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ON PLAGUE IN A TIME OF EBOLA

Cormac Ó Gráda

University College Dublin

Dublin 4

[cormac.ograda@ucd.ie]

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ABSTRACT

Ebola and plague share several characteristics, even though the second and third plague epidemics dwarfed the 2014-15 Ebola outbreak in terms of mortality. This essay reviews the mortality due to the two diseases and their lethality; the spatial and socioeconomic dimensions of plague mortality; the role of public action in containing the two diseases; and their economic impact.

KEYWORDS: plague, mortality, health, economic history

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ON PLAGUE IN THE TIME OF EBOLA

In 2014-15 the Ebola epidemic attracted global attention for its highly infectious character, its high fatality rate, the lack of any known cure for it, and the grave risks that it posed for health workers and third parties. The fears that it generated and the strict public health measures that it prompted echoed responses to the medieval Black Death and the third plague pandemic that originated in China in 1855.

Ebola and plague have rather similar incubation periods and both cause painful and distressing deaths. Indeed, for a time in the 2000s, some experts were convinced that plague was a form of Ebola (Little 2011; Haensch et al. 2010). Ebola is transmitted from person to person; whereas that is literally true of only one rare and highly lethal form of plague, pneumonic plague, the relatively recent finding (Drancourt et al. 2006; Ayyadurai et al. 2010; Hufthammer and Walløe 2013) that plague can be transmitted by body lice—and does not require the presence of rats and rat fleas—implies a route approximating transmission from person to person. Both diseases have led to searches for a medical remedy: rapid in the case of plague in the late 1890s in the wake of the discovery of the bacillus responsible, and similarly rapid in 2014-15 with the preparation for use of the (previously discovered) VSV-EBOV vaccine. These common characteristics prompt the following reflections about

See Kool 2005. Giovanni Boccaccio (author of the Decameron) was convinced that ‘the malady was communicated by speech or association with the sick… or by touching the clothes of the sick’, which led people ‘to shun and abhor all contact with the sick and all that belonged to them, thinking thereby to make each [their] own health secure’.
plague and Ebola, even though the second and third plague epidemics dwarfed the 2014-15 Ebola outbreak in terms of mortality. I review the mortality due to the two diseases and their lethality; the spatial and socioeconomic dimensions of plague mortality; the role of public action in containing the two diseases; and their economic impact.  

1. Deaths from Plague and Ebola

Around the time that this Forum was conceived quite alarming predictions were being made about the likely spread of Ebola in western Africa and beyond. Econometric simulations were forecasting that if the disease spread, one million or more would die ‘in the next six months’, forecasts soon echoed by the U.S. Centers for Disease Control and Prevention (CDC). The World Health Organization’s mid-October 2014 forecast of 10,000 new cases weekly was more modest, though it still dwarfed the actual cumulative out-turn of about 30,000 cases and over 11,000 deaths (roughly 0.05 per cent of the combined population of the three affected countries—although this does not include the likely impact on non-Ebola mortality). 

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3 For an excellent earlier essay relating the two diseases see Green 2014.

shaky assumptions and extraordinarily unrealistic in retrospect, were published as much for their ‘shock value’ and as a stimulant to public and private action as anything else.⁵

Whereas demographic data on Ebola are plentiful, historians of the Black Death have very little solid data to work on. There is no unanimity on the Black Death’s toll. After a careful scrutiny of the evidence for England nearly four decades ago medievalist John Hatcher (1977: 21-25) declared the ‘most judicious’ estimate of excess mortality from the first outbreak of plague in Europe in 1348-51 to be 30-45 per cent. Paolo Malanima’s analysis of country-level data implies that the Black Death and secondary epidemics reduced Europe’s population by 28 per cent between 1300 and 1400 (Malanima 2012: 314), but with considerable variation across countries, ranging from only 10 per cent in Austria (including Bohemia and Hungary) and 14 per cent in Belgium to 44 per cent in Scandinavia and 50 per cent in Ireland. More recent estimates also range widely, from a third to over three-fifths (Rawcliffe et al. 2015; Campbell 2015). In aggregate these guesstimates suggest that the first outbreak of the Black Death reduced Europe’s pre-plague population of about 80 million ⁶ by between 25 and 50 million.


⁶ Excluding Russia. See Malanima 2012: 312.
The Norwegian historian Ole Benedictow bases his case for the upper-bound estimate of at least three-fifths on what he interprets as ‘the remarkable similarity of the levels of mortality in ... widespread and diverse regions’, although some critics have linked this ‘similarity’ to Benedictow’s dismissal of any inconvenient data implying lower mortality (Benedictow 2004: 381-83; Cohn 2005; Horrox 2006: 199; Noymer 2007: 623-24; Mengel 2011: 22fn74, 32fn110). More recent work on Central Europe suggests that the Black Death’s impact there has been ‘greatly exaggerated’ (Mengel 2011: 31-32). But in the absence of anything remotely resembling civil registration the evidence is very thin and selective.

One way of evaluating the plausibility of a population decline of three-fifths is to ask how would it have impacted on the size and distribution of income. The economic impact of the Black Death in its wake was certainly dramatic. Clark reckons that the real wages of agricultural labourers in England rose by 55.5 per cent between 1339-48 and 1349-58, while real rents fell by 30 per cent (Clark 2007: 133; 2001: 25); Humphries and Weisdorf (2015) reckon that the real wages of unskilled women rose by 20 per cent between the 1340s and the 1350s; according to Malanima (2007: 157-58) the real wages of Italian urban workers by about 35 per cent and those of rural workers by 63 per cent over the same period (see too Pamuk 2007). These are big changes, but how do they square with Benedictow’s 60 per cent decline in population?

