The Spread of Malaria to Southern Europe in Antiquity: New Approaches to Old Problems

ROBERT SALLARES, ABIGAIL BOUWMAN
and CECILIA ANDERUNG*

Introduction

The discoveries in the late nineteenth century that malaria is caused by protozoan parasites, which are transmitted by mosquitoes, quickly led to intense speculation about its history in antiquity. The historiography of malaria has passed through three distinct phases during the last hundred years or so. The first generation of historians to consider the effects of malaria did exaggerate its significance in some respects. The argument by W H S Jones that the Greek doctrine of fevers was based on malaria was generally and rightly accepted. However, it is not surprising that his view that malaria was a major reason for the degeneration of the moral character of the ancient Greeks attracted little sympathy.1 The eradication of malaria from southern Europe in the 1930s and 1940s contributed to a decline of interest in the subject. Subsequently medical historians and even professional malariologists tended to minimize the historical significance of malaria.2 The revisionist tendencies of this second phase of research led to attempts to reassess some of the details of the evidence upon which Jones had relied. For example, Leonard Bruce-Chwatt and Julian de Zulueta rejected Jones’s belief that Plasmodium falciparum, the most dangerous of the four species of human malaria, was already active in Greece in the fifth century BC. They suggested that it started to spread in southern Europe only during the time of the Roman Empire and attributed all the references to intermittent tertian fevers in Hippocratic texts dating to the fifth and fourth centuries BC to the less virulent P. vivax.3 Although the literature produced during this second phase of scholarship was in many ways more sophisticated, it still suffered from some of the same weaknesses; in particular, analysis proceeded in a purely qualitative manner, without any consideration of the effects of malaria on historical human populations in quantitative terms. A second weakness was a tendency to make generalizations covering the whole of Mediterranean Europe. Since many types of mosquito are

---

* Robert Sallares, MA; PhD, Dept of Biomolecular Sciences, UMIST, PO Box 88, Manchester, M60 1QD, United Kingdom (e-mail: rsallares@aol.com); Abigail Bouwman, MSc; PhD, Dept of Biomolecular Sciences, UMIST; Cecilia Anderung, MSc; Dept of Evolution, Genomics and Systematics, EBC, Uppsala University, Sweden.

We wish to thank David Soren for providing the photos of Lugnano and the bones.


incapable of transmitting malaria to humans, mosquito breeding sites do not occur everywhere, and many mosquitoes do not fly further than a few hundred yards from their breeding sites, malaria can only be really understood by micro-analyses, conducted at a very local level, of geography, hydrology, climate, competition between different species of mosquito for breeding sites, and human activities.

Some recent studies, opening up a third wave of research, have focused on the effects of malaria on historical European populations in quantitative terms. These studies have conclusively demonstrated that malaria did have severe effects in terms of both morbidity and mortality on human populations in early modern Europe, but only within strictly circumscribed geographical areas. Malaria did not occur everywhere. Consequently, it is now possible to move towards a modern synthesis which affirms by analogy (given the scarcity of quantifiable demographic data from the ancient world) that malaria did have considerable effects in Mediterranean Europe in antiquity—killing or debilitating people to the extent that it altered the age structure of human populations, as well as changing human settlement patterns and influencing the nature of agricultural systems—but usually only at a localized level. The effects of malaria on human populations and societies in Mediterranean Europe during the time of the Roman Empire are now well understood. However, the debate about the time of introduction of *P. falciparum* to southern Europe and indeed the date of its origin as a specifically human parasite are still very much matters of controversy. The main purpose of this article is to consider some new approaches, some new evidence, and present a new hypothesis about the early history of malaria in southern Europe.

### Research in Molecular Evolution

Comparative analysis of the genomes or complete DNA sequences of pathogenic microorganisms has introduced a new way of studying the evolution of human pathogens. Comparisons of 18S ribosomal DNA gene sequences suggest that the divergence of *P. falciparum* from its only known close relative *P. reichenowi*, a parasite of chimpanzees, ran parallel to the divergence of hominids from chimpanzees, approximately five to ten million years ago. This is an example of parasite:host co-speciation. In contrast, the other three species of human malaria (*P. vivax*, *P. malariae*, *P. ovale*) were lateral transfers from other primates to humans. However, there is intense controversy about the more recent molecular evolution of *P. falciparum*. One controversial hypothesis claims that all currently existing *P. falciparum* populations are descended from a most recent common ancestor (MRCA) which lived only a few thousand years ago during the Neolithic period.

---


6. *P. ovale* is not considered any further in this paper since it was never common in Mediterranean countries.

The Spread of Malaria to Southern Europe in Antiquity

scientists insist that *P. falciparum* has maintained a large effective population size with a high degree of genetic diversity for several hundred thousand years. Such divergent conclusions are possible because different scientists use different datasets and different methods for analysing the molecular data, which are in any case often difficult to date.

