

1 **Large-scale comparative analyses of tick genomes elucidate their genetic**
2 **diversity and vector capacities**

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1 **Summary**

2 Among arthropod vectors, ticks transmit the most diverse human and animal
3 pathogens, leading to an increasing number of new challenges worldwide. Here, we
4 sequenced and assembled the high-quality genomes of six ixodid tick species and
5 further resequenced 678 tick specimens to understand three key aspects of ticks:
6 genetic diversity, population structure and pathogen distribution. We explored the
7 genetic basis common to ticks, including heme and hemoglobin digestion, iron
8 metabolism, and reactive oxygen species, and unveiled for the first time that both
9 genetic structure and pathogen composition in different tick species were mainly
10 shaped by ecological and geographic factors. We further identified species-specific
11 determinants associated with different host ranges, life cycles and distributions. The
12 findings of this study provide an invaluable resource for research and control of ticks
13 and tick-borne diseases.

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1 Ticks (Acari: Ixodida), which are obligate blood-feeding arthropods, are distributed
2 all over the world from tropic to subarctic regions, with the oldest records dating back
3 to the mid-late Cretaceous (Anderson and Magnarelli, 2008; Peñalver et al., 2018).
4 Ticks are the most versatile vectors, capable of transmitting the broadest spectrum of
5 pathogens, including bacteria, protozoa, fungi, nematodes and viruses, to humans,
6 livestock and wildlife. More than 28 tick species are known to cause a variety of
7 human diseases, such as Lyme disease and spotted fever group rickettsiosis (Jongejan
8 and Uilenberg, 2004), even causing deaths due to misdiagnosis and delayed treatment.
9 Persistent and relapsing infections, as well as long-term sequelae caused by tick-borne
10 pathogens, further worsen the quality of human health (Krause et al., 2008; Mac et al.,
11 2019). Furthermore, the global economic burden in animal husbandry due to
12 tick-borne infections is very large. For instance, the most notorious veterinary
13 ectoparasite, *Rhipicephalus microplus*, is estimated to lead to an annual loss of
14 US\$ 2.5 billion throughout tropical and subtropical regions (Barker and Walker,
15 2014).

16 The threats of tick-borne diseases (TBDs) to human health have unpredictably
17 increased with contemporary urbanization, deforestation, climate change and the
18 rapidly changing interactions between people, animals and their respective habitats. A
19 recent example is the exotic disease vector *Haemaphysalis longicornis*, which has
20 infested multiple states in the United States (Beard et al., 2018) and caused great
21 concern. Even worse, the surging number and geographic expansion of emerging
22 TBDs have caused social anxiety due to unknown health consequences and the lack of
23 approaches to control their transmission. Therefore, fundamental knowledge of tick
24 genomes and genetic diversity is urgently needed, which will undoubtedly open new
25 avenues for research on tick biology, vector-pathogen interactions, disease
26 transmission and control strategies.

27 The first tick genome sequenced, that of *Ixodes scapularis*, offered a glimpse into
28 the genetic architecture and genomic features of the tick (Gulia-Nuss et al., 2016).
29 However, different tick species adapt to diverse environmental niches, feed on diverse

1 hosts ranging from reptiles to mammals and birds, and exhibit complex and distinct
2 life cycles. The dominant tick species across China, including *Ixodes persulcatus*,
3 *Haemaphysalis longicornis*, *Dermacentor silvarum*, *Hyalomma asiaticum*,
4 *Rhipicephalus sanguineus*, and *Rhipicephalus microplus*, have their species-specific
5 characteristics. For example, *Hae. longicornis* is a widely distributed tick species
6 indigenous to eastern Asia, whereas *Hy. asiaticum* prefers to live in desert or
7 semidesert environments (Figure 1A). *R. microplus* has a typical one-host cycle, while
8 most others are three-host ticks depending on the number of host animals they attach
9 themselves to during their life cycle (Figure 1B). Therefore, to better understand their
10 genetic complexity and reveal the links between the genomic variation and
11 geographic distribution, ecological adaptation and vector capacity of ticks, we
12 performed large-scale comparative analyses of 684 ixodid tick genomes, representing
13 six dominant tick species across China (Figures S1-S2).

14 **Six high-quality ixodid tick reference genomes**

15 We used larvae of above-mentioned six representative ixodid ticks for *de novo*
16 genome sequencing. We first constructed ≥ 15 Kb DNA libraries for the PacBio
17 Sequel System and generated 162~303 Gb of subreads with high sequencing depth
18 (approximately 67~95 \times) (Table S1). Considering the relatively high error rate of
19 PacBio sequencing, we further constructed short-fragment libraries (350 bp) and
20 sequenced them using the Illumina HiSeq X-Ten platform, which generated 106~134
21 Gb of clean reads (Table 1). We used these high-quality short reads to perform K-mer
22 frequency analyses to estimate the genome sizes (Table S1) and to correct the short
23 indels and substitutions in the PacBio assembly. To further improve the continuity of
24 the assembled tick genomes and anchor the assemblies into chromosomes, we used
25 Hi-C data to order and orient the contigs as well as to correct misjoined sections and
26 merge overlaps (Figure 1C). Finally, we assembled six tick genomes, achieving
27 8,620~15,174 contigs with scaffold N50 lengths of 533~208,696 Kb and contig N50
28 lengths of 340~1,800 Kb (Table 1; Table S1). Subsequently, we used Benchmarking
29 Universal Single-Copy Orthologs (BUSCO) and the proportion of properly aligned

1 Illumina paired-end reads to evaluate the completeness of these assemblies, which
2 further demonstrated their high completeness and accuracy (Table 1).

3 By combining *de novo* and homology-based approaches, 52.6~64.4% of the
4 repetitive elements were identified from these six assembled tick genomes (Table 1),
5 which is comparable to that from the latest available genome of *I. scapularis*
6 embryonic 6 (ISE6) cell line (~63.5%) (Miller et al., 2018). Among the annotated
7 repeats, LINE and LTR constituted the most abundant known repeat families,
8 representing 8.6~18.3% and 6.5~16.1% of the repetitive sequences, respectively
9 (Table S1). By combining transcriptome-based, homology-based, and *ab initio*
10 approaches, 25,718~29,857 protein-coding genes were predicted from these tick
11 genomes (Table 1). The gene numbers are slightly larger than those predicted in *I.*
12 *scapularis* and two closely related species, namely, *Centruroides sculpturatus* (bark
13 scorpion) and *Parasteatoda tepidariorum* (common house spider) (Thomas et al.,
14 2018) (Table S1), which could be explained by the high completeness and accuracy of
15 the assembled genomes as well as the pairwise homology searches among these six
16 tick species. The average gene length varied greatly among the six tick species, from
17 the smallest (6,466 bp) in *Hae. longicornis* to the largest (15,067 bp) in *I. persulcatus*,
18 with 3.0~4.8 exons per gene and an average intron length of 2,754~3,760 bp (Table
19 S1), indicating the substantial differences in genetic structure among these ticks.

20 To further elucidate the genetic diversity of these tick species, we compared the
21 chromosome size, abundance of repetitive elements, gene content, GC content,
22 noncoding RNA content and synteny of these six tick genomes (Figure 1D). *D.*
23 *silvarum* had the largest genome size and the largest chromosome 1 (> 452 Mb), ~100
24 Mb larger than those of the other species (Table S1). In contrast, the genome size and
25 gene content of *I. persulcatus* were the lowest, while its repetitive elements and
26 noncoding contents were the highest. The GC content was relatively similar across
27 different tick species. Among the six sequenced tick genomes, *I. persulcatus* exhibited
28 very low conserved synteny, which reflects its high genetic divergence from the other
29 tick species. To calculate the evolutionary distances of the six tick species and *I.*

1 *scapularis* from arachnids, orthologous protein sequences were obtained from these
2 species and two outgroup species, *C. sculpturatus* and *P. tepidariorum*, and used to
3 construct a maximum likelihood tree. The divergence time was estimated based on the
4 coding sequences of 464 single-copy orthologous genes. As shown in Figure 1E, the
5 phylogenetic analysis divided the ticks into two main clades, with the two ixodids (*I.*
6 *scapularis* and *I. persulcatus*) closely related to each other and sharing a common
7 ancestor ~200 million years ago (MYA) with the other five ticks. *Hae. longicornis*, *R.*
8 *microplus*, *R. sanguineus*, *Hy. asiaticum*, and *D. silvarum* were clustered together and
9 differentiated from a common ancestor about 137.8 MYA. This genome-based
10 phylogeny constitutes mutual confirmation with the morphological evolutionary tree
11 for ticks (Hoogstraal and Aeschlimann, 1982).

