Conflicts and the spread of plagues in pre-industrial Europe

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One of the most devastating environmental consequences of war is the disruption of peacetime human-microbe relationships, leading to outbreaks of infectious diseases. Indirectly, conflicts also have severe health consequences due to population displacements, with a heightened risk of disease transmission. While previous research suggests that conflicts may have accentuated historical epidemics, this relationship has never been quantified. Here, we use annually resolved data to probe the link between climate, human behavior (i.e. conflicts), and the spread of plague epidemics in pre-industrial Europe (AD 1347–1840). We find that AD 1450-1670 was a particularly violent period of Europe’s history, characterized by a mean twofold increase in conflicts. This period was concurrent with steep upsurges in plague outbreaks. Cooler climate conditions during the Little Ice Age further weakened afflicted groups, making European populations less resistant to pathogens, through malnutrition and deteriorating living/sanitary conditions. Our analysis demonstrates that warfare provided a backdrop for significant microbial opportunity in pre-industrial Europe.
Introduction

Historians, scientists, and wider society have generally paid little attention to bygone epidemics, with the marked exceptions of the Black Death and the Great Plague of London (Duffy, 1977). This narrow outlook has recently changed due to the coronavirus pandemic and its profound impacts on human health, the global economy, and the geography of travel. For instance, the ongoing Covid-19 crisis has sparked renewed interest in Albert Camus’ novel “The plague”, originally published in 1947. The fascist “plague” that inspired the novel may no longer be a reality, but many other varieties of “pestilence” mean that this theme still has relevance today (Franco-Paredes, 2020). Pandemics are the most dramatic manifestation of the rapid and efficient spread of infectious pathogens, capable of influencing the course of world history. Understanding why, when, and how past epidemics/pandemics spread is therefore key to contextualizing current outbreaks.

The 2019/2020 coronavirus (Covid-19) pandemic has sharpened focus on the role of human population movements in rapidly spreading pathogenic microbes from a local hotspot to the global scale (Bedford et al., 2020; Chinazzi et al., 2020). In the past 40 years, outbreaks of infectious diseases have also been underpinned by population exoduses. Scourges have often emerged in forcibly displaced populations, invariably linked to a breakdown of health and social services (Murray et al., 2002). The Office of the United Nations High Commissioner for Refugees (UNHCR) reported that the “global forced displacement population” who have escaped conflict, persecution, or human rights violations totaled ~40 million people at the end of 2016 (The UN Refugee Agency, 2016). In 2018, it is estimated that 25 people were forced to flee insecure areas every minute (The UN Refugee Agency, 2018). War produces a multitude of opportunities for pathogenic microbes and constitutes an extremely effective way to promote microbial traffic and increase human morbidity and mortality. Migrants can act as vectors for infectious disease, leading to severe epidemics in receiving areas, where displaced populations are often housed in cramped refugee camps (see the Darfur region of Sudan; Degomme and Guha-Sapir, 2010). As early as 1995, the UNHCR stated that measles, diarrheal diseases, acute respiratory infections, and malaria account for between 60% and 80% of reported deaths in refugee camps (The UN Refugee Agency, 1995).

The health consequences of wars are nowadays circumvented by basic health care services, which alleviate the spread of epidemics, even in countries engaged in armed conflicts and where interventions are challenging (Spiegel et al., 2010; Leaning and Guha-Sapir, 2013). During the Late Middle Ages to the Early Modern Age, when persistent conflicts marred the European continent, the spread of plague (caused by the bacteria Yersinia pestis) was probably aggravated and enhanced through population fleeing war zones, increasing the geographical range of epidemics.

Fatigue, malnutrition, wounds, and stress are known to lower immune responses in human populations. Furthermore, camp life in crowded and unsanitary conditions favors the spread of contagious diseases and creates ideal ecological niches for both native and imported parasites. It has recently been suggested that there is an urgent need for a quantitative framework for modeling modern conflicts and epidemics (Banerjee, 2019). Within this context, there is potentially much to learn from the past and historical data are key to calibrating models. Here, we quantify the fundamental link between conflicts, plagues, fatalities, and the evolution of world population for the period from the Late Middle Ages to the Early Modern Age (AD 1340–1900). We also analyze how climate deterioration aggravated past epidemics/pandemics.