The most recent major study has synthesized these two polar views, concluding that the MRCA of *P. falciparum* is approaching 100,000 years old, although it has undergone a major population expansion within the last 10,000 years. Consequently the view adopted here is that *P. falciparum* is indeed a very ancient human parasite. Mario Coluzzi has cogently argued that its evolution as a human parasite is intricately bound up with the evolution of its most important and efficient vector in tropical Africa today, the mosquito species complex *Anopheles gambiae*. Since the forest ecotype is the oldest ecotype of *A. gambiae*, highly anthropophilic behaviour probably evolved in a closed forest environment in which the mosquitoes had few other large animals apart from humans to bite, rather than on open grasslands with a large variety of alternative prey. This is a highly plausible hypothesis, but the date at which humans entered the African rain forests then becomes a critical question. The current trend in research on human evolution is to stress that early hominids lived in environments which included some forests, not in completely open grasslands, and it is now known that early hominids were present in West Africa as well as East Africa as early as six million years ago. Moreover, recent archaeological research indicates that human hunter-gatherer populations have been continuously present in rain forest environments for a very long time. As a result, the association in tropical Africa between the forest ecotype of *A. gambiae* and *Homo sapiens* need not be as recent as the Holocene period, but could easily be much older and involve earlier species of hominids. Consequently, the hypothesis about the role of *A. gambiae* does not necessarily contradict the results of the molecular studies of malarial DNA which suggest that *P. falciparum* is a very ancient organism. The Holocene population expansion of *P. falciparum* in Africa would have been facilitated by the evolution of other ecotypes of *A. gambiae* from its original forest ecotype. The MRCA approximately 100,000 years ago is probably associated with the expansion of early modern humans. It was part of a malaria population which preyed on the human population to which “mitochondrial Eve”, the ancestor of all currently existing human mitochondrial genotypes, belonged.

---


Ancient History

In the light of the uncertainties generated by current scientific research on the molecular evolution of malaria, it is still well worth considering the alternative way of exploring the early history of human malaria, namely working from the present as far back as possible using the evidence of historical documentary sources and archaeological materials. It is very likely that all species of human malaria (temperature-dependent diseases) and their vector mosquitoes were absent from Europe during the Quaternary Ice Ages. Nevertheless, there are plenty of references to the intermittent fevers characteristic of malaria in ancient literature such as the works of the Hippocratic Corpus in Greece in the fourth and fifth centuries BC, in ancient Indian texts (difficult to date), and in Chinese literature of the first millennium BC. These accounts leave no doubt whatsoever that benign tertian fever (caused by *P. vivax*) and quartan fever (caused by *P. malariae*) were endemic in the Old World from Greece to as far as India and China by 500 BC. Evidently these types of human malaria arrived in Greece some time between the end of the last Ice Age and 500 BC.

The history of *P. falciparum* (responsible for malignant tertian fever) is even more problematic, as has already been noted. It is generally agreed that it evolved in tropical Africa. Nevertheless, the earliest direct evidence for its history comes from Europe. The Roman author Celsus, writing during the reign of the emperor Tiberius in the first half of the first century AD, differentiated the clinical symptoms of semiterminal fever from tertian fever in an account whose accuracy as a description of the differences between falciparum and vivax malaria has been recognized since the time of Étto Marchiafava and Angelo Celli, the first scientists to identify falciparum parasites under the microscope, in 1889. There are two types of tertian fever. The first type begins and finishes in the same manner as quartan fever. Its characteristic feature is that it disappears for one complete day and returns on the third day. The second type is much more dangerous. It too recurs on the third day. However, out of the forty-eight hours, the paroxysm lasts for almost thirty-six hours (sometimes more or less), and there is no total cessation during the remission, although it becomes less severe. Most doctors call this type of fever semiterminal.

Marchiafava described the text of Celsus as follows: “the clearest and most exact description of the two clinical types of tertian fevers are to be found in the works of Celsus”. There are two important points to note here. Firstly, Celsus regarded all types of intermittent malaria is traced by mitochondrial DNA’, *Mol. Biochem. Parasitol.*, 2000, **111**: 163–71.

13 Celsus, *De medicina*, III.3.2: “*tertianarum vero duo genera sunt. Alterum eodem modo, quo quartana, et incipienti et desinens, illo tantum interposito discriminate, quod unum diem praestat integrum, tertio redit. Alterum longe perniciosius, quod tertio quidem die revertitur, ex quadraginta et octo horis fere triginta et sex per accessionem occupat (interdum etiam vel minus vel plus), neque ex toto in remissione desistit, sed tantum levius est. Id genus plerique medicici ignotum spectatorem appellant.*”