12 **Essential genetic basis of tick hematophagy and the related phenotype**

13 The six sequenced genomes provide a unique resource for understanding the
14 genetic basis of tick hematophagy through comparative genomics and transcriptomics
15 analysis. Through protein family (Pfam) domain-based comparison of the six ticks
16 with *I. scapularis* tick (Miller et al., 2018), three other blood-feeding arthropods
17 (*Anopheles gambiae*, *Aedes aegypti*, and *Glossina morsitans*) and two arachnids (*P.*
18 *tepidariorum* and *C. sculpturatus*), we found that protein families implicated in
19 peptidase activity, transferase activity, transcription regulator activity, transmembrane
20 transporter activity and immunity have notable expansions in ticks (Figure 2A; Table
21 S2). Most of these protein families are relevant to the blood-sucking process. For
22 example, 3~15-fold proliferation of peptidase family M13, ABC-2 family transporter
23 protein, serine protease inhibitor, and glutathione S-transferase occurred in tick
24 genomes (Table S2); these families are involved in hemoglobin digestion, heme
25 transport, blood coagulation, fibrinolysis, detoxification, and oxidative stress
26 (Dickinson and Forman, 2002; Horn et al., 2009; Lara et al., 2015; Rubin, 1996).

27 Long attachment time to the host (several days to weeks), large volume of blood
28 meal (hundreds of times its unfed weight), and broad meal source range (the blood of
29 almost all terrestrial animals) are unique traits of hematophagous ticks and should be

1 involved in many physiological processes, including detoxification of xenobiotic
2 factors, host questing, blood meal digestion, nutrient metabolism, and immune
3 response (Figure 2B). The six tick genomes sequenced in this study provided strong
4 evidence that unlike most eukaryotes (Braz et al., 1999; Gulia-Nuss et al., 2016;
5 Perner et al., 2016), blood-dependent ticks have lost most genes encoding heme
6 biosynthesis and degradation, making them strictly dependent on exogenous sources
7 of heme from the host (Table S3). Thus, ticks are likely to have evolved to acquire
8 and transport heme and iron for vitally important physiological processes and at the
9 same time to maintain redox homeostasis, where free heme and iron can catalyze the
10 generation of reactive oxygen species (ROS). To investigate the potential mechanism
11 associated with iron homeostasis, we surveyed the gain and loss of iron
12 metabolism-related genes in tick genomes and found that the transmembrane protease
13 serine 6 family of matrilysin-2 (TMPRSS6) was significantly expanded (Table S3). In
14 addition, genes associated with antioxidant enzymes, radical scavengers, or
15 heme-mediated activators associated with ROS were mostly conserved across all tick
16 species (Figure 2C, Table S3). This further indicated the importance of maintaining
17 antioxidant systems for ticks, on the one hand to avoid oxidative stress and on the
18 other hand to affect pathogen transmission indirectly by changing its balance with
19 other microbes, as reported in mosquitoes (Cirimotich et al., 2011; Kumar et al., 2010;
20 Oliveira et al., 2011). Furthermore, genes related to immune systems and interactions
21 with pathogens were relatively conserved (Figure 2D; Tables S3), which suggests that
22 ticks may have evolved multiple cellular and humoral immunities to achieve success
23 at the tick-host interface and to maintain a balance at the tick-pathogen interface. In
24 addition, we observed the absence of many genes (*Imd*, *Fadd*, *Dredd*) in the immune
25 deficiency pathway (Table S3), which is essential for recognition and response to
26 Gram-negative bacteria in *Drosophila* (Palmer and Jiggins, 2015), indicating a
27 different strategy of immunological defense against microbes between ticks and fruit
28 flies.

29 We further performed comparative transcriptomic analysis between unfed and fed

1 ticks and found that the differentially expressed genes in various ticks were all
2 enriched in functions of heme and iron ion binding, oxidoreductase activity, and chitin
3 metabolic process (Figure 2E). For example, the upregulated genes in TMPRSS6
4 family exhibited 3~97 fold change during blood sucking in all ticks. The results
5 further elucidate the common genetic basis for tick blood feeding and highlight the
6 importance of these mechanisms for their parasitic lifestyle. Considering that genes
7 after duplication tend to be nonfunctionalized, neofunctionalized or subfunctionalized
8 (Sandve et al., 2018), we explored their expression changes between unfed and fed
9 ticks and found that duplicated genes in larger gene families exhibited a significantly
10 larger standard deviation of fold change than those in smaller gene families
11 (Spearman's rank correlation test, $p < 0.001$), indicating the diversification of these
12 homologous genes in blood-feeding after gene expansion.

13 We next explored the genomic features associated with the species-specific traits
14 that are critical for vector control, including evolutionary distance, host range,
15 geographic distribution and life cycle. *I. persulcatus* in the Prostriata clade evolved
16 much earlier and parasitizes a more diverse range of host groups than the other five
17 tick species (Beati and Klompen, 2019; Hoogstraal and Aeschlimann, 1982). A
18 notable expansion of gene families associated with blood meal digestion,
19 detoxification of xenobiotic factors (such as acaricides, poisons, and environmental
20 pollutants), and nutrient metabolism including serine carboxypeptidase, TMPRSS6,
21 cytochrome P450, and alcohol dehydrogenase etc., was found in *I. persulcatus*
22 (Figures 2C-2D; Table S3). These expansions may confer to *I. persulcatus* additional
23 advantages for nutrient acquisition and endogenous/exogenous detoxification during
24 blood feeding. *Hae. longicornis* has the widest geographic distribution (Figure 1A)
25 and was recently detected in the United States (Beard et al., 2018). We discovered the
26 expansion of known gene families implicated in blood feeding by comparative
27 genomic analyses in *Hae. longicornis* (Figures 2C-2D), which may account for its
28 adaptation to colonize diverse habitats and ecological niches.

29 Another distinguishing trait of ticks is their life cycle. *R. microplus* has a typical

1 one-host cycle. The expanded chemosensory gene family, e.g., the ionotropic
2 receptors (IRs) (Figures 2F-2G; Table S3), which have been associated with a variety
3 of sensory functions (Eyun et al., 2017), may facilitate the strict parasitization by *R.*
4 *microplus* of the same host in each developmental stage. In addition, cytochrome
5 P450 genes, encoding a major family of enzymes involved in the detoxification of
6 xenobiotics, were strikingly reduced (Figure 2G; Table S3). The down-regulations
7 genes after blood meal in RNA-seq differential expression analysis were also enriched
8 in P450 gene families of *R. microplus* (Fisher's exact test, $p = 0.03$). Those may be
9 potentially attributed to *R. microplus* one-host life cycle and a lack of selection
10 pressure.

11 **Population structure and genetic diversity of six tick species**

12 Population evolution is particularly challenging for ticks, as their life cycle
13 consists of long off-host periods (months to years) in changing environments and
14 because of their great reproductive potential, with thousands of eggs being laid after
15 repletion. The genetic diversity of ticks is largely unknown due to the lack of genomic
16 data from different habitats. With the advantage of having acquired six high-quality
17 genomes, we resequenced 678 wild-caught specimens of the six tick species across 27
18 provinces, metropolises or autonomous regions of mainland China, spanning eight
19 ecogeographical faunas and a variety of ecological settings, including coniferous
20 forest, steppe, farmland, desert, shrubland and tropical forest (Figure 1A). Maximum
21 likelihood trees based on full mitochondrial sequences and nuclear single nucleotide
22 variants within single-copy genes were constructed to explore the population structure
23 and genetic diversity among these tick individuals.

24 Through comparison of the six population structures, we found that different tick
25 species have evolved a common dispersal strategy. An ecogeographical distribution
26 pattern was observed for *I. persulcatus*, *D. silvarum*, *Hy. asiaticum*, and *R. sanguineus*
27 (Figure 3A; Figure S3). *I. persulcatus* was relatively restricted in the boreal
28 coniferous forest and temperate forest; *D. silvarum* detected in Shanxi formed a
29 subdivision; The morphologically indistinguishable *R. sanguineus* could be mainly

1 subdivided into two clades, one thriving in tropical forest or shrubland and the other
2 in farmland; and *Hy. asiaticum* was distributed in the same ecological fauna but was
3 geographically differentiated between Xinjiang and Inner Mongolia. Although further
4 investigations of diverse ecosystems, different hosts and larger datasets are needed for
5 broader generalization of these results, our findings suggest that the local adaptation
6 to different ecological niches coupled with geographic distance by restriction of active
7 tick movement can explain the observed patterns of population subdivision in ticks.