Results

Plague outbreaks. Arguably the most infamous plague outbreak in human history was the second plague pandemic (AD 1346–1720) for western Europe; ending around AD 1840 in Eastern Europe, North Africa, and the Near East; Dols, 1979; Hays, 2005), which started with the Black Death (AD 1346–1353). This pandemic swept through pre-industrial Asia and Europe (Benedictow, 2006). Scholars believe that the outbreak originated in China in the early 1330s, before spreading along trade routes and reaching Europe via Mediterranean ports in the late 1340s (Herlihy, 1997; Panum, 2007). However, the ultimate origin of the Black Death still remains uncertain (China, Mongolia, India, central Asia, and southern Russia; Norris, 1977). Recurring plagues lingered on for centuries, particularly in cities, and, for instance, the Great Plague of London (AD 1665–66; Roberts, 1966) is blamed for around 79,000 fatalities. While numerous epidemics, probably induced by different pathogens [the Plague of Athens (430–426 BC), the Antonine plague (AD 165–180), the Cyprian plague (AD 250)] and the first plague pandemic [Plague of Justinian (AD 541–750)] severely affected Europe (Little, 2007; Cohn, 2008; McMichael, 2012; Wagner et al., 2014) or not (Morduchai et al., 2019), the second plague pandemic remains the most devastating event in human history, killing some 30–60% of Europe’s population (75–200 million individuals). A number of recent studies have focused on the geography of the second plague pandemic in pre-industrial Europe (Büntgen et al., 2012; Schmid et al., 2015; Yue and Lee, 2018), but none have looked to quantify the link between human behavior, through the lens of conflicts, and the spread of epidemics (Banerjee, 2019).

Figure 1 illustrates that conflict dynamics, which engendered population displacements, manifest a strong correlation with the spread of plagues and increased fatalities between AD 1450 and 1670. The strongest correlations, highlighted by the Mantel scalograms, frame a particularly deadly period during which all datasets follow the same trend, spanning the end of the Middle Ages and the onset of the early Modern Period (AD 1450–1670). Conflicts were frequently associated with plagues before AD 1450. The Black Death (AD 1346–1353) struck during a violent phase in European history, notably symbolized by the Hundred Years’ War (AD 1337–1453) and the Reconquista (which ended in AD 1492). The plague was so devastating that it briefly interrupted these conflicts due to the significant death toll (Nolan, 2006). During the Hundred Years’ War, the epidemic reached France in 1347 AD before striking England in the summer of AD 1348 (Cusack, 2019). The pathogen reached the British Isles through the port of Melcombe Regis in Dorset County before hitting Bristol and later London, in the autumn (Hawkins, 1990; Bolton, 1996). The Black Death (AD 1346–1353) reached England, like other countries, via the terrestrial and maritime trade routes that interlaced Asia and Europe (Schmid et al., 2015; Yue et al., 2016). After this, the disease spread rapidly amongst military populations.

Conflicts and plague outbreaks. The rise in conflicts played out first, with a shift in the number after AD 1450 (Fig. 2), followed, around AD 1465/1470, by a growth in towns and cities affected by the plague. The acme occurred between AD 1465 and AD 1670, when the dynamic of all signals points to multiple concurrent events (Fig. 2). Cross-correlations applied to the conflicts versus plagues and plagues versus fatalities time series show a positive correlation \( P_{\text{value}} < 0.001 \), suggesting a chronological relationship between the three variables. A composite sequence (termed CPF, based on the sum of conflicts, plague incidences, and fatalities) further demonstrates that this period is linked to a
plateau in world population figures (Fig. 3). The rate of demographic growth (50-year average) slowed from AD 1470 to 1620 and became negative during the period AD 1620–1670 before strongly increasing from the late 17th century onwards. The increasing spread of plagues during the period AD 1400–1500 (Fig. 3) is probably related to the growing population density (Supplementary Fig. 1), as the pathogen was able to reach more
hosts rapidly. The connection between conflicts, plagues, population density, and fatalities is furthermore underlined by the neighbor-joining clustering (Supplementary Fig. 1), which places the conflicts at the beginning of the dynamic.

