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MULTIPLE HYBRID GENOTYPES OF *LEISHMANIA* (*VIANNIA*) IN A FOCUS OF MUCOCUTANEOUS LEISHMANIASIS

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Abstract. The principal agent of mucocutaneous leishmaniasis (MCL) is the South American protozoan parasite *Leishmania* (*Viannia*) *braziliensis*. This organism is generally considered to be clonal, that is, it does not undergo genetic exchange. Nevertheless, apparent hybrids between several *Leishmania* species have been reported in the New World and the Old World. When we characterized isolates of *Leishmania* (*Viannia*) from a single focus of cutaneous leishmaniasis (CL) and MCL, we found a remarkable phenotypic and genotypic diversity, with 12 zymodemes and 20 microsatellite genotypes. Furthermore, 26 of the 59 isolates were *L. braziliensis*/*L. peruviana* phenotypic hybrids that displayed 7 different microsatellite genotypes. A hybrid genotype was the only organism isolated from 4 patients with MCL. Thus hybrids must be included among the potential agents of MCL. Despite the propensity for clonality, hybrids are also an important feature of *Leishmania* (*Viannia*) and may give rise to epidemiologically important emergent genotypes.

INTRODUCTION

Leishmaniasis is a major public health problem in much of Latin America, where *Leishmania* of the subgenus *Leishmania* are agents of visceral leishmaniasis (VL), cutaneous leishmaniasis (CL), and diffuse cutaneous leishmaniasis (DCL). The subgenus *L. (Viannia)*, which is restricted to the New World, causes CL and metastatic mucocutaneous leishmaniasis (MCL).

In Peru, both CL and MCL are endemic. *Leishmania (Viannia) braziliensis* and *L. (V.) peruviana* are most frequently associated with CL, although *L. (V.) guyanensis*, *L. (V.) lainsoni* and *L. (Leishmania) amazonensis* have also been reported.¹ MCL is attributed to *L. braziliensis*. However, lesions from *L. braziliensis* and *L. peruviana* are not distinct in the early stages of CL, and species have often been incriminated on the basis of known geographical range, with *L. peruviana* found mostly in the western Andes and inter-Andean valleys and *L. braziliensis* occurring predominantly at lower altitudes in the Amazonian region. Mucosal leishmaniasis (ML), with involvement of the mucosae by contiguity from a primary lesion, has been described for all species causing CL in Peru.¹ However, ML is distinct from MCL, which involves metastatic spread to mucosal sites some time after a primary infection. Control of MCL depends predominantly on passive or active case finding, diagnosis, and effective treatment.² Dogs are commonly infected with *L. braziliensis* and/or *L. peruviana*, and they may act as a peridomestic reservoir.^{3,4} The sylvatic reservoir hosts of *L. peruviana* and *L. braziliensis* are incompletely known, although terrestrial small rodents have been implicated for some strains.^{4,5}

Where the Amazonian forest and Andean regions meet, as in the Department of Huánuco, both *L. braziliensis* and *L. peruviana* may occur sympatrically. *Leishmania (Viannia)* isolates that appear to be hybrids between *L. braziliensis* and *L. peruviana* have been reported from this region of Peru.⁶

Herein we describe the phenotypes (obtained by multilocus

enzyme electrophoresis, MLEE), and genotypes (obtained by microsatellite multilocus typing, MLMT) of 59 isolates from the Department of Huánuco. The results show that putative hybrid phenotypes and genotypes are common. A remarkable degree of diversity is revealed, suggesting that propagation is not entirely clonal and indicative of some form of genetic exchange in the Huánuco *L. (V.)* population.

MATERIALS AND METHODS

Isolate collection. *Leishmania* were isolated in 1994–1995 from 45 humans and 14 dogs, from cutaneous lesions on the ear or nose, in villages around Huánuco City (2000–3000 m above sea level). Eleven patients had MCL, and 34 had CL. Full ethical permission was given by the Universidad Peruana Cayetano Heredia with informed consent obtained from human subjects for parasite diagnosis. Households in these areas routinely keep dogs, donkeys, pigs, and chickens. A full list of the isolate codes is available from the authors upon request.