8 *Hae. longicornis* is particularly interesting because it is capable of rapidly
9 invading new areas and explosively proliferating in established ranges (e.g., recent
10 invasion to the USA). A very close genetic distance of the *Hae. longicornis*
11 population was observed in the phylogenetic analysis, although this species had a
12 wide geographic distribution occupying diverse ecosystems (Figures 3A-3C).
13 Population structure models supported the division of *Hae. longicornis* into one major
14 population and one minor population (Figure 3A). The major domestic population
15 lacked clear geographic structuring, which suggested that this species was selected for
16 dispersion rather than local competitiveness, which prevented selection for locally
17 adapted phenotypes. The minor population was mainly from three provinces (Fujian,
18 Shanghai and Jiangsu) along the southern coastline of China (Figure 3B). Compared
19 with the major population, the minor was close to the ancestral root of the
20 phylogenetic tree and shared a high similarity with strains from New Zealand
21 (Guerrero et al., 2019) (Figure 3C). Understanding the contribution of migrating birds
22 to the domestic and overseas movement of *Hae. longicornis* is warranted for further
23 dissection of the dispersion of this vector population.

24 As a tick with a typical one-host cycle, *R. microplus* has a distinct population
25 structure and gene flow compared with three-host ticks. We found that *R. microplus*
26 can be clustered into three major clades which largely correspond to their
27 geographical subdivisions: Clade 1 includes specimens from Southwest China
28 (Yunnan), Clade 2 from Southeast China (Hainan and Guangdong to Jiangxi and
29 Fujian) and Clade 3 from South Central China (Guizhou and Chongqing to Hubei,

1 Hunan, Anhui and Zhejiang) (Figure 3B; Figure S4). Comparison of the branches
2 from different provinces showed high F_{ST} values (>0.50), indicating the high genetic
3 differentiation among various *R. microplus* populations in China. Interestingly,
4 phylogenetic analysis based on mitochondrial sequences showed some differences in
5 the tree topology compared with that based on nuclear genome sequences (Figures 3A,
6 3C), indicating distinct paternal and maternal population structures and migration
7 patterns within this species. We speculated that the host specificity within this species
8 may drive local selection patterns of *R. microplus* and greatly alter its population
9 structure (Araya-Anchetta et al., 2015). We also detected extensive gene gain-and-loss
10 events among three subdivisions of *R. microplus* and found that the discriminated
11 genes were enriched in pathways related to the regulation of epithelial cell
12 proliferation and NF- κ B (Figures 3D-3E). The top discriminated genes, such as
13 ubiquitin protein ligase and mucin-6-like protein, indicated some differences of
14 immune response among the three clades.

15 **Key drivers of pathogen distribution in ticks**

16 The complex genomic diversity among tick species implies complicated
17 tick-pathogen interactions, which prompted us to further understand the tick-borne
18 pathogen ecology and evolution. We evaluated the impacts of host gene flow on
19 pathogen distribution by metagenomic analysis of the six tick species. Host DNA
20 contamination could be effectively removed by using the six tick genomes obtained.
21 After filtering the host sequences by mapping the sequencing reads to tick genome
22 assemblies, microbial composition analysis and pathogen identification were
23 performed for each of the 678 specimens.

24 The tick taxonomy is an important factor in defining the potential of a tick to
25 transmit pathogens. Our study for the first time unveiled the landscape of pathogens
26 carried by six tick species collected from a wide range of geographical sources. In
27 general, the relative abundance of certain pathogens was quite different across the six
28 tick species (Figure 4A). *I. persulcatus* and *Hae. longicornis*, traditionally the most
29 important vectors of human and animal diseases (Fang et al., 2015), were found

1 bearing various bacterial species of *Anaplasma*, *Babesia*, *Borrelia*, *Coxiella*, *Ehrlichia*,
2 and *Rickettsia* (Figure 4A). In contrast, *R. sanguineus* had the lowest abundance of
3 bacterial pathogens. *R. microplus*, which transmits *Babesia* and *Anaplasma* in
4 livestock and wild ruminants, possesses a *Coxiella*-like endosymbiont as the most
5 abundant bacterial taxon (Figure 4A). Notably, *D. silvarum* presented the largest
6 relative abundance of *Rickettsia* (Figure 4A). *Hy. asiaticum* carried the highest
7 relative abundance of *Coxiella burnetii* and *Francisella tularensis* (Figure 4A), the
8 causative agents of Q fever and tularemia, respectively.

9 The interplay among humans, animals and ecosystems is well acknowledged.
10 However, the driving factors of interactions among the environment, pathogens,
11 vectors and hosts have not yet been clearly addressed for TBDs. Each geographical
12 fauna has specific ecological features and thus favors different forms of animal life.
13 We observed that the bacterial distribution had an overall correlation with the
14 ecogeographical faunal region for a given tick species (Figure 4A). For example, the
15 relative abundances of *Anaplasma* and *Ehrlichia* in *R. sanguineus* were lower in
16 tropical forest and shrubland areas than in farmland faunal regions ($p < 0.05$, Mann–
17 Whitney U test); for *D. silvarum*, nonpathogenic *Anaplasma* was prevalent in North
18 China, and for *R. microplus*, *Rickettsia* was prevalent in Southwest China (Figure 4A)
19 ($p < 0.001$, Mann–Whitney U test). To quantify the microbial divergence across
20 regions, we compared the Bray-Curtis (BC) dissimilarities of the tick microbiota
21 between different geographic faunas and within the same geographic fauna (Figure
22 4B), and found that the calculated BC dissimilarities varied by geographic distance
23 for each tick species (Figure 4C). We found that the more the geographic fauna or
24 distance diverged, the larger the tick microbiota dissimilarity was, and such a pattern
25 may consequently impact the pathogen distribution. In addition to above key drivers,
26 we also found that different subtypes of *R. microplus* and *Hae. longicornis* exhibited
27 different positive rates of *Rickettsia* ($p < 0.001$ for *R. microplus* and $p < 0.05$ for *Hae.*
28 *longicornis*, Kruskal-Wallis test) (Figure 4A), further indicating the necessity of
29 determining and monitoring the tick subspecies or subpopulations with more pathogen

1 load.

2 We further summarized all the reported human cases with TBDs in China from
3 1980 to 2020 (Figure 4D; Figure S5; Table S4). During the past 40 years, at least 22
4 diseases caused by tick-borne bacteria or protozoa have been reported. The
5 northeastern China is a high-risk area where about 15 pathogens, half of which were
6 emerging agents, have caused human infections (Jia et al., 2018; Jia et al., 2013; Jia et
7 al., 2014; Jiang et al., 2018; Jiang et al., 2015; Li et al., 2015). We mapped the
8 abundance and proportion of pathogens of different tick species onto their collection
9 sites (Figure 4D). By overlapping the distributions of TBDs and detected pathogens in
10 ticks, we found that pathogenic *Rickettsia* had both the high prevalence and large
11 abundance in the ticks from the Northeastern China, where spotted fever group
12 rickettsioses were frequently diagnosed. However, the abundance of tick pathogens
13 does not strictly correlate with their transmission rate to human in general. Besides the
14 reason that the identified pathogens in ticks of this study may not be at infectious
15 stage when they were sampled, another possible explanation is that there might be
16 under-reported cases of TBDs due to lack of etiologic diagnosis tests in many
17 endemic areas. It should be noted that although the abundance of *Borrelia* was only 3%
18 of that of *Rickettsia* in the ticks in Northeastern China, it has caused the disease
19 incidence as high as *Rickettsia* (Figures 4E-4F). Taken together, these findings
20 suggest that pathogen abundance may not be the sole factor in determining the risk of
21 human infection, which highlights the necessity of more sensitive approaches to
22 identify the low abundance pathogens in ticks.

23 In conclusion, the genomes of six representative species generated in this study
24 provide novel insights into tick-specific blood feeding life, tick-pathogen interactions
25 and the development of genetic tools for tick control. The large-scale genomic
26 re-sequencing of 678 wild-caught tick specimens further unveils the high genetic
27 heterogeneity of ticks, reflecting their local adaptation to diverse ecological niches.
28 Based on metagenome profiling and pathogen screening of these tick specimens, we
29 described the landscape of microbial pathogens, including some emerging human

1 pathogens, carried by six tick species collected from a wide range of geographical
2 sources. The pathogen composition in different tick species is mainly shaped by
3 ecological and geographic factors, and different subpopulations may have diverse
4 tick-borne pathogen profiles. We believe the tick genomes and their associated
5 pathogen profiles generated in this study will undoubtedly benefit the community on
6 global tick and TBD control.

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7 **AUTHORS CONTRIBUTIONS**

8 W.C.C, F.Z., N.J. designed and supervised research. Y.S., J.F.J., X.M.C., B.G.J.,
9 Q.C.C., S.J.D., X.J.W., J.G.Z., X.D.R., T.C.Q., C.H.D., J.X.C., P.F.D., X.H.H., E.J.H.,
10 J.Z.L., H.Z.S., X.W., C.C.W., T.C.Y., Q.B.H., W.L., H.Y.C., L.G.Z., J.H.T. collected
11 samples. Q.W, T.T.Y, L.F.L., W.W., L.Y.X., J. L., prepared materials for sequencing.
12 Q.W., L.Z., Y.S., W.B.G., X.B.N. set up the database, W.Z., W.F.Y., Y.C.G, T.L,
13 performed genome sequencing. W.Z., W.F.Y., Y.C.G, T.L, W.S, performed genome
14 assembly and annotation. W.S., L.D., J.W, N.J., F.Z. performed genome analysis and
15 interpretation, J.W. W.S, L.D., N.J., Y.H,Z. R.Z.Y prepared figures and tables. L.B.S.
16 provided the tick cell line and edited the manuscript, N.J., J.W., F.Z. W.C.C. wrote the
17 paper.

