International beeswax trade facilitates small hive beetle invasions
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International trade can facilitate biological invasions, but the possible role of beeswax trade for small hive beetles (SHBs), Aethina tumida Murray (Coleoptera: Nitidulidae) is poorly understood. SHBs are parasites of social bee colonies native to sub-Saharan Africa and have become an invasive species. Since 1996, SHBs have established in all continents except Antarctica. Here, we combine mitochondrial DNA analyses (COI gene, N = 296 SHBs, 98 locations) with previously published beeswax trade data (FAO) for 12 confirmed SHB invasions. Our genetic data confirm previous findings and suggest novel SHB African origins. In nine out of 12 invasion cases, the genetic and beeswax trade data match. When excluding one confirmed pathway (bee imports) and two cases, for which no FAO data were available, the genetics and beeswax trade data consistently predict the same source. This strongly suggests that beeswax imports from Ethiopia, South Africa, Tanzania and the USA, respectively, have mainly been responsible for the past invasion success of this beetle species. Adequate mitigation measures should be applied to limit this key role of beeswax imports for the further spread of SHBs. Combining genetics with trade data appears to be a powerful tool to better understand and eventually mitigate biological invasions.

Invasive species are a major threat to food security and conservation of natural biodiversity 1–2. Typically, such biological invasions follow a jump dispersal pattern reflecting human assisted transmission often across the entire globe 3–4. In particular, there is growing evidence that international trade can play a major role in spreading alien species 5–6, but in many cases the actual trading routes and goods involved remain poorly understood. Aiming at mitigating the global impact of invasive species, a better understanding of the role of specific international trading routes as transmission pathways appears therefore crucial. Since genetic tools are routinely used to reconstruct routes of invasion 7, one feasible approach seems to be a combination of genetics data with public data on international trade 8 for specific cases of invasive species. This also holds true for invasive pests and pathogens associated with managed honey bees, Apis mellifera Linnaeus (Hymenoptera: Apidae) 9.

One invasive species associated with honey bees is the small hive beetle (SHBs), Aethina tumida Murray (Coleoptera: Nitidulidae) 10. SHBs are parasites of honey bee colonies native to sub-Saharan Africa 11,12. Larval and adult SHBs feed on honey, pollen, bee larvae, as well as dead or live adult bees 13–15. SHB larvae can cause severe damage to honey bee colonies 14, often resulting in the full structural collapse of the entire nest 12. Since 1996, SHBs have emerged as an invasive species and have now established on all continents except Antarctica 10,16–18. At present, it is still unclear whether a single or multiple introductions into the USA have occurred 19–21. SHBs in Australia appear to have a different origin than beetles in North America and the initial North American beetles shared the same source 11. The outbreaks in Quebec (Canada) 19 appear to have originated from the USA 16,21 and the SHBs confirmed in Alberta (Canada) in 2006 16 were probably a novel introduction via Australian package bees 21. SHBs confirmed in the Calabria region (Italy) in 2014 21 appear to have been introduced from Cameroon 11. This introduction into Calabria was followed by man-mediated migration to Sicily 11. SHBs detected and originally reported to be widespread in Egypt 13 were not confirmed by a latter extensive survey 20, thereby suggesting a possible false positive diagnosis in the first place. In 2004, SHB larvae were intercepted in a shipment of queens from the USA to Portugal, but pest establishment was successfully prevented by rigorous sanitation 27. Recently, introductions into Hawaii (2010) 10 and Brazil (2015) 16, were traced back to South Africa 28 and the USA 29, respectively. None of the confirmed SHB introductions reported in Portugal (2004) 27, Jamaica (2005) 30, Mexico (2007) 10, and

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K (Hd) haplotypes were detected, displayed the highest genetic diversity (pi ± = 0.00256, 2.522)

Table 1. Number of analysed individuals (N), polymorphic sites (S), parsimony informative sites (PIS), number of haplotypes (h), haplotype diversity (Hd), nucleotide diversity (pi) and average number of differences (K) in mt-DNA sequences computed for Aethina tumida collected from five geographical regions.

| Regions | Sample | Polymorphic | Parsimony | Number of | Haplotype | Nucleotide | Average number of |
|---------|--------|-------------|-----------|------------|-----------|-------------|------------------|
|         | size (N) | sites (S) | informative sites (PIS) | haplotypes (h) | diversity (Hd) | diversity (pi) | nucleotide differences (K) |
| Australia | 30 | 19 | 10 | 4 | 0.510 ± 0.109 | 0.00285 ± 0.00076 | 2.522 |
| Americas | 115 | 16 | 13 | 5 | 0.301 ± 0.056 | 0.00183 ± 0.00039 | 1.518 |
| Europe | 10 | 45 | 43 | 4 | 0.889 ± 0.075 | 0.02070 ± 0.00226 | 20.289 |
| Asia | 21 | 36 | 36 | 2 | 0.381 ± 0.101 | 0.01404 ± 0.00370 | 13.714 |
| Africa | 120 | 105 | 73 | 80 | 0.983 ± 0.005 | 0.02490 ± 0.00148 | 13.646 |
| All | 296 | 118 | 89 | 90 | 0.857 ± 0.018 | 0.02280 ± 0.00103 | 12.450 |

Cuba (2012)8, El Salvador (2013)10, Nicaragua (2014)10, Philippines (2014)10, South Korea (2016)12 and Mauritius (2016)18 have so far been studied genetically.