Isolates were phenotyped and genotyped against the following *L. (V.)* reference strains: *L. (V.) braziliensis* (MHOM/BR/84/LTB300); *L. (V.) peruviana* (MHOM/PE/94/LC1152; MHOM/PE/84/LC26; MHOM/PE/84/LC39); *L. (V.) panamensis* (MHOM/PA/71/LS94); *L. (V.) guyanensis* (MHOM/BR/75/M4147); *L. (V.) shawi* (MHOM/BR/94/M15065); *L. (V.) lainsoni* (MHOM/BR/81/M6426); and *L. (V.)* sp. n.⁷ (ISQU/BR/86/IM2832).

Isolation and in vitro cultivation of parasites. Reference strains were retrieved from liquid nitrogen storage onto biphasic 4N blood slopes.⁸ Peruvian stocks were dispatched from Lima on 4N blood slopes and were passaged onto fresh slopes and incubated at 23°C upon receipt. Isolates from this first-passage culture were stored under liquid nitrogen; subsequent passages were kept to a minimum to reduce culture selection of parasite strains. Promastigote cultures were expanded in alpha-modified minimal essential medium (Sigma-Aldrich Ltd., Gillingham, Dorset, UK) supplemented with 10% heat-inactivated fetal bovine serum, 50 µg/mL gentamicin, 30 mM NaHCO₃, 40 mM HEPES, 20 mM D-glucose, 4 mM L-glutamine, 10 µM hemin, 30 µM adenine, 10 µM folic acid, and 10 µM D-biotin (all supplements from Sigma Chemi-

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cal Co., Gillingham, Dorset, UK). Enzyme lysates were prepared from logarithmic-phase bulk cultures according to the method of Evans and others.⁸

Isoenzyme electrophoresis. Diversity of the *Leishmania* isolates was initially analyzed by MLEE using thin-layer starch gels.⁹ The enzymes applied were mannose phosphate isomerase (MPI, EC 5.3.1.8); glucose phosphate isomerase (GPI, EC 5.3.1.9); proline dipeptidase (PEPD, EC 3.4.13.9); phosphoglucomutase (PGM, EC 2.7.5.1); nucleoside hydrolyase using 2 different substrates, inosine (NHi, EC 3.2.2.1; 2 loci: NHi1 and NHi2) and deoxyinosine (NHd, EC 3.2.2.x); 6-phosphogluconate dehydrogenase (6PGD, EC 1.1.1.44); glucose-6-phosphate dehydrogenase (G6PD, EC 1.1.1.49); esterase (ES, EC 3.1.1.1); aspartate aminotransferase (ASAT, EC 2.6.1.1); and alanine aminotransferase (ALAT, EC 2.6.1.2).

Microsatellite genotyping and DNA sequencing. Subsequent analyses involved multilocus microsatellite typing (MLMT)¹⁰ and DNA sequencing. Microsatellites AC01, AC16, and AC52¹¹ were amplified from promastigote genomic DNA. The microsatellites were amplified with primers AC01F and AC01R-FAM (GAGAGGCCACCAGACACGTCAGCACAC and CCCCTTCCTTCGCCTTCAACACCTTTAC, respectively), AC16F and AC16R-TET (CTTCTTCTCATGCTGCACGGTCTCCTCCTT and CCATGGGCGGGCTTGTTCGTTACTTTTTTA, respectively), and AC52F and AC52R-HEX (CCACCGCCGGCTTCACTAC and GCGGCAATCGTCTGGCTAAA, respectively). Reverse primers were fluorophore-labeled as indicated (Perkin-Elmer, Beaconsfield, UK). Amplification reactions were carried out according to a protocol modified from that of Russell and others.¹¹ PCR amplification for each sample was done in a 10 µL reaction mix containing 10× NH₄ reaction buffer (160 mM (NH₄)₂SO₄, 670 mM Tris-HCl, pH 8.8, 0.1% Tween-20 [Bioline, London, UK]), 1 mM (AC01, AC52) or 2 mM (AC16) MgCl₂, 0.2 mM each of dATP, dCTP, dGTP, and dTTP (Pharmacia LKB, Upsala, Sweden), 5 pmol of each primer, 0.5% formamide (v/v), 1 U of *Taq* DNA polymerase (Bioline), and 25 ng of genomic DNA. PCR amplification was carried out in microtiter plates with sealed lids using the heated-lid option in an MJ Research PTC-200 Peltier thermocycler (Genetic Research Instrumentation Ltd., UK) using the following parameters: 35 cycles of 95°C for 30 s, 62°C (AC01 and AC52) or 60°C (AC16) for 30 s, 72°C for 1 min, followed by a final extension period of 10 min at 72°C. The multiplexed microsatellite products were sized on an ABI 377 automated sequencer by Genescan® and Genotyper® software (Applied Biosystems, Warrington, UK).

