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| Original Citation:   |
|--|
| Availability: This version is available http://hdl.handle.net/2318/98855 since   |
| Published version: DOI:10.1016/j.biocon.2010.03.001  |
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Host taxon-derived Sarcoptes mite in European wild animals revealed by microsatellite markers, Rasero R, Rossi L, Soglia D, Maione S, Sacchi P, Rambozzi L, Sartore S, Soriguer R, Spalenza V, Alasaad S. DOI: 10.1016/j.biocon.2010.03.001

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# Host taxon-derived Sarcoptes mite in European wild animals

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- 16 **Abstract** Microsatellite genotyping was applied on individual Sarcoptes mites from 15
- wild mammalian populations belonging to 10 host species in three European countries,
- 18 using 10 Sarcoptes mite specific markers. The results showed that the geographical
- 19 separations had real biological significance for the definition of mite sub-populations,
- 20 and that the degree of genetic exchange occurring between mites from different
- 21 localities was related to the geographical distance between locations. Wild host-derived
- 22 mite populations were clustered into three main groups: herbivore-, carnivore- and
- 23 omnivore-derived Sarcoptes mite populations. Omnivore-derived was halfway between
- 24 herbivore- and carnivore-derived Sarcoptes mite populations. The separation between
- 25 the three mite groups (herbivore-, carnivore- and omnivore-derived mite) was more

supported than that by the geographical separations; nevertheless a kind of subclustering was detected within each group (carnivore-, omnivore- and herbivore-), scattering mite populations up to their geographical localities (countries). The lack of gene flow between Sarcoptes populations may have improved parasitic adaptations and led to, what we called host taxon-derived (carnivore host-, herbivore host- and omnivore host-derived) Sarcoptes mite populations in European wild animals. Our results demonstrated that Sarcoptes is not a single panmictic population even within locations, which will have important ramifications on the study of population genetic structure, life cycle, diagnosis and monitoring protocols, and could contribute to the better understanding of the epidemiology of the ubiquitous Sarcoptes mite.

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37 **Keywords:** Sarcoptes scabiei; Genetic epidemiology; Genetic structure; Microsatellite

38 markers; Omnivore-derived; Carnivore-derived; Herbivore-derived; Host taxon-derived.

#### Introduction

Predicting the spread of a disease to wildlife is critical in order to identify populations at risk, target surveillance, and design proactive management programmes (Blanchong et al. 2008). Recently, there has been an increased interest in diseases of free-living animals and the ecological role of diseases in populations, particularly their ability to regulate animal abundance (Scott 1988; Lyles & Dobson 1993; Robinson 1996; Daszak et al. 2000). There is also an awareness that free-living species may act as reservoirs of diseases of man and domestic animals (Robinson 1996; Daszak et al. 2000).

Although neglected as a pathogen, the ectoparasite Sarcoptes scabiei continues to affect humans and a wide range of mammalian hosts on a worldwide scale (Bornstein et al. 2001; Pence & Ueckermann 2002; Walton et al. 2004a). The introduction of infected domestic animals and the succeeding adaptation of Sarcoptes mite to a new highly susceptible and receptive wild host have been proposed to give rise to sarcoptic mange epizootics in previously mange-free wildlife populations (Arlian 1989).

In several European wild mammal populations, Sarcoptes mite infections are endemic and cause devastating mortality as reported especially for Alpine and Pyrenean chamois, Iberian ibex, aoudad, and red fox (Fandos 1991; Mörner 1992; Pérez et al. 1997; León-Vizcaino et al. 1999; González-Candela et al. 2004; Rossi et al. 2007). Notwithstanding, only few cases have been reported in other sympatric host species, like stone marten, badger, lynx, and roe deer (Ryser-Degiorgis et al. 2002; Oleaga et al. 2008).

Morphological studies have failed to identify any significant differences among mite populations (Fain 1978), and experimental cross contamination of Sarcoptes mite between hosts of different species is commonly unsuccessful (Arlian et al. 1984; Arlian

1989). Apparently no epidemiological relationship exists, in Europe, between mange foci affecting wild ruminants, wild boars, and carnivores (Berrilli et al. 2002).

The question as to whether Sarcoptes mites might be divided into different species or whether they are, in fact, monospecific has been the subject of an ongoing debate (Zahler et al. 1999; Burgess 1999; Berrilli et al. 2002; Gu & Yang 2008; Alasaad et al. 2009c). Zahler et al. (1999) and Berrilli et al. (2002), using the ITS-2 sequences as genetic markers, did not detect clear-cut evidence of genetic separation related to host species or geographic location. As well as, in our previous study, we have shown that ITS-2 rDNA does not appear to be suitable marker for examining genetic diversity among Sarcoptes mite populations from different wild host species and/or geographical localities (Alasaad et al. 2009c). In phylogenetic analyses bootstrapping support for the closest relationships may be relatively poor due to reduced time to accumulate informative changes in the sequences examined. Further resolution is therefore provided in faster evolving hypervariable sequences such as nuclear polymorphic microsatellite loci (Walton et al. 2004b).

