d_S ratio among sites, we also compared a model (M3) with three distinct classes of d_N/d_S to model 0. Significance was determined by comparing the negative of twice the log-likelihood difference ($-2\Delta l$) with the χ^2 distribution, with degrees of freedom equal to the difference in the number of parameters estimated between the two nested models. A Bayesian analysis was used to calculate the posterior probabilities that sites with $d_N/d_S > 1$ were influenced by positive selection.²⁶ We used the new Bayes empirical Bayes approach, to have greater confidence in the prediction.²⁷ Convergence was checked by repeating the analyses with different initial d_N/d_S values, and, in all cases, identical likelihoods and parameter estimates were obtained.

To test for correlations with mating systems, we analyzed for variation in the d_N/d_S ratio between lineages. We first compared a model with one d_N/d_S estimated for all lineages with a "free-ratio" model in which we estimated d_N/d_S for each lineage. We next compared a one-ratio model with a model in which each lineage was assigned a class on the basis of mating system, as

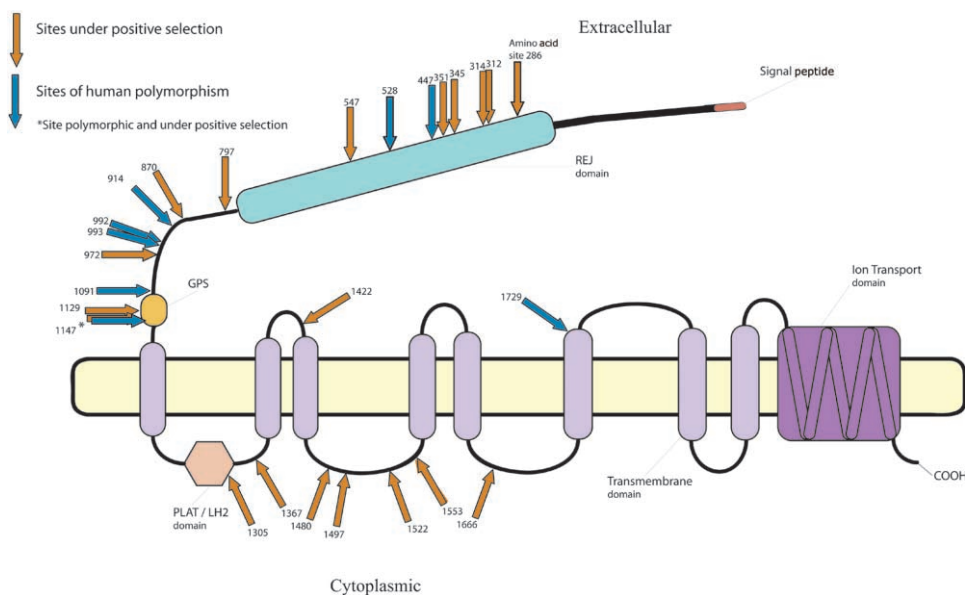


Figure 2. The PKDREJ protein with sites predicted to be subject to positive selection under model 8 (orange arrows) and amino acid polymorphic sites (blue arrows). The majority of polymorphic sites fall in the same domains as sites predicted to have evolved by positive selection.

described by Dorus et al.²⁸ Significance was determined by comparing the negative of twice the log-likelihood difference ($-2\Delta l$) with the χ^2 distribution, with degrees of freedom equal to the difference in the number of parameters estimated between the two nested models.

Chromatograms from the human panel were automatically base called, assembled, and scanned for SNPs with use of the Phred/Phrap/polyPhred programs (v. 14.0) and were visually inspected using Consed.^{29–32} The final consensus sequence was a single protein-coding region of 6,762 bp. Finished sequence data from the human panel were exported, and haplotypes were inferred using PHASE.³³ The default-phase certainty parameters $p = q = 90\%$ were used. Haplotypes were all well resolved. Two haplotypes per individual were obtained for this autosomal gene from the human panel of 48 individuals. DnaSP version 4.0³⁴ was used to estimate population genetic parameters and genetic distances and to perform tests of neutrality. The chimpanzee sequence that we determined experimentally was used as the outgroup. The protein architecture and domain identities were inferred using the SMART program.³⁵ The human amino acid sequence and polymorphism data were then used in the programs SIFT³⁶ and PolyPhen,³⁷ to infer the possible consequences of amino acid changes on the function of the PKDREJ peptide.