In its new ranges, SHBs can have a strong impact on local honey bees and can also infest colonies of other social bees10 as well as nests of solitary bees45. It appears therefore high time to slow down the ongoing spread of SHBs until better mitigating options will become available31. Central to that is clearly the identification of the actual transmission means. So far, a whole range of actual and possible transmission pathways has been identified, with import of bees and bee products being the major confirmed contributors so far10. Accordingly, countries have put in place legislation and quarantine measures to reduced chances for entry of SHBs as well as other bee pests and diseases37. While most of these efforts have focused on the control of living bee imports, hive products have so far been rather neglected, even though one introduction of SHBs into Canada could be traced back to beeswax imports22. Therefore, it seems obvious that at least some additional SHB introductions may have occurred in association with international trade of beeswax.

Beeswax is a creamy coloured substance used by worker bees to build the comb that forms the structure of their nest38. It is widely used by humans, e.g. in cosmetics, pharmaceutical preparations and food production39. In particular, beekeepers use large quantities of beeswax for making beeswax comb foundation42,43. Beeswax can significantly range in quality34 and respective international trade can occur in various forms, ranging from non-processed “crude” wax over foundations to highly processed ones (i.e. melted and purified)32,35. For beeswax in the form of “honeycomb”, there are trade regulations with respect to SHBs available38. Either the beeswax originates from a country free of A. tumida infestation or precautions, such as “thoroughly” cleaning, have been taken to prevent infestation/contamination with SHB life stages36. However, for processed beeswax there are currently no trade restrictions with respect to SHBs4,36, probably because wax processing is assumed to kill all SHB life stages (e.g. melting). Interestingly, countries in sub-Saharan Africa, the endemic range of SHBs4,12, only participate in international trade with the sale of beeswax and honey4, the latter not being confirmed as a SHB transport means so far10. Nevertheless, correlations between global beeswax trade routes4 and confirmed SHB introductions with respective areas of origin have yet to be taken into consideration. This is mainly due to few data sets existing on molecular markers tracing back the origin and possible transmission pathways of SHB19-21,24,28,29. These previous studies could obviously not have covered all confirmed introductions so far and/or only considered comparatively few native locations of SHBs.

The aim of this study is to shed further light on possible SHB invasion pathways to mitigate its further spread. For that purpose, we here combine for the first time genetic data of 12 confirmed SHB invasions with FAO data on international trade of beeswax7. If the latter constitutes a major means of SHB transport globally, we suspect that in the majority of cases genetic traces of origin match with respective data on the imports of beeswax.

Results
Mitochondrial COI haplotypes. Analysis of mitochondrial COI DNA sequence data from 296 individuals revealed 90 unique haplotypes representing 0.857 ± 0.018 and 0.02280 ± 0.00103 (mean ± SD) haplotype and nucleotide diversity, respectively. Among the sequences analysed, 118 segregating (polymorphic) sites were detected (Table 1), 89 of which were parsimony-informative and 29 were singleton sites.

81 haplotypes were found in a single country (Australia, Benin, Burkina-Faso, Burundi, DR Congo, Ethiopia, Italy, Kenya, Madagascar, Malawi, Nigeria, Central African Republic, South-Sudan, Sudan, Tanzania, Togo, Uganda, and the USA) while the nine others were present in at least two different countries. The most widely distributed haplotype (Hap_11) was found with a global frequency of 35.1% and was detected in Africa (Tanzania and the Americas (Brazil, Canada, Cuba, Jamaica, Mexico, and USA) (Fig. 1). The two other most frequent haplotypes were Hap_1 and Hap_23 with global frequencies of 11.5 and 4.7%, respectively. Whereas Hap_1 occurred in association with international trade of beeswax8, if the latter constitutes a major means of SHB transport globally, we suspect that in the majority of cases genetic traces of origin match with respective data on the imports of beeswax.
Phylogenetic reconstruction. The Bayesian phylogenetic reconstruction based on COI sequences separates the samples into six different clades, labelled A, A1, A2, B, B1, and B2 (Fig. 2). These different clades can be grouped in two main African populations labelled Pop1 and Pop2 (Fig. 2). Pop1 is consisted of Western (Benin, Nigeria, Togo, and Burkina Faso) and Central African (Cameroon, Central African Republic, and Democratic Republic of Congo) populations, but also some Eastern regions (Uganda, South Sudan, Sudan, Burundi, and Ethiopia). Pop2 includes some Eastern (Tanzania, Kenya, South Sudan, and Madagascar), Southern (South Africa, Malawi, and Zimbabwe) and Western African (Burkina Faso) populations. Pop2 shows 93% similarity with Burkina Faso samples from Fada-Ngourma. In Pop1 and Pop2 different clades are well-supported by posterior probabilities (>90%).

Most of the Italian specimens belong to clade A, mainly A1 and A2. The other Italian specimens, which are not included in clade A, show similarity with each other. In clade A1, the Italian samples (ItalyCal8, and ItalyCal9) cluster with one Ugandan sample (UgandaAl2) with a Bayesian posterior probability of 86%. In the same clade, other Italian samples (ItalyCal2, Italy12, Italy3, Italy4 2015) show a high similarity with the Portuguese sample (96%). In clade A2, Italian samples, which contains the first detected specimen in Calabria (ItalyCal1_KT380624), cluster with Ethiopian samples with a Bayesian posterior probability of 97%.