18 **DECLARATION OF INTERESTS**

19 The authors declare no competing interests.

20

1 **Figure Legends**

2 **Figure 1 Basic information and genomic comparison of six tick species.** (A) Map
3 of sample collections. The size of the circle represents the number of tick samples
4 collected in the area. Geographical fauna were recorded as follows: Northeast China
5 (I); North China (II); Neimenggu-Xinjiang (III); Qinghai-Xizang (IV); Southwest
6 China (V); Central China (VI); South China (VII). Ecological fauna are also shown on
7 the map with different colors. (B) Illustration of ticks with a 3-host life cycle, in
8 which larvae and nymphs feed on blood once before molting, the adults feed once,
9 and then, the fully engorged tick drops from the host and lays thousands of eggs to
10 continue the life cycle. (C) Hi-C interactive heatmap of the genome-wide organization
11 of 11 chromosomes for five ticks. For auxiliary assembly of chromosomes, assemblies
12 were cut into bins of the same length. The effective mapping read pairs between two
13 bins were used as a signal of the strength of the interaction between the two bins.
14 With the numbered chromosomes as the coordinates, the color of each dot represents
15 the log value of the interaction intensity of the corresponding bin pair of the genome,
16 and the interaction intensity increases from yellow to red. Chr represents
17 chromosomes. (D) Comparative genomic analysis of six tick species. From the outer
18 circle to the inner circle, nine types of information, namely, chromosome size,
19 Illumina data coverage, PacBio data coverage, Hi-C data coverage, repeat abundance,
20 gene abundance, GC content, ncRNA, and gene synteny, are labeled successively with
21 the letters a-i. In the synteny analysis, the blue and red lines denote *R. microplus* and
22 *Hae. longicornis*, serving as the reference genome, respectively. (E) Maximum
23 likelihood phylogeny of all sequenced ticks with two species of Arachnoidea as
24 outgroups. The estimated divergence time between clades is labeled on the branch
25 nodes. See also Figures S1-S2.

26

27 **Figure 2 Genetic basis of tick hematophagy and the related phenotype.** (A)
28 Species-specific and shared Pfam family among ticks and other arthropod species.
29 Each cell in the heatmap represents the normalized gene count (across all species on

1 the left side) of a Pfam family. Only the Pfam that are specific to ticks or common
2 with other blood-feeding arthropod species are shown. Pfams are further grouped
3 according to their functions in biological processes or activities. (B) Unique
4 hematophagous traits of ticks, including detoxification of xenobiotic factors (a), host
5 questing (b), blood meal digestion (c), nutrient metabolism (d), and immune response
6 (e). (C) Gene counts of four gene categories in six tick species: detoxification of
7 xenobiotics (yellow), iron metabolism (deep blue), hemoglobin digestion (green) and
8 oxidative stress (purple). (D) Gene counts of six tick species related to five
9 hematophagous traits of ticks. (E) Gene ontology (GO) enrichment analysis based on
10 the transcriptomic data of unfed and fed ticks. The biological process, cellular
11 component, and molecular function categories are referred to as BP, CC and MF,
12 respectively. From the inner circle to the outer circle, three levels of GO enrichment
13 are displayed with nodes. The sector of the nodes in outermost circle represents the
14 proportion of DE genes in three ticks, namely, *I. persulcatus*, *Hae. longicornis* and *R.*
15 *microplus*. The sector of the nodes in inner circles represents the absence or presence
16 of DE genes. (F) Gene counts of four different perception pathways to quest preferred
17 hosts in six tick species. (G) Phylogenetic analysis of the IR25a gene (left) and P450
18 gene group I family (right). The colors of the nodes on the tree represent different tick
19 species. See also Tables S2-S3.

20

21 **Figure 3 Genetic diversity and population structure of six tick species.** (A)
22 Phylogenetic structure of tick populations based on the mitochondrial genome. The
23 subtitle of each tree indicates the species name and the number of specimens. The
24 color of the tree tip represents the ecological fauna type of the sample location. (B)
25 Geographical population structure of *Hae. longicornis* and *R. microplus*. In the top bar
26 plot, each vertical line shows the membership probability of a specimen inherited
27 from each of the inferred ancestral populations (K=5) for *Hae. longicornis*, and
28 specimens are grouped by the sampled province as annotated by the line segment on
29 the top. The bottom plot shows the same information for *R. microplus*. Pie charts on

1 the map aggregate the same membership probability of ancestral populations for all
2 specimens in each province. Neighboring provinces are connected according to the
3 F_{ST} value between the two provinces. (C) Phylogenetic structures of *Hae. longicornis*
4 (left) and *R. microplus* (right) populations based on their nuclear genomes. The strain
5 previously reported in New Zealand and its close relative were highlighted. (D)
6 Circos plot of genes with elevated copy numbers in the three clades of *R. microplus*.
7 (E) GO enrichment analysis of genes with elevated copy numbers in the three clades
8 of *R. microplus*. The heatmap color represents the adjusted p-value (-log 10). The
9 biological process, cellular component, and molecular function categories are referred
10 to as BP, CC and MF, respectively. See also Figures S3-S4.

11

12 **Figure 4 Potential pathogen profiling of six tick species.** (A) The distribution and
13 abundance of known tick-borne pathogens and their related species in the six tick
14 populations. The relative abundance of the microbes in each sample was estimated by
15 read counts per 100,000 reads. Subtypes of each tick species were classified based on
16 the phylogenetic analysis of the resequenced genomes. Geographic fauna and
17 ecological fauna were selected according to the Chinese fauna classification and were
18 annotated in the corresponding colors. Bacterial species of twelve human pathogenic
19 genera are shown, and each genus name is indicated below the heat map. Human
20 pathogens are annotated in deep gray, and nonhuman pathogens are annotated in light
21 gray. (B) Bray-Curtis dissimilarity between each pair of samples, grouped within the
22 same geographic fauna or between different geographic fauna. (C) Bray-Curtis
23 dissimilarity between each pair of samples varied by geographic distance. (D)
24 Epidemiological distribution of tick-borne disease (TBD) patients and tick pathogens.
25 The cases of human infection were reported between 1980 and 2020. The pies
26 indicate pathogen composition, with the color of circle outline representing tick
27 species. The circle size indicates the relative abundance of all pathogens per 10^5
28 microbial reads, and the color and area of pies indicate the species and relative
29 abundance of each pathogen, respectively. The Northeastern China is highlighted in

1 dark gray. (E) The relative abundance (node color) and positive rate of 33 human
2 pathogenic bacteria or protozoa species of the ticks in the Northeastern China. (F) The
3 reported incidences of TBD among the risk population in the Northeastern China. See
4 also Figure S5 and Table S4.

Table 1. Summary of the Assembly and Annotation Information of the Sequenced Tick Genomes

	<i>I. persulcatus</i>	<i>Hae. longicornis</i>	<i>D. silvarum</i>	<i>Hy. asiaticum</i>	<i>R. sanguineus</i>	<i>R. microplus</i>	<i>I. scapularis</i> ^a
Data statistics							
Illumina clean data (Gb)	118.4	115.1	134.1	121.1	105.9	110.1	49.6
Pacbio subreads (Gb)	165.1	303.1	202.7	162.3	183.6	170.6	192.5
Hi-C clean data (Gb)	-	306.0	210.1	201.3	185.5	168.8	-
Assembly statistics							
Contig span (Mb)	1,901.7	2,554.5	2,473.0	1,713.1	2,364.5	2,529.8	2,691.1
Contig N50 (Kb)	532.9	740.0	340.0	555.4	541.9	1,800.7	269.7
Chromosome size (Mb)	-	2,230.7	2,384.8	1,539.3	2,210.2	2,140.8	-
Scaffold N50 (Kb)	532.9	204,922.3	189,477.5	137,335.1	208,696.2	183,350.9	835.7
GC content (%)	46.0	47.4	46.9	46.6	46.8	45.8	46.0
Genome completeness							
Mapping rate (%)	97.5	93.6	98.1	97.9	92.7	97.7	98.7
Coverage rate (%)	98.1	96.7	98.8	99.2	98.1	98.3	96.6
BUSCO (%)	93.2	91.8	91.6	93.3	92.3	90.3	95.0
Annotation statistics							
Repeat content (%)	64.4	59.3	60.2	52.6	61.6	63.1	63.5
Gene numbers	28,641	27,144	26,696	29,644	25,718	29,857	24,501
Mean gene length (bp)	15,067	6,466	12,166	10,574	11,201	8,818	26,459
Mean CDS length (bp)	1,091	892	1,097	960	1,016	1,009	1,348

5 a. Scaffold N50, GC content, and annotation statistics were calculated using the latest available genome of *I. scapularis* ISE6 cell line (Miller et al., 2018).