Results

At least 5,867 of 6,762 bp of *PKDREJ* were amplified and sequenced for 14 primates. The first 895 bp contains a GC-rich region that was could not be PCR amplified. A portion of this region encodes the signal sequence that is cleaved off the mature protein, and the remainder is of unknown function. A neighbor-joining tree was constructed and resulted in a well-supported topology that agreed with the accepted phylogeny^{38,39} of humans, great apes, and Old

and New World monkeys (fig. 1). A maximum-likelihood tree produced the same topology.

An analysis of the multispecies panel with use of CODEML indicates that positive selection has acted on *PKDREJ* (table 1). Model 3, which incorporates three categories of d_N/d_S values, was a significantly better fit to the data than was model 0, which estimates a single average d_N/d_S value over all sites. Whereas a comparison of model 3 and model 0 indicates a significant variation in d_N/d_S between sites, M3 is not a robust test of adaptive evolution. Thus, we compared more-general models that use a beta distribution (M7), which ascribes values of d_N/d_S between 0 and 1 across all sites with a model (M8) that includes an additional category with a class of sites that have d_N/d_S estimated from the data. This latter selection model M8 was a significantly better fit to the data than was the neutral M7 model whose d_N/d_S values are constrained between 0 and 1. This comparison is a robust test of positive selection.⁴⁰ Approximately 3% of amino acids or 19 codons appear to have evolved by positive selection with an average d_N/d_S (ω) of 2.703, as estimated by model 8. To control for false-positive results due to ancestral recombination, we performed the analyses without the closely related species (we excluded chimpanzee, bonobo, gorilla, orangutan, and all but one macaque). The rationale for this is that these species are so closely related that there could still be sorting of ancestral polymorphisms with recombination, which could lead to false-positive results. The analysis of this smaller data set was completely consistent with the analysis of the broader data set containing all 14 primates, indicating positive selection.

On the basis of predictions of protein architecture of

Table 2. SIFT and PolyPhen Prediction of Amino Acid Polymorphisms

Position	AA		Frequency of Derived Allele (%)	Prediction		
	Ancestral	Derived		PolyPhen	SIFT	Location
914	P	L	63	Damaging	Tolerated	Extracellular
1091	N	S	34	Benign	Tolerated	Extracellular
992	T	P	33	Benign	Tolerated	Extracellular
1147	I	M	15	Benign	Damaging	GPS unit, extracellular
993	V	A	8	Benign	Tolerated	Extracellular
528	R	E	3	Benign	Tolerated	REJ module, extracellular
1729	V	I	2	Benign	Tolerated	Transmembrane, extracellular
447	R	G	1	Benign	Tolerated	REJ module, extracellular

PKDREJ,³⁵ 6 of the 19 sites predicted to have evolved by positive selection fall within the REJ module (fig. 2). Five sites fall in or between the GPS region and the REJ module. These sites are all predicted to be exposed to the extracellular environment and could therefore influence sperm-egg interaction(s) with other extracellular molecules. Seven sites fall on predicted low-complexity regions adjacent to the highly conserved and functionally important PLAT/LH2 domain that is potentially involved in mediating membrane attachment via binding to other proteins and transmembrane domains on the cytoplasmic side of the molecule. One additional site falls between transmembrane regions 2 and 3 in the extracellular side (fig. 2).

To search for a correlation between the intensity of sperm competition and the amount of positive selection in all primate lineages studied (fig. 1), we compared d_N/d_S values along different branches with a neutral model that estimates a single d_N/d_S for all branches. For the branch-length model, a single d_N/d_S per branch is estimated on the basis of a set number of categories that we predetermined on the basis of the degree of sperm competition that might be expected, given the mating systems of the different species.²⁸ In contrast with a study of the primate seminal-fluid proteins SEMG2²⁸ or SEMG1,⁴¹ no correlation was discovered between the amount of molecular evolution in a lineage of *PKDREJ* and the degree of sperm competition.