Clade B1 includes SHB from the USA (USA2, US NthAm2), Mexico, Jamaica, Cuba, Canada (British Columbia, Quebec) and Brazil. Interestingly, within this clade all samples cluster with Kenyan, Zimbabwean, and Tanzanian samples with a Bayesian posterior probability of 95%. In clade B, all samples from Australia, USA (USA1, USNthAm1), Canada (Alberta, Ashville, and Concordia), Costa Rica, Hawaii and South Korea show similarity with Tanzanian and South African samples.

FAO data. We considered beeswax imports from countries with known native or introduced SHB populations up to two years before the introduction into new ranges (Table 2). Based on prior invasion history (USA, Australia), this appears to be a suitable time window. SHB populations probably have to increase first to cause clinical symptoms of infestation, thereby fostering their detection by beekeepers.

Beeswax imports into the Americas. From 1994 to 1996, Tanzania was the first African beeswax trade partner to the USA. The USA imported 39, 29 and 70 tons (=T) in 1994, 1995 and 1996 respectively. From 2000 to 2002, Canada imported 363.84 T of beeswax per year, of which ~87% was from the USA. During this period, Canada did not import any beeswax from Africa. In the suspected period of introduction of SHBs into Jamaica (i.e. 2003–2005), FAO data clearly show that this country imported 3 T of beeswax only in 2003 and from the USA. From 2005 to 2007, the USA and Germany (no confirmed SHB cases) were the main beeswax suppliers of Mexico with an average of 81.33 and 75.33 T per year respectively (with a maximum import of 108 and 78 T in 2007). Costa Rica is a small beeswax importer with an average of 3.33 T per year from 2012 to 2014. During this period, Costa Rica imported only from two countries: Argentina (2 T; no confirmed SHB cases) and USA (1.33 T). In 2012 and 2013, Costa Rica imported beeswax exclusively from the USA with an average of 1 T per year. The
USA, after India (no confirmed SHB cases), was the second largest beeswax exporter to Brazil with an average of 5.67 T per year from 2013 to 2015. There are no data available on the quantity of beeswax imported to Cuba.

**Beeswax imports into Australia.** For the period 1999–2001, data are not available in the FAO database. However, from 2005 to 2016 South Africa was the second largest beeswax supplier (after China) to Australia with an average of 80.6 T per year.

**Beeswax imports into Italy, Portugal and South Korea.** From 2012 to 2014, Italy imported from Asia (283.33 T), Europe (242.01 T), Africa (20 T) and the Americas (2 T). In 2012, Italy imported beeswax from Africa exclusively from Ethiopia (18 T). During this period, Italy did not import any beeswax from Cameroon. From 2002 to 2004, Portugal exclusively imported from Europe: Spain (20 T), Germany (5.67 T), UK (3.67 T), France (0.33 T) and Netherlands (0.33 T). None of these countries had any confirmed SHB cases. Portugal did not import any beeswax from the USA. South Korea imported beeswax from Americas exclusively from the USA with 11 and 16 T in 2015 and 2016 respectively.

**Combination of genetics and beeswax trade data.** To shed light on the role of beeswax imports for the spread of SHBs, we combined our genetic data with beeswax trade data (FAO) for 12 confirmed SHB invasions. For that purpose, we considered imports from countries in the endemic range of SHBs and with prior confirmed cases. In nine out of 12 invasion cases, the genetics and beeswax data matched, i.e. countries of origin suggested by genetics data matched with beeswax trade records (Table 2, Fig. 3).

**Discussion**

Our genetic data combined with FAO data strongly suggest that imports of beeswax play a previously underestimated key role in propagating SHBs globally. Indeed, in nine out of 12 cases, beeswax trade data and our genetic data match. When excluding one confirmed introduction pathway (bee imports, Portugal) and the two cases, for which no FAO data were available, the genetics and FAO trade data consistently predict the same source of...
which is a logic consequence of genetic bottlenecks due to invasion events. Among these areas, the Americas are also in line with recent reports on the origin of SHBs in Hawaii and Brazil, suggesting that SHBs discovered in these areas could have been introduced from the USA. In nine out of 12 invasion cases, the genetics and beeswax data match as indicated in bold. In the Portugal case, import of queen bees was shown and for two further cases, beeswax trade data are not available.

### Table 2. Combined genetics data (this study) and beeswax trade data (FAO) for 12 confirmed small hive beetle (SHB) invasions.

| Invasion | Genetics | Beeswax | Origin |
|----------|----------|---------|--------|
| USA (mainland) | Kenya, Tanzania, Zimbabwe, South Africa | Tanzania | Tanzania |
| Mexico | Kenya, Tanzania, Zimbabwe | USA (no SHBs) | USA |
| Jamaica | Kenya, Tanzania, Zimbabwe | USA | USA |
| Cuba | Kenya, Tanzania, Zimbabwe | — | USA |
| Canada | Kenya, Tanzania, Zimbabwe, South Africa | USA | USA |
| Brazil | Kenya, Tanzania, Zimbabwe | USA, India (no SHBs) | USA |
| Costa Rica | Tanzania, South Africa | USA | USA |
| USA (Hawaii) | Tanzania, South Africa | — | USA, South Africa |
| South Korea | South Africa, Tanzania | USA | USA |
| Australia | South Africa, Tanzania | South Africa, China (no SHBs) | South Africa |
| Portugal | Unknown (same as Italy) | Spain, Germany, UK, France, Netherlands (all no SHBs) | USA |
| Italy | Unknown, Ethiopia, Uganda | Various countries, incl. Ethiopia | Ethiopia |

SHBs. This creates an urgent demand to better consider the role of international beeswax trade for the spread of this invasive species.