Table 1 reports the predicted outcomes for a range of population losses and the elasticity of substitution associated with the CES production function:
\[ Q = E(aL^\rho + b(T+K)^\rho)^{1/\rho} \]

where \( E \) is a measure of efficiency, \( Q \) is output, \( L \) (labour), \( T \) (land), and \( K \) (capital) are the factors of production and \( a \), and \( b \) are factor shares (here both assumed to be 0.50). In this specification the elasticity of substitution, \( \sigma \), equals \( 1/(1-\rho) \). Let initial \( E, L, K, \) and \( T \) equal 1, and assume that the Black Death affected only \( L \), which is a constant proportion of the population (compare Hirshleifer 1966); the effect of a reduction in \( L \) then depends on factor shares and \( \rho \). In that case the post-Black Death wage, \( w_i \), equals \( [1/L_i]^{1-\rho} \). As seen in Table 1, the actual changes in wages are much more modest than those implied by Benedictow’s 60 per cent decline (i.e. \( L_i=0.4 \)) for a plausible range of \( \sigma \) (\( 0.8<\sigma<1.5 \)).

—Table 1 about here—

2. Relative Lethality

Although some contemporaries believed in selective recovery from the Black Death (Crespo and Lawrenz 2014), most highlighted its lethality, i.e. the high probability that those who were infected would die from it. Hard data on the lethality of the Black Death in the fourteenth century are lacking. Paul Slack states that ‘in the absence of appropriate treatment, between 40% and 60% of the victims die’, but Benedictow, realizing that his mortality estimate would require a very high fatality rate indeed to be plausible, invoked that of
nearly four-fifths recorded during the Bombay Presidency pandemic of 1896-1900 in support (Slack 2012: 5; Benedictow 2004: 9; compare Nathan 1898: 13-14; Fee 1900). Benedictow might equally have invoked the case-fatality rates recorded for the Chinese population of Hong Kong between 1894-1907—92 per cent. But then would he not also have had to take into account the much lower fatality rate of Hong Kong's non-Chinese population—59 per cent (Brown 1913; Pryor 1975; Benedict 1996: 142, 206fn10)—and the accompanying implication that the health and nutritional status of the populations at risk may have mattered?

An added complication—also pertinent to Ebola—is whether lethality varied during the course of an epidemic. Hatcher (2008: 208) suggests that as the first outbreak in 1348 abated ‘the battle between the frailty of the victims and the potency of their affliction began to become less unequal’. Similarly, towards the end of the Moscow outbreak of 1771-72 ‘several persons who had the plague were but slightly indisposed, and walked about though they had buboes upon them’ (Alexander 2003: 2006). Moreover, there is the following tantalizing claim by the pope's surgeon regarding the lethality of successive outbreaks of plague in fourteenth-century Avignon (as cited in Cohn 2008):

In 1348, two thirds of the population were afflicted, and almost all died; in 1361, half the population contracted the disease, and very few survived; in 1371, only one tenth were sick, and many survived; while in 1382, only one twentieth of the population became sick, and almost all of these survived.

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7 For a similar assertion referring to the outbreak of 1665 in London see Payne (1894: 94).
The death rates recorded in Italian lazzaretti (pest houses) in the seventeenth century were much lower than Benedictow’s four-fifths. However, as Cipolla has made clear, rates such as the one-half recorded in the lazzaretto (or pest-house) of Pistoia in 1630-31 may underestimate the plague’s lethality, for two reasons. First, the survivors included many people who were quarantined because they displayed symptoms mistaken for plague or had resided with plague victims and, second, an unknown number had already died of the plague before they could be sent to the lazzaretto (Cipolla 1973: 105; 1981: 65). Whether such considerations are enough to bridge the gap between 50 and 80 per cent remains moot. Note too the remarks of Russian physician Gustav Orraeus, who whilst accompanying troops wrote during the 1770-72 plague: ‘Of our sick infected by the plague, one can calculate that a third recover; but of the local inhabitants, much the greater part die of it, because their relatives in this case immediately abandon them, besides which they do not take any medicine’ (cited in Alexander 2003: 105). Today the WHO reckons that half of those struck by plague recover without the aid of any medical treatment, a further indication that factors such as nutritional status and health offer some resistance against Yersinia pestis. During the 2014 outbreak of bubonic plague in Madagascar 119 confirmed cases resulted in 40 deaths; an outbreak in August-September 2015, this time of the pneumonic form, killed 10 of 14 victims.

8 Nearly two-thirds of those quarantined in Moscow in 1771-72 did not survive (Alexander 2003: 226).
The paucity of standard documentary evidence enhances the value of some recent archaeological research on the lethality of the fourteenth-century Black Death. Sharon DeWitte and her colleagues have compared skeletal evidence from London’s East Smithfield plague cemetery and non-epidemic samples from medieval cemeteries in the Danish towns of Viborg and Odense in order to test for adverse selection among those who succumbed to plague (DeWitte and Wood 2008; DeWitte and Hughes-Morey 2012; DeWitte and Slavin 2013). Their findings are further ammunition against the case for no selection. They report that ‘people who experienced physiological stressors, and who developed stress markers in response to those stressors, at some point (perhaps even long) before the arrival of the epidemic were subsequently more likely to die during the Black Death compared to their peers who lacked the stress markers’ (DeWitte 2014: 114). The finding is striking, although whether the economic and environmental pressures endured by Londoners, of whom there were about one hundred thousand on the eve of the Black Death, approximated those facing the people buried in Odense and Viborg, both towns with populations of a few thousand inhabitants at most, is questionable.