The AC01 products were sequenced by dye-terminator cycle sequencing and aligned using Sequence Navigator (Applied Biosystems).

Population genetics analysis. Resultant data from MLEE and MLMT were tested for genetic recombination and segregation using five population-genetics analyses: Hardy-Weinberg (HW) equilibrium,¹² fixation index (Fis),¹² *D'* index,¹³ *R*² index,¹³ and index of association (*I*_A).¹⁴

RESULTS

The isoenzyme phenotypes of the isolates were diverse. Twelve zymodemes (LON217-228) were encountered among the 59 Huánuco stocks by MLEE. One human isolate was

identified as *L. lainsoni*, and 3 dog isolates as *L. (V.)* sp. n.⁷ For the remaining 55 isolates (Table 1), the enzymes GPI, G6PD, ASAT, and ALAT were monomorphic while MPI, PEPD, PGM, NHi, NHd, 6PGD, and ES were polymorphic. The 55 isolates were initially classified according to their defining MPI profile, diagnostic for *L. braziliensis* and *L. peruviana*; 25 isolates were *L. braziliensis* (inferred MPI genotype 1/1), 4 were *L. peruviana* (2/2), and 26 had a double-banded MPI profile indicative of *L. braziliensis/L. peruviana* hybrids (1/2). Mixed cultures were excluded both by the occurrence of the same phenotype in biologic clones⁶ and by distribution of band intensities, including the presence of the classic triple-banded hybrid phenotype (with a central band of greater intensity) for dimeric enzymes.^{6,9} All putative hybrids had a triple-banded profile for the dimeric enzyme NHd. In contrast to the *L. braziliensis/L. panamensis* hybrids from Nicaragua,⁹ the Huánuco hybrids did not have the same overall phenotype (Tables 1 and 2).

Twenty multilocus microsatellite genotypes were identified, including 6 different multilocus genotypes among isolates from MCL lesions (Tables 1 and 2). Double peaks indicated heterozygosity. Twenty-three *L. braziliensis/L. peruviana* hybrids were heterozygous at the AC01 locus, as were many *L. braziliensis*. AC01 DNA sequence was obtained for all 55 isolates that generated a PCR product for the AC01

TABLE 1
Zymodeme and microsatellite genotypes of 55 *Leishmania (Viannia)* isolates, Huánuco, Peru

Species	Zymodeme ³⁸	Microsatellite genotype			No. of isolates*
		AC01	AC16	AC52	
<i>L. braziliensis</i>	LON222	227/237	244/246	265/295	2 (1)
		227/237	244/246	NS	1 (1)
	LON223	229/233	248/262	NS	3 (1)
		LON224	227/231	244/244	265/295
	LON225	227/231	244/250	NS	1 (1)
		227/233	244/274	265/295	1
		227/231	NS	265/295	1 (1)
		227/231	244/244	265/295	1
		227/231	248/248	265/295	12 (1)
		227/231	248/248	NS	1
	LON226	237/237	244/244	NS	1
		LON217	235/235	244/266	NS
241/241	250/250		NS	1 (1)	
X	X		X	2	
<i>L. peruviana</i>	LON218	227/231	240/240	X	1
		227/231	240/240	NS	1
		227/231	240/244	265/295	1
		227/231	244/244	265/295	12 (2) (V)
		227/231	244/244	NS	1 (1)
		227/231	244/248	NS	1
		227/231	244/250	265/295	1
		227/231	NS	NS	1
		229/231	250/252	NS	1
		237/237	250/252	NS	1
		ND	ND	ND	2 (1)
LON219	227/231	240/240	X	1	
LON220	227/231	240/244	265/295	1	
LON221	227/231	244/244	265/295	1	