Walton et al. (1999; 2004b), using multi-locus genotyping applied to microsatellite markers, substantiated previous data that gene flow between scabies mite populations on human and dog hosts is extremely rare in northern Australia. As well as, genetic differences were detected between geographically distinct populations, even between householders. Microsatellite markers were used by Alasaad et al. (2008b) to describe a new phenomenon of genetic structuring among S. scabiei at individual host skin-scale.

Taking into account all the above-mentioned information, the aim of the present study was to test the extent of genetic relationship between sympatric wild host-derived Sarcoptes mite populations, and to study the influence of the geographical isolation on

the genetic structuring of Sarcoptes. This is pivotal for wildlife health management in order to understand the geographic variation among bordered mite populations, and to measure the patterns of host specific differences, especially in sympatric hosts.

### Materials and methods

## Collection of S. scabiei

Using Postponed Isolation (Post-frozen Isolation) and Direct Isolation (Live Isolation) methods, as described by Alasaad et al. (2009b), 251 Sarcoptes mites were collected from the crusted skin of 100 animals belonged to 15 populations of 10 European wild mammalian species, as listed in Table 1, which were sampled in Italy, France and Spain (Fig. 1). Rupicapra rupicapra rupicapra, Cervus elaphus, Martes martes, Ovis musimon, Capra ibex, and Vulpes vulpes were sympatric in Northeast Italian Alps. V. vulpes, Martes foina and Sus scrofa were sympatric in Northwest Italian Alps. Taking into account the topography of Sierra Nevada Mountain and that the first case of Sarcoptes mite infection was reported in Dílar Valle (East Sierra Nevada) (Pérez et al. 1997), mites from Sierra Nevada were divided into two different groups, East and West populations. All mites were identified as S. scabiei based on known morphological criteria (Fain 1968).

In Table 1, term 'Code' refers to all mites belonging to the same geographical and/or host species-derived population, from now onwards called 'component population' or, simply, population (Bush et al. 1997).

### Preparation of Sarcoptes gDNA

114 115 The DNA of individual Sarcoptes mites was extracted with the NucleoSpin Tissue kit 116 procedure (Macherey-Nagel, Düren, Germany) with some modifications proposed by 117 Soglia et al. (2009), and recently with HotSHOT Plus ThermalSHOCK technique 118 (Alasaad et al. 2008a). 119 120 Fluorescent-based polymerase chain reaction analysis of microsatellite DNA 121 122 From the panel described by Walton et al. (1997), ten microsatellites (Sarms 33-38, 40, 123 41, 44, and 45) were selected and analysed with one 10× multiplex PCR. Each 15 μl 124 PCR reaction mixture consisted of 3 µl of the single mite DNA, together with the PCR 125 mixture containing all primer pairs (ranged from 0.04 to 0.1 µM per primer), 200 µM of 126 each dATP, dCTP, dGTP, and dTTP, 1.5 µl of 10× PCR buffer (200 mM KCl and 100 mM Tris-HCl, pH 8.0), 1.5 mM MgCl2, and 0.15 µl (0.5 U/reaction) HotStartar Taq 127 128 (QIAGEN, Milano, Italy). Samples were subjected to the following thermal profile for 129 amplification in a 2720 thermal cycler (Applied Biosystems, Foster City, CA, USA): 15 130 min at 95°C (initial denaturing), followed by 37 cycles of three steps of 30 s at 94°C 131 (denaturation), 45 s at 55°C (annealing) and 1.5 min at 72°C (extension), and a final 132 elongation of 7 min at 72°C. 133 134 Microsatellite analysis 135

Using 96-well plates, aliquots of 12  $\mu$ L of formamide with Size Standard 500 Liz (Applied Biosystems, Foster City, CA, USA) and 2  $\mu$ l PCR product were prepared. Then, the plates were heated for 2 min at 95°C and chilled to 4°C. Fluorescent PCR

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amplification products were analyzed by ABI PRISM 310 Genetic Analyzer with pop4.

Allele calling was performed using the GeneMapper v. 4.0 software (Applied

Biosystems, Foster City, CA, USA). To track and minimize the amount of error

associated with genotyping, the genetic data were collected twice by SA and DS.

Descriptive statistics and cluster analysis

CONVERT 1.31 software (Glaubitz 2004) was used to reformat files for the statistical softwares. Descriptive statistics and diversity analyses were carried out with GenAlEx v. 6.2 (Peakall & Smouse 2006), Genepop v. 4.0 (Raymond & Rousset 1995), Fstat v. 2.9.3 (Goudet 1995), and Arlequin v. 3.1 (Excoffier et al. 2005) softwares, i.e. allelic richness (R), number of private alleles, allele frequencies, unbiased expected (He) and observed (Ho) heterozygosity, test for Hardy-Weinberg equilibrium (HWE), test for linkage equilibrium (LE), and F statistics. All pairs of the component populations were compared for homogeneity of genetic variation using the Wilcoxon's matched-pairs signed-rank test (GraphPad InStat software).