My reading of the scant evidence is that both the mortality and lethality rates associated with bubonic plague are likely to have varied across Europe. Yet even though Benedictow’s estimate of mortality, if not also that of lethality, seems too high, his depiction of the first outbreak of the Black Death as ‘the greatest catastrophe ever’ is apt. No other disaster in European history—war, famine, or epidemic—matches it. The Black Death was unparalleled because it was so easy to transmit and because of its power to kill.
Before the 1890s the only half-effective cure against the bubonic plague was to lance the buboes, an extremely painful procedure which apparently reduced the fatality rate (but compare Boghurst 1894: 87, 89). The fatality rate from Ebola in 2014-15 was much lower than from untreated *Yersinia pestis* (Table 2). The fatality rate was also very variable between the three countries, but lower everywhere than in the Republic of the Congo in 2002-4 (where out of 178 reported cases, 157 resulted in death). World Health Organization data suggest that the overall fatality rate during the recent outbreak was 40 per cent, ranging from 29 per cent in Sierra Leone to 67 per cent in neighbouring Guinea (see Table 2 and Figure 1b). There were 36 cases and 15 deaths outside the three countries. Neighbouring Nigeria and Mali accounted for 28 of those cases and 14 of the deaths.

—Figures 1a and 1b about here—

A second striking and disturbing implication of Table 2 is the very high proportion—nearly five per cent—of health workers among those who died during the Ebola outbreak. Nearly all of these were native-born: five health workers from high-income countries contracted the disease, and only one died. The proportion of all Ebola deaths ranged from 3.9 per cent in Guinea to 5.6 per cent in Sierra Leone. Nor were the victims confined to the lowest-paid workers:

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in Sierra Leone they included several physicians. These are very high proportions, given that transmission requires physical contact (WHO 2015). Perhaps the cumbersome procedures involved in ‘donning and doffing’ the highly uncomfortable Ebola protective suits worn by over-stretched health workers led some to take short cuts?  

Equally striking are the fatality rates among health workers struck by Ebola and the big gap between fatality rates in Guinea (less than half) and Sierra Leone (over seventy per cent). In Liberia and Sierra Leone health workers who contracted Ebola were more likely to die of it than others, even though presumably they were more likely to be diagnosed early. In all three countries, moreover, WHO data imply the fatality rate among health workers grew worse as the outbreak progressed. In Guinea it rose from 59 per cent before 26 November 2014 to 77 per cent thereafter; the percentages in Liberia were 42 and 51, respectively, and in Sierra Leone 21 and 36.

Despite this, the risk to health workers seems to have decreased over time, because WHO data imply that they were less likely to contract it over time. Thus, up to 26 November 2014, health workers represented nearly six per cent of all deaths, but thereafter—when presumably they were much more numerous—they represented about three per cent. Conditions ‘improved’ in all three countries in this regard.

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3. Did the Plague Distinguish Between Rich and Poor?

Malcolm Casadaban, who died of *Yersinia pestis* in Chicago in 2009, was a most unlikely plague victim. A biology professor, he succumbed to accidental exposure to a strain of the virus in his lab. But although the plague bacillus was blind, the most likely victims of plague have always been disproportionately the poor; ‘top people’ such as the Queen Eleanor of Aragon, the parents and siblings of Florentine merchant Francesco Datini, and the two archbishops of Canterbury who died in rapid succession during the first outbreak of the Black Death, were exceptional among their peers. In Noorthouck’s account of the last London outbreak, ‘the distemper was left chiefly to prey upon the common people; which it did to a degree, as to obtain the name of the *poor’s plague*’ (Noorthouck 1773: 220).

Cipolla’s study of plague in Prato in 1630 revealed that among those who fought it, the elite (health officers, physicians) were hardly touched while ‘the gravediggers and the attendants in the pest-house died like flies’ (Cipolla 1973: 107-08). During the last outbreaks of plague in London searchers and especially corpse bearers were presumably at similar risk, but according to court publisher Roger L’Estrange the attack of 1665 took away no ‘person of prime authority and command’. Again, in Moscow in 1770-1 the plague, ‘as is generally the case, raged chiefly among the common people; the nobles and better sort of
inhabitants escaped the contagion, a few only excepted, who fell victims to their rashness and negligence... Amid so great a number of deaths, ... there were only three persons of family, few of the principal citizens, and not more than 300 foreigners of the common class, who fell victims to the plague; the rest consisted of the lowest order of the Russian inhabitants’ (de Mertens 1799: 34-35). During the third pandemic in Hong Kong the white population escaped almost unscathed, and the small number who contracted plague were much less likely to die of it than the Chinese. An American public health specialist found this ‘interesting and strange’, given that ‘the white man has mixed freely with the Chinaman, going in and out of Chinatown at all hours, and many of them have their places of business within the infected areas; and yet they escape infection’ (Brown 1913: 555-56).