Bold shows the phenotypes (zymodemes) and the multilocus microsatellite genotypes associated with mucosal disease (MCL); X = not amplifiable; ND = not done; NS = not scorable by Genotyper®. ***Bold** in parentheses shows numbers of isolates from MCL, roman numerals (I, V) show number of isolates from dogs. Reference strains used in phenotyping and genotyping were *L. (V.) braziliensis* (MHOM/BR/84/LTB300); *L. (V.) peruviana* (MHOM/PE/94/LC1152); MHOM/PE/84/LC26; MHOM/PE/84/LC39); *L. (V.) panamensis* (MHOM/PA/71/LS94); *L. (V.) guyanensis* (MHOM/BR/75/M4147); *L. (V.) shawi* (MHOM/BR/94/M15065); *L. (V.) lainsoni* (MHOM/BR/81/M6426); and *L. (V.)* sp. n. (ISQU/BR/86/IM2832).

TABLE 2
Summary of zymodemes and multilocus microsatellite genotypes

Species	No. of isolates [MCL associated]	No. of zymodemes (10 loci) [MCL associated]	Minimum+ no. of microsatellite genotypes (3 loci) [MCL associated]
<i>L. braziliensis</i>	26 [6]	5 [4]	8 [4]
<i>L. peruviana</i>	4* [1]	1 [1]	2 [1]
<i>L. b/pe</i> hybrids	26† [4]	4 [1]	7 [1]
<i>L. lainsoni</i>	1 [0]	1 [0]	1 [0]
<i>L. (V.) sp. n.</i>	3§ [0]	1 [0]	2 [0]
Totals	59 [11]	12 [6]	20 [6]

* 1 from dog; †10 from dog; §all from dog; +X and NS are not considered distinct genotypes here (see Table 1).

locus. In most cases, the number of dinucleotide repeats from sequence analysis correlated exactly with allele size by Genotyper®. When isolates were scored as heterozygous, the superimposed downstream sequences were clearly interpretable as 2 alleles differing by number of repeats. For example, all *L. braziliensis* isolates scored using Genotyper® as 227/231 were found to have overlapping sequence reads that could be interpreted as 2 alleles differing by 2 repeats (i.e., 8 and 10), giving allele sizes that varied by 4 base pairs. Only 6 of the *L. braziliensis/L. peruviana* hybrids were heterozygous at the unlinked AC16 locus. The complexity of AC52 profiles only allowed limited allele scoring.

Five population-genetics analyses, Hardy-Weinberg (HW) equilibrium,¹² fixation index (Fis),¹² *D'* index,¹³ *R*² index,¹³ and index of association (*I*_A),¹⁴ were applied to the data for 58 isolates, including (due to apparent sharing of alleles) *L. (V.) sp. n.*⁷ but excluding *L. lainsoni*. The results were as follows:

- Four of 10 polymorphic enzyme loci (MPI, ES, PEPD, and NHd) showed no significant deviation from HW equilibrium (*P* > 0.05, HW exact test), whereas 4 enzyme loci (PGM, GPI, G6PD, NHi1) and both microsatellite loci (AC01, AC16) deviated from HW equilibrium (NHi2 and ASAT were monomorphic, ALAT and 6GPD had null alleles).
- Excess heterozygosity was indicated by negative Fis values at 3 of 10 polymorphic loci MPI (-0.133), NHi (-0.425), NHd (-0.281), and AC01 (-0.288), whereas 4 loci showed deficits of heterozygosity, 3 of which were entirely homozygous.
- D'* indices were calculated for pairwise combinations of loci, with "1" denoting complete linkage and "0" indicating no linkage.¹³ Calculations were based on maximum-likelihood estimates of gametic frequencies.¹³ Reference to the *L. major* genome sequencing project indicates at least 8 different chromosomal assignments, although the loci are not yet mapped for *L. (Viannia)*. Seven out of 13 pairwise combinations showed a value of *D'* < 0.7. Three of these pairwise values (MPI × NHi1, AC16 × NHi1, and AC16 × NHd) were statistically significant (*P* ≤ 0.01, *Q* test).¹³ Corresponding *r*² values of linkage disequilibrium were lower, as is normal for this index.¹³
- The Maynard Smith index of association, *I*_A, which assesses linkage disequilibrium over all loci, was calculated. *I*_A values for all enzyme loci (0.59) and for all loci (0.97) did not indicate panmixia (*I*_A = 0). Values are comparable to those obtained with a smaller number of loci for *Try-*

panosoma brucei populations exhibiting epidemic clonality.¹⁵

In summary, overall the results from these tests indicated that the Huánuco *L. (Viannia)* population diverges from clonality or linkage disequilibrium. Firstly, the partial lack of significant deviation from HW equilibrium and, secondly, the significant *D'* values imply that some form of genetic exchange has occurred among the Huánuco *L. (Viannia)* population.