6

7 SUPPLEMENTAL INFORMATION

8 **Figure S1. Illustrations of the Six *de novo* Sequenced Tick Species, Related to**
9 **Figure 1.**

10 **Figure S2. Life Cycles (A-E) and Parameters (F) under Laboratory Rearing**
11 **Conditions of Six Tick Species, Related to Figure 1.**

12 **Figure S3. Phylogenetic Structures for Tick Populations Based on Mitochondrial**
13 **(A) and Nuclear (B) genome, Related to Figure 3.**

14 **Figure S4. Geographical Population Structures of (A) *Hae. longicornis* and (B) *R.***
15 ***microplus*, Related to Figure 3.**

16 **Figure S5. Epidemiological Distribution of the Human Cases Infected with**
17 **Tick-borne Diseases and of Pathogen Profiles in Six Tick Species, Related to**
18 **Figure 4.**

19

20 **Table S1. Details of the Assembly and Annotation Information of the Sequenced**
21 **Tick Genomes and Their Homolog Species, Related to Table 1.**

22 **Table S2. Pfam Comparison across Seven Tick Species, Three Other**
23 **Blood-feeding Arthropods and Two Arachnids, Related to Figure 2A.**

24 **Table S3. Comparative Analysis of the Genes Associated with Tick**
25 **Hematophagy and the Related Phenotype, Related to Figure 2.**

26 **Table S4. Epidemiological Distribution of the Human Cases Infected with**
27 **Tick-borne Diseases from 1980 to 2020, Related to Figure 4D.**

28 **Table S5. Detailed Commands, Parameters and Configurations Used in Tick**
29 **Genome Assembly, Repeat Identification and Gene Annotation, Related to**
30 **STAR★METHODS.**

31

32

33	STAR★METHODS
34	● KEY RESOURCE TABLE
35	● RESOURCE AVAILABILITY
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37	○ Materials Availability
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57 **STAR★METHODS**

58

59 **KEY RESOURCE TABLE**

60

61 **RESOURCE AVAILABILITY**

62 **Lead contact**

63 Further information and requests for resources and reagents should be directed to and
64 will be fulfilled by the Lead Contact, Wu-Chun Cao (caowc@bmi.ac.cn).

65

66 **Materials Availability**

67 The study did not generate any new reagents.

68

69 **Data and code availability**

70 The genome assemblies and annotations generated in this study are available at BIGD
71 (<https://bigd.big.ac.cn>, project accession ID PRJCA002240). We have also submitted
72 the genome assemblies to GenBank (accession ID:
73 JABSTQ000000000-JABSTV000000000) with the project accession ID:
74 PRJNA633311. The raw data of re-sequenced samples are available at BIGD
75 (accession number PRJCA002242). We provided a detailed list of software,
76 commands, parameters and configuration files used in genome data analyses in Table
77 S5.

78

79 **METHODS DETAILS**

80 **Sample collection**

81 From November 2017 to January 2019, ticks were collected from 28 provinces,
82 metropolises or autonomous regions of mainland China. The collection sites were
83 selected according to their ecological environments, including coniferous forest,
84 steppe, farmland, desert, shrubland and tropical forest. Ticks were collected by
85 dragging a standard 1-m² flannel flag over vegetation or from domestic or wild
86 animals such as cattle, dogs, sheep, goats, cats, rabbits, camels, deer, and boars. The
87 latitude and longitude of each collection site were recorded. The species, sex and
88 developmental stage of each tick were identified by entomologists. Adult ticks were
89 used for tick genome resequencing to understand their genetic diversity, population

90 structure and pathogen distribution. Most of the *R. sanguineus* and *R. microplus* ticks
91 were collected from animal hosts. A majority of the *I. persulcatus*, *Hae. longicornis*,
92 *D. silvarum*, and *Hy. asiaticum* specimens were free questing ticks. Live ticks were
93 transported to the laboratory, and dead ticks were directly stored at -80 °C. A total of
94 678 specimens were used for tick genome resequencing (Figure 1A).

95 Live adult ticks of *I. persulcatus*, *Hae. longicornis*, *D. silvarum*, *Hy. Asiaticum*,
96 *R. sanguineus*, and *R. microplus* collected from the Heilongjiang (129.22°E,
97 44.96°N), Shandong (122.32°E, 36.89°N), Shanxi (110.93°E, 38.70°N), Tibet
98 (91.09°E, 30.68°N), Guangxi (109.96°E, 22.41°N) and Guizhou (107.96°E, 26.56°N)
99 provinces, respectively (Figure 1A), were laboratory reared to obtain larvae and then
100 used for *de novo* genome sequencing. Laboratory mice (for *I. persulcatus*), rabbits
101 (for *Hae. longicornis* and *D. silvarum*) and goats (for *Hy. asiaticum*) were used for
102 blood feeding to obtain engorged females. Engorged *R. sanguineus* and *R. microplus*
103 ticks were directly collected from dogs or cattle on site. Engorged female ticks were
104 reared separately under a 12-hour light/12-hour dark photoperiod at 25 °C in
105 desiccators in which a saturated aqueous solution of K₂SO₄ was used to maintain
106 relative humidity. Larvae hatched from a single female were used for the subsequent
107 *de novo* genome sequencing (Illumina, PacBio sequencing and Hi-C experiment),
108 considering their lower contamination of environmental bacteria than those directly
109 collected from natural environments, and their single maternal source which may
110 reduce genetic complexities. In addition, to reduce the genetic heterozygosity of *R.*
111 *microplus*, the embryo-derived cell line BME/CTVM23 (Alberdi et al., 2012) of *R.*
112 *microplus* was also subjected to deep sequencing and then used for genome
113 scaffolding.

114

115 ***De novo* sequencing, assembly and annotation**

116 ***Genomic DNA preparation and genome sequencing***

117 Larvae hatched from a single female were used for *de novo* sequencing.
118 Approximately 50-100 larvae of each tick species were collected, thoroughly
119 surface-sterilized (two successive washes of 70% ethanol, 30 s each) and then used
120 for genomic DNA extraction using the DNeasy Blood & Tissue Kit (Qiagen, USA).
121 The integrity of the DNA was determined using an Agilent 4200 Bioanalyzer (Agilent
122 Technologies, Palo Alto, California, Genomic DNA Analysis ScreenTape and

123 Genomic DNA Reagents). Two high-throughput sequencing platforms, namely, the
124 Illumina HiSeqX-Ten and Pacific Bioscience Sequel, were used to generate
125 sequencing data. First, more than 1 μ g of DNA was used to construct short
126 fragmented libraries with an insertion size of 350 bp, which were then sequenced on
127 the Illumina HiSeqX-Ten platform. For each tick species, approximately ~110Gb
128 Illumina sequencing data were generated. Second, 8 mg of DNA was sheared using
129 g-Tubes (Covaris, Woburn, MA) and concentrated with AMPure PB magnetic beads.
130 Each single-molecule real-time (SMRT) bell library was constructed using the Pacific
131 Biosciences SMRTbell Template Prep Kit 1.0. The constructed libraries were
132 size-selected on a BluePippinTM system for molecules \geq 15 kb, followed by primer
133 annealing (Sequencing Primer v3) and the binding of SMRTbell templates to
134 polymerases with the Sequel Binding and Internal Control Kit 3.0. Sequencing
135 (Sequel Sequencing Kit 3.0 Bundle, SMRT Cell 1M v3 Tray) was performed on the
136 Pacific Bioscience Sequel platform by Annoroad Gene Technology Beijing Co. Ltd.

