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*PNAS* 2007;104:6278-6282; originally published online Mar 26, 2007;  
doi:10.1073/pnas.0607426104

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# Germ-line chimerism and paternal care in marmosets (*Callithrix kuhlii*)

C. N. Ross<sup>\*†</sup>, J. A. French<sup>‡</sup>, and G. Ortí<sup>\*</sup>

<sup>\*</sup>School of Biological Sciences, University of Nebraska, Lincoln, NE 68588; and <sup>‡</sup>Departments of Psychology and Biology, University of Nebraska, Omaha, NE 68182

Edited by John C. Avise, University of California, Irvine, CA, and approved February 14, 2007 (received for review August 25, 2006)

**The formation of viable genetic chimeras in mammals through the transfer of cells between siblings *in utero* is rare. Using microsatellite DNA markers, we show here that chimerism in marmoset (*Callithrix kuhlii*) twins is not limited to blood-derived hematopoietic tissues as was previously described. All somatic tissue types sampled were found to be chimeric. Notably, chimerism was demonstrated to be present in germ-line tissues, an event never before documented as naturally occurring in a primate. In fact, we found that chimeric marmosets often transmit sibling alleles acquired *in utero* to their own offspring. Thus, an individual that contributes gametes to an offspring is not necessarily the genetic parent of that offspring. The presence of somatic and germ-line chimerism may have influenced the evolution of the extensive paternal and alloparental care system of this taxon. Although the exact mechanisms of sociobiological change associated with chimerism have not been fully explored, we show here that chimerism alters relatedness between twins and may alter the perceived relatedness between family members, thus influencing the allocation of parental care. Consistent with this prediction, we found a significant correlation between paternal care effort and the presence of epithelial chimerism, with males carrying chimeric infants more often than nonchimeric infants. Therefore, we propose that the presence of placental chorionic fusion and the exchange of cell lines between embryos may represent a unique adaptation affecting the evolution of cooperative care in this group of primates.**

callitrichid | genetic chimerism | genomic conflict | social behavior

Genetic chimerism, the mingling of two or more genomic lineages within an individual (1), is rare in mammals, but chimerism is prevalent in the hematopoietic tissues of marmosets and other callitrichid primates (2, 3). In these species, fraternal twins exchange cell lines through chorionic fusion during early development (2, 4, 5). On the basis of karyotypic evidence from *Callithrix jacchus* (2, 3), estimates are that 95% of pregnancies result in the birth of hematopoietic chimeric twins. Chorionic fusion of the twins' placentas begins on day 19 and is complete by day 29, forming a single chorion with anastomoses connecting the embryos, which are still at a presomite stage in development (4–7). The fusion of the chorions and a delay in embryonic development at this stage allows the exchange of embryonic stem cells via blood flow between the twins (2, 8). As a result, the infants are genetic chimeras with tissues derived from self and sibling embryonic cell lineages (2, 3, 8).

Although there is little doubt that tissues derived from hematopoietic origin are universally chimeric (9), the existence of chimeric cells in nonhematopoietic tissues, including germ-line cells, has not been established. Karyotypic analysis of *C. jacchus* revealed that testes cells express unusual orientation during meiosis, and this evidence suggested that the germ-line cells might include female cells present because of chimerism (8, 10). However, further karyotypic analysis refuted these findings, and an analysis of sex ratios in captive colonies of *C. jacchus* suggested that germ-line chimerism was not present (11). To investigate whether chimerism occurs in tissues other than those derived from the hematopoietic system, species-specific micro-

satellite markers were used to examine the extent and distribution of chimerism.