Climate and plagues. The role of climate in the reemergence of plagues in pre-industrial Europe has been previously debated in the literature (Schmid et al., 2015; Yue and Lee, 2018). Focusing on two time series of annually resolved temperature anomalies (European summer and Northern Hemisphere temperatures: Neukom et al., 2014; Luterbacher et al., 2016), it appears, in each case, that plague outbreaks mostly coincided with colder periods (Fig. 3 and Supplementary Fig. 2). The cross-correlations indicate a positive link between plague outbreaks and both climate sequences (P_{value} < 0.001) but with a higher correlation for the reconstruction based on Northern Hemisphere temperature anomalies. While variations in temperatures seem to have influenced the scourges, we find that precipitation anomalies (Cook et al., 2015) are not significant in explaining the plague data. Figure 4 summarizes (i) the link between the increase in conflicts and the rise of plague incidences in Europe, (ii) the role of cooler temperatures in the development of the plague, and (iii) the impact of conflicts and plagues on fatalities. The wavelet analyses (Fig. 5) demonstrate that the time series are characterized by equal periodicities, namely statistically significant cycles of 55 and 23 years. The associated Mantel scalograms show that while temperature anomalies are clearly associated with the CPF, drought is not a significant factor in explaining plague incidences (Figs. 3 and 5).

Discussion

Geography of population movements. In AD 1343–1347, the siege of Caffa (present-day Feodosia, Crimea) illustrates the devastating effect of the conflict–plague alliance. Caffa was the main port for Genoese merchant ships (Gardiner, 1995) and, in AD 1343, the Mongols/Tartars besieged the city (Dols, 1977). With the plague decimating the Mongol armies, they ordered the corpses to be placed in catapults and launched into the city, hoping that the disease would subsequently spread 

intra muros

(narrative of Gabriele De’ Mussi from AD 1343–1346). The plague was transmitted to Europeans by this stream of sick corpses inside Caffa. Italians fleeing the city brought the plague to Mediterranean ports (Wheelis, 2002). For instance, the plague first broke out at Messina in October 1347 AD. Three major centers of Mediterranean contagion then developed: Sicily, Genoa, and Venice (Nolan, 2006). This view was recently revised with the addition of the Venetian community of Tana, which may have played an equally important role in the transmission of the plague as the Genoese colony of Caffa (Barker, 2021). Later, during the English Civil Wars (AD 1639–1651) outbreaks of plague in AD 1644 and 1645 largely decimated armies and cities (Slack, 1990; Jennings, 2003). Disease, and in particular typhus and plague, killed more of the local population than the fighting due to the displacement of civilians (Jennings, 2003). During the Great Northern War (1700–1721 AD), a severe plague epidemic, peaking between 1708 and 1712 AD, ravaged the Baltic region (Frandsen, 2010). Most probably introduced via Constantinople, the plague first spread to southern Poland (Pinczów), and was then introduced by the army and traders into the Baltic Sea. The plague reached the Baltic coast of Prussia in AD 1709. It spread and affected all parts of the Baltic Sea in AD 1711, reaching Hamburg in AD 1712. The plague caused many deaths in the army as well as the depopulation of towns and rural areas due to the fleeing of civilians (Frandsen, 2010). In AD 1799, during the capture of Jaffa fortress (present-day Tel Aviv, Israel), Napoleon Bonaparte’s troops contracted the bubonic plague. Following Jaffa, Napoleon Bonaparte expected to defeat the Turks at Acre (present-day Acre, Israel). While Bonaparte’s army was defeated, the plague, which had accompanied the French from Jaffa, persisted, spread, and was introduced at Acre (Harris, 2006).