137 To further improve the continuity of the assembled genomes, approximately 100
138 ~ 200 larvae of five tick species were used for chromosome conformation capture
139 (Hi-C) experiments (*I. persulcatus* was not included due to its limited sample size).
140 Cells/tissues were crosslinked using 40 ml of 2% formaldehyde solution at room
141 temperature for 15 min. A total of 4.324 ml of 2.5 M glycine was added to quench the
142 crosslinking reaction. The supernatant was removed, and the tissues were ground with
143 liquid nitrogen and resuspended in 25 ml of extraction buffer I containing 0.4 M
144 sucrose, 10 mM Tris-HCl (pH 8.0), 10 mM MgCl₂, 5 mM β -mercaptoethanol, 0.1 mM
145 phenylmethylsulfonyl fluoride (PMSF), and 13 protease inhibitors (Sigma) and then
146 filtered through Miracloth (Calbiochem). The filtrate was centrifuged at 4000 rpm and
147 4 °C for 20 min. The pellet was resuspended in 1 ml of extraction buffer II (0.25 M
148 sucrose, 10 mM Tris-HCl (pH 8), 10 mM MgCl₂, 1% Triton X-100, 5 mM
149 β -mercaptoethanol, 0.1 mM PMSF, and 13 protease inhibitors) and centrifuged at
150 14,000 rpm and 4 °C for 10 min. The pellet was resuspended in 300 ml of extraction
151 buffer III (1.7 M sucrose, 10 mM Tris-HCl (pH 8), 0.15% Triton X-100, 2 mM
152 MgCl₂, 5 mM β -mercaptoethanol, 0.1 mM PMSF, and 1 μ l of protease inhibitor),
153 loaded on top of an equal amount of clean extraction buffer III and then centrifuged at
154 14,000 rpm for 10 min. The supernatant was discarded, and the pellet was washed
155 twice by resuspending in 500 μ l of ice-cold 1 \times CutSmart buffer and then centrifuged

156 for 5 min at 2,500 ×g. The nuclei were washed with 0.5 ml of restriction enzyme
157 buffer and transferred to a safe-lock tube. Next, the chromatin was solubilized with
158 dilute SDS and incubated at 65 °C for 10 min. After quenching the SDS with Triton
159 X-100, overnight digestion was performed with a 4-cutter restriction enzyme (400
160 units of MboI) at 37 °C on a rocking platform. The next step was Hi-C specific,
161 including marking of the DNA ends with biotin-14-dCTP and performing blunt-end
162 ligation of crosslinked fragments. The proximal chromatin DNA was religated using
163 the ligation enzyme. The nuclear complexes were reverse-crosslinked by incubating
164 with proteinase K at 65 °C. DNA was purified by phenol-chloroform extraction, and
165 biotin-C was removed from nonligated fragment ends using T4 DNA polymerase.
166 Fragments were sheared to 100-500 bp by sonication. The fragment ends were
167 repaired using a mixture of T4 DNA polymerase, T4 polynucleotide kinase and
168 Klenow DNA polymerase. Biotin-labeled Hi-C samples were specifically enriched
169 using streptavidin magnetic beads. A-tails were added to the fragment ends by
170 Klenow (exo-), and then the Illumina paired-end sequencing adapter was added via a
171 ligation mix. Finally, the Hi-C libraries were amplified by 10-12 cycles of PCR and
172 sequenced on an Illumina HiSeqX-Ten (HiSeq X Ten Reagent Kit v2.5).

173 ***Genome size estimation***

174 Before *de novo* assembly, we estimated the genome size of each tick species. For each
175 tick species, we built an Illumina short-read library using the DNA material from the
176 same source as the PacBio sequencing library, and ~110 Gb Illumina sequencing data
177 were generated. Based on the Illumina data, Jellyfish (v2.1.3) (Marçais and Kingsford,
178 2011) was employed to calculate the frequency of each K-mer (k=21). Then, the
179 genome size was estimated using a previously described method based on K-mer
180 distribution (Liu et al., 2013).

181 ***Genome assembly and quality assessment***

182 PacBio reads were first assembled using four *de novo* assemblers: Canu (Koren et al.,
183 2017), Falcon (Chin et al., 2016), SMARTdenovo (Istace et al., 2017) and wtdbg
184 (Ruan and Li, 2020). The best assembly was selected according to the optimal
185 continuity and completeness, and the final version of the genome assembly was
186 polished by Arrow and error-corrected by Pilon (Walker et al., 2014) using Illumina
187 reads. The completeness of the final assembly was evaluated using two criteria: (1)
188 BUSCO (v3.0, arthropoda_odb9) (Simão et al., 2015) based on the evolutionarily

189 informed expectations of gene content from near-universal single-copy orthologs; (2)
190 mapping rate and coverage of Illumina reads on the assembled genomes.

191 Scaffolding was performed using Hi-C-based proximity-guided assembly for five
192 tick species, excluding *I. persulcatus*. Hi-C reads were first aligned to the draft
193 genome using the bowtie2.2.3 algorithm (Langmead and Salzberg, 2012). According
194 to the Hi-C protocol and the fill-in strategy, unmapped reads were mainly composed
195 of chimeric fragments spanning the ligation junction. HiC-Pro (V2.7.8) was used to
196 identify ligation sites and align back to the genome using the 5' fraction of the read
197 (Servant et al., 2015). The assembly package Lachesis (Burton et al., 2013) was used
198 to perform clustering, ordering and orienting. Based on the agglomerative hierarchical
199 clustering algorithm, we clustered the scaffolds into 11 chromosome groups based on
200 the karyotypes of chromosomes from a previous report (Qin et al., 1997). Contigs
201 from the polished and corrected assembly were anchored to chromosome groups with
202 a length ratio of 80% ~ 95%.

203 ***Additional assembly procedures for Hae. longicornis***

204 The initial genome size of *Hae. longicornis* was estimated to be 5.4 G based on the
205 Illumina sequencing data of 100 larvae, which was much larger than those of the other
206 five tick species. Considering its nontypical K-mer Poisson distribution, we assume
207 that the elevated genome size could be attributed to the heterozygosity of the larvae
208 used for *de novo* sequencing. Therefore, we resequenced additional *Hae. longicornis*
209 specimens from three provinces (Beijing, Shandong and Zhejiang), with one male and
210 one female from each province. The genome sizes of three males and two females
211 were approximately 2.4-2.8 Gb. Interestingly, the genome size of the female from
212 Shandong was approximately 3.6 Gb. The larger genome size of this female may be
213 related to the additional chromosomes in the parthenogenetic lineage, which was
214 supported by the detected genetic markers of the parthenogenetic lineage in the
215 female sample (Chen et al., 2014).

216 The overestimated genome size of *Hae. longicornis* indicated its high genome
217 heterozygosity in the PacBio library. Therefore, additional assembly procedures were
218 adopted beyond the conventional pipeline to improve the assembly quality. First,
219 before assembly, we used the Illumina reads of a single female sample to correct the
220 PacBio reads using LorDEC version 0.8 (Salmela and Rivals, 2014). Second, we
221 filtered a subset of the PacBio reads that showed a low LorDEC correction ratio

222 (<25%, i.e., proportion of PacBio reads covered by Illumina reads). After filtering, the
223 corrected PacBio reads were fed into the assembler. Third, we obtained a core genome
224 by removing the genome sequences from 7 redundant homologous chromosomes of
225 the core female genome by using redundans (Pryszcz and Gabaldón, 2016) (with
226 parameters including an identity of 80% and overlap of 50%). Finally, contigs of the
227 core male *Hae. longicornis* assembly were anchored in 11 chromosomes using the
228 Hi-C data.

229 ***Repeat annotation***

230 Repetitive sequences and transposable elements (TEs) in each tick genome were
231 identified using a combination of *de novo* and homology-based approaches. Briefly,
232 RepeatMasker (open-4.0.6) (Chen, 2004) and RepeatProteinMask (v.4.0.6) were used
233 to identify and classify different TEs by aligning genome sequences against Repbase
234 version 23.12 (Jurka et al., 2005) with default parameters. To identify tandem repeats,
235 TRF v4.0.6 (Benson, 1999) was used with the following parameters: Match = 2,
236 Mismatch = 7, Delta = 7, PM = 80, PI = 10, Minscore = 50, MaxPeriod = 500, -d, -h.

237 ***Genome annotation***

238 Gene annotation was accomplished by integrating evidence or predictions from
239 transcriptome-, homology- and *ab initio*-based approaches. In the transcriptome-based
240 approach, RNA was extracted from six tick species. In brief, ticks were quickly
241 washed in RNase-free water twice and homogenized in RLT solution under liquid
242 nitrogen. The homogenate was then incubated at 55 °C for 10 min with proteinase K
243 (Qiagen, USA) and centrifuged for 30 s at full speed. The homogenized lysate was
244 used for further RNA extraction using the RNeasy Mini Kit (Qiagen, USA). RNA
245 quality was assessed using an Agilent Bioanalyzer 2200 (Agilent Technologies, Inc.).
246 RNA-seq libraries were generated by using RiBo-Zero Gold rRNA Removal Reagents
247 (Human/Mouse/Rat) (Illumina). Paired-end (150 bp) sequencing of the RNA library
248 was performed on an Illumina HiSeq 4000 platform. RNA-seq reads generated from
249 each tick species were assembled by Trinity (v2.4.0,
250 <https://github.com/trinityrnaseq/trinityrnaseq>) with default parameters (Haas et al.,
251 2013). The assembled transcripts were aligned to each assembled genome and were
252 used to predict gene structure by PASA (v2.3.3
253 http://wfleabase.org/release1/PASA_gene_annotation.html) (Haas et al., 2008). The
254 protein sequences of homologous species, including *I. scapularis*