fevers, including what he called semitermian fever, as very widespread and common in his own time.\(^\text{17}\) Falciparum malaria was not a rare disease in the Roman world in the early first century AD. Secondly, although there was explicit discussion in antiquity of the possibility of new diseases arising, there is no indication whatsoever that Celsus (or any other ancient author) regarded falciparum malaria (or any other type of malaria) as a new disease.\(^\text{18}\) On the contrary, by using the Greek form of the word “semitermian”, he recalled the earlier Hippocratic texts which also mention this type of fever. It was explicitly observed by another Roman author, Quintus Serenus, that the Romans had not coined a Latin name of their own for the disease, borrowing the Greek expression instead.\(^\text{19}\) This suggests that the disease was known in Greece before it became endemic in central Italy. The semitermian fevers described by Celsus are indeed mentioned in the Hippocratic Epistles, for example in the second “constitution” (description of a particular set of atmospheric conditions thought to be responsible for disease) on the island of Thasos in northern Greece. The account of the first “constitution” at Thasos offers a one sentence definition of semitermian fever which is perfectly compatible with the statements of Celsus.\(^\text{20}\) We are also given the important detail that patients suffering from semitermian fevers in the autumn frequently ended up in a coma.\(^\text{21}\) Cerebral malaria is a major syndrome of falciparum malaria, which is not caused by the other species of human malaria. This syndrome was also noticed later in antiquity, for instance by Archigenes of Apamea c. 100 AD, as cited by Aetius of Amida.\(^\text{22}\) The uneven distribution of extant classical Greek literature, which mostly originates from Athens (never a hotspot for malaria), creates various problems, which cannot be discussed in detail here, regarding the prevalence of malaria across classical Greece as a whole. Nevertheless, it is clear that semitermian fevers, like other intermittent fevers, were common in at least some localities in northern Greece in the fifth century BC.\(^\text{23}\) The second “constitution”

---

\(^{\text{17}}\) Celsus, De medicina, III, 3.1.

\(^{\text{18}}\) Plutarch, Moralia, VIII, 9.731b–734c, and Pliny, Historia naturalis, XXVI, 1–9, both discussed leprosy as a new disease in Hellenistic and Roman times.


\(^{\text{21}}\) Hippocrates, op. cit., note 13 above, pp. 624–5: ἄρρητος τὸ ὀψιλος καὶ μάλαιστα ὁστος καὶ πλαίν κωμάτωθες. Julian de Zulueta’s argument (mentioned by Corbellini and Merzagora, op. cit., note 16 above, p. 22), that sufferers from semitermian fever died too slowly for falciparum malaria to have been the cause, does not take into account the likelihood, discussed below in this article, that by the fifth century BC at least some populations in northern Greece already had a high frequency of human genetic mutations conferring some resistance to malaria. M Gmek, Diseases in the ancient Greek world, Baltimore, Johns Hopkins University Press, 1989, p. 281, accepted the identi- fication of falciparum malaria in the Hippocratic texts.

\(^{\text{22}}\) Aetius of Amida, Libri medicinales, VI, 3.

\(^{\text{23}}\) For the problem of whether Macedonia was severely affected by malaria during the time of its rise to power under Philip and Alexander the Great in the fourth century BC, see E Borza ‘Some observations on malaria and the ecology of central Macedonia in antiquity’, Am. J. Ancient Hist., 1979, 4: 102–24.
Robert Sallares, Abigail Bouwman and Cecilia Anderung

states that many people on Thasos were affected by semitertian fevers.24 The details about semitertian fever given in the Hippocratic texts are congruent both with Celsus’ description and with modern knowledge of falciparum malaria. There is no evidence that the Greek word changed its meaning between the fifth century BC and the first century AD. Consequently, there is no good reason for not accepting that falciparum malaria was well known and common in at least some parts of Greece by the fifth century BC. The Hippocratic texts simply set a *terminus ante quem* for its appearance, since there is a scarcity of relevant earlier literature.25 Indeed, as will be seen later, there is another, completely different, type of argument which suggests that it was active in Greece well before the fifth century BC.

However, a rather different picture emerges for Italy. One can discern three stages in the northwards spread of malaria in Italy (Figure 1). Firstly, literary sources suggest that it was already active in Sicily by the fifth century BC, when it was probably responsible for the epidemic which affected the Athenian army outside Syracuse during the Peloponnesian War, as Mirko Grmek once argued.26 Recent research on the physical environment of the territory of Syracuse has concluded that the area of marshland near the city was much larger in antiquity than it was in early modern times.27 This strengthens Grmek’s arguments by showing that the natural environment around Syracuse was indeed highly favourable for malaria 2500 years ago. Similarly, numerous Greek colonies were established along the coasts of southern Italy c. 800–600 BC in regions that later suffered from intense malaria. Paestum in Campania, where famous Greek temples were constructed in the sixth and fifth centuries BC, is one well known example. Later on the city declined because of neighbouring marshes which became unhealthy, as Strabo records in the first century BC.28

The spread of malaria in western central Italy (the second stage of its spread in that country) seems to have been mainly a phenomenon of the period c. 400–100 BC. In the archaic and early classical periods, up to c. 400 BC, Etruscan cities such as Vulci and Vetulonia flourished in the Maremma, the coastal region of Tuscany, which was described as pestential by Pliny the Younger in the first century AD.29 Vulci occupies a lowland site in the Maremma in an area where there was intense malaria in the early modern period. The archaeological remains, particularly the large numbers of imported Greek vases, testify to prosperity in the archaic period, while the famous wall paintings of the François tomb record contacts with Rome. It has even been suggested that the place name “Vulci” signified “prosperous place” in the ancient Etruscan language.30 It is difficult to believe that there