Our study paints a more complete picture of the devastation wrought by conflicts and plagues. From the Late Medieval to Early Modern Era, Europe and neighboring countries were hit by several waves of conflicts (Fig. 2 and Supplementary Fig. 3), causing peaks in mortality and disease. This promoted the spread of plagues in both military and civilian populations, a trend further accentuated by a growing population density in Europe during the 15th century AD (Fig. 3) and by population displacements far beyond the initial zones of hostility. Europe suffered from the 

European wars of religion,

but also from revolts, territorial ambitions, and great power conflicts (Nolan, 2006).
The same occurred in Eurasia, before and during the Ottoman Empire (Sugar, 1977; Aksan, 2015; Varlik, 2015). Civilian populations were significantly affected because armies tended to devastate civilian areas in an effort to feed themselves, causing food shortages, deplorable sanitary conditions, diseases, and population displacement (Ramsey, 2016). This phenomenon was exacerbated by the increasing length of conflicts, such as the Thirty Years’ War (Parker, 1997) and the Eighty Years’ War (Israel, 1995), which played out in areas subjected to repeated devastation. It led to more populations being affected by conflicts, with a greater health impact and more diseases. The wars of this era were considered to be amongst the most lethal before the modern period (Ramsey, 2016). This phase was devastating because it also corresponded to the start of the widespread use of gunpowder and firearms (sometimes called “Gunpowder Empires”; McNeill, 1993). These new weapons caused greater devastation on civilians and forced populations to flee the zones of hostility (Streusand, 2011).

Throughout history, it has been shown that, in most conflicts, infectious diseases kill more soldiers and civilians than weapons (Wilson, 1995). Conflicts themselves can act as vectors of epidemics, as they result in mixing of both military and civilian populations, increasing the probability of disease (Smallman-Raynor and Cliff, 2004). Wars displace large population masses that flee conflict zones in search of refuge in more geopolitically stable areas (Kendall et al., 2013). These populations are potentially vectors of infectious diseases which can spread in host areas (Kett, 2005; Degomme and Guha-Sapir, 2010).

**Spread of plagues.** For the period AD 1450–1670, recurrent plague outbreaks may be explained by other scenarios. It has been suggested that reintroductions of the bacterium into European ports occurred from reservoirs located in Asia, with a delay of 15 ± 1 years, and that no permanent plague reservoirs persisted inside Europe (Schmid et al., 2015). This concept has recently been challenged, suggesting that natural reservoirs may also have been located within Europe, in the southern Alpine region (Carmichael, 2014). Rasmussen et al. (2015) has also shown that the early branching of *Y. pestis* appeared 5783 years ago and that the ancestor of *Y. pestis* strains was widely distributed across Eurasia from the Bronze Age onwards. Large-scale population movements (Allentoft et al., 2015; Haak et al., 2015) may have facilitated the first plague outbreaks (Rasmussen et al., 2015).

A second and complementary hypothesis suggests that the spread of plagues, once the bacterium reached Europe, occurred via major trade routes which maintained sufficient contagion density to sustain plague transmission (Yue et al., 2016). In addition to these hypotheses, our analysis suggests that conflicts, through population displacements from war-affected areas or through a plague-endemic region, favored the spread of plagues via major but also secondary routes. This scenario could explain how peripheral towns were affected by plagues (shown in Supplementary Fig. 3), and why fatalities and the plateau in world population growth are both correlated with conflicts and plague outbreaks (Figs. 3 and 4).

A final hypothesis is that Europe was primed for plague outbreaks (De Witte, 2015). Food crises were ubiquitous in Europe because of a population boom set against the backdrop of land shortages. Europe was ravaged by the Great Famine (AD 1315–1317), resulting from both torrential rains that ruined harvests and institutional factors, including market failure, and a

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**Fig. 4 Relationships between conflicts, plague outbreaks, fatalities, and temperature anomalies.** The climate data are Northern Hemisphere temperature anomalies (Neukom et al., 2014). Linear and polynomial models, as well as a smoothing, were added for each curve.
disproportionate allocation of crop resources across different social strata (De Witte and Slavin, 2013). Famine strongly discriminated between social echelons, affecting mainly peasants and urban paupers. Other crises, such as the Great Bovine Pestilence in England (AD 1319–1320; Slavin, 2012), aggravated the situation. The gap between the poor and the elite was huge, with significant disparities in diets; peasant lives were extremely difficult. Food shortages, malnutrition, and degrading sanitary conditions paved the way for recurrent plagues.