The results confirm the African origin of this invasive species and that SHBs found in Australia and Canada are most likely originating from South Africa and USA respectively. Our data are also in line with recent reports on the origin of SHBs in Hawaii and Brazil, suggesting that SHBs discovered in these areas could have been introduced from the USA. In nine out of 12 invasion cases, the genetics and beeswax data match as indicated in bold. For the Portugal case, import of queen bees was shown and for two further cases, beeswax trade data are not available.

The highest COI diversity was found in Africa with 80 out of 90 identified mitochondrial haplotypes. This high genetic diversity confirms previous findings as well as the African origin of SHBs. Furthermore, the phylogenetic reconstruction suggests the existence of at least two large African populations, in which several haplotypes are present. The population Pop1 is more heterogeneous than population Pop2, thereby suggesting that the large region (West, Central and Eastern Africa) is probably the SHBs nucleus of origin. The comparatively high mobility of adult SHBs and resulting panmixia could explain the non-grouping of the different sub-populations by country in the phylogenetic tree. In addition, Pop2 showed a considerable similarity with samples from Burkina Faso (Fada-Ngourma) belonging to Pop1. This population may have invaded the southern and eastern regions. This could explain the homogeneity and low diversity observed within population Pop2. Based on our data, it is not clear if these two SHB populations actually overlap or not. A more exhaustive sampling in the neighbouring geographic areas between the two populations and more genetic markers will be required to shed light on the causes for this observed divergence.

In the invaded areas (Americas, Australia and Asia) the COI diversity was low when compared to Africa, which is a logic consequence of genetic bottlenecks due to invasion events. Among these areas, the Americas showed the lowest genetic diversity probably reflecting comparatively fewer introduction events. Alternatively, but not mutually exclusive, an original genetic diversity may have been lost secondarily.

Our results clearly show that one of the haplotypes (Hap_11) present in the Americas (USA, Brazil, Canada, Cuba, Jamaica and Mexico) is the same as that found in Tanzania. Combining our genetics data with the FAO data base, the most likely invasion scenario is that the first specimens found in South Carolina in November 1996 were introduced via infested beeswax containers from Tanzania. Since it can take years for mass reproduction with obvious clinical symptoms to occur, SHBs could have easily been introduced one to two years before their confirmation. Indeed, the USA imported 39 and 29 T of beeswax respectively in 1994 and 1996 from Tanzania. These SHBs almost certainly crossed the USA border to invade Quebec in 2008 (<25 km).

Our data do not confirm the previous finding of an Australian origin of Canadian SHBs. Most likely, the SHBs introduced from Australia back then were not able to establish a local population in their northern distribution limit in Canada (but see Essex county, Ontario for an exception). It appears therefore most likely that novel introductions from the USA have occurred in the meantime. Similarly, according to our data, the USA is the most probable source of SHBs in other countries of the Americas (Brazil, Cuba, Jamaica and Mexico), which is also reflected in respective beeswax trade activities.

The results of the haplotype analysis show that the second haplotype (Hap_23) found in the USA is also present in Canada, Costa Rica and South Korea. According to FAO statistics, Costa Rica and South Korea have not imported beeswax from Africa. However, both countries have mostly imported beeswax from the USA, thereby supporting that the SHBs detected in Costa Rica and more recently in South Korea originated from the USA.
introduction from the USA via bee import\textsuperscript{7}, but our data suggest the same African origin as for Italy; yellow lines suggested by genetics; dashed red lines (24) 2016, Guanacoste, Costa-Rica (this study); (red line into Portugal from any country with known SHB populations\textsuperscript{8}. Indeed, the beeswax was not imported with some Italian samples from Calabria, thereby strongly suggesting the same yet unidentified African origin are among the largest producers of beeswax with respective productions of 5,542 and 1,308 T in 2016\textsuperscript{8}. Therefore, second introduction into the same region most likely originated from Uganda (clade A1). Ethiopia and Uganda (1) 1996, Charleston, South Carolina, USA, (2) 2000, Itay-Al-Baroud, Egypt, (3) 2001, Richmond, NSW, Australia, (4) 2002, Manitoba, Canada, (5) 2004, Lisbon, Portugal, (6) 2005, Jamaica (2010), (7) 2006, Alberta and Manitoba, Canada, (8) 2007, Coahuila, Mexico, (9) 2007, Kununurra, North Australia, (10) 2008, Perth Australia, (11) 2008, 2009, Quebec, Canada, (12) 2008, 2013 Ontario, Canada, (13) 2010, Panàwèa, Big Island, Hawaii, (14) 2012, Cuba, (15) 2012, Naracoorte in Eastern South Australia; (16) 2013, El Salvador, (17) 2014, Nicaragua), (18) Sovereto, Calabria, Italy, (19) 2014, Renmark, Australia, (20) 2014, Lupon, Philippines, (21) 2015, Piracicaba, São Paulo State, Brazil\textsuperscript{16}, (22) 2016, Miryang-si, GN, Korea\textsuperscript{17}, (23) 2016, Mauritius island\textsuperscript{18}, (24) 2016, Guanacoste, Costa-Rica (this study); (red line = our genetic data; solid red lines = invasion pathways suggested by genetics; dashed red lines = origin from Australia and/or South Africa; dotted red line = confirmed introduction from the USA via bee import\textsuperscript{1}, but our data suggest the same African origin as for Italy; yellow line = matching beeswax trade data\textsuperscript{a} with our genetic data).