Parish register data shed further light on the socioeconomic gradient of plague mortality in sixteenth- and seventeenth-century London. Figure 2 compares the number of deaths, distinguishing between areas within and outside the walls, relative to their non-plague norm, in four plague years when deaths reached 5 to 6 times their habitual levels. One of the most striking features is the sharp fall between the 1560s and 1660s in mortality in the richer intramural parishes. An increasing tendency for the better off to flee from the city during plague years may help account for this. In 1665 the exodus included

---Figure 2 about here---

the King and his court (with Charles II manifesting a ‘paternal regard’ for his subjects from a safe distance in Oxford), ‘almost all the rich’, and ‘a great many of the clergy who were in circumstances’ (Harvey 1769: 415-16). Note conversely the significant increase over time in mortality in London’s rapidly urbanizing out-parishes.

Another likely reason for the socioeconomic gradient is the increasing self-segregation of the rich in London over time. This is the clear implication of Figure 3, which compares the spatial distribution of wealth by quartile in 1582 and 1638 (Cummins et al. 2015). Improvements in the quality of housing—‘bricks’ instead of ‘sticks’ (Keene 2001: 28)—within the walls may also have played a role.

—Figure 3 about here—

Alfani and Bonetti’s (2015) analysis of an unusually rich dataset from the town of Nonantola in northwestern Italy during the plague epidemic of 1630-31 paints a somewhat different picture. In Nonantola the rich appear to have been at as much risk as the poor and, indeed, the richest part of the town is where the risk of dying from plague was highest.

4. How Did the Plague Spread?
The erratic advance and retreat of Ebola in 2014-15 prompts a reflection on the spread of plague during its last visitations. The first named casualty of the last outbreak in London died in the low-income and densely populated parish of St. Giles-in-the-Fields, located to the northwest of the old walled city, on Christmas Eve 1664 (Moote and Moote 2006). This case raises two issues. First, it was claimed at the time that there was an apparent gap in plague deaths between then and 9 February 1665 and thereafter until 22 April 1665. Was the long hiatus between the first death due to ‘a hard frost which set in this winter and continued till March 1665; when its virulence was revived by the advance of the spring’ (Noorthhouk 1773: 217)? Or was it due to people concealing the truth from the officials responsible for recording the data, ‘as people were very loth at first to have neighbours believe their houses were infected, by money, or by other means they procured the dead to be returned as dying of other distempers’ (Harvey 1769: 439)? The London Bills of Mortality, a weekly record of deaths by cause, have always been considered a useful but fallible source on the demography of the plague. That is why the attempt by Cummins et al. (2015) to track London’s last plague epidemics over time and across parishes supplements what can be safely inferred from the bills with burial data from surviving parish registers.

The second is the location of the deaths. A contemporary described the London outbreaks of 1603 to 1636 as beginning ‘the first time by a surfeit in White Chapell, the second time, by Seamen, about the same place, the third by reason of rotten mutton at Stepney, the fourth with a packet of carpets from Turkey, the fifth with a Dogge that come over from Amsterdam’ (cited in Brett-
James 1935: 205). The London apothecary William Boghurst\(^\text{12}\) (1894: 26), however, noted that ‘the Plague hath put itself forth in St. Giles’s, St. Clement’s, St. Paul’s, Covent Garden, and St. Martin’s this 3 or 4 years, as I have beene certainly informed by the people themselves that had it in their houses in those Parishes’. This suggests a combination of local and exogenous factors, as does Payne’s introduction to Boghurst (1894: xiv-xv), although the consensus on London’s last epidemics is that ‘the plague is never originally bred with us but always brought accidentally from abroad’ (Harvey 1769).

The link between the outbreak of 1665 and ‘the transport of infection from Smyrna to Amsterdam in 1663’ (Slack 1981: 470) is plausible. But there is a conundrum; if indeed plague reached England from the continent (i.e. Holland), the first outbreak would most likely have occurred in some docklands parish. Why Saint-Giles-in-the-Field in the West End? Can we credit the story that it originated there with ‘some Levant goods that came from Holland... carried to a house in Long Acre near Drury Lane, where they were first opened’, whereupon ‘two Frenchmen’ died there and another who fled from Drury Lane to Bearbinder Lane introduced the plague into the city (Noorthhouck 1773: 217), a story which according to Payne (in Boghurst 1894: xi-xii) is ‘not supported by any authentic data’?

In their study of the spread of infection across the city week-by-week Cummins \textit{et al.} (2015) infer the location of the last outbreaks of plague from an epidemiological model applied to parish burial data. They find that these

\(^{12}\) In the \textit{Dictionary of National Biography} entry on Boghurst, Paul Slack, the greatest living expert on the history of plague, deems Boghurst ‘the most reliable eyewitness account of bubonic plague in 1665’. 
outbreaks were indeed more likely to start in the poor northern suburbs, before spreading around the walls and then within the City proper. In no case did mortality first appear in dockland parishes, implying that—perhaps—plague could have been generated internally in response to local conditions (see Figure 4). This would support a telluric or ‘localist’ rather than a ‘contagionist’ interpretation of the spread of plague in 1665. Interestingly, l’Estrange’s Intelligencer noted that the plagues of 1625 and 1636 had broken out ‘among the Butchers of White-Chapel (the likeliest Place for it always to shew itself here in) where there could not be least Suspicion of Foreign Goods.’