---

24 Hippocrates, op. cit., note 13 above, pp. 618–9; ἡμιτραπέσια, τραπέσια, ἄκριβες, τεταρταίοι, πλανῆτες, ἄκριτοι de τῶν ὑπογεγραμμένων πυρετῶν πολλάίσιν ἐγένοτο.
25 This is a better approach to the problem than assuming, as W H S Jones (op. cit., note 1 above, p. 76) did, that the silence of very scarce earlier sources proves that malaria was unknown in Attica until c. 430 BC.
28 Strabo, *Geographia*, V.4.13.251C.
Figure 1: Map of Italy, adapted from Luigi Torelli’s map of the distribution of malaria in Italy published in 1882. The arrows show the spread of the disease in the past, with very approximate dates (TAQs), following its vector mosquitoes.
was endemic malaria there during the city’s period of prosperity, but it is barely mentioned in literary sources dating to the first and second centuries BC, when a significant quantity of written evidence first becomes available from Rome. Consequently it is likely that malaria, and by implication its mosquito vectors, spread widely along the coast of Tuscany in the third and second centuries BC. The earliest record of endemic malaria at a specific location in this region by a contemporary author, Cato the Elder, relates to Graviscae, the port of the Etruscan city of Tarquinia, c.150 BC, but this mention, of course, only provides a \textit{terminus ante quem} for its arrival.\footnote{31} Both the malaria of Graviscae and the mosquitoes that transmitted it probably arrived by sea, either from Sardinia or Sicily or directly from North Africa. In the late second century BC Asclepiades of Bithynia described severe intermittent fevers with the cerebral symptoms frequently found in falciparum malaria as common in the city of Rome itself, and Galen repeated this opinion in the second century AD.\footnote{32} There is no doubt that the countryside around Rome had become a zone where malaria was endemic by the time of the Roman Empire. Mike McCormick has recently demonstrated by a study of papal correspondence that the seasonal pattern of travel that is well known for the early modern period, in which people from northern Europe tended to go to Rome in winter to avoid the “bad air” of summer, was already becoming a custom as early as the eighth century AD.\footnote{33}

The third and final stage in the spread of malaria in Italy was its advance to northern Italy. Vitruvius and Strabo state that the Po delta region, around Ravenna in northeastern Italy, was healthy about 2,000 years ago.\footnote{34} This was evidently regarded as an unusual state of affairs for a marshy region. It cannot be attributed to any failure of malaria itself to reach the area, since the constant movements of merchants, administrators, soldiers and slaves during the Roman Empire would have spread malarial parasites within their bodies all over Europe. Nor was there a shortage of mosquitoes, since Roman authors such as Sidonius Apollinaris explicitly mention the mosquitoes of Ravenna.\footnote{35} However, only certain anthropophilic species of mosquito are effective vectors of malaria. The most likely inference is that the mosquitoes of Ravenna in antiquity were not efficient vectors of the disease. Endemic malaria became possible in this region only during the medieval period, when a new anthropophilic species of mosquito, \textit{Anopheles sacharovi}, colonized the coasts of northeastern Italy. The spread of malaria in Italy did not happen suddenly. It took at least 1500 years, from \textit{c}.500 BC to \textit{c}.1000 AD. The hypothesis that this process depended on the spread of certain types of mosquito is highly plausible, just as speciation in the \textit{A. gambiae} complex played an important role in the spread of malaria in tropical Africa, as Mario Coluzzi has argued.\footnote{36} It has already been noted that Bruce-Chwatt and de Zulueta believed that \textit{P. vivax} and \textit{P. malariae} were present in Mediterranean Europe before

The Spread of Malaria to Southern Europe in Antiquity

*P. falciparum*. However, all three of these species of human malaria seem to have been present together, according to the earliest available literary evidence for both Greece and Italy. Logically, the prior presence of *P. vivax* and *P. malariae* would have required the availability as vectors anyway of suitable anopheline mosquito species, which could equally well have been transmitting *P. falciparum* at the same time. All three were ready to move together simultaneously into new regions, but had to wait for their mosquito vectors to move first.

The Infant Cemetery at Lugnano in Teverina, Umbria, Italy

So far only the evidence of historical documentary sources has been considered. However, scientific evidence for the ancient spread of malaria is now starting to become available through research on ancient biomolecules. Between 1988 and 1992 David Soren and his colleagues (from the University of Arizona) excavated a fascinating archaeological site at Lugnano in Teverina, about seventy miles up the Tiber valley from Rome. One of the largest infant cemeteries found in Roman Italy, dating to c.450 AD, was unearthed in the ruins of an abandoned Roman villa (see Figure 1 for the location). This site has yielded important new evidence for the spread of malaria.

Infants rarely received proper burial in Roman times. Several features of the finds led the archaeologists to conclude that the infant deaths had been caused by a disease epidemic. All the burials occurred in a short period of time within one year, since joins between potsherds from different levels of the fill indicate that the cemetery represents a single archaeological stratum. The spatial distribution of the burials within the cemetery suggests the increasing activity of an epidemic during that period of time. Plant remains indicate that the time of year was the summer, the season when malaria (a temperature-dependent disease) was active in Italy in the past. The number of burials is larger than the number of young children in the adjacent town of Lugnano today. This gives an indication of the size of the adult female breeding population that gave birth to the infants c.450 AD. A disease is required that was capable of sweeping throughout the entire population and infecting virtually all the pregnant women at the same time. Moreover 22 out of the 47 excavated infants were premature births, while most of the others were neonates. Consequently, a

---

37 On the question of the reactivity of modern tropical strains of *P. falciparum* to European mosquitoes, see Sallares, op. cit., note 5 above, pp. 33–6. Refractoriness was probably overcome in North Africa and the Near East in prehistory before *P. falciparum* had even reached southern Europe.