Alternatively, but not mutually exclusive, the SHBs in Costa Rica simply reflect the ongoing SHB invasion front in Central America\textsuperscript{10}.

Our results confirm previous findings that SHBs found in Australia are most likely originating from South Africa\textsuperscript{21}. Interestingly, the SHBs found in Hawaii are also similar to South African haplotypes\textsuperscript{25}, thereby implying that Australian and Hawaiian beetles both originated from the same African source population. Alternatively, but not mutually exclusive, the Hawaiian SHBs may have originated from the Australian populations. Unfortunately, there are no FAO data available concerning beeswax imports from South Africa to Australia and to Hawaii, respectively.

When compared to the Americas, SHB populations found in Europe show a higher haplotype diversity, thereby supporting multiple introduction events\textsuperscript{24}. Phylogenetic analysis shows that specimens from Italy belong to a different clade than the one in the USA and Australia. Our data therefore support earlier results that the occurrence of SHB in Italy is due to an independent introduction from Africa and not from the USA or Australia\textsuperscript{10}. The phylogenetic tree clearly shows two clades, A1 and A2, to which most SHB specimen from Italy belong, therefore, suggesting two separate introductions into the Calabria region. The first specimen detected in September 2014 in Gioia Tauro in the Calabria region\textsuperscript{23} most likely originated from Ethiopia (clade A2). A second introduction into the same region most likely originated from Uganda (clade A1). Ethiopia and Uganda are among the largest producers of beeswax with respective productions of 5,542 and 1,308 T in 2016\textsuperscript{8}. Therefore, SHBs detected in the region of Calabria (only a few kilometres from the port of Gioia Tauro) most probably came from infested wax containers from Ethiopia and/or Uganda. Interestingly, Italy however only imported beeswax from Ethiopia until 2012\textsuperscript{8}. This suggests that SHBs were probably present in Calabria at least two years before their official confirmation similar to the USA and Australia\textsuperscript{15}. Moreover, it also shows that those SHBs in Calabria most likely originating from Uganda must have arrived via another yet unidentified pathway, e.g. bee imports as in the case of Portugal 2004\textsuperscript{27}. Fascinatingly, the Portuguese sample shows a very high similarity (96%) with some Italian samples from Calabria, thereby strongly suggesting the same yet unidentified African origin (assuming that SHBs were not present in Italy back in 2004). The FAO data show that beeswax was not imported into Portugal from any country with known SHB populations\textsuperscript{8}. Indeed, the \textit{A. tumida} larvae were detected in a shipment of queens from Texas (USA)\textsuperscript{25}. Since the Portuguese sample therefore unequivocally originated from the USA, this strongly implies that at least a second, previously not detected, introduction into the USA must have occurred independently from the Tanzanian one (see above). Therefore, we can here clarify the question whether a single or multiple introductions into the USA have occurred\textsuperscript{21}. Based on our genetic data, it seems safe to assume that at least two independent introductions into the USA must have occurred.

The combination of our genetic and beeswax trade data\textsuperscript{a} enabled light to be shed on a previously overlooked key factor for SHB spread. Since the origin of the Portugal SHB introduction in 2004 can unambiguously be traced back to shipment of queen bees from the USA\textsuperscript{36}, only for 11 out of the 12 confirmed introductions, which were investigated here, sources could not certainly be traced back so far. Correlations are not causations, however, given that two independent data sets match in their correlations (genetics and FAO), this further supports a causal relationship. Indeed, in the nine remaining cases, for which both genetic and FAO data were available,
both data sets consistently suggest the same country of origin. This strongly suggests that international beeswax trade is the major factor explaining the past invasion history of small hive beetles. It seems therefore most unfortunate that nobody paid really attention up to now on the role of beeswax for SHB spread. Interestingly, the international transport of beeswax usually occurs via ships\textsuperscript{35}\textsuperscript{40} and some SHB introductions so far occurred near major harbors (Charleston, USA\textsuperscript{35}; Giria Tauro, Italy\textsuperscript{40} or on islands (Jamaica, Hawaii, Cuba, Philippines)\textsuperscript{40}. Since the international shipping industry is responsible for the carriage of \textasciitilde 90\% of world trade\textsuperscript{40}, it appears unsurprising that the surroundings of harbours were starting points of several SHB invasions. It seems therefore prudent to take into account beeswax trade for adequate mitigation measures to further limit the spread of this pest species.