While the usual mode of transmission of the bubonic plague to humans is via black rats (*Rattus rattus*) that host infective fleas (Kugeler et al., 2015), alternative modes include either pneumonic transmission (Kool, 2005) or via an intermediate human ectoparasite vector (e.g. human body louse or human fleas; Little, 2007). The pneumonic mode results from secondary involvement in bubonic cases after the spread of plague bacilli to the lungs before direct transmission (Pechous et al., 2016). The third mode of transmission, septicemic, was quite rare but highly fatal, killing within hours, before buboes had time to form (Cusack, 2019). Transmission of the bacilli also involved fleabite and produced fatal sepsis by different routes: through the lymphatic system, directly through the circulatory system, or both (Sebbane et al., 2006). The spread of conflict-linked plagues would advocate in favor of the pneumonic or human body louse/flea transmission of the bacilli. While the black rat population has been proposed as a plague reservoir in harbor cities (Keeling and Gilligan, 2000), via...
the fleas that it hosts (Xenopsylla cheopis and Nosopsyllus fasciatus), its role in the transmission of plague is debated (Davis, 1986; Hufhammer and Walloe, 2012). Rats were absent in large areas of northern Europe during the second plague pandemic and plague outbreaks often peaked when temperature and humidity were unfavorable for the arthropod vectors of plague to proliferate (Cohn, 2008). This would have hampered the rapid dissemination of the disease (Schmid et al., 2015). A spread from person-to-person therefore seems more plausible.

Cold climate. Another component that seems to have favored the persistence and recurrent spread of plagues is climate instability before and during the Little Ice Age (Neukom et al., 2014; Figs. 3 and 4 and Supplementary Fig. 2). It has been shown that warmer and wetter conditions favored the flea burden and host abundance (black rats or great gerbil) in the case of bubonic plague (Stenseth et al., 2006). While regional conditions may have promoted the second plague outbreak in Central Asia (Stenseth et al., 2006), the “pre” Little Ice Age (AD 1300–1550) was wetter and cooler in both Europe and Eurasia (Jones et al., 2001, 2006; Griggs et al., 2007; Kaniwsky et al., 2011; Xopliaki et al., 2018; Lüning et al., 2019). Because rat/gerbil and flea blooms are favored by climate conditions not recorded from Eurasia to Europe (Lionello, 2012; Kushner and Stein, 2019), and as rats were scarce in northern Europe during the plague outbreaks (Davis, 1986), the epidemics may have mostly spread via human-to-human transmission. Previous studies have mainly demonstrated that climate change has important health-related consequences because it can weaken afflicted populations by inducing food shortages, malnutrition, and even starvation (McMichael, 2012; Patz et al., 2000). These factors can alter human–microbe relationships and promote the (re)emergence of infectious diseases. Climate change did not initiate the emergence of the scourge during the period AD 1450–1670, but our data analysis suggests that it favored subsequent outbreaks by weakening European populations before and during the Little Ice Age. The “extreme wet years” in Europe and Eurasia during the pre-Little Ice Age had a strong impact on harvests. Fluctuations in harvests are clearly linked to environmental factors, mainly an increase in rainfall regime. In southern England, the lowest grain and straw yields are observed in wet-cold as well as in wet-warm years, whereas the highest grain yields are more frequent in dry or cold years and the highest straw yields in dry or warm years (Chmielewski, Potts, 1995). Kettlewell et al. (2003) have also noted that high summer precipitation in England and Wales leads to low grain growth. The link between heavy rain and poor harvests was evoked by Beveridge (1921, 1922) as early as the 1920s, who clearly showed a correlation between the rise in grain prices and increasing precipitation in England, France, Belgium, Germany, and Austria. In ancient China, Tian et al. (2017) have suggested that cooler climate conditions indirectly accentuated the prevalence of epidemics through famines during the Little Ice Age. Munzar (1995) describes a similar situation in Finland for the years AD 1695–1697. Food crises also engulfed Europe at the same time (Appleby, 1980). Our data support previous research on temperature anomalies (Schmid et al., 2015; Yue and Lee, 2018) and imply that colder conditions and conflicts affected food-production systems and human nutrition. Furthermore, colder conditions engendered indoor crowding and a greater likelihood of human-to-human infection, either through the medium of the human louse/flea or via the pneumonic form (Little, 2007). It is worth noting that, between the 1400s and 1800s, Europe was one of the most densely populated and urbanized areas on the planet (Klein Goldewijk et al., 2010).