DISCUSSION

The trypanosomatids that cause human diseases (trypanosomiasis and leishmaniasis) were for many years considered to be incapable of genetic exchange, with reproduction confined to binary fission, in which one parental form divides to produce two identical offspring. Sexual dimorphism is not apparent, and chromosomes do not condense and therefore cannot be visualized. As a result, meiosis and mitosis cannot be observed directly.

Perceptions that *T. brucei* and *Trypanosoma cruzi* are entirely clonal have changed dramatically. With the aid of drug-resistance markers, genetic exchange was demonstrated to occur in the tsetse fly salivary glands, both within and between the *T. brucei* subspecies.¹⁶ The mechanism appears to be Mendelian, with occasional aneuploid progeny.¹⁷ Dependent on the locality under study, the population genetics of *T. brucei* ranges from panmixia in some undisturbed natural hosts and habitats through epidemic clonality, to true clonality.¹⁵ Among *T. cruzi* populations, phylogenetic analysis revealed genetic exchange¹⁸ and hybrids were produced experimentally in the laboratory.¹⁹ The *T. cruzi* experimental hybrids were derived from mammalian cells, not from the triatomine bug vector¹⁹ (although this does not exclude occurrence of genetic exchange within the vector). The mechanism of hybridization in *T. cruzi*, involving fusion, genome erosion, and recombination, was unusual but compatible with hybrid genotypes and the extensive range of DNA content found among natural populations. Heitman²⁰ has drawn attention to striking parallels between *T. cruzi* and the fungus *Candida albicans*: cell fusion in *C. albicans* yields tetraploid progeny, which in appropriate growth conditions undergo random chromosome loss to revert to diploidy.²¹

Similarly, perceptions of *Leishmania* as entirely clonal have been questioned by reports of several instances of naturally occurring hybrid strains, especially from the New World. On the basis of phenotypic and genotypic markers, hybrids have been described between *L. braziliensis* and *L. panamensis* in Nicaragua⁹; between *L. braziliensis* and *L. guyanensis* in Venezuela²²; between *L. braziliensis* and *L. guyanensis* in Ecuador²³; and between *L. braziliensis* and *L. peruviana* in Peru.⁶ In the Old World, *L. major/arabica* hybrids were described.²⁴ Putative parental and hybrid phenotypes of the *L. donovani* complex (*L. donovani*; "*L. archibaldi*") occur, sympatrically in East Africa, and sequencing of housekeeping genes encoding enzymes shows mosaic characters across such strains.²⁵ Preliminary genetic analysis suggests that genetic exchange may occur among *L. tropica* populations in the Middle East.²⁶ Most recently, genetic hybrids between *L. infantum* and *L. major* have been described, from immunocompromised patients in Portugal.²⁷ Mixed *Leishmania* infections occur in natural hosts and in vectors, and infections in mammals may

last for decades, providing ample opportunities for interactions between distinct genotypes. Overall, this is substantial circumstantial evidence that an extant mechanism of genetic exchange remains to be described for *Leishmania*, with epidemiologic implications, for example, in the spread of emergent strains or drug resistance.

L. peruviana was recorded in the Department of Huánuco prior to the epidemic from which the isolates in this study were derived. However, prior to this epidemic, CL was seldom encountered and no cases of MCL were recorded (Llanos-Cuentas, unpublished data). It is likely that introduction of *L. braziliensis*⁶ resulted in the increased prevalence of CL, and outbreak of MCL. *L. braziliensis* may have been introduced by human immigration from another region or by human or canine intrusion into an unidentified sylvatic transmission cycle.