Analysis of structure and relationships among host-specific mite populations were studied using two different approaches:

(i) Multilocus proportion of shared alleles (Dps) was computed between all possible pairs of individual mites using the Microsat software (Minch 1997) ignoring any preliminary information on origins of parasites. One thousand datasets were generated by resampling the input data (bootstrapping), the Neighbor-Joining algorithm was used as implemented by the Phylip v. 3.6 package (Felsenstein 1989), and a consensus dendrogram was obtained. The dendrogram was visualized using the Dendroscope v.

2.2.2 software (Huson et al. 2007).

(ii) The analysis of relationships among mites was then improved by a Bayesian assignment test using the method implemented by the Structure v. 2.2 software (Pritchard et al. 2000). We performed 50000 MCMC (Markov chain Monte Carlo) replicates following a burn-in period of 10000 steps. This parameter set was run 20 times for each of different numbers, K, of genetic clusters of multilocus genotypes; all values of K from 1 to 20 were tested. The probability of the multilocus genotype of any individual mite to occur in each of the K clusters was computed. We used the admixture model (each mite drew some fraction of its multilocus genotype from each of the K clusters) allowing the allele frequencies to be correlated among clusters. This configuration has been considered the best in the case of subtle population structure (Falush et al. 2003). We used the height of the modal value of the distribution of  $\Delta K$  in order to estimate the uppermost number of clusters capturing the overall mite sample structure, as suggested by Evanno et al. (2005). We then associated any individual mite with the cluster that corresponded to its greatest membership, q, that is fraction of its multilocus genotype; a threshold value  $q \ge 0.9$  was used. Finally, each of the inferred clusters was associated with the component populations of its mites. If a cluster was labelled with multiple mite populations, an additional substructure analysis for K values from 1 to 5 was performed testing only the mites assigned to that cluster.

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## **Results**

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# Descriptive statistics

Ten marker loci were analysed on 251 mites belonging to 15 populations of 10 European wild mammalian species from Italy, France and Spain; 101 alleles were detected. Allele count for each of the 10 loci ranged from six (Sarms 37) to 15 (Sarms

34). Proportion of missing genotypes was as low as 0.04 and it did not affect single locus or population. Forty two private alleles were detected in 11 wild host-derived mite populations, ranging from 1 (ItNWMf, SpNEVv and SpNWRp) to 10 (FrNESs). Whereas in ItNECe, SpEMf, ItNEOam, and ItNEMm no private alleles were identified (Table 2). The highest within-population genetic variability was observed for the two S. scrofa mite populations, in spite of their small size, whereas little variation was found in ItNWVv and SpNWRp (Table 3).

Allelic richness (R) and heterozygosity (He) were used as most informative parameters of diversity. In particular, allelic richness provided a measure of the number of alleles independent of sample size, hence allowing comparison among different populations. Level of genetic diversity varied both across loci and among populations. Wilcoxon's test stated (P<0.001) that C. ibex mites showed more variability (R = 2.6, He = 0.339) than mites from the other ruminants whereas S. scrofa mites were the most variable at all (R = 3.4, He = 0.667).

LE test (Lewontin 1964; Slatkin 1994; Slatkin & Excoffier 1996) was performed for all loci and significant linkage disequilibrium (P<0.05) was observed for 18 pairs when all the mite populations were pooled. In no cases disequilibrium was observed at the same loci in more than two populations individually analysed. HWE estimates were assessed of 70 locus-by-population comparisons, 40 (57%) showed significant heterozygosity deficiencies. Deviations from HWE did not point at any locus in particular. All populations deviated from HWE across loci after sequential Bonferroni correction (P<0.001).

Population differentiation based on allele frequencies for all 15 populations gave an overall Fst = 0.721. Each locus significantly (P<0.001) contributed to distribution of variability among populations with per-locus values ranging from 0.290 to 0.821. This

very high estimate means that most of the global Sarcoptes genetic variability resided among rather than within component populations different in geographical and host-derived distribution of mites.

Structure and relationships among mite populations

(i) Multilocus proportion of shared alleles (Dps) as a measure of genetic similarity between all pairs of mites.

Genetic variability among populations of Sarcoptes mites collected from the same host species from different localities.

The proportion of shared alleles between pairs of individual mites from the two C. pyrenaica mite populations in Spain (SpSWCp and SpSECp) were scattered randomly with no evidence of distribution based on geographical location of hosts. Individual Sarcoptes mites belonging to the three V. vulpes mite populations from the Northeast and Northwest Italian Alps as well as from Northeast Spain showed clear clustering up to their original populations. V. vulpes mite population from Spain was the most different one, supported by 980/1000 bootstraps. The V. vulpes mites from Northeast and Northwest Italian Alps were relatively more similar between each other, their distribution across two distinct clusters being very poorly supported (169/1000 bootstraps) (Fig. 2). Regarding mites from S. scrofa populations from Northwest Italian Alps and from Northeast France, strong separation was detected between them (1000/1000 bootstraps, data not shown).

Genetic variability among populations of Sarcoptes mites collected from sympatric host

239 species.