War-plague alliance. Our study, based on both short and long-term trends, suggests that warfare significantly transformed Europe’s human–microbe environments, and the spread of infectious disease. It is clear that conflicts may have fostered the spread of infectious diseases and, in turn, pandemics may have led to social unrest leading to conflict. All of these factors were juxtaposed and constantly influencing each other. The devastating war–plague alliance led to a strong increase in fatalities producing a plateau in world demographics. Conflicts seem to have favored the spread of plagues by the movement of military troops and by pushing civilian populations to flee in unsanitary conditions. The climate of the pre-Little Ice Age and the Little Ice Age further reinforced outbreaks due to its impact on food resources—generating malnutrition that weakened the human immune system—and on the congregation of people indoors. Plagues caused by Y. pestis have greatly reduced or disappeared thanks to the invention of quarantine and efforts to develop prevention and intervention measures, namely improved hygiene and sanitation (Bramanti et al., 2019). In Europe, the disease was supplanted by smallpox (orthopox virus variola—VARV; Li et al., 2007; Duggan et al., 2016). For many centuries, smallpox devastated mankind, causing significant deaths (Behbehani, 1983; Geddes, 2006; Davenport et al., 2018). The bubonic plague has not fully disappeared and is still active in some countries (cf. Madagascar, Democratic Republic of Congo and Peru).

Concluding remarks

The 2019/2020 coronavirus pandemic has highlighted how the geography of population movements can drive the rapid escalation of infectious diseases. Within this context, recent events in Syria and Yemen show that warfare creates opportunities for pathogenic microbes to spread rapidly in displaced human populations (Abbara et al., 2020; Daw, 2020). Nonetheless, the role of armed conflict in driving recurrent plague outbreaks in the Late Middle Ages to Early Modern Age of Europe, one of the most violent in the continent’s history, is largely unknown despite its potential importance in constraining current human–microbe models of war-affected regions. Previous studies have underscored the absence of plague reservoirs in medieval Europe and have emphasized climate deterioration as a possible driver for the re-emergence of outbreaks. They have also highlighted the role of terrestrial and maritime trade routes in spreading historical outbreaks of the plague.

To the best of our knowledge, this study is the first to assess the role of conflicts in promoting the spread of plagues in pre-industrial Europe. First, we demonstrate that conflicts, plagues, and fatalities were significantly correlated during the period AD 1450–1670. Second, we find that the latter period corresponds to a plateau in world population figures. Finally, we suggest that the cooler climate conditions of the Little Ice Age affected food resources, weakening immune responses in populations through malnutrition and stress.

While recent research has focused on the role of warfare in promoting the emergence of epidemics in refugee camps, the effects of historical conflicts on human health are largely unexplored. Our study, based on high-resolution historical data from pre-industrial Europe, finds that recurrent conflicts between AD 1450 and 1670 entailed the displacement of military troops and civilian populations that acted as key vectors for disease transmission. This historical perspective reinforces the fundamental need for good healthcare in war-affected populations, in order to circumvent epidemics. Our study also demonstrates that there is potentially much to learn from the past regarding infectious disease diffusion and that such data are key to calibrating current quantitative models. These data are finally a warning for present-day
overpopulation, global warming, geopolitical tensions, the constant spread of industry, escalating poverty, and our compromised diets.

**Methods**

We collated data for European plagues, conflicts, and climate covering the end of the medieval period to the onset of the Modern Era.