—Figure 4 about here—

Benedictow’s claim that the incidence of plague was similar across ‘widespread and diverse regions’ is not borne out by the spatial variation in mortality in early modern London and in sixteenth- and seventeenth-century Italy. Indeed, the considerable variation in mortality across relatively small distances is a conundrum remarked on and studied by Italian scholars (Cipolla 1981; Alfani 2013). Factors such as the timing of when the plague struck, geography and topography, the weather, and governance are all likely to have played a role. After surveying the data for a range of central-northern Italian cities—and in particular the neighbouring cities of Prato and Pistoia in 1630-

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13 The terms in inverted commas are Payne’s (foreword to Boghurst (894: xiv)).
Cipolla cautiously concluded ‘one has to be cautious about generalizations’ (1981: 84-85, 108).

5. Did Public Action Matter?

Why did plague recede in Western Europe during the sixteenth and seventeenth centuries? Why had it not done so earlier? After rejecting as likely factors the disappearance of the black rat (*rattus rattus*), environmental improvement, and changes in the nature of the disease, Slack (1981; see too Slack 2012) opted for increasingly effective public action in the form of quarantining, removing foul-smelling refuse, and draconic measures against infringement. Such action, which reduced the likelihood of transmission from fleas (and lice) to humans, required credible sanctions and a degree of public support. The effectiveness of public health measures depended on being ready, eliminating corruption in the forms of breaching quarantines, concealing deaths, deliberate misdiagnoses of plague cases, and anti-social behaviour in general (Cipolla 1979: 20). A local study of the last outbreaks of plague in Leicester, pointing out that the town had fared better than neighbouring towns such as Loughborough and Melton, attributed this to ‘the stringent regulations enforced during the time of plague... a benefit as well as a credit to her inhabitants’ (Wilshere 1968-69: 64).

Even Benedictow (2004: 95) concedes the role of Milan’s ruthlessly effective Visconti rulers in saving that city from plague in 1348-50—though plague would return there with a vengeance later. In Milan, the conviction—
not prevalent at the time—that plague was contagious led to efforts at controlling the movements of contacts. But public action can claim few other victories in 1348-50. In Florence (in Boccaccio’s words) ‘huge amounts of filth were removed from the city by officials charged with that task; sick people were forbidden to enter the city; advice was given on how to stay healthy; devout persons made humble supplication to God not once but many times, in processions and by other means’, but the plague had its way in an ‘almost miraculous manner’.

Quarantining and maritime cordons came later; the first lazzaretti were those in Ragusa (1377) and Venice (1423) (Tognotti 2013). The policy of isolating suspected plague victims in lazzaretti was based on the (false) presumption that the disease was spread from person to person, but may well have worked to the extent that lazzaretti were located in areas that were less flea- and rat-infested than the streets on which plague raged; but the English policy of shutting ‘sound people’ in their homes when a death had occurred instead of allowing them escape infection may well have been counterproductive (Payne 1894: 99; Slack 2012: 83-84). Cipolla has described in some detail how municipal action—through the enforcement of quarantines, the funding of lazzaretti, improvements in public hygiene, the banning of processions, and so on—reduced the threat of plague in parts of seventeenth-century Italy. The improvements were by no means comprehensive or universal: while the cities of Tuscany escaped rather lightly in 1629-30, plague killed one in three in Venice and three in five in Verona. Half the population of Genoa and Naples succumbed in 1656-57, while public action limited the mortality to one in ten in
Rome (Alfani 2013). Other institutional responses to plague included the London Bills of Mortality (continuously from 1603) and bans on processions and other large congregations of people.\textsuperscript{14}

Although what caused plague was not understood at the time, Cipolla depicts the acceptance of the measures associated with effective public action in northern and central Italy as a victory of science over superstition. How very different it was in Moscow at the height of the plague outbreak of 1771 when ‘an outrageous mob broke open the pest-houses and the quarantine-hospitals, renewing all the religious services which it is customary with them to perform at the bed-side of the sick, and digging up the dead bodies and burying them afresh in the city... They began again to embrace the dead, despising all manner of precaution’ (de Mertens 1799: 23). These unfortunates believed that precaution and resistance against the plague, a divine punishment, was futile.

This is reminiscent of reports from western Africa in 2014-15. Ebola took hold in Sierra Leone in May 2014 just after hundreds of people attended the death of a local faith healer who had succumbed to the disease. Traditional burial customs were also blamed for the early spread of the disease in Guinea (WHO); indeed, a WHO official declared that ‘fear and resistance from locals were more powerful drivers of the epidemic than any mistakes by the WHO or anyone else’.\textsuperscript{15}

\textsuperscript{14} In 1604 the authorities in London ordained that theatres should close once the number of plague deaths in the Bills exceeded thirty. See James Shapiro, ‘How Shakespeare’s great escape from the plague changed theatre’, \textit{The Guardian}, 24 Sept 2015.

\textsuperscript{15} WHO, ‘Sierra Leone: a traditional healer and a funeral: more than 300 Ebola cases link back to one funeral’ [http://who.int/csr/disease/ebola/ebola-6-months/sierra-leone/en/];
On the eve of the Ebola crisis Sierra Leone had 0.022 physicians per 1,000 inhabitants, Liberia 0.014, and Guinea 0.115. Compare this to Ireland’s 0.35 physicians and surgeons per 1,000 inhabitants on the eve of the Great Famine of the 1840s. Although the medics behind these numbers did not know how to cure either plague or Ebola, the numbers also reflect broader medical infrastructures. All three affected countries currently have per capita income levels about one per cent of Ireland’s; by the same token they are much poorer than, say, seventeenth-century Italy or England, though they have also been growing much faster in recent years. Given the economic context, some will ask how come the epidemic was so limited and vanquished so quickly!