The dendrogram of individual mites from six sympatric host-derived populations from Northeast Italian Alps, and three sympatric host-derived populations from Northwest Italian Alps allowed the clustering of the mites into three groups (Fig. 3). The first group was formed by all carnivore-derived mites from East and West Italian Alps, mainly mites from V. vulpes and, in additon, M. foina (Figure 3, a) and M. martes (Figure 3, b), nevertheless it included also a mite from C. ibex (Figure 3, c). The second group included the herbivore-derived mite populations from Northeast Italian Alps, mainly mites from C. ibex and R. Rupicapra, all scattered across the cluster. In addition, the cluster included O. aries musimon (Figure 3, d) and C. elaphus (Figure 3, e) mites. The separation of herbivore- and carnivore-derived mites was then quite clear-cut (470/1000 bootstraps). The S. scrofa mites from Northwest Italian Alps were near the carnivore-derived mites but distinct from them (572/1000 bootstraps).

species and geographical localities.

Five clusters resulted from the analysis of the overall 15 wild host-derived mite populations (Fig. 4). Cluster I included almost all the herbivore-derived mites from Spain (SpNWRp, SpSWCp and SpSECp) and the C. ibex mite which clustered with carnivore-derived parasites in Figure 3 (c). Although some mites scattered around, evidence of separation between SpNWRp and SpSWCp-SpSECp was observed. Clusters IIa and IIb contained all the omnivore-derived mites, ItNWSs and FrNESs respectively, and one ItNECi (included in cluster IIb, Figure 4, arrow). Cluster III included all the carnivore-derived mites, i.e. ItNEMm, ItNWMf, ItNWVv, ItNEVv as well as SpNEVv (Figure 4, a) and SpEMf (Figure 4, b). Cluster IV contained almost all

Genetic variability among Sarcoptes populations distributed according to both host

the herbivore-derived mites from Italy (ItNERr, ItNECi, ItNECe, and ItNEOam,) and one SpNWRp.

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(ii) Analysis of mite population structure by the Bayesian method.

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The modal value of the statistic  $\Delta K$  for the overall dataset (251 mites) stated that the uppermost cluster value was K = 4 (Evanno et al. 2005). Each of the four inferred clusters was then associated with the information of its mites. For each cluster the average membership and number of mites assigned with the greatest membership were computed (Fig. 1). Cluster I shared all mites with the cluster I in Fig. 4, i.e. nearly all mites of Spanish ruminants (SpNWRp, SpSECp and SpSWCp) and one ItNECi. High proportions of membership were always obtained (q > 0.97). Cluster II grouped all the S. scrofa mites (ItNWSs and FrNESs) (see Fig. 4, clusters IIa and IIb) and their membership fraction was q > 0.98. Two ItNECi (for one of them see Fig. 4, cluster IIb) and three SpNEVv mites were also added but they lacked to show high membership in this cluster (q < 0.70). Cluster III shared most mites with the cluster III in Fig. 4 grouping all mites of Italian V. vulpes (ItNEVv and ItNWVv), M. martes (ItNEMm), and M. foina (ItNWMf and SpEMf) with very robust membership (q > 0.95) for all mites but SpEMf parasite with q = 0.86). One SpNEVv mite showed some similarity to this cluster although with poor membership (q = 0.56). Cluster IV grouped the majority of the mites of Italian ruminants (ItNERr, ItNECi, ItNECe, and ItNEOam) and one SpNWRp with very high membership (q > 0.97). Its members corresponded to those of the cluster IV in Fig. 4.

The computation of the statistic  $\Delta K$  was repeated separately for four subsets of samples made up of the main geographical and host-specific mite groups, i.e. mites

belonging to S. scrofa (ItNWSs and FrNESs), Italian ruminants (ItNERr, ItNECi, ItNECe, and ItNEOam), V. vulpes (ItNEVv, ItNWVv, and SpNEVv), and Spanish ruminants (SpNWRp, SpNECp and SpNWCp). No evidence of substructure was detected in any case.

Spanish V. vulpes mites received ambiguous assignment as in case of mixed ancestry. However, they were collected from a single host animal and lacked a substantial component population as a reference. Two ItNECi were misplaced in S. scrofa mite cluster with low membership. One of them was also assigned to the sympatric V. vulpes mite cluster (Fig. 3, c) or to the allopatric Spanish ruminant mite cluster (Fig. 4, Cluster I) depending on the populations used for the comparison. Such individual parasites seemed to be randomly assigned since they carried multilocus genotypes infrequent in their population, so the algorithm could not recognize their ancestry.

In synthesis, the proportion of shared alleles as a similarity measure among mites and the assessment of structure using the Bayesian method provided patterns in agreement with each other. When the full data set was used, four distinct genetic clusters of mites were inferred, i.e. omnivore-, Italian herbivore-, carnivore-, and Spain herbivore-derived parasites.