To survey polymorphisms within human populations, the *PKDREJ* gene was amplified and sequenced from a panel of 48 individuals (96 chromosomes). A total of 24 polymorphic sites were found in the human panel, 8 of which resulted in amino acid changes. All polymorphisms were diallelic. Two of the eight sites are located in the REJ module, and five more fall between the REJ module and the GPS site, corresponding to the regions that have several sites predicted to be under positive selection. One additional site falls at the end of transmembrane region 6. It is notable that all amino acid polymorphisms fall within parts of the protein that are predicted to be extracellular. The polymorphism at amino acid 447 (arginine→glycine) is at a site predicted by PAML to be under positive selection among species. The effect of the amino acid-changing polymorphisms was predicted using SIFT³⁶ and PolyPhen.³⁷ These programs compare the amino acid poly-

morphisms with amino acid variation in other members of the family, the location in the protein, and other factors, to predict whether the polymorphism would affect protein function (in particular, be deleterious). For genes under positive selection, we suggest it is possible to use SIFT and PolyPhen to detect polymorphisms that may have a functional consequence and therefore might be targets for positive selection. Results showed that two (R447G and R528E) of the eight nonsynonymous polymorphisms alter charge. PolyPhen reported changes to amino acid position 914 (L914P) as probably damaging to PKDREJ function, and SIFT reported changes to position 1147 (I1147M) as potentially affecting PKDREJ protein structure, indicating that these may have functional significance. Some of these derived alleles are at intermediate frequencies (table 2). In the case of amino acid 914, the nonsynonymous SNP is predicted to be possibly damaging, but the derived SNP occurs at a frequency of 0.63 overall and is similar in frequency in both populations. Slightly deleterious mutations are generally maintained at low frequencies in populations. Positive selection can act to increase the frequency of a mutation. The relatively high derived allele frequency, clustering of nonsynonymous SNPs by sites undergoing adaptive evolution, and indication of functional differences suggest that positive selection may be acting on these SNPs to alter protein function or specificity within humans as well as among primate species.

Twenty-six human haplotypes were inferred using PHASE.³³ The most common haplotype appeared in 35% of the total population and in 63% of the European American population but in only 22% of the African American population. The European American population was dominated by a smaller number of haplotypes shared by a greater number of individuals. Among European Americans, there were four shared haplotypes among seven singletons, compared with eight shared haplotypes and 11 singletons in the African American population. The African American population had greater haplotype diversity (HD = 0.904) compared with European Americans (HD = 0.640). Nonsynonymous changes were largely shared by both populations, with only a single additional nonsynonymous change found exclusively in the African American population. The differences in diversity between

Table 3. Summary Statistics

Population	Sample Summary					Parameter Estimate						Test Statistic			
	<i>N</i>	Synonymous	Nonsynonymous	Total	No. of Haplotypes	Φ	π	HD	<i>k</i>	F_s	F_{ST}	<i>D</i>	<i>F</i>	<i>H</i>	Tajima's <i>D</i>
African American	50	15	7	22	19	4.912	1×10^{-3}	.904	7.265	-1.745725	1.23638	-.408	1.546
European American	46	10	6	16	11	3.641	6×10^{-4}	.64	4.254	.2533	.457	-2.381	.529
Total sample	96	16	8	24	26	...	1×10^{-3}	.815	6.452	-3.822	.178	-.135	.43736	-.928	1.135

NOTE.—Chimpanzee is the outgroup. Φ (per sequence) is from S ($\Phi - W$); *k* = average number of nucleotide differences; F_s ; *D* = Tajima's *D*; F_s = Fu's F_s statistic; *H* = Fay and Wu's *H* with outgroup; *F* = Fu and Li's *F* with outgroup⁴⁷; *D* = Fu and Li's *D* with outgroup.⁴⁷

these two population samples are primarily due to the abundance of singletons in the African American population, a pattern typical of that population.⁴² The difference in genetic makeup is reflected in the large $F_{ST} = 0.178$ compared with the $F_{ST} = 0.157$ human average⁴² and a significant χ^2 value ($P = .014$) for genetic differentiation.

We compared the levels of polymorphism within humans with divergence between humans and chimpanzee or gorilla, using the McDonald-Kreitman (MK) test,⁴³ which measures deviations from the expected ratio of nonsynonymous: synonymous polymorphisms to divergence. The MK test produced significant *P* values, such that neutrality can be rejected. We suggest two explanations for why neutrality is rejected. First, there could be an excess of synonymous polymorphisms within humans. An excess of synonymous polymorphisms could arise because of a balancing polymorphism at this locus. Alternatively, there could be a significant accumulation of nonsynonymous divergence since the human and chimpanzee lineages diverged. We favor the former idea, since there is additional evidence of balancing selection on the basis of analyses of the frequency spectrum (see below). The significant analyses of d_N/d_S suggests that balancing selection is most likely acting on nonsynonymous sites. Although the MK tests is generally considered to be robust to demographic effects,^{44,45} the test was run on subsamples of the human polymorphism data. Results from the European American sample proved to be nonsignificant, whereas results from the African American sample were highly significant.