**Dataset conflicts.** The definition of conflicts encompasses all events that can cause population displacements, from wars to revolts, including civil wars, insurgencies, rebellions, or battles. The data for conflicts mainly derive from the *Conflict Catalog* compiled and curated by P. Brecke (2020). The datasets were started to be compiled in 1998 (Brecke, 1998). All conflicts were sorted by date, duration, and cities(regions/states/countries) engaged in war. We summed events by year to create a linear time series covering the period AD 1347–1840 (Fig. 1). While conflicts may be missing, this dataset was controlled and used to detect general trends, to avoid uncertainties or misinterpretations.

**Dataset plagues.** Plagues in this study correspond to all the recorded epidemics caused by the pathogen *Y. pestis* (Bos et al., 2011). The initial dataset originates from the work of Biraben (1976). This seminal dataset was digitized (Atanasiu et al., 2008; Büntgen et al., 2012; Voigtländer and Voth, 2013) and improved with the additions from Russia, Constantinople, and Turkey (Schmid et al., 2015). We used this improved dataset for plague outbreaks. It has been shown that this dataset does not fully capture all historical plague activity across Europe, and that Biraben’s dataset has led to an important overestimation of plagues in cities and an underestimation in towns and villages (Rosen and Curtis, 2018). According to Rosen and Curtis (2018), the data collected by Biraben only denote the availability of sources mentioning plague and not the severity or pervasiveness of the disease in any given year. To avoid misinterpretation, we focus on overall trends, not on individual events that may be poorly expressed in the time series.

**Dataset fatalities.** Fatalities were gleaned from the *Conflict Catalog* (Brecke, 1998). We summed events by year to create a linear time series covering the period AD 1347–1840. We converted this dataset into fatalities ratioed to world population. The resulting time series was log transformed. The world population data were converted this initial matrix into an annually resolved dataset. Climate time series for the period AD 1347–1840 were then employed to test the similarities between the composite sequence (conflicts, plagues, and fatalities) and the climate proxies (Fig. 5).

**Statistical analyses.** All data were analyzed using XL-Stat2017 and PAST, version 2.17c. A simple smoother (smoothing transform with moving average as basic function; smoothing 9-year and 31-year) was first applied to assess long-term trends in conflicts, plague outbreaks, and fatalities (Fig. 1). Mantel scalograms (distance measure: Bray–Curtis), based on smoothed series (3-year smoothing), were subsequently used to test the similarities between the time series (Fig. 1). The apex of the triangle is the similarity between the first and last point. The base of the triangle shows similarities between pairs of consecutive points (Hammer and Harper, 2006).

We then assessed all datasets (Fig. 2) for long-term trends (3-year smoothing). Three homogeneity tests (Pettitt, SNHT, and Buishand) were applied to the time series to detect shifts in long-term dynamics. Each discordant period was categorized and its average denoted by “mu”.

We z-score transformed the time series and created a composite sequence (termed CPF) based on the sum of conflicts, plague incidences, and fatalities (Fig. 3). The results are shown with the full dataset and a long-term trend (3-year smoothing). A sinusoidal model was then applied to detect the long-term periodicities ($P_{value} < 0.001$). In tandem, we calculated growth rates for world population (Fig. 3), using a 50-year average (with standard deviation). The data are based on a subtraction between each consecutive data point and these differences were used to calculate 50-year averages. Climate proxies were transformed into long-term trends and a sinusoidal signal ($P_{value} < 0.001$) was fitted (Fig. 3).

To analyze the relationships between conflicts, plagues, fatalities, and temperature anomalies, we sorted the time series by ascending values (for the x-axis) and plotted the resulting curves with the standard error for each data point (Fig. 4). We added the results of the linear and polynomial models for each curve. A smoothing function was subsequently applied (Fig. 4).

We tested the periodicity of each time series (Fig. 5) using wavelet analyses (with Morlet as the basis function). Mantel scalograms (distance measure: Bray–Curtis; smooth 3-year) were subsequently applied to assess long-term trends in conflicts, plagues, and fatalities (Fig. 3).

**Data availability**

All the data are available at https://data.mendeley.com/datasets/ysrta8478w/1.

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Author contributions
D.K. and N.M. gathered the data, performed the analysis, and wrote the paper.

Competing interests
The authors declare no competing interests.

Additional information
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