### Discussion

Differentiation of host-specific mites using morphological traits, apart from being very difficult and time consuming, proved to be impossible to implement when mites of the same host-specific variant, but belonging to different geographical component populations, have to be compared (Arlian et al. 1984; Arlian 1989). Short fragments of

mitochondrial or ribosomal DNA spacer regions have been shown not to be suitable markers for examining genetic diversity among Sarcoptes mite populations (e.g. Alasaad et al. 2009c). Further resolution is therefore provided in faster evolving hypervariable sequences such as nuclear polymorphic microsatellite loci. Microsatellites have previously demonstrated to provide strong support for geographically discrete populations, they showed congruence with evolutionary patterns at the population level, and reported genetic differentiation at the skin-scale of individual mangy hosts (Bowcock et al. 1994; Walton et al. 2004b; Alasaad et al. 2008b).

The number of the mites utilized in this study (251 samples) has to be considered high as compared with previous studies in this field (see Alasaad et al. 2009a for review).

All the component populations showed a strong deficiency of heterozygosity over all loci and mites belonging to the same component population showed to be scattered through the same cluster more than subdivided across individual host animals.

Sarcoptes mites lack free-living stages. Individual hosts, depending on their susceptibility and behaviour, are more or less ephemeral habitats and may provide patchy environments which hamper random mating (Price 1980; Criscione et al. 2005). All mites on an individual host could form an 'infrapopulation' (Bush et al. 1997) with some recurrent generations on that host. Number of generations is affected by short generation interval of the parasite, about three weeks, as well as by life expectancy of the infested host, depending on its susceptibility.

In our data set, the reduced gene pools made mites alike each other and hid possible equilibrium between dispersive process and gene flow among infrapopulations. This may be due to rapid diffusion of few genotypes as for an epidemic population structure (Oura et al. 2005).

Another evident feature of our results is the lack of homogeneity of genetic diversity across populations. S. scrofa mites were the most variable at all. Wild boar populations are widespread and growing, and generally show higher resistance to parasites than other mammalian species (Rodrigues & Hiraoka 1996; Nejsum et al. 2009). Consequently, a single host can be affected by repeated infestation events through mites from other infrapopulations or, even, from other component populations.

The other major determinants of gene flow among mites are the degree of host specificity and geographical structure of host populations. Previous investigations showed that approaches to individual clustering provide appropriate characterisation of population structure at high Fst values (Rosenberg et al. 2001; Manel et al. 2002; Latch et al. 2006). In the presence of very diverging taxa, few loci are needed to achieve high performance, regardless of the sample sizes (Manel et al. 2002; Tadano et al. 2008). In fact, the ideal marker locus for our purposes should be monomorphic within any taxon and polymorphic across taxa (Reed 1973).

In our data set, the unusually high value of Fst and the high number of private alleles, in most populations, indicated that the mite component populations were very unlike each other. All the 10 loci provided a significant component of among-population diversity. As a consequence, our marker panel provided good accuracy for analysis of the genetic characteristics of Sarcoptes populations.

Sarcoptes scabiei (1) from different host species belonging to different geographical localities, (2) from the same host species belonging to different geographical localities, and (3) from closely related host species belonging to different geographical localities clustered up to their original populations. Clear genetic diversity among mite populations from different geographical localities exists. The differences show to be as stronger as the geographical separation between host populations is

larger. In the case of short geographical separations (East and West Sierra Nevada) mites from C. pyrenaica scattered randomly in the dendrogram and no clear separation was detected. The differentiation between V. vulpes mites from Spain and Italy was highly supported whereas the genetic separation between V. vulpes mites from East and West Italian Alps was poorly supported. This finding suggests that gene flow occurring among mites from different localities is related to the geographical distances.

The individual mites belonged to the six sympatric host-derived mite populations from East Italian Alps, and the three sympatric host-derived mite populations from West Italian Alps clustered into three main groups (Fig. 1, 3, and 4): herbivore-derived mites (ItNECi, ItNERr, ItNEOm and ItNECe), carnivore-derived mites (ItNEVv, ItNEMm, ItNWMf and ItNWVv), and omnivore-derived mites (ItNWSs). In particular, mites from S. scrofa were distinct from both herbivore- and carnivore-derived mites and they did not cluster with the sympatric Northwest Italian populations. In other words, the host-specific separation among the three clusters was stronger than that by the geographical separation between East and West Italian Alps.

Similar results were obtained when the overall mite samples of our investigation were analysed (Fig. 1 and 4). For example, Cluster III contained all the carnivorederived mites regardless of their geographical origins from different European countries under study.

Our results from the sympatric wild animals in Italy and from the general analysis of all mite populations show unambiguously lack of gene flow or recent admixture among Sarcoptes populations carnivore-, herbivore-, and omnivore-derived. Mite transmission may occur within each mite cluster herbivore-, carnivore-, and omnivore-derived but it seems to be extremely rare or absent among them. This might

improve parasite adaptations and led to, what we called, host taxon-derived (carnivore, herbivore, and omnivore host-derived) Sarcoptes mite populations.