The goal of international relief in 2014-15 was to compensate for frail public health infrastructures and widespread destitution. However, given the huge transfers involved, corruption was inevitable. In November 2014 in Sierra Leone, when health workers protested violently at not being paid, all a spokesman for the National Ebola Response Centre could offer was that ‘somebody somewhere needs to be investigated (to find out) where these moneys have been going’.

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16 In 2013 Gross National Income per capita, adjusted for PPP, was $720 in Sierra Leone; $480 in Guinea; and $400 in Liberia (World Bank). Nigeria’s more advanced health infrastructure helped it stop Ebola from spreading in late 2014 (‘How Nigeria stopped Ebola, The Economist, 20 October 2014).

17 Sara Jerving, ‘Why Liberians thought Ebola was a government scam to attract western aid: decades of corruption have left Liberians suspicious of their government’, The Nation, 16 Sept 2014.
The disease that had already killed about sixty people in Guinea since late 2013 was identified as Ebola on 22 March 2014. At that time the WHO characterized it as ‘an outbreak of limited geographic area and only a few chains of transmission’, and this assessment tallied with evidence on earlier outbreaks of Ebola in central Africa. However, by late May the disease had reached Sierra Leone and on 8 August 2014, by which time the death toll had reached nearly one thousand, the WHO declared the outbreak an ‘international health emergency’. In the following weeks alarming forecasts of the likely death toll from Ebola spurred the international community to action. By May 2015 such transfers exceeded $3 billion, more than 30 per cent of the combined GDP of the three recipient economies.

Still, there was plenty criticism of the weak and delayed response of the WHO and of the international community. Médecins sans frontières, who provided most of the frontline defence against Ebola at the outset, accused the authorities in Guinea and Sierra Leone of seeking to conceal the outbreak; the biotech company Metabiota, which was employed to monitor the crisis in Sierra Leone, of refusing to share crucial data about the progress of the outbreak; and the WHO of prevarication. The international community was rather slow to ‘wake up’ about Ebola. For example, there were only 8 references to ‘Ebola’ in the Irish Times before the end of June 2014. The first reference on

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April 1st referred to an epidemic claiming eighty lives in Guinea, while a feature on Médecins sans frontières on June 21st mentioned Ebola only in passing. Not surprisingly, the speed with which epidemic was eventually controlled led to speculation regarding what might have been achieved had large-scale intervention begun a few months earlier.  

6. The Cure

Like plague in the past, the recent Ebola outbreak inspired fear and panic because of its lethality, its long incubation period (WHO Ebola Response Team 2014; Van Kerkhove et al. 2015), and the lack of a medical cure. The eradication of plague across most of the globe relied on preventive rather than curative measures, and it took centuries for those preventive measures to become fully effective. One of the main defences against its spread, quarantine, involved isolating victims for a biblical forty days—a big multiple of plague’s incubation period of 2 to 6 days (which, of course, was not known or understood at the time).

Quarantine has also been a key weapon against Ebola; a precautionary 21-day quarantine is stipulated for those who have been in close contact with an Ebola victim. In October-November 2014 the publication of a paper by Haas (2014) arguing that a 21-day isolation period was too short led to a flurry of

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(mainly uninformed) hysterical commentary—‘Fear-bola’ (Jantz 2014)—regarding the dangers posed by U.S. health workers returning home from western Africa. Policy makers rightly ignored this panic, recognizing the danger that draconian quarantines would deter volunteers and increase avoidance.  

Alexandre Yersin’s identification of the plague bacillus, *pasteurella pestis*, in 1894 enabled the Russian-born microbiologist Waldemar Haffkine to produce his anti-plague vaccine after just three months of intensive work in a makeshift laboratory in Bombay in January 1897. Haffkine had previously produced an effective anti-cholera vaccine in Pasteur’s laboratory in Paris and so, once the *Yersinia pestis* bacillus had been identified, producing a vaccine based on heat-killed *Y. pestis* cultures was well within his capacity. After minimal trials, the new vaccine was pronounced ready for use. Granted, its unpleasant side effects lessened its appeal, but it was effective in reducing the incidence and lethality of disease (*British Medical Journal* 1900; Meyer *et al.* 1974).

As with plague before Haffkine’s discovery, there was no therapeutic cure for Ebola during the epidemic of 2014-15. One of the truly dramatic by-

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22 Breaking news in mid-October 2015 that the virus persisted in some survivors of Ebola long after their recovery raised new fears [e.g. The Guardian, 17 October 2015 http://www.theguardian.com/world/2015/oct/16/how-pauline-cafferkeys-ebola-relapse-tears-up-everything-doctors-thought-they-knew]. The WHO’s response, rightly, was that there was no need for panic [http://www.who.int/csr/disease/ebola/virus-persistence/en/]: ‘While it is now clear that virus persists longer in semen than previously thought, the risk of people being infected with Ebola by those who have survived the disease is probably low. Although sexual transmission by survivors with persistent virus is a possibility, it appears to be rare. In areas of Sierra Leone – Kailahun and Kenema - that had very large outbreaks and have high numbers of survivors, there have been no new cases of Ebola for 300 days.’
products of that epidemic was the final release of a highly effective vaccine. In fact, the clinical discovery of VSV-EBOV dates back to 2003, when scientists working for the Canadian Public Health Agency produced a drug that was effective in protecting rhesus monkeys. Had Ebola struck in the 1890s it is possible—and this is not to advocate the same response today—that a similar drug would have been discovered, in which case the authorities would have gambled on its immediate use, as happened in the case of Haffkine’s vaccine.