40 Sallares, op. cit., note 5 above, pp. 61–3, on the seasonality of malaria in Italy.

41 Unfortunately no corresponding adult cemetery has been discovered.
Robert Sallaress, Abigail Bouwman and Cecilia Anderung

disease is required which produces not only high death rates but also a high rate of miscarriages in pregnant women.

Falciparum malaria fits the bill. It causes intrauterine growth retardation, which is associated with neonatal mortality, in immune or semi-immune pregnant women in tropical countries where malaria is holoendemic today. 42 However in the past it also produced a high rate of miscarriages in non-immune pregnant women in temperate or subtropical regions where it had an epidemic rather than an endemic presence. 43 For example, a series of cases in the United States in the first half of the twentieth century yielded a foetus death rate of 60 per cent, while there were also numerous cases of miscarriages during the Monte Cassino epidemic in 1944, the last major malaria epidemic in modern Italian history, and similarly at Amritsar in India in the early twentieth century. 44 There is evidence that maternal malaria increases susceptibility to malaria in infants in the first year of life. Maternal malaria in the last week of pregnancy is particularly associated with neonatal mortality. 45

The archaeological site is situated 185 metres above sea level in gently undulating countryside about 3.5 km away from the river Tiber, which is visible from the site. River valleys were a prime location for malaria in Italy because river floods frequently created mosquito breeding sites when the floodwaters retreated leaving behind small pools. The vicinity of Orte, on the Tiber a few kilometres from Lugnano, was particularly vulnerable to flooding in the past. A local doctor, Angelo Sorgoni, described the pernicious symptoms of intermittent fevers among itinerant agricultural labourers around Narni, near Lugnano, in 1832. 46 This confirms that the local environment of Lugnano is indeed highly favourable to malaria. Relying on these circumstantial arguments, the archaeologists proposed the hypothesis that the deaths of the Lugnano infants were caused by an epidemic of falciparum malaria during the summer of one year in the middle of the fifth century AD. 47

43 A Barbosa and B López Arjona, El paludismo en el primer año de la vida, Madrid, Sáez Hermanos, 1935, pp. 11–18.

320
The Spread of Malaria to Southern Europe in Antiquity

Direct Evidence for Falciparum Malaria

This hypothesis was tested by attempting to extract *P. falciparum* DNA from bone samples from the infant skeletons. A fragment of *P. falciparum* ribosomal DNA was successfully amplified from two separate DNA extractions from the skeleton of burial no. 36, the oldest child found on the site (a female 2–3 years old at death).\(^48\) The fact that malarial DNA could be amplified by the polymerase chain reaction from such an old skeleton suggests a massive infection, since it is unlikely in practice that an asymptomatic infection with a low level of parasitaemia could be detected after such a long passage of time. Consequently it is likely that falciparum malaria did indeed cause the death of this particular child. The results prove that falciparum malaria was present and tend to confirm the archaeologists’ hypothesis that it was the cause of the Lugnano epidemic.\(^49\) Although positive results were obtained from only one out of forty skeletons that were sampled, it should be noted that it is not in the nature of the epidemiology of falciparum malaria for it to occur sporadically. If environmental conditions were favourable for it to have been present at all, many people would have been infected. The low success rate can be safely attributed to poor DNA preservation in the bones.\(^50\) The significance of the Lugnano epidemic is that it shows that falciparum malaria and by inference its principal vector in central Italy, *Anopheles labranchiae*, had attained their maximum geographical distributions in that region by the middle of the fifth century AD. *A. labranchiae* occurred as far up the Tiber valley as Orte, near Lugnano, as recently as the Second World War. Further north it was replaced by *A. atroparvus*, a secondary vector.\(^51\)

Documentary evidence shows that in late antiquity malaria was frequently conceptualized as a demon. In fact the term *abracadabra*, used in magic, originated as a spell against semitertian fever recorded by the Roman author Quintus Serenus.\(^52\) There is a remarkable synthesis of modern science and ancient pagan rituals in the archaeological finds from Lugnano, since the feet and hands of the infant who has yielded some malarial DNA were weighted down with stones and tiles, presumably to prevent the demons of malaria from escaping and causing further devastation (Figure 2). The magical rituals attested at Lugnano

---

\(^{48}\) For details of the methods used, see R Sallares and S Gomzi, ‘Biomolecular archaeology of malaria’, *Ancient Biomolecules*, 2001, 3: 195–213 (with EMBL/Genbank accession nos. AJ426487–8 for the malaria sequences that were obtained).

\(^{49}\) Determination of the sex of a sample of these infants (see also next footnote) indicated that the infant population had a fairly normal sex ratio (five female and four male). Consequently there is no question of selective exposure of female infants in this cemetery.