Ideally, beeswax shipments are not contaminated prior to sending. However, this appears notoriously difficult in regions with SHB populations due to the highly mobile and cryptic behaviour of the adult beetles\textsuperscript{35,40}. Regardless of the actual beeswax processing stage\textsuperscript{33,34}, adult SHBs are therefore very likely to invade and successfully hide in any beeswax trade package. Indeed, even small commercial honey bee queen cages, which are obviously much easier to visually screen by the beekeepers compared to containers and large boxes, were already responsible for two independent SHB invasions (Portugal\textsuperscript{22}; Northern Territory, Australia\textsuperscript{40}). It therefore appears inevitable that adult SHBs can and eventually will invade any beeswax trade containers during packaging. Considering adult SHB survival under starvation (at least 12 days without any food and water; Peter Neumann, unpublished data), chances for adult SHB invasion of and hiding\textsuperscript{39} in beeswax trade containers as well as the impact of this invasion on bees in the new ranges, it seems as if beeswax imports from countries with known SHB populations should be banned. Alternatively, rigorous quarantine measures in the importing countries might help, but in light of our rather fragmentary knowledge of SHB physiology, this appears risky.

In conclusion, our genetic results (1) confirm the African origin of SHBs, (2) show a high genetic diversity in Africa, (3) suggest two main native populations, (4) confirm previous findings and (5) suggest novel origins. Since our genetic data match well with data on international trade of beeswax\textsuperscript{4}, a closer investigation of international beeswax trade may enable to slow down the further spread of this invasive species. In general, it appears worthwhile to use this combination of genetics and public trade data in more cases of invasive species.

Methods

Sample collection. Adult SHBs (N \(=\) 1542) were manually sampled\textsuperscript{41} from 101 naturally infested A. mellifera colonies from 98 locations both in the native range of SHBs in sub-Saharan Africa (Benin, Togo, Nigeria, Burkina-Faso, Central African Republic, Democratic Republic of Congo, Uganda, Sudan, South Sudan, Ethiopia, Kenya, Tanzania, Malawi, Burundi, Madagascar and South Africa) as well as from 11 confirmed SHB introductions (USA, Australia, Canada, Philippines, Mexico, Jamaica, Hawaii, Cuba, Italy, Brazil and South Korea). Samples from Guanacaste, Costa Rica taken in 2016 by J. Pettis were also included because detailed morphometric analysis\textsuperscript{41} confirmed that the samples do belong to A. tumida. The last remaining SHB larvae from the Portugal introduction in 2004\textsuperscript{41} was also included. Aethina convolor specimen collected in Australia were used as outgroup. Please refer to Table 3 for details of the sampling. All samples were preserved in 70\% Ethanol, transported at room temperature and stored at \(-20\textdegree\text{C}\) in a laboratory until further analyses.

Genomic DNA extraction. A modified protocol from Neumann et al.\textsuperscript{35} was used. In brief, all samples were crushed individually in 2 mL microcentrifuge tubes containing 5 mm metal beads and 100 \(\mu\text{L}\) TN buffer (10 mmol/L Tris, 10 mmol/L NaCl, pH 7.6). Crushed samples were homogenized with a Tissuemizer (QiAGEN, Hombrechtikon, Switzerland) for 25 sec at 20 \(\text{Hz/}text{sec}\) frequency using a Qiagen Retsch\textsuperscript{6} MM 300 mixer mill (Thermo Fisher Scientific, Zurich, Switzerland) and centrifuged for 60 sec at 2000 rpm. 50 \(\mu\text{L}\) of supernatant was taken from the homogenate and used for DNA extraction using innuPrep DNA Mini Kit (Analytik Jena, Jena, Germany) by following the manufacturer’s recommendations. The DNA yield and purity of each sample was checked using a Spectrophotometer Thermo Scientific\textsuperscript{79} Nanodrop 2000 (Axon lab, Baden-Dättwil, Switzerland).

PCR amplification and COI gene sequencing. A fragment of Cytochrome C Oxidase subunit I (COI) gene was amplified using primers AT1904S (5\textsuperscript{'}-GGTGGATCTTCAGTTGATTAGC-3\textsuperscript{'}\text{)} and AT2953A (5\textsuperscript{'}-TCAGCTGGGGATATAATTG-3\textsuperscript{'}\text{)} for SHB specimens\textsuperscript{39}. Additionally, another COI fragment was also amplified with LCO1490 (5\textsuperscript{'}-GGTCACAAATGATAAGATATGG-3\textsuperscript{'}\text{)} and HCO2198 (5\textsuperscript{'}-TAACTTCAGGGTGACCAA AAAATCA-3\textsuperscript{'}\text{)} for A. convolor specimen\textsuperscript{41}. PCR tests were carried out in 25 \(\mu\text{L}\) (15.88 \(\mu\text{L}\) millipore water, 5 \(\mu\text{L}\) 5x reaction buffer, 1 \(\mu\text{L}\) of (0.4 \(\mu\text{mol/L}\) each primer (reverse and forward), 0.125 \(\mu\text{L}\) (0.63 units) Taq DNA polymerase, 2 \(\mu\text{L}\) tenfold-diluted DNA) in a Biometra\textsuperscript{10} Thermal Cycler (Thermo Fisher Scientific, Switzerland) as follows: 95\textdegree\text{C} for 2 min, 35 cycles of 95\textdegree\text{C} for 20 sec, 56\textdegree\text{C} for 30 sec, 72\textdegree\text{C} for 30 sec, and 72\textdegree\text{C} for 2 min. Positive and negative controls were included in each PCR (DNA of previously identified SHB specimen from Nigeria (based on both morphometrics and DNA sequence data) or millipore water). Aliquots of PCR products were run by electrophoresis on a 2\% (w/v) agarose gel.