However, before the latest Ebola outbreak the Canadian government had merely patented the VSV-EBOV vaccine and presented vials of it to the WHO to try out on volunteers. It took the current crisis for the Merck Group to buy the commercial rights to a VSV-EBOV from a small biotech company in November 2014. By April 2015 Merck were organizing Phase 3 trials in Guinea. Although the crisis was almost over by the time the vaccine was ready for use (July 2014), the speed with which it was developed is nonetheless rather impressive.

6. The Economic Consequences of the Black Death and Ebola

In October 2014 the World Bank predicted that Ebola would cost the three affected countries US$25 billion in economic losses in 2015. That would have been equivalent to almost twice their combined GDPs, surely enough to cripple them economically for years to come. By January 2015 that prediction

23 The initial discovery, by Canada’s Public Health Agency, predated the latest crisis; they had sold the commercial rights to a small U.S. biotech company NewLink in 2010.
had been reduced to a still significant US$1.6 billion, equivalent to about 12 per cent of combined GDPs. The most recent estimates (see Figure 5) are much more sanguine, implying that the impact of Ebola on the level of economic activity has been minor and will be temporary. The significant drop in GDP in Sierra Leone, by far the most dynamic of the three economies, in 2014 was almost entirely due the global collapse in iron ore prices rather than to Ebola.  

Analogous estimates of the economic impact of the Black Death are impossible, but that impact has nonetheless been the subject of extended debate among economic historians. Elementary Malthusian economics predicts that an exogenous shock such as the Black Death, that reduces population but leaves the capital stock and other resource endowments intact, will result in reduced output but an increase in wages relative to other factor payments (Hirshleifer 1968; Le Roy Ladurie 1974: 40-50; Clark 2001). And there can be no doubt but that the first attack of the Black Death in western Europe resulted in significantly improved living standards for most survivors, while reducing urbanization levels and shifting the balance between crop cultivation and pasture in the countryside (e.g. Campbell 2006; Malanima 2012). Labour’s gain ‘was to prove a potent driving force behind revolutionary changes in economic and social institutions, including the decline of serfdom and feudalism, and a golden age for peasants and labourers’ (Hatcher 2008: 321).

Some go further, holding that high wages led to labour saving technologies such as the Gutenberg printing press; or crediting the Black Death with leading to, or being indirectly responsible for, an ‘industrious revolution’ or an industrial revolution (e.g. Pamuk 2007; Koyama 2012; Voigtländer and Voth 2013). Evidence of the impact of the Black Death on income or wealth distribution is very thin: the only study I have come across is that Alfani and Ammanati (2014), which reports a significant albeit temporary reduction in wealth inequality in north-central Italy in the wake of the Black Death.

There is also general agreement that it took the population of Europe a long time to recover its pre-plague level, partly because plague kept returning but also because, in some countries at least, of incessant warfare in the decades and centuries that followed. According to Broadberry et al. (2015) England’s population reached its post-1348 nadir a century later (c. 1450), three-fifths below its peak on the eve of the Black Death; it would not reach that peak again until the late 1620s.

The economic and demographic impact of subsequent outbreaks could much be less dramatic, as highlighted by the case of London. This is seen in the absence of any significant ‘plague’ effect on wages between the mid-sixteenth and mid-seventeenth centuries (Cummins et al. 2015; Boulton 1996). Such was the power of the city to attract labour from the countryside that the last epidemic of 1665 had no appreciable impact on wage levels. The remarkable power of the metropolis to attract labour in the wake of plague, noted by Graunt (1676 [1899]: 367) at the time, is indicated by its ability to
make good so quickly after 1665 one hundred thousand lives lost, even though the total number of young adults living outside London at that time that cannot have numbered much more than a million. The elasticity of labour supply is also indicated by the speed with which new apprentices replaced those who had succumbed (Table 3 below). The last outbreaks of plague in England left no appreciable traces because they were largely confined to urban areas, particularly the metropolis. Still, the welfare gains from the eradication of plague were substantial (Ó Gráda 2015).

Two important reservations temper the traditional Malthusian account of the post-Black Death era. First, the landed elite everywhere naturally resisted improvements in labour's status, sometimes violently, and with varying success. In eastern Europe this led to what Friedrich Engels dubbed a ‘second serfdom’. Second, the de-urbanization that followed mortality and emigration reduced the volume of both internal and international trade. This may well have led to some diseconomies of scale, with the result that although the lot of the landless labourer improved, the overall state of the economy may not have.

This is closer to the scenario that has recently been proposed for Italy, where the impact of plague in the seventeenth century was heavy and pervasive, and affected both rural and urban areas. Plague has been blamed for turning ‘one of the wealthiest areas of Europe’, and one with a ‘solid economy’, into an economic backwater (Alfani 2013; see too Alfani and Percoco 2014). In Italy the plague outbreaks of 1629-30 and the 1650s, unlike that of 1348, brought no increase in real wages in its wake (Malanima 2007). Why the difference?
One hypothesis is that the seventeenth-century population loss slashed internal aggregate demand at a time when increasing protectionism was reducing foreign demand. The resultant fall in labour supply thus generated a negative productivity shock by depriving Italy of scale economies that would prove impossible to recover. Given the small size of production units in this era, such economies at the level of the individual firm can hardly have mattered much: but the loss of external economies in realms such as communications and urban services is plausible, given the decline in both the size of several of Italy’s major cities and their share of total population (Malanima 2005: 103).