\(^{51}\) Sallares, op. cit., note 5 above, pp. 64–8.

\(^{52}\) Quintus Serenus, *Liber medicinalis*, 51.935–940, op. cit., note 19 above.
provide archaeological evidence for the survival of pagan traditions connected to medicine in the countryside of central Italy in the fifth century AD despite the advance of Christianity.\textsuperscript{53}

**Indirect Evidence for the Spread of Malaria**

The development of molecular population genetics over the last twenty years or so now provides a new, indirect, method of tracking the effects of malaria in the ancient Mediterranean world that was not available to previous generations of historians, by searching for evidence of human genetic mutations associated with resistance to malaria. The lack of haplotype diversity associated with most of the malaria resistance mutations found in modern Mediterranean populations suggests that they have evolved within the last few

\textsuperscript{53} Literary evidence for the interaction of magic and religion in Umbria at this time has been discussed by G Bartocci, ‘The cultural construction of the Western conception of the realm of the sacred: co-existence, clash and interbreeding of magic and sacred thinking in fifth- and sixth-century Umbria’, *Anthropology and Medicine*, 2000, 7: 373–88.
Figure 3: An example of an amphora burial from the infant cemetery at Lugnano in Teverina.

This timescale is plausible, since it was too cold for malaria in Europe during the glacial periods. The most ancient osteological evidence for thalassaemia comes from the Near East, from a skeleton excavated at the now submerged village of Atlit Yam off the coast of Israel, dating to about 10,000 years ago. Various different sorts of evidence indicate that all the major types of inherited genetic resistance were present in the Mediterranean world by the time of the Roman Empire, about 2000 years ago. For example, the skeleton of a child with osteological modifications attributed to thalassaemia was excavated at Pisa in Italy in strata dated to the fifth to seventh centuries AD. A skeleton of a woman approximately seventeen years old, with morphological changes attributed to thalassaemia, was excavated at the Roman villa of Settefinestre, on the coast of Tuscany, dating to the third century AD. Similar evidence occurs earlier in the Greek world of Magna Graecia in southern Italy. Eleven skeletons with the porotic hyperostosis characteristic


of thalassaemia (presumably all homozygotes) were excavated in the Pantanello necropolis at the Greek colony of Metapontum, dating to the fourth and fifth centuries BC.\(^{58}\) This suggests that the Greek colonists had a significant population frequency of thalassaemia mutations and by inference a population history of contact with endemic falciparum malaria before the inhabitants of Latium and Etruria in western central Italy, around Rome, came into contact with it. Grmek examined seven terracotta statuettes from Smyrna in Asia Minor, dating to the Hellenistic period, which have "swollen faces with hypertrophied zygomatic arches and symmetrical protuberances on the fronto-parietal parts of the skull", with the base of the nose crushed. He concluded that these figurines are classic illustrations of homozygous thalassaemia.\(^{59}\)

The sickle-cell trait was also present by Roman times. The skeleton of an approximately twenty-year-old man with porotic hyperostosis was excavated on the island of Failaka in the Persian Gulf. The bones were radiocarbon dated to 2130 ± 80 BP. Scanning-electron microscopy of the bones directly revealed fossilized erythrocytes with the characteristic sickle shape.\(^{60}\) Research in experimental archaeology has confirmed that it is possible for human red blood cells to leave a recognizable print on archaeological materials.\(^{61}\) Today haemoglobin S occurs with a frequency of about 2 per cent in some Mediterranean populations. Haplotype analysis has demonstrated that the sickle-cell trait in Sicily, northern Greece, and western Arabia is in linkage disequilibrium with the Benin haplotype in western central Africa.\(^{62}\) This constitutes direct evidence for gene flow linked to human migration from central Africa to Mediterranean Europe in historical times. The Failaka individual might have been a descendant of a Macedonian soldier from northern Greece with the Benin haplotype. Alternatively he might have been one of the local inhabitants, since there is molecular evidence for an independent origin of the sickle-cell trait in India and eastern Arabia, with a different haplotype association. The Indian variant of homozygous sickle-cell disease displays a mild clinical phenotype, associated with high levels of foetal haemoglobin which inhibits sickling.\(^{63}\) This could explain how the Failaka individual survived to adulthood.

To complete this picture some biomolecular evidence is now available for glucose-6-phosphate dehydrogenase (G6PD) deficiency, the third major type of human genetic mutation commonly occurring in Mediterranean populations which confers some resistance to malaria. G6PD deficiency is caused by any one of about 140 different mutations or combinations of mutations that affect the functioning of a gene that plays a critical role in a biochemical pathway for removing oxidants that would otherwise damage erythrocytes.\(^{64}\)


\(^{64}\) A list of G6PD mutations is given by E Beutler and T J Vollmy, ‘Haematologically important
The Spread of Malaria to Southern Europe in Antiquity

Such damage shortens the life span of red blood cells, causing premature phagocytosis or apoptosis, before malarial parasites inside the cell have completed their development. The large number of mutations, differential inactivation arising from the gene’s location on the X chromosome, and other unknown factors, produce a wide range of expressions of G6PD deficiency in homozygous and heterozygous females and hemizygous males.