Additional tests of neutrality on *PKDREJ* (Tajima's *D*,⁴⁶ Fu and Li's *D* and F_s ,⁴⁷ and Fay and Wu's H ⁴⁸) were performed on the total human sample and the two subsamples, with chimpanzee as the outgroup for Fay and Wu's *H* (table 3). Significance was determined by coalescent simulations, with only Tajima's *D* showing significance ($P = .04$) and for only the African American population, suggesting an excess of intermediate-frequency polymorphisms. Additionally, the value of Tajima's *D* ($D = 1.5$) for the African American sample was the second highest value seen in the SeattleSNPs database of 247 genes from the same African American samples, which places it in the 99th percentile of values (median Tajima's *D* for the African American sample is -0.52). Thus, the magnitude and direction of Tajima's *D* for *PKDREJ* is highly unusual for the African American population. The value of Tajima's *D* for European Americans was 0.591, a typical value for this population based on comparisons with the Seattle-

SNPs database (median value is 0.379 for the CEPH population).

Discussion

We compared PKDREJ sequences from 14 primates plus two population samples of humans. Multiple statistical tests reject equilibrium-neutral expectations and suggest that *PKDREJ* has evolved by positive selection in the primate lineage (table 1). Nineteen codons are identified that show signs of positive selection. These fall into several regions predicted to be important for function, including the REJ domain, the GPS domain, and a region between them. Of the 19 sites predicted to be subject to positive selection, 12 are in the predicted extracellular part of the molecule. An additional seven sites occur on the first three intracellular loops between transmembrane domains. These are low-complexity regions but contain a potential lipid-interaction site.⁴⁹

We also surveyed PKDREJ for amino acid polymorphisms within humans. Several nonsynonymous polymorphic sites fell within or close to codons for which we predicted adaptive evolution. For example, 87% of human nonsynonymous polymorphisms fall within the REJ domain and the region before the GPS domain, the REJ module, and the extracellular region around the putative GPS domain representing ~50% of the protein (fig. 2). All eight of the amino acid polymorphic sites are in predicted extracellular regions. The African American population appears to have an excess of intermediate-frequency variants compared with the expectations of an equilibrium-neutral model.

Although the function of PKDREJ remains unknown, it has been suggested as a candidate sperm receptor for the ZP.^{13,18,50} The rapid evolution of fertilization proteins is hypothesized to result from male-male competition in promiscuous mating systems in which sperm compete to fertilize the egg. Alternatively, there could be conflict between males and females, where the egg is selected to avoid fertilization by multiple sperm (which is usually fatal to the egg) but sperm are selected to fertilize rapidly.⁵¹ We were unable to document any correlation between rate of *PKDREJ* evolution and mating system, such as that reported for the primate seminal-fluid protein SEMG2.²⁸ Whereas this result is consistent with female-male rather than male-male competition having driven the evolution of *PKDREJ*, the lack of correlation may instead be due to

Table 4. MK Test: Chimpanzee/Gorilla Comparison

Comparison and Species	Findings by Sample		
	Total	European American	African American
Synonymous substitutions:			
No. of polymorphic sites (human)	16	10	15
Fixed differences between species:			
Chimpanzee	10	11	10
Gorilla	18	18	18
Nonsynonymous substitutions:			
No. of polymorphic sites (human)	8	6	7
No. of fixed differences between species:			
Chimpanzee	21	21	21
Gorilla	27	27	27
NI:			
Chimpanzee	.238	.314	.222
Gorilla	.333	.4	.311
Fisher's exact test <i>P</i> (two tailed):			
Chimpanzee	.015	.122	.013
Gorilla	.045	.151	.039

NOTE.—Significant values are shown in bold.

inaccuracies on the estimates of the degree of sperm competition. Additionally, sperm competition and sperm-egg conflict are not mutually exclusive, since intense sperm competition can select for the same characteristics that increase the likelihood of polyspermy and increased egg-sperm conflict.