The amplicons were purified with ExoSAP-IT\textsuperscript{TM} (Thermo Fisher Scientific, Zurich, Switzerland) and sequenced in both directions (forward and reverse). For a better coverage of the 1091 bp fragment to be sequenced, two internal primers, Aet-int-F (5\textsuperscript{'}-CTTCCTGCTACAATTAATTGCG-3\textsuperscript{'}\text{)} and Aet-int-R (5\textsuperscript{'}-TTTGTTGATCATGAAGATTCGC-3\textsuperscript{'}\text{)} were added to the sequencing reactions\textsuperscript{42}. All samples were analyzed using the Big Dye\textsuperscript{8} Terminator v3.1 cycle sequencing kit (Thermo Fisher Scientific, Zurich, Switzerland). Sequencing was performed using a 96-capillary ABI PRISM\textsuperscript{8} 3730XL Genetic Analyzer (Thermo Fisher Scientific, Zurich, Switzerland). The nucleotide sequences were deposited in the GenBank database under accessions MK024945-MK025231.

Please refer to Table 3 for details of the sampling. All samples were preserved in 70\% Ethanol, transported at room temperature and stored at \(-20\textdegree\text{C}\) in a laboratory until further analyses.

PCR amplification and COI gene sequencing. A fragment of Cytochrome C Oxidase subunit I (COI) gene was amplified using primers AT1904S (5\textsuperscript{'}-GGTGGATCTTCAGTTGATTAGC-3\textsuperscript{'}\text{)} and AT2953A (5\textsuperscript{'}-TCAGCTGGGGATATAATTG-3\textsuperscript{'}\text{)} for SHB specimens\textsuperscript{39}. Additionally, another COI fragment was also amplified with LCO1490 (5\textsuperscript{'}-GGTCACAAATGATAAGATATGG-3\textsuperscript{'}\text{)} and HCO2198 (5\textsuperscript{'}-TAACTTCAGGGTGACCAA AAAATCA-3\textsuperscript{'}\text{)} for A. convolor specimen\textsuperscript{41}. PCR tests were carried out in 25 \(\mu\text{L}\) (15.88 \(\mu\text{L}\) millipore water, 5 \(\mu\text{L}\) 5x reaction buffer, 1 \(\mu\text{L}\) of (0.4 \(\mu\text{mol/L}\) each primer (reverse and forward), 0.125 \(\mu\text{L}\) (0.63 units) Taq DNA polymerase, 2 \(\mu\text{L}\) tenfold-diluted DNA) in a Biometra\textsuperscript{10} Thermal Cycler (Thermo Fisher Scientific, Switzerland) as follows: 95\textdegree\text{C} for 2 min, 35 cycles of 95\textdegree\text{C} for 20 sec, 56\textdegree\text{C} for 30 sec, 72\textdegree\text{C} for 30 sec, and 72\textdegree\text{C} for 2 min. Positive and negative controls were included in each PCR (DNA of previously identified SHB specimen from Nigeria (based on both morphometrics and DNA sequence data) or millipore water). Aliquots of PCR products were run by electrophoresis on a 2\% (w/v) agarose gel.

The amplicons were purified with ExoSAP-IT\textsuperscript{TM} (Thermo Fisher Scientific, Zurich, Switzerland) and sequenced in both directions (forward and reverse). For a better coverage of the 1091 bp fragment to be sequenced, two internal primers, Aet-int-F (5\textsuperscript{'}-CTTCCTGCTACAATTAATTGCG-3\textsuperscript{'}\text{)} and Aet-int-R (5\textsuperscript{'}-TTTGTTGATCATGAAGATTCGC-3\textsuperscript{'}\text{)} were added to the sequencing reactions\textsuperscript{42}. All samples were analyzed using the Big Dye\textsuperscript{8} Terminator v3.1 cycle sequencing kit (Thermo Fisher Scientific, Zurich, Switzerland). Sequencing was performed using a 96-capillary ABI PRISM\textsuperscript{8} 3730XL Genetic Analyzer (Thermo Fisher Scientific, Zurich, Switzerland). The nucleotide sequences were deposited in the GenBank database under accessions MK024945-MK025231.
DNA sequences alignment and analyses. The sequencing data were assembled and edited with Sequencher® software v5.4.1 (Gene codes corporation, Michigan, USA). The consensus sequences obtained were aligned using the MEGA 7 package and the MUSCLE alignment algorithm43, and compared to reference nucleotide sequence available in the GenBank database (http://www.ncbi.nlm.nih.gov/genbank/). All sequences with bad quality or ambiguous electropherograms were sequenced one in each direction twice and completely removed if still unclear. All SHB COI sequences from non-sampled countries available in GenBank database were added for further analyses. The alignment was performed on sequences of different lengths, ranging from 739 bp (A. concolor) to 980 bp (A. tumida). The program MrModelTest v2.328 was used to test models of evolution on SHB COI sequences for Bayesian Inference analyses. The best-fit model of nucleotide substitution for all sequences was SYM + G selected by Akaike Information Criterion (AIC). Bayesian inference (BI) was implemented with MrBayes v3.2.6 44. The evolutionary model employed six substitutions types (“nst = 6”), with the stationary state frequencies set to be equally fixed (“statefreqpr = fixed (equal)”). Rate variation across sites was modeled using a gamma distribution (“rates = gamma”). The Markov chain Monte Carlo (MCMC) search was run with 4 chains for 25,000,000 generations, with trees sampled every 200 generations and the first 6,250,000 trees discarded as ‘burn-in’. The 739 bp COI gene sequence from A. concolor was used as an outgroup to root the trees. ITOL (Iterative Tree of Life) v4.2 (https://itol.embl.de/) was used to visualize and annotate the phylogenetic tree.