The Lugnano infants were screened for the presence of the two commonest G6PD mutations in Mediterranean populations. None of those skeletons from which human DNA could be extracted yielded any evidence for the 844 G→C mutation in exon 8 of the G6PD gene (the asymptomatic Seattle variant), but one infant (burial no. 22) tested positive for the 563 C→T mutation in exon 6 (G6PD Med variant). This mutation, the most frequent variant in Mediterranean populations, accounts for about 61 per cent of cases of G6PD deficiency in mainland Italy. It is associated with severe deficiency, potentially fatal neonatal jaundice in infants, and favism in older children. The disease favism was not formally described in medical literature until 1856. However, the taboo on eating broad beans (ingestion of which precipitates favism in some G6PD deficient individuals) among the followers of Pythagoras in antiquity is probably an ancient reference to it, as Grmek argued, even if Elinor Lieber’s suggestion that “the Pythagorean community was primarily designed for the benefit of persons handicapped by favism” is rather speculative. The sex of the infant at Lugnano was determined as male by the amelogenin sex determination system. It is uncertain whether the G6PD Med variant actually provides any protection against malaria to male hemizygotes (as well as female heterozygotes). These experimental results confirm that G6PD deficiency, like thalassaemia and the sickle-cell trait, was already spreading in Mediterranean Europe during the time of the Roman Empire, presumably as a human evolutionary response to the spread of malaria.

It is generally thought that each individual mutation for traits such as thalassaemia and G6PD deficiency evolved only once and was then spread by human migration at a regional scale. The extraction of DNA from human bones from 18th to mid-19th century London, has revealed the G6PD Med variant, and other genetic markers, in ancient populations throughout Europe and the Mediterranean. The spread of these mutations, and the diseases they cause, may have had significant impacts on human health and evolution in the region.

---


---


level, since human populations in each part of the world where malaria occurs or occurred in the past have their own distinctive mutations which do not occur elsewhere, and most of the mutations exist in the context of restricted haplotype diversity. For example, linkage disequilibrium with a silent mutation at nucleotide 1311 in exon 11 of the G6PD gene suggests that the G6PD Med variant probably evolved only once in the Mediterranean region and was then spread by human migrations around the Mediterranean from a single point of origin, although it did have a second quite separate point of origin in India lacking the association with the exon 11 mutation.\(^{71}\) One of the two most frequent β-thalassaemia mutations in Mediterranean populations today, the B + IVS nt 110 mutation (G → A), which occurs in areas of Greek colonization in Italy, attains its highest frequencies today in eastern Mediterranean countries. It is very common in Greece, and is also associated with a high level of haplotype diversity in Turkey.\(^{72}\) Consequently, Italian scientists suggested that this particular mutation originated in Greece or Anatolia and was spread westwards to Italy by Greek colonization from the eighth century BC onwards.\(^{73}\) This hypothesis implies that falciparum malaria became an important agent of natural selection on human populations in the eastern Mediterranean before it became important in Italy. (If, for the sake of argument, the Greek colonists in southern Italy had not experienced falciparum malaria in their homeland of Greece before migrating, they should have evolved their own distinctive mutations afterwards when they encountered it later on in Italy, on the principle that each region evolves its own mutations.) This hypothesis is also congruent with the ancient literary evidence discussed earlier which suggests that the classical Greeks were familiar with falciparum malaria before the people of central Italy had experienced its effects.

The second common β-thalassaemia mutation in Mediterranean populations today, the CD39 mutation, has a quite different geographical distribution, concentrated in areas of Phoenician colonization in the western Mediterranean. Although it has been suggested that it evolved in the Levant and was spread from there to the western Mediterranean by the Phoenicians,\(^{74}\) its present distribution is almost entirely western. It achieves its highest level of associated haplotype diversity by genetic recombination in North Africa.\(^{75}\) Consequently it now seems more probable that the CD39 mutation actually evolved in North Africa, rather than the Levant, and was then spread northwards to Sardinia, Sicily and Spain by Phoenician colonization from North Africa during the first millennium BC. North Africa is the main centre of abundance today and the presumed main Ice Age refuge of the mosquito species Anopheles labranchiae, the most important vector of malaria in Italy in recent times.


72\) Tadmouri, *et al.*, op. cit., note 54 above.

73\) A Cao, M Gossens, and M Pirastu, ‘β-thalassaemia mutations in Mediterranean populations’, *Br. J. Haematology*, 1989, 71: 309–12. Earlier discussions by medical historians and palaeopathologists (e.g. Grmek, op. cit., note 21 above, pp. 254–64) did not employ the evidence of DNA sequence data, which was not yet available.

74\) Cao, *et al.*, op. cit., note 73 above.

The Spread of Malaria to Southern Europe in Antiquity

Adding up all this evidence, a picture emerges of two quite separate routes whereby malaria itself, its vector mosquitoes, and the associated human genetic mutations which give some degree of resistance to malaria, reached southern Europe in the past. One route passed from Tunisia in North Africa, via Sicily and Sardinia, to Italy. The second route commenced in the eastern Mediterranean and led from the Levant or Anatolia to Greece and then on to Italy. Previous scholarly literature generally speaks of malaria in Italy and in Greece in the same breath, without considering the possibility that malaria had different histories in these two countries. However, we have just seen that there is some evidence that falciparum malaria became an important agent of natural selection on human populations in the eastern Mediterranean before it became an important agent in mainland Italy.