255 (<https://www.vectorbase.org/>), *C. sculpturatus*
256 (<https://i5k.nal.usda.gov/content/data-downloads>) and *P. tepidariorum*
257 (<https://i5k.nal.usda.gov/content/data-downloads>), were retrieved from public
258 databases. In addition, as the six tick species sequenced are closely related species, the
259 genes of all five species annotated only by PASA were also added to the homologous
260 gene dataset. Homologous protein sequences were aligned to the tick genome
261 assemblies using TBLASTN v2.2.28+
262 (<https://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/2.2.28/>) with e-value=1e-5
263 (Camacho et al., 2009), and the gene structure was predicted by GeneWise v 2.2.0
264 (Birney et al., 2004). *Ab initio* gene prediction was performed using Augustus v3.3
265 (Stanke et al., 2004), GlimmerHMM v 3.0.4 (Majoros et al., 2004), SNAP (Korf,
266 2004), and GeneMark v3.51 (Besemer and Borodovsky, 2005). Based on the above
267 evidence, we used EvidenceModeler (EVM) v1.1.1(Haas et al., 2008) to integrate the
268 gene models predicted by the above approaches into a nonredundant and more
269 complete gene set. Finally, the functions of the protein-coding genes were predicted
270 by searching against multiple gene annotation databases, including SwissProt
271 (<http://www.ebi.ac.uk/interpro/search/sequence-search>), NT
272 (<https://www.ncbi.nlm.nih.gov/nucleotide/>), NR
273 (<https://www.ncbi.nlm.nih.gov/protein/>), Pfam (<http://xfam.org/>), Eggnog
274 (<http://eggnogdb.embl.de/>), GO (<http://geneontology.org/page/go-database>), and
275 KEGG (<http://www.genome.jp/kegg/>).

276 ***Noncoding RNA annotation***

277 Four types of noncoding RNAs (ncRNAs), namely, microRNAs (miRNAs), transfer
278 RNAs (tRNAs), ribosomal RNAs (rRNAs) and small nuclear RNAs (snRNAs), were
279 identified. The tRNA genes were identified using tRNAscan-SE v1.3.1 (Lowe and
280 Eddy, 1997) with default parameters. The rRNA fragments were predicted by aligning
281 human rRNA sequences to the assembled genome sequences by BLASTN with the
282 parameter e-value <1e-5. The miRNA and snRNA genes were searched using BLAST
283 against the Rfam v13.0 database using INFERNAL v1.0 with a family-specific
284 “gathering” cutoff of Rfam (Griffiths-Jones et al., 2005).

285 ***Collinearity analysis***

286 Collinear segments were detected between assembled genomes using JCVI software
287 (v0.8.4, <https://github.com/tanghaibao/jcvi>) (Tang et al., 2015) with default
288 parameters.

289

290 **Gene family and phylogenetic analysis**

291 To infer tick evolutionary history, a maximum likelihood phylogenetic tree was built
292 based on the protein sequences of nine species, including the six tick species
293 sequenced in this study, *I. scapularis* and two outgroup species (*C. sculpturatus* and
294 *P. tepidariorum*) (Thomas et al., 2018) . First, single-copy genes within the nine
295 species were identified, and all-to-all BLAST was performed for all protein sequences
296 (E-value <10⁻¹⁰ and identity >30%). Gene families (i.e., ortholog or paralog groups)
297 were identified using OrthoMCL (Li et al., 2003) with the parameters -I=1.5.
298 Single-copy gene families (n=464) were used for subsequent phylogenetic analysis.
299 The protein sequences of these single-copy genes were aligned using MUSCLE (v3.6)
300 (Edgar, 2004) and then used to construct a maximum likelihood tree by PhyML (v3.0)
301 (Guindon et al., 2010).

302

303 **Divergence time estimation**

304 The divergence time within the nodes of the phylogenetic tree was estimated by the
305 MCMCTREE program of PAML (v4.4) (Yang, 2007) with parameters RootAge=500,
306 model=4, alpha=0, clock=3, sample frequency=2, burn-in=20000, nsample=100000,
307 and finetune="0.00876 0.03724 0.06828 0.00789 0.44485". The divergence time was
308 corrected using calibration points from the TimeTree website (<http://timetree.org/>)
309 (Kumar et al., 2017).

310

311 **Gene family analysis and comparison**

312 The expansion and contraction of gene families were determined by comparing the
313 cluster size differences between the ancestor and each of the six investigated tick
314 species and *I. scapularis* using the CAFE program
315 (<http://sourceforge.net/projects/cafehahnlab/>) (De Bie et al., 2006). CAFE used a
316 random birth-and-death model to infer gene family size across the tree. To calculate
317 the probability of the transitions in each gene family size from parent to child nodes in
318 the tree, a probabilistic graphical model was introduced. According to the conditional

319 likelihoods, we calculated the possible p-value in each lineage. A p-value of 0.05 was
320 used to identify significantly expanded/contracted families.

321 **Comparative genomics**

322 *Pfam analysis*

323 We searched the potential Pfam domains from 12 species of three groups, including
324 six ticks sequenced in this study, *I. scapularis* (Miller et al., 2018), other
325 blood-feeding arthropods *A. aegypti* (Matthews et al., 2018), *A. gambiae* (Holt et al.,
326 2002), and *Glossina* (International Glossina Genome Initiative, 2014), and a
327 non-blood-feeding outgroup *C. sculpturatus* and *P. tepidariorum* (Thomas et al.,
328 2018). Briefly, amino acid sequences of each species were scanned using all profiles
329 from Pfam database version 31 (El-Gebali et al., 2019) by hmmscan version
330 hmmer-3.1b1. The scanned results were filtered with an e-value cutoff of 1e-3, and
331 overlapping/redundant hmm matches were removed. Genes assigned to Pfam were
332 counted within each species. To identify Pfams that differed between the three groups,
333 we used a fold change >2 of the group median value as the selection criteria. Two sets
334 of Pfams were identified using the two-fold criteria: (1) Pfams that were abundant in
335 ticks compared with other blood-feeding arthropods and the outgroup; (2) The Pfams
336 showed similar abundances (fold change ≤ 2) among ticks and other blood-feeding
337 arthropods but were more abundant in these organisms than in the outgroup.

338 *Orthology analysis*

339 We performed orthology analysis for our six genomes and *I. scapularis* (Miller et al.,
340 2018) genome. First, the protein sequences of gene families with various functions,
341 including iron metabolism, carbohydrate metabolism, amino acid metabolism,
342 chemosensory functions, gustatory functions, immune functions, heme and
343 hemoglobin digestion, detoxification of xenobiotic factors, opsin-related functions,
344 lipid metabolism, oxidative stress, purine metabolism, and mechanosensation, were
345 retrieved and divided into subgroups according to their specific functions (Anderson
346 et al., 2008; Antunes et al., 2012; Bohbot et al., 2014; Cabezas-Cruz et al., 2017;
347 Della Noce et al., 2019; Eyun et al., 2017; Galay et al., 2013; Graça-Souza et al.,
348 2002; Graça-Souza et al., 2006; Gulia-Nuss et al., 2016; Hajdušek et al., 2013;
349 Hajdusek et al., 2016; Hajdusek et al., 2009; Horn et al., 2009; International Glossina
350 Genome Initiative, 2014; Iovinella et al., 2016; Josek et al., 2018; Liu et al., 2012; Liu
351 et al., 2011; Merino et al., 2011; Pal et al., 2004; Perner et al., 2016; Salem et al.,

2014; Sanders et al., 2003; Sonenshine and Macaluso, 2017; Sultana et al., 2010; Weisheit et al., 2015; Whiten et al., 2017; Winzerling and Pham, 2006). The gene families in each subgroup are shown in Table S3. Second, using collected sequences in each subgroup as query sequences, a second step of BLASTp were performed to search ortholog protein sequences in our assembled genomes (e-value <1e-5, identity $\geq 50\%$, match percentage of shorter sequence between query and subject $\geq 25\%$). For each subgroup, multiple-sequence alignments were performed using MUSCLE v3.8.31 with the default parameters, and PhyML v3.3.20190321 was employed to construct a phylogenetic tree. Based on the phylogenetic tree, genes with high reliability and the gene number for each subgroup were determined after filtering distantly related genes.