Population structure of Sarcoptes would be that of a species subdivided into genetically small populations with restricted gene flow among local demes (Price 1980; Martínez et al. 1999; Nadler & Hafner 1990). Strong specialisation could be the result of a host taxon-derived shift and, even if two host taxon-derived species are sympatric as for their host species, they should be considered as allopatric if the parasites have no possibility of host choice. In other words, the host sympatry is not the same as the parasite sympatry.

The probability of disease transfer between sympatric host taxon-derived species could be reduced by evolved intrinsic mechanisms, this means that the behaviour has been selected to impede crosses between individuals from two different host taxon-derived, and this could represent the first step of sympatric speciation (McCoy 2003). Host taxon-derived effect is stronger than the geographical separation in the definition of speciation events.

The existence of host taxon-derived Sarcoptes mites could be the explanation of the mange-free wildlife populations in sympatry with other mangy wild animals, like the mange-free C. ibex and R. rupicapra of West Italian Alps which are in sympatry with the endemically mangy population of V. vulpes. This effect could be the reason behind the successful of the cross-transmission/infection in some Sarcoptes varieties e.g. S. scabiei var vulpes/canis readily infect dogs and other canids as well as felids including domestic European cats, as all of them belong to the same host taxon-derived (Bornstein 1995).

We have no clear explanation of this taxonomic affiliation. Further studies on dispersal capability of host animals and their disposition to interact with each other, host

behaviour and parasite adaptation are needed to explain the host taxon-derived Sarcoptes. Characterization of host genetic structure in addition to mite population genetic structure would contribute additional valuable information.

## Acknowledgements

We would like to thank A.R. Molinar Min (Università degli Studi di Torino-Italy) for assistance with mite collections; S. Lavin (Universitat Autonoma de Barcelona-Spain), S. Rossi (ONCFS, Paris-France), G. Capelli (Istituto Zooprofilattico Sperimentale delle Venezie, Padua-Italy), W. Mignone (Istituto Zooprofilattico Sperimentale del Piemonte, Liguria e Valle d'Aosta, Imperia-Italy), J.M. Pérez (Jaén University, Jaén-Spain) and J.E. Granados (Espacio Natural Sierra Nevada, Granada-Spain) are thanked for providing mite samples and the last two for supporting S.A. investigation stay in Italy. S. Angelone (Swiss Federal Institute WSL, Zürich-Switzerland) and X.Q. Zhu (South China Agriculture University, Guangzhou-China) are thanked for helpful comments. The research was supported by MURST contract year 2004, Prot. 2004078701\_001 (LR)

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**Table 1** Countries, geographical locations and host species used in this study, together with the number of host animals and Sarcoptes mite samples.

| Code    | Countries | Geographical | Host taxon | Host species        | N°. of animals N | N°. of mites |
|---------|-----------|--------------|------------|---------------------|------------------|--------------|
| ItNERr  | Italy     | Northeast    | Herbivore  | Rupicapra rupicapra | 20               | 63           |
| ItNECi  | Italy     | Northeast    | Herbivore  | Capra ibex          | 10               | 25           |
| ItNECe  | Italy     | Northeast    | Herbivore  | Cervus elaphus      | 1                | 1            |
| ItNEOam | Italy     | Northeast    | Herbivore  | Ovis aries musimon  | 1                | 2            |
| ItNEVv  | Italy     | Northeast    | Carnivore  | Vulpes vulpes       | 7                | 23           |
| ItNEMm  | Italy     | Northeast    | Carnivore  | Martes martes       | 1                | 3            |
| ItNWVv  | Italy     | Northwest    | Carnivore  | Vulpes vulpes       | 11               | 30           |
| ItNWMf  | Italy     | Northwest    | Carnivore  | Martes foina        | 1                | 2            |
| ItNWSs  | Italy     | Northwest    | Omnivore   | Sus scrofa          | 1                | 3            |
| FrNESs  | France    | Northeast    | Omnivore   | Sus scrofa          | 4                | 5            |
| SpNEVv  | Spain     | Northeast    | Carnivore  | Vulpes vulpes       | 1                | 4            |
| SpNWRp  | Spain     | Northwest    | Herbivore  | Rupicapra pyrenaica | 9                | 26           |
| SpSECp  | Spain     | Southeast    | Herbivore  | Capra pyrenaica     | 21               | 33           |
| SpSWCp  | Spain     | Southwest    | Herbivore  | Capra pyrenaica     | 11               | 30           |
| SpEMf   | Spain     | West         | Carnivore  | Martes foina        | 1                | 1            |

**Table 2** Private alleles detected at the 10 microsatellite loci for the host-associated mite populations, together with their frequencies.