Indices of sequence diversity, including number of mt-DNA haplotypes (Nh), haplotype diversity (h), nucleotide diversity (pi) and average number of pairwise nucleotide differences (K), were estimated using DnaSP v645. These parameters were calculated for samples grouped into five geographical regions: (1) Africa; (2) Americas; (3) Australia; (4) Europe and (5) Asia. In addition, the genealogical and geographical relationships among COI haplotypes were analyzed using the TCS network as implemented in PopArt46.

FAO data. Based on SHB invasion history, it can take two years after introductions before official pest confirmation (USA: 1996–1998)15. We therefore took advantage of published data8 to collect information about imports of beeswax into countries with recent introductions at least two years before the notification of the presence of the SHB.

| Country                          | Site (N of sequenced individuals) | Year |
|----------------------------------|-----------------------------------|------|
| Australia                        | Townsville (5), Cairns (6), Nambour (5), Victoria-Melbourne (4), Bathurst (3), S.E. Queensland (2) | 2016 |
| Benin                            | Keta (3), Abomey (3)              | 2015 |
| Brazil                           | Piracicaba (30), Sao Pedro (6)    | 2016 |
| Burkina-Faso                     | Fada Ngourma (2), Tenkodogo (3), Ziniaré (2), Bobo Dioulasso (3) | 2015 |
| Burundi                          | Rusiga (1)                        | 2016 |
| Canada                           | Canary (6), Mckenzie Road (1), LeFevtre Road (1), Eco-dairy (1) | 2015 |
| Central African Republic         | Keleng (2), Ndara (3), Sibut (2), Teremon (3) | 2015 |
| Costa Rica                       | Guanacoste (3)                   | 2016 |
| Cuba                             | Jovenallos (5), San Nicolas de Bari (5), Jaruco (5) | 2016 |
| Democratic Republic of Congo     | Maboya 1 (2), Maboya 2 (1), Maboya 3 (1), Maboya 4 (1), Mulo 1 (1), Mulo 2 (2), Mulo 3 (1), Mulo 4 (1), Kabomo 1 (1), Kabomo 2 (1), Kabomo 3 (2), Kabomo 4 (2) | 2016 |
| Ethiopia                         | Gedeo zone (5), Sidama zone (5)   | 2016 |
| Italy                            | Candinoni (1), Rizziconi (1), Melicucco (1), Polistena (1), Feroleto Della Chiesa (1), Taurianova (1), Ahtila (3) | 2016 |
| Jamaica                          | Old Harbour (5)                   | 2016 |
| Kenya                            | Meru (2)                          | 2016 |
| South Korea                      | Miryang (5)                       | 2017 |
| Madagascar                       | Ambat (3), Anjozorobe (1), Beomby (2), Isatra (3), Mantaosao (3) | 2016 |
| Malawi                           | Chawala (1), Chakhuntha (1)       | 2015 |
| Mexico                           | San Ignacio (4), Llera (4), Linares (4), Tancitaro (4), Carretera (4), Candelaria (4), Benito Juarez (4), Tepich (4), Yucatan (4) | 2016 |
| Nigeria                          | Osogbo (2), Ayangha (2), Otan (2), Oyan (2), Taraba (1) | 2015 |
| Philippines                      | Davao 2 (6), Davao 3 (5), Davao 4 (5) | 2015 |
| Portugal                         | Lisboa region (1 Larvae)          | 2004 |
| Republic of South Africa         | Northern Cape (9)                 | 2016 |
| South Sudan                      | Yei City (2)                      | 2015 |
| Sudan                            | Khartoum (1)                      | 2015 |
| Tanzania                         | Kilimanjaro (8), Arusha (6)       | 2015 |
| Togo                             | Seva (2), Notise (3)              | 2015 |
| Uganda                           | Jangano (2), Keirere (2), Rubona (1), Kyanyamutale (1), Oryang-Ojuma (1), Busiwu (2), Alik (2), Wasswa/Kavunza (2), Kanyonza (2) | 2015 |
| USA                              | Baton Rouge 1 (2), Baton Rouge 2 (3), Baton Rouge 3 (3), Oahu kunia (2) | 2016 |

Table 3. Sampling overview. The country, site and year of sampling are shown as well as the number of sequenced individuals per site.
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**Author Contributions**

F.O.I., Q.H., O.Y. and P.N. conceived and designed the study. F.O.I. performed the experiments. F.O.I., Q.H. and O.Y. analysed the data. F.O.I. prepared figures. F.O.I. and P.N. wrote the main manuscript text. All authors reviewed the manuscript.

**Additional Information**

**Competing Interests:** The authors declare no competing interests.

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