A second argument is that the sheer size of the seventeenth century epidemics in Italy meant that skilled and comfortably off inhabitants were not immune, so that there was an attendant adverse impact on human capital. This is by no means implausible; the starkly different outcomes in the fourteenth and seventeenth centuries might be explained by the universality of the former outbreak compared to the local but very severe character of the Italian outbreaks of the seventeenth century, which placed Italy at a serious competitive disadvantage in their aftermath.

—Table 3 about here—

7. A Final Reflection

The classic criticism of an independent role for 'demand' in the Industrial Revolution in England is Mokyr (1977).
Despite the major differences between them, *Yersinia pestis* and Ebola share many resonances. The campaign to contain and eradicate Ebola—and the attendant red tape and corruption—recalls the varied attempts by the authorities at ridding Western Europe of plague. Later efforts to control plague had an international dimension: the work of Haffkine, a Jew born in Odessa, in Bombay was funded by the local authorities and by the Aga Khan and plague’s virtual eradication in pre-independence India owed much to colonial policies (Harrison 1994: 152-58; Dasgupta 2005). Today the resources and knowledge available campaigns against epidemics like plague and Ebola are global rather than local. In the case of Ebola, NGOs such as Médecins sans frontières, institutions such as the WHO, and the governments of the countries affected combined in bringing the 2014-15 epidemic under control.

According to WHO data the epidemic had caused 11,313 deaths up to mid-October 2015, by which time the crisis had been stayed, with only 23 deaths since the end of August. The number is very modest compared to, say, estimates of famine deaths in Somalia in 2011-12 or of deaths from malaria in sub-Saharan Africa in 2014 (0.4 million), yet the global impact of Ebola was far greater.

At the height of the crisis the Harvard global health specialist Paul Farmer (2014) insisted that ‘if patients are promptly diagnosed and receive aggressive supportive care—including fluid resuscitation, electrolyte replacement and blood products—the great majority, as many as 90 per cent,

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26 Put at 0.26 million by Checchi and Robinson (2013); for a few caveats see Ó Gráda (2015: 181-3).
should survive’. Easier said than done, given the fears generated by Ebola\textsuperscript{27}, and the primitive health infrastructures (e.g. in Sierra Leone ‘Most hospitals and clinics don’t have running water, and some don’t have electricity’\textsuperscript{28}) and rickety economies of the counties in question. Yet how many lives might have been spared by a prompter response or by extra funding remains to be discovered (DuBois et al. 2015).

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\textsuperscript{27} These included both the fear of being infected and that of being isolated. As in the case of plague in western India in the late 1890s some those living in Ebola-affected areas in 2014 believed that health workers were spreading the disease. In the US, where an Ebola outbreak was never a serious possibility, one poll in October 2014 suggested that nearly two-thirds of the population feared such an outbreak while another indicated that nearly half feared that a family member would contact Ebola (Jantz 2014).

\textsuperscript{28} Amy Maxmen, ‘To prevent the next plague, listen to Boie Jalloh’, *NPR*, 8 October 2015 [http://www.npr.org/sections/goatsandsoda/2015/10/08/446631677/to-prevent-the-next-plague-listen-to-boie-jalloh].


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Figure 1a. Cumulative cases in Guinea, Liberia and Sierra Leone
Figure 1b. Cumulative fatality rates in Guinea, Liberia and Sierra Leone

Figure 2. Mortality in plague years compared with average mortality over the preceding five years
Figure 3. The spatial distribution of wealth in London, with parishes grouped by quartile, 1582 and 1638
Figure 4. The spread of plague in London
Figure 5. GDP Growth Per Annum 2009-2017 in the Affected Economies
Table 1. Predicted wage increase for different values of $L_1$ and $\sigma$

<table>
<thead>
<tr>
<th>$L_1$</th>
<th>$\sigma=1.5$</th>
<th>$\sigma=1$</th>
<th>$\sigma=0.8$</th>
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<tbody>
<tr>
<td>0.8</td>
<td>16</td>
<td>25</td>
<td>32</td>
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<td>0.6</td>
<td>41</td>
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<td>89</td>
</tr>
<tr>
<td>0.4</td>
<td>89</td>
<td>150</td>
<td>214</td>
</tr>
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</table>

Table 2. The Recent Ebola Epidemic: Cases and Deaths

<table>
<thead>
<tr>
<th>All</th>
<th>Health Workers</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cases</td>
</tr>
<tr>
<td>Guinea</td>
<td>3,800</td>
</tr>
<tr>
<td>Liberia</td>
<td>10,672</td>
</tr>
<tr>
<td>Sierra Leone</td>
<td>13,982</td>
</tr>
<tr>
<td>Total</td>
<td>28,454</td>
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</tbody>
</table>


Table 3. Average number of apprentices enrolled annually before and after three mortality crises in London

<table>
<thead>
<tr>
<th>Influenza, 1557-9</th>
<th>Plague, 1593</th>
<th>Plague, 1603</th>
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<tbody>
<tr>
<td>5 years before</td>
<td>3 years after</td>
<td>% change</td>
</tr>
<tr>
<td>314</td>
<td>520</td>
<td>66</td>
</tr>
<tr>
<td>1,039</td>
<td>1,574</td>
<td>51</td>
</tr>
<tr>
<td>1,439</td>
<td>2,505</td>
<td>74</td>
</tr>
</tbody>
</table>

Source: Rappaport 1989: 75
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