| Pop (N°mites) | Locus | Allele      | Freq       | Pop (N° mites)   | Locus       | Allele     | Freq  |
|---------------|-------|-------------|------------|------------------|-------------|------------|-------|
|               | ms33  | 224         | 0,008      | ItNEVv (23)      | ms35        | 150        | 0,065 |
|               | ms33  | 244         | 0,025      | ILINEVV (23)     | ms41        | 232        | 0,022 |
|               | ms34  | 170         | 0,016      | ItNWVv (30)      | ms35        | 146        | 0,200 |
| ItNERr (63)   | ms34  | 192         | 0,190      | 1111 00 0 0 (30) | ms41        | 264        | 0,033 |
|               | ms41  | 214         | 0,008      | SpNEVv(23)       | ms38        | 205        | 1,000 |
|               | ms41  | 250         | 0,083      |                  | ms33        | 266        | 0,800 |
|               | ms38  | 290         | 0,008      |                  | ms33        | 268        | 0,100 |
|               | ms34  | 188         | 0,104      |                  | ms33        | 270        | 0,100 |
|               | ms34  | 208         | 0,042      |                  | ms34        | 182        | 0,200 |
| ItNECi (25)   | ms35  | 138         | 0,022      | FrNESs(5)        | ms35        | 126        | 0,200 |
| ItiNECI (23)  | ms37  | 176         | 0,045      |                  | ms35        | 128        | 0,300 |
|               | ms41  | 244         | 0,026      |                  | ms36        | 287        | 0,400 |
|               | ms38  | 223         | 0,043      |                  | ms37        | 178        | 0,900 |
|               | ms35  | 160         | 0,333      |                  | ms41        | 228        | 0,600 |
|               | ms36  | 263         | 0,017      |                  | ms44        | 274        | 0,700 |
| SpSWCp(30)    | ms36  | 273         | 0,017      |                  | ms33        | 274        | 1,000 |
|               | ms40  | 217         | 0,100      | ItNWSs(3)        | ms34        | 200        | 0,250 |
|               | ms40  | 225         | 0,067      | 1014 44 25 (3)   | ms40        | 235        | 1,000 |
|               | ms35  | 158         | 0,015      |                  | ms45        | 176        | 0,500 |
| SpSECp(33)    | ms36  | 277         | 0,015      | ItNWMf(2)        | ms38        | 219        | 0,250 |
|               | ms45  | 164         | 0,030      | SpNWRp (26)      | ms45        | 198        | 0,679 |
| ItNECe(:      | 1);   | l); ItNEOar | n(2) and I | tNEMm(3): no pr  | ivate allel | e detected |       |
|               |       |             |            |                  |             |            |       |

Table 3 Descriptive statistics for the main mite populations (R=allelic richness;

He=expected heterozygosity; Ho=observed heterozygosity)

|    | ItNERr | ItNECi | ItNEVv | ITNWVv | SpNWRp | SpSWCp | SpSECp | FrNESs-<br>ItNWSs |
|----|--------|--------|--------|--------|--------|--------|--------|-------------------|
| R  | 1.9    | 2.6    | 2.0    | 1.5    | 1.6    | 1.8    | 1.8    | 3.4               |
| Не | 0.215  | 0.339  | 0.232  | 0.119  | 0.119  | 0.217  | 0.216  | 0.667             |
| Но | 0.048  | 0.097  | 0.126  | 0.020  | 0.051  | 0.077  | 0.103  | 0.340             |

**Fig. 1** Europe map showing approximate sites for sample collection, together with structure clusters. The colours within bars show the proportion of membership of each individual to the genetic clusters for each Sarcoptes population separately. The pie charts give the genetic membership per Sarcoptes population. 1=SpNWRp, 2=SpSWCp, 3=SpSECp, 4=ItNWSs, 5=FrNESs, 6=SpNEVv, 7=ItNEVv, 8=ItNWVv, 9=ItNEMm, 10=ItNWMf, 11=SpEMf, 12=ItNERr, 13=ItNECi, 14=ItNECe, and 15=ItNEOam. For site abbreviations see Table 1.

**Fig. 2** Unrooted Dps consensus dendrogram for individual Sarcoptes mites from three V. vulpes-derived mite populations from Northwest and Northeast Italian Alps, and from Northeast Spain. Numbers at the nodes are percentage values of 1000 bootstraps supporting the same branching structure. Codes in this figure (bold for Northwest Italian Alps, italic for Northeast Italian Alps, grey for Northeast Spain) represent the sample codes in Table 1 and Fig.1.

Fig. 3 Unrooted Dps consensus dendrogram for individual Sarcoptes mites from six sympatric host-derived mite populations in Northeast Italian Alps, and three sympatric host-derived mite populations in Northwest Italian Alps (Table 1). Numbers at the nodes are percentage values of 1000 bootstraps supporting the same branching structure. Thick branches for all carnivore-derived mites, thin branches for all herbivore-derived mites, grey branches for S. scrofa-derived mites. Carnivore-derived mites: V. vulpes and Herbivore-derived mite: C. ibex and R. Rupicapra. a: M. Foina, b: M. martes, c: C. ibex, d: O. aries musimon and e: C. elaphus.