The 59 Huánuco isolates analyzed here showed a remarkable degree of isoenzyme diversity considering that they originated from such a small geographical area. *L. guyanensis* and *L. amazonensis*, previously reported from Peru,¹ were not identified. However, 4 species—*L. peruviana*, *L. braziliensis*, *L. lainsoni*, and *L. (V.) sp. n.*⁷—were found to occur sympatrically in Huánuco. In addition, *L. braziliensis/L. peruviana* phenotypic hybrids were common and almost as abundant as *L. braziliensis*. This local prevalence of hybrid strains is reminiscent of the predominance of the genetic hybrids of *T. cruzi* among human cases of Chagas disease in Paraguay and adjacent regions.²⁸ The gene encoding the enzyme MPI has recently been sequenced from both *L. braziliensis* and *L. peruviana*, and phenotypic differences have been shown to correspond with a single nucleotide polymorphism (SNP), changing a threonine to an arginine, which has been used as the basis of a PCR identification assay.²⁹ It would be of interest to confirm that the current *L. braziliensis*, *L. peruviana*, and *L. braziliensis/L. peruviana* hybrid isolates, and those from a wider geographical range, conform to the predicted SNP genotypes.

It was surprising to find 4 MLEE phenotypes and 7 microsatellite genotypes among the *L. braziliensis/L. peruviana* hybrids. This might be a consequence of the rapid evolution of microsatellites, or it could be consistent with the occurrence of more than one hybridization event. It is very unlikely that these genotypes are explicable by mutation, with no hybridization event. By analogy, multilocus sequence typing (MLST) of the *L. donovani* complex has revealed multiple heterozygous sites within a gene and at several loci, with sharing of alleles within and across genetic groups: recombination,^{25,30} rather than mutation,³¹ is considered to be the most parsimonious explanation.

The majority (12) of the *L. braziliensis/L. peruviana* hybrids had a single genotype (Table 1). The expansive clonal propagation of one of the putative hybrid genotypes suggests that a hybrid agent has emerged with increased fitness relative to the parental strains, although a neutral event, such as a population bottleneck, unlikely in view of the diversity within the population, may also have been involved. One comparison of promastigote growth rates found no evidence that *L. braziliensis/L. peruviana* hybrids had an enhanced growth rate in vitro.³² Nevertheless, it would be of interest to compare the metastatic potential of the hybrid and nonhybrid genotypes in the hamster model of MCL³³ or perhaps in the mouse ear model for dissemination.³⁴ The occurrence of *L.*

braziliensis/L. peruviana hybrids in dogs and humans (Tables 1 and 2) from the same area indicates that both are exposed to the same infective sand fly population. This suggests that dogs might act as a reservoir host and enhance propagation of the emergent hybrid genotype.

Six zymodemes and at least 6 microsatellite genotypes were isolated from patients with MCL. Isolation of *L. peruviana* from a single case of MCL is an interesting finding but, in the absence of more cases, must be interpreted with caution. Previously in Peru, a single case of mucosal leishmaniasis (ML) was attributed to *L. peruviana*.¹ Four of the MCL patients yielded a hybrid isolate, with no evidence of mixed infection, so patients carrying such isolates must be considered at risk of developing MCL. Thus, the finding of unequivocal genotypic markers for isolates that carry the risk of progression to MCL is still an elusive goal.³⁵

As mentioned above, MLST applied to the *L. donovani* complex has already revealed the genetic basis of MLEE and provided a higher resolution approach to resolving genetic groups and to understanding relationships between them.^{25,30} MLMT has given even higher resolution of intraspecific population structure.¹⁰ Diversity of the subgenus *L. (Viannia)* is seen in some endemic foci,³⁶ although the level observed in Huánuco is extraordinary. Genetic diversity has also been recorded from individual patients.³⁷ Furthermore, it is apparent that there is some overlap of boundaries between the perceived species of the subgenus *L. (Viannia)*. In combination, MLST and MLMT will be a powerful approach to understanding the complex molecular epidemiology and population genetics of *L. (Viannia)* and for further investigation of the extent of genetic exchange in natural or experimental populations. For the future MLMT of *L. (Viannia)*, a much wider panel of microsatellite markers is required, and such a panel is in preparation (R. Oddone, G. Schönián, and K. Kuhls, personal communication and unpublished data). In localities where *L. (Viannia)* species overlap, we anticipate the discovery and possible emergence of other human infective hybrid genotypes, some with potential to generate severe disease.

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