Results of the MK test were significant, which we interpret as an excess of synonymous polymorphisms in human *PKDREJ* (table 4), although an excess of amino acid fixation between species may also contribute. Because the significance of the MK test result could be due, in part, to divergence in the chimpanzee outgroup, the MK test was repeated using gorilla as the outgroup. Results were again significant for the total population. The test was run on the two subpopulations, and results were found to be significant only for the African American sample. In all cases, the *P* value was diminished in relation to tests using the chimpanzee outgroup, suggesting that the choice of outgroup contributed to the significance of the result. A comparison of the neutrality index (NI) values (table 4), which provides a qualitative measure of the direction and extent of amino acid changes, were both <1, where a value of 1 indicates neutrality, >1 is purifying selection, and <1 is positive selection. The value of NI is closer to 1 when human is compared with gorilla than when human is compared with chimpanzee.

Consistent with the finding of excess synonymous substitutions in the African American population by use of the MK test, analysis of the frequency spectrum also suggests that balancing selection acts on *PKDREJ*. For *PKDREJ* in the African American population, the value of Tajima's *D* (1.5) was significantly positive and an extreme outlier compared with the values of 247 other loci from the SeattleSNPs data set. Positive values of Tajima's *D* are due to an excess of alleles with intermediate-frequency variants.⁴⁶ This pattern is seen under balancing selection as

well as a narrow window of time during the recovery after population bottlenecks. Demographic changes affect all genes, whereas selection typically acts on a single locus. Large positive values of Tajima's *D* are rare in the African American population, and other genes in the African American population show no evidence of a bottleneck. The distribution of polymorphisms and signs of selection on different human haplotypes in the African American population suggest that selection has driven the diversification of human reproductive alleles.

One of the striking features of the analysis presented here is the evidence of rapid, adaptive divergence of *PKDREJ* among primates, coupled with human polymorphism data indicating the presence of balancing selection. Thus, there is high divergence among species and high levels of variation within species. Such an observation is inconsistent with multiple complete selective sweeps and suggests variable selective pressures (e.g., balancing selection coupled with occasional selective sweeps) or perhaps a succession of partial selective sweeps. This pattern has been documented in a variety of other genes involved in reproduction. For example, both the *Drosophila* seminal-fluid protein *Acp26Aa*^{52–54} and the sea urchin sperm protein *bindin*^{55,56} show this pattern of high levels of differences both within and among species. Such consistent patterns across multiple taxonomic groups suggest a potentially fruitful area of population genetics for modeling the evolutionary processes that could lead to these patterns. For example, some models of sexual conflict predict diversification within species of genes involved in reproduction.⁵⁷

Coevolution between sperm proteins and their egg receptors can result in the divergence of alleles. If a mutation in a sperm protein creates sufficient advantage in fertilizing eggs that display a particular variant of the egg-coat protein, it will be selected for despite the expense of being

inferior at fertilizing eggs displaying other egg-coat alleles. This sperm protein may then become specialized, responding to changes in that egg protein while growing ever more divergent from other sperm alleles. Mismatch between mating types is more likely to occur between populations that infrequently exchange gametes, since there is a likely to be a trade-off between the benefit of increased effectiveness of fertilization and the odds of encountering a new egg-coat protein for which the sperm protein is an ineffective receptor.

The pattern of variability and departures from neutrality in the sequence evolution of PKDREJ strongly suggest positive selection on the evolution and diversification of PKDREJ. The putative structure of the PKDREJ protein and its testis-specific expression suggests that this gene is functionally analogous to the sea urchin sperm protein suREJ, which plays a role in fertilization. We have found evidence of adaptive changes in the evolution of PKDREJ among species as well as evidence of diversifying selection within species. There appears to have been selection for increased diversity in the number of haplotypes in the African American population sample. It is unknown whether there are functional differences between any of the haplotypes, but the existence of multiple alleles with functional differences could have consequences for human fertilization.

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Web Resources

Accession numbers and URLs for data presented herein are as follows:

GenBank, <http://www.ncbi.nlm.nih.gov/Genbank/> (for *PKDREJ* [accession number NM_006071] and nonhuman primate *PKDREJ* sequences [accession numbers EF517278–EF517291])
 Oxford Evolutionary Biology, <http://evolve.zoo.ox.ac.uk/software.html?id=seal> (for Se-AL sequence alignment editor)
 SeattleSNPs, <http://pga.gs.washington.edu/>

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