**Fig. 4** Unrooted Dps consensus dendrogram for individual Sarcoptes mites from the 15 wild host derived populations (Table 1) using a similarity matrix based on the proportion of shared alleles. Thick branches for all carnivore-derived mites, thin branches for all herbivore-derived mites, grey branches for S. scrofa-derived mites. Cluster I (SpNWRp, SpSWCp and SpSECp), Clusters IIa and IIb (ItNWSs and FrNESs), Cluster III (ItNEMm, ItNWMf, ItNWVv and ItNEVv) and Cluster IV (ItNERr, ItNECi, ItNECe, and ItNEOam). a: SpNEVv, b: SpEMf and arrow: ItNECi.

Fig. 1

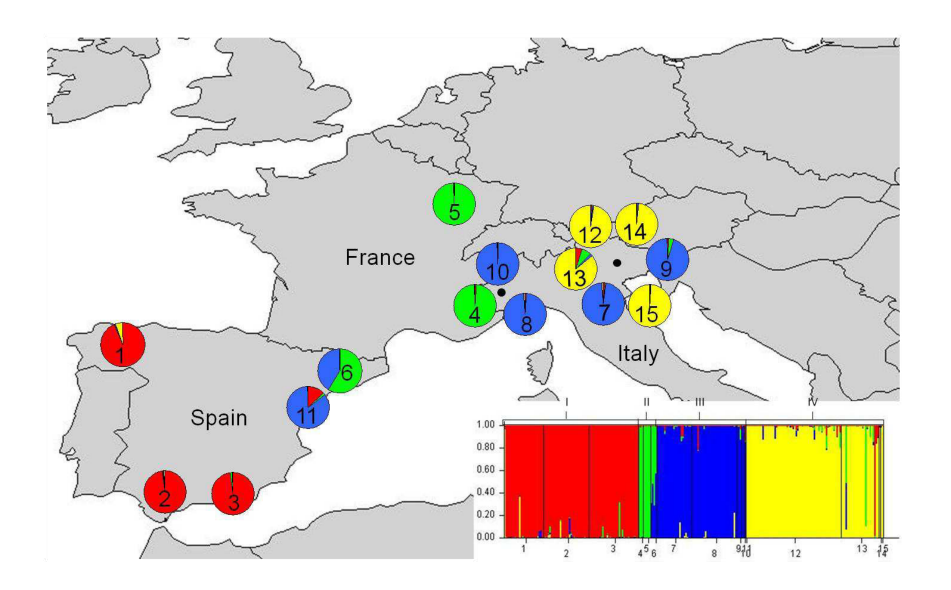


Fig. 2

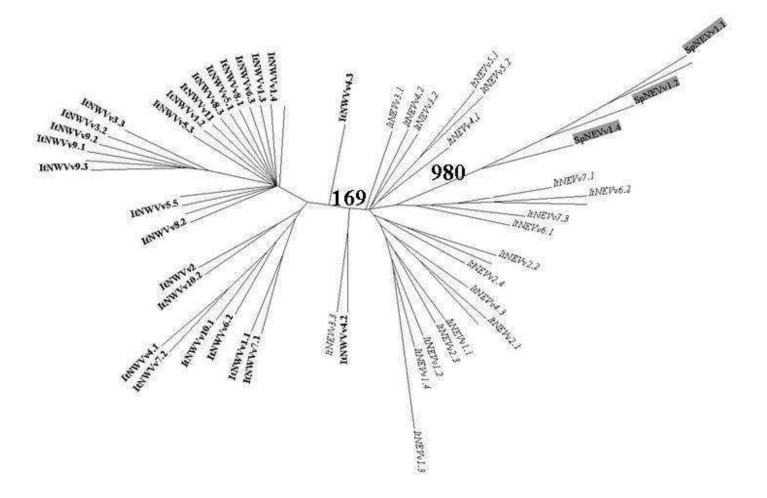


Fig. 3

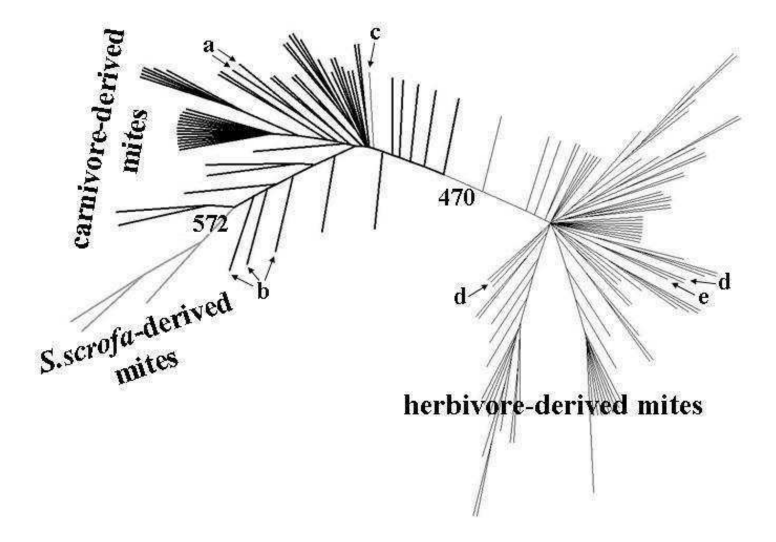


Fig. 4