The existence of chimerism throughout somatic and germ-line tissues may have important implications for the evolution of paternal and alloparental care characteristics of this taxon, through genomic conflict or altered perceptions of relatedness between members of a family group (12, 13). Genomic conflict in individuals with genetic heterogeneity has been identified as a possible evolutionary mechanism, influencing behavioral and developmental traits (14–17). Conflict within an individual may influence the development of kin recognition mechanisms. Specifically, how do chimeric organisms identify an individual and determine relatedness to another chimeric individual (12)? Somatic chimerism may provide individuals with self-matching kin recognition cues, causing an overestimate of their relatedness to chimeric offspring. Although the exact mechanisms of kin recognition are unknown in primates, baboons appear to be capable of recognizing paternal offspring, which may involve phenotype matching (18). Phenotype matching has been conclusively demonstrated to occur in at least one mammal species (19). If chimerism in marmosets involves more than hematopoietic tissues, then we predict differential parental behavior toward chimeric and nonchimeric infants and altered estimates of relatedness from those expected for nonchimeric mammals.

## Results

We examined the prevalence of chimerism in tissues derived from different embryonic origins by analyzing genotypes of microsatellite loci with a probability of detecting chimerism of 98% based on parental genotypes for these loci. A total of 92 intergenerational individuals that included 36 twin sets of *Callithrix kuhlii* (Wied's black tufted-ear marmosets) and their parents were assessed. The samples were genotyped in an appropriate blind fashion such that the identity of the individual and the tissue type were unknown. All alleles were noted for each locus, and samples were identified as potentially chimeric if they contained three or four allelic variants at a single locus. The samples were then matched to identity, and twins were noted to be chimeric at a tissue only if the alleles were found to match both the parents as well as their twin. Further, a majority rule approach was used to assign alleles as "self" (i.e., diploid and inherited vertically from the parents) and "sibling" (inherited horizontally from the twin *in utero*) [see example in supporting information (SI) Fig. 3]. Of the 36 twin sets surveyed, 26 (72.2%) were determined to carry chimeric tissues. Exchange of alleles

Author contributions: C.N.R., J.A.F., and G.O. designed research; C.N.R. performed research; C.N.R. and G.O. analyzed data; and C.N.R., J.A.F., and G.O. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

<sup>†</sup>To whom correspondence should be addressed at: University of Texas Health Science Center, 15355 Lambda Drive, Suite 3.325, South Texas Centers for Biology in Medicine, San Antonio, TX 78245. E-mail: rossc4@uthscsa.edu.

This article contains supporting information online at [www.pnas.org/cgi/content/full/0607426104/DC1](http://www.pnas.org/cgi/content/full/0607426104/DC1).

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**Table 1. The number of *Callithrix kuhlii* individuals chimeric for each tissue type**

Tissue	Tissue type	Genotyped, no.	Chimeric, no.	Chimeric, %
Samples from deceased animals				
Placenta	H	7	7	100.0
Blood	H	2	2	100.0
Spleen	H	28	14	50.0
Liver	H	39	15	38.5
Heart	S	30	7	23.3
Hair	S	35	6	17.1
Lung	S	30	4	13.3
Kidney	S	33	4	12.1
Gonad	G	21	2	9.5
Skin	S	36	2	5.6
Brain	S	31	1	3.2
Muscle	S	34	1	2.9
Samples from living animals				
Sperm	G	7	4	57.1
Saliva	S	31	16	51.6
Blood	H	45	22	48.9
Hair	S	50	13	26.0
Fecal	S	22	2	9.09

H, hematopoietic; S, other somatic; G, germ line.

between twins was not always bidirectional. In 14 twin sets, only one twin was found to have chimeric tissue types, whereas the other 12 chimeric twin sets revealed chimerism in both twin's tissues.