Role of Different Species of Mosquito

Such a situation can be explained if it is remembered that the most important vectors of malaria in the western Mediterranean and the eastern Mediterranean were two different species of mosquito, *A. labranchiae* and *A. sacharovi* respectively, both members of the *A. maculipennis* complex. Not only are they different species, but they also have different geographical distributions and origins. *A. labranchiae*, the most important vector in central and southern Italy, originated in North Africa. It is a non-diapausing species of mosquito which remains active throughout the year and indeed breeds all the year round in North Africa, but cannot do so in mainland Italy. It had to stay inside houses or man-made animal shelters to survive the winter in southern Europe. In contrast *A. sacharovi*, the most important vector of malaria in Greece, tends to hibernate, a useful habit for surviving southern European winters. The Near East, the centre of its geographical distribution today, was presumably its refuge during the glacial periods.

The hypothesis is proposed here that *A. sacharovi* spread from the Near East to Greece (together with other Near Eastern vectors such as *A. superpictus*) a considerable time before *A. labranchiae* spread from North Africa to mainland Italy. As a result, there was intense malaria in some parts of Greece and the Near East before there was intense malaria in Italy. Consequently some of the most important malaria resistance mutations in Mediterranean populations probably evolved in the eastern Mediterranean in prehistory and were then spread westwards to Italy by Greek migrations in the eighth century BC. At about the same time Phoenician activity in the western Mediterranean opened up the second route, from North Africa to Italy. Although there were humans on Sardinia as early as c. 20000 BP, the spread of falciparum malaria to Sardinia was probably a result of Phoenician colonization. There was a considerable Phoenician presence not only on the coast but in the interior of Sardinia in the seventh century BC, with archaeological evidence for trade with the Etruscans, followed by a Carthaginian occupation from North Africa. The two routes

---

interacted in Sicily, the scene of both Phoenician and Greek colonization, where both of the two major β-thalassaemia mutations attain high frequencies. However, despite the availability of these two routes of travel, these mutations did not actually reach very high frequencies in Latium and Tuscany in central Italy because these areas were not directly colonized by the Greeks or Phoenicians. Subsequently, the Benin haplotype of the sickle-cell trait travelled with migrants (slaves?) from central Africa, probably exploiting the two routes, since it occurs in some human populations in both the western and the eastern Mediterranean. This hypothesis integrates, firstly, the molecular evidence for the origin and distribution of human genetic mutations giving resistance to malaria; secondly, data on the behaviour, origin and distribution of the mosquito vectors; and, thirdly, the historical documentary evidence for the late spread of falciparum malaria in central Italy and its earlier presence in Greece.

The gradual spread of malaria in mainland Italy clearly occurred in historical times. The final step in this process did not happen until the medieval period, when endemic malaria emerged in the Po delta region of northeastern Italy, presumably as a result of the arrival of A. sacharovi, the dominant vector in recent times in that region. The spread of malaria to northeastern Italy occurred at a time when Ravenna and the emerging commercial centre of Venice were closely associated with the Byzantine Empire, as McCormick has pointed out. It is significant that it was the predominantly eastern Mediterranean A. sacharovi, rather than the western Mediterranean A. labranchiae, which became dominant in northeastern Italy.

The chronology of the spread of malaria in mainland Italy is reasonably clear. However, the chronology of its earlier spread in the eastern Mediterranean remains very uncertain. Before the texts of the Hippocratic Corpus begin to appear in the fifth century BC there is a lack of relevant literary sources. Furthermore, malaria itself does not cause any unambiguous morphological lesions on human bones, and no biomolecular evidence is currently available from Greece. Mirko Grmek and Danielle Gourevitch interpreted depictions of Heracles slaying the multi-headed serpent of Hydra on Corinthian pottery dating to the period c. 630–570 BC as a reference to attempts to drain the malarial marshes of Lerna in the Argolid. This is quite plausible, but once again it only sets a terminus ante quem. The fact that the Greeks who founded colonies in the south of Italy from the eighth century BC onwards carried with them the commonest eastern Mediterranean β-thalassaemia mutation indicates that falciparum malaria was having severe effects in at least some parts of Greece by then. Beyond that point, investigation of porotic hyperostosis on human skeletal remains from archaeological sites is probably still the most promising approach to malaria in the prehistory of Greece, as argued by J Lawrence Angel.

81 This is the explanation of the problem posed by Bruce-Chwatt and de Zulueta, op. cit., note 3 above, p. 15.
82 Sallares, op. cit., note 5 above, pp. 78–85.
85 J L Angel, ‘Porotic hyperostosis or osteoporosis symmetrica’, in D Brothwell and A Sandison (eds), Diseases in antiquity, Springfield, IL, C C Thomas, 1967, pp. 378–89, argued that there was malaria at Lerna in the Middle Bronze Age.