363

364 **Differential transcriptome analysis**

Unfed and fed ticks of *I. persulcatus* (3 vs 6 ticks were pooled as unfed vs fed group, respectively), *Hae. longicornis* (10 vs 10) and *R. microplus* (14 vs 24) were used for RNA extraction and transcriptome sequencing. The high-quality transcriptomic data were aligned to the reference genome using HISAT2 v2.1.0 (Kim et al., 2019). The read count of each gene was calculated for each sample by HTSeq v0.6.0 (Anders et al., 2015), and fragments per kilobase per million mapped reads (FPKM) values were then determined. DE genes were analyzed using EdgeR(v3.28.1) (Robinson et al., 2010) with false discovery rate (FDR) ≤ 0.05 and $|\log_2(\text{fold change})| \geq 1$. The dispersion parameter of DE model was estimated using the estimateCommonDisp() function in the EdgeR package. Enriched GO terms (<http://geneontology.org/>) of the DE genes were identified using Fisher's exact test in the topGO package (Alexa and Rahnenfuhrer, 2007) (FDR <0.05). Enriched pathways were tested based on the KEGG database (Kyoto Encyclopedia of Genes and Genomes, <http://www.kegg.jp/>) using clusterProfiler (Yu et al., 2012) (FDR <0.05).

379

380 **Population structure analysis**

381 ***Genomic DNA extraction and library preparation for resequencing***

All 678 adult ticks collected from the wild were thoroughly surface-sterilized, and genomic DNA for resequencing was isolated using the AllPrep DNA/RNA Mini Kit

384 (Qiagen, USA). The DNA concentration was measured using the Qubit dsDNA HS
385 Assay Kit in a Qubit® 2.0 fluorometer (Life Technologies, CA, USA). Sequencing
386 libraries were constructed using the NEBNext® Ultra™ DNA Library Prep Kit for
387 Illumina (NEB, USA) following the manufacturer's recommendations, and index
388 barcodes were added to attribute sequences to each sample. The library preparations
389 were sequenced on an Illumina NovaSeq platform (NovaSeq 6000 SP Reagent Kit),
390 and paired-end reads were generated.

391 ***Variant calling and population structure models***

392 Illumina reads of 678 tick samples were aligned to the corresponding reference
393 genome using BWA (version 0.7.17-r1188) (Li and Durbin, 2009). Variants were
394 called following the recommended GATK 4.0 pipeline (Van der Auwera et al., 2013).
395 Variant sites with quality scores ≥ 30 were kept for subsequent analysis. Based on the
396 called variants, we generated the full mitochondrial sequence of each specimen and
397 built maximum likelihood trees by MEGA7 (Kumar et al., 2016) using the GTR+F+I
398 substitution model. The tree was rooted using mitochondrial sequence of
399 *Ornithodoros hermsi* (NC_039832.1) as outgroup. For variant calling on the nuclear
400 genome, we selected variants with sufficient reads ($8 \leq$ read depth ≤ 12 , genotype
401 rate $> 70\%$), as the mean genome read coverage was $\sim 8\times$. To build the phylogenetic
402 tree of the nuclear genome, we used SNPs (minor allele frequency $\geq 5\%$) in 464
403 single-copy genes that are supposed to be conserved across tick species. An external
404 dataset from New Zealand (SRR9226159) (Guerrero et al., 2019) was added to the
405 phylogenetic analysis of *Hae. longicornis* and processed using the same pipeline as
406 that used for the six tick genomes sequenced in this study.

407 Geographical population structure was analyzed using fastSTRUCTURE (Raj et al.,
408 2014) using SNPs in the mitochondria. For each tick species, fastSTRUCTURE
409 was run for K (number of ancestral populations) from 2 to 10 with fivefold
410 cross-validation. The fastSTRUCTURE model selected the best value of K=2 for
411 *Hae. longicornis* and K=3 for *R. microplus* by maximizing the marginal likelihood of
412 the fastSTRUCTURE model. However, to enable fair comparison between the two
413 species, we chose a more detailed population structure (K=5), as shown in Fig. 3B.
414 The population structure was plotted using Pophelper (2.3.0) package (Francis, 2017)
415 and CLUMPAK (<http://clumpak.tau.ac.il>) (Kopelman et al., 2015). To measure
416 population differentiation, we calculated the F_{ST} between all pairs of populations in

417 each province for *Hae. longicornis* and *R. microplus* based on the SNPs within their
418 mitochondria. First, the numerator and denominator of the Hudson F_{ST} estimator were
419 calculated for each SNP. Then, across all SNPs, the ratio of the average numerator
420 and denominator was calculated as the final F_{ST} estimator between two populations
421 (Bhatia et al., 2013). F_{ST} calculations were conducted using the python scikit-allele
422 package (version 1.2.1, <https://github.com/cggh/scikit-allele>) (Alistair Miles and
423 Harding, 2016).

424 ***Copy number variation detection in the genomes of R. microplus***

425 We found that *R. microplus* can be clustered into three major clades. First, genes
426 with read counts > 2 in at least half of the samples were selected to calculate the copy
427 number changes in the three clades of *R. microplus*. Second, the read counts of the
428 genes were normalized to gene length. In each sample, the normalized gene read
429 count was divided by the median of all genes to calculate the fold change (cf) of the
430 copy number (CN). Third, the cfCN of the gene was compared with each sample
431 median cfCN by the function of t.test (paired=T) in R to calculate the significance in
432 each clade. The p-values were adjusted for multiple testing correction using
433 Benjamini-Hochberg correction as a function of p.adjust (method = "BH") in R. In
434 each clade, genes with adjusted p-values < 0.001 and median cfCN ≥ 2 were referred
435 to as increased CN genes. According to the gene annotation results, GO enrichment
436 analysis was limited to the 4-level GO terms and implemented by a hypergeometric
437 test with the phyper() function in R. The enrichment p-value was adjusted by the
438 p.adjust function (method = "BH") in R.

439

440 **Metagenomic analysis and pathogen detection**

441 Tick sequences were filtered by SAMtools (version 0.9.24) (Li et al., 2009) after
442 mapping the reads of 678 specimens to tick genomes by BWA (version: 0.7.17), and
443 all unmapped reads were retained for subsequent analysis. Taxonomic classification
444 was performed by aligning the filtered reads to the NR database using DIAMOND
445 (version 0.9.24, parameters: -f 102 -top 10) (Buchfink et al., 2015). To estimate the
446 relative abundances of different bacterial species, we extracted all taxonomic IDs
447 according to the NCBI taxdump
448 (<ftp://ftp.ncbi.nlm.nih.gov/pub/taxonomy/taxdmp.zip>) (*Rickettsia*: TaxID780,
449 *Anaplasma*: TaxID768, *Ehrlichia*: TaxID 943, *Borrelia*: TaxID 138, *Babesia*: TaxID

450 5864, *Theileria*: TaxID 5873, *Francisella*: TaxID262, *Bartonella*: TaxID 773,
451 *Coxiella*: TaxID 776, *Hepatozoon*: TaxID 75741, *Toxoplasma*: TaxID 5810,
452 *Candidatus Neoehrlichia*: TaxID 467749). Sequence similarity (>70%) were used as
453 the threshold to screen the alignment results. The classification of species pathogenic
454 to human or not to was based on currently available literatures. After normalizing all
455 classified sequences to 100,000 microbial reads, the relative abundance of each
456 pathogen was estimated by calculating the sequences classified to this species. We
457 also adopted a widely-used tool, Metaphlan2 (Segata et al., 2012), for metagenomic
458 taxonomic profiling, but only a very limited number of pathogens could be found in
459 different tick species. Therefore, we used the results of NR-blast-based method for
460 downstream analyses.

461 **Epidemiological data search strategy**

462 We searched PubMed and ISI (Web of Science) for articles published in English, and
463 WanFang database, China National knowledge Infrastructure, and Chinese Scientific
464 Journal Database of articles published in Chinese between Jan 1, 1980 and April 30,
465 2020. We used the following search terms: “tick-borne disease”, “tick-borne
466 zoonosis”, “tick-borne zoonotic disease”, “tick-associated agent”, “tick-associated
467 microbe”, and “China”. The articles about tick-borne viral diseases were excluded.
468 We did a secondary manual search of the references cited in these articles to find
469 relevant articles. We investigated all the articles related to detection, identification, or
470 case reports of tick-borne microbes in human beings. Each case was geo-referenced to
471 a Chinese map in the prefecture-level with ArcGIS 10.2 (Johnston et al., 2004) (ESRI,
472 Redlands, CA, USA) according to the patient’s living location or visiting hospital.

473

474 **QUANTIFICATION AND STATISTICAL ANALYSIS**

475 Quantification and analysis procedures of genome, transcriptome and metagenome
476 data were provided in the relevant sections of Method Details. To test the correlation
477 between gene family size and standard deviation of gene expression (fold change),
478 Spearman’s rank correlation coefficient was calculated. Fisher’s exact test was used to
479 test the enrichment of down-regulated genes in P450 families. Mann–Whitney U test
480 was used to compare the prevalence of pathogen in different faunal or geographical
481 regions. Kruskal-Wallis test was used to compare the positive rate of *Rickettsia* in
482 different subtypes of ticks. All these tests were performed in R environment and p

483 value below 0.05 was considered statistically significant. For all analyses, the
484 meaning and value of n and the measures of center, dispersion, and precision used can
485 be found in the relevant main text or in Method Details.

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