Chimerism was found to be present in every tissue that was analyzed, and the occurrence of chimerism in tissues harvested from marmoset cadavers differed significantly across liver, spleen, kidney, heart, lung, brain, muscle, skin, and hair ("Samples from deceased animals" in Table 1;  $n = 25$ , Cochran's  $Q = 51.6$ ,  $df = 8$ ,  $P < 0.001$ ). To determine which tissue type accounted for the variance among the tissues, the tissues were ranked according to percentage of occurrence of chimerism and then grouped. A comparison of liver and spleen revealed a nonsignificant difference between the tissue types ( $n = 25$ ,  $Q = 4$ ,  $df = 1$ ,  $P = 0.15$ ). The addition of hair samples revealed a significant difference between the tissue types ( $n = 25$ ,  $Q = 14.3$ ,  $df = 2$ ,  $P < 0.001$ ). The grouping of all other tissues (heart, hair, lung, kidney, skin, brain, and muscle) resulted in no significant difference between the tissue types for the presence of chimerism ( $n = 25$ ,  $Q = 10.3$ ,  $df = 6$ ,  $P > 0.1$ ). Hematopoietic tissues were significantly more likely to be chimeric than all other tissue types ( $\chi^2 = 4.88$ ,  $df = 1$ ,  $P < 0.05$ ). The assessment of chimerism in tissues collected from living marmosets revealed nonsignificant differences between the tissue types ("Samples from living animals" in Table 1).

The presence of sibling-derived alleles in multiple tissues suggested that all embryonic cell lineages in *C. kuhlii* might be affected by chimerism, including gametic tissue. In fact, gonadal tissue was found to be chimeric (2/21), and sperm samples were also chimeric (4/7). Additionally, the 36 twin sets analyzed for chimerism comprised multiple generations within 15 family lines. We determined that individuals in 5 of the 15 families passed on alleles to their offspring that represented gene lineages inherited horizontally from the sibling (see examples in Fig. 1 and SI Fig. 4). One breeding female, whose uterine twin was a male, produced offspring that inherited her sibling's alleles. This documents the possibility that an XY primordial germ cell is capable of maturing and producing viable eggs in a female, a phenomenon that has not been documented for primates. Although we are not currently able to document the fate of the Y chromosome during development of the female's oocytes, our

data suggest the intriguing possibility that a female may pass on a Y chromosome to her offspring.

The presence of cells derived from different lineages within an individual may impact behavioral decisions. Genetic chimerism may give rise to genomic conflict such that an individual's decision to cooperate within a group and care for members of the group may depend on the true, or perceived, genetic relatedness between the individuals (12, 16). To illustrate this with a simple example, we consider the increased proportion of shared alleles, because of genetic chimerism, between male twins produced by nonchimeric parents. A chimeric individual's coefficient of relatedness to his twin could increase from the expected fraternal twin value of  $r = 0.5$  to as much as  $r = 1$  in certain tissues. Based on the prevalence of chimerism, the proportion of cells within a tissue that carry sibling alleles, and the probability of the direction of exchange obtained from our data, we estimate that male twins are on average related by  $r = 0.574$  (see SI Text for calculations). More specifically, in a case of unidirectional exchange in which the soma of the donating twin is nonchimeric, he is related to the sperm of the recipient twin by an average  $r$  of 0.625 (see SI Text). The relatedness calculations suggest that chimeric callitrichid siblings are more closely related to each other than typical nonchimeric mammalian siblings. Calculations of relatedness under more complex scenarios and involving parental chimerism are beyond the scope of this report; thus, at this stage, it is not known how parental-offspring relatedness may be affected by chimerism.

A different issue, but also with potential behavioral implications, is perceived relatedness through kin recognition. To investigate whether the presence of chimeric epithelial tissues, those most likely to mediate kin recognition, are associated with changes in parental care, maternal and paternal infant carrying effort for the first 2 weeks of life was compared between infants determined to be epithelial chimeras ( $n = 10$ ) and those determined to be nonchimeric ( $n = 20$ ). Alloparents do not typically provide care during this early period (20) and thus were not included in the analysis. Females carried chimeric infants significantly less than nonchimeric infants ( $F_{1,28} = 7.61$ ;  $P = 0.01$ ), but fathers carried chimeric infants significantly more than nonchimeric infants ( $F_{1,28} = 12.08$ ;  $P = 0.002$ ) (Fig. 2). No significant differences in carrying time by mothers or fathers





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