**Yersinia pestis** DNA Sequences in Late Medieval Skeletal Finds, Bavaria

To the Editor: *Yersinia pestis*, the causative agent of plague, is held responsible for 3 human pandemics: the Justinian plague (5th–7th century), the Black Death (13th–15th century), and modern plague (1870s to present). In 1894, Alexandre Yersin identified *Y. pestis* during an epidemic of plague in Hong Kong (1). However, whether *Y. pestis* was indeed responsible for the medieval epidemic is still controversial. *Y. pestis* specific DNA has been detected in medieval skeletal finds (2,3), although some investigators have failed to do so (4), leading to the suggestion that a viral hemorrhagic fever was the agent of these medieval pandemics (5).

Against this background, we investigated a mass burial site that was discovered under the sacristy of the St. Leonhard Catholic church in Manching-Pichl, near Ingolstadt in Bavaria, Germany. In 1984, during the renovation of this church, 75 human skeletons and several scattered skeletal elements were discovered under the sacristy. The skeletons lay densely packed in 4 layers. Apparently, no grave pit had been dug at all, but the bodies were obviously deposited side by side and covered with dirt layer by layer. An approximation of the time of burial was possible only by means of accompanying building structures, which were dated to the Gothic period (1250–1500 CE).

In the course of research preceding this study, our research group found *Y. pestis* DNA in 10 of 33 examined individual skeletal remains from the mass grave beneath the sacristy (6) by using the primer pair YP12D/YP11R (3). In the current study, the remains of 6 persons from the mass burial site that had positive *Y. pestis* DNA results before were further investigated. DNA from additional tooth samples of these persons was extracted, and more markers mapped on the *Y. pestis* high copy number plasmid pPCP1 were included. In addition to the primer systems YP12D/YP11R and YP11D/YP10R (3), the following primer systems was used. The primer pair YP14F (5′- TCCGGGTCAGGTAATATGGA-3′)/YP13R (5′-ACCAGCCTTTCATGAGG-3′) amplifies another sequence section (positions 6953–7082, reference: *Y. pestis* strain CO92 plasmid pPCP1 sequence AL109969.1) on the *Y. pestis* pla gene (encoding plasminogen activator). The plasminogen activator belongs to the virulence factors of *Y. pestis*. The primer pair pst-F (5′-GGTAA ATCGCTGAACCGAA-3′)/pst-R (5′-AACAACCTCCTGACGCT TT-3′) amplifies a 129-bp sequence section (positions 5026–5154, GenBank accession no. AL109969.1) on the *pst* gene (encoding pesticin activity protein). Pesticin is a bacteriocin that is active against only a few closely related microorganisms. The primer pair PCP-F (5′- CATCCACATGCTCAACCCCTA-3′)/ PCP-R (5′-CTGAACGCATTTCAG TGTTG-3′) can amplify a 128–130-bp sequence section (positions 8428–8555, GenBank accession no. AF053945.1; positions 8428–8557, GenBank accession no. AL109969.1) on plasmid pPCP1.

These 3 new primer sets, designed with the aid of the software component Primer3 (7), were applied by using a suicide PCR method (3,8); i.e., new primer pairs targeting sequences not previously amplified in the laboratory were used. At no time during all examinations was any modern *Y. pestis* DNA included.

The sample preparation, DNA extraction, PCR setup, electrophoretic separation, and sequencing of amplicons are described elsewhere (6,9); however, we used 0.20 μmol/L of each primer, an annealing temperature  

<table>
<thead>
<tr>
<th>Identification no.</th>
<th>Sex, age at death, y/sex</th>
<th>Tooth sample</th>
<th>DNA extract†</th>
<th>YP12D/YP11R‡</th>
<th>YP11D/YP10R</th>
<th>YP14F/pst-R</th>
<th>PCP-F/PCP-R</th>
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<tbody>
<tr>
<td>17-I</td>
<td>≈11/M</td>
<td>1st</td>
<td>I</td>
<td>–</td>
<td>+</td>
<td>NT</td>
<td>NT</td>
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<td></td>
<td>1st</td>
<td>I</td>
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<td>NT</td>
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<td>2nd</td>
<td>I</td>
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<tr>
<td>22</td>
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<td>2nd</td>
<td>I</td>
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<td>–</td>
<td>NT</td>
<td>NT</td>
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<tr>
<td>34-I</td>
<td>Young adult/M</td>
<td>1st</td>
<td>I</td>
<td>+</td>
<td>+</td>
<td>NT</td>
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<td>–</td>
<td>NT</td>
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<tr>
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<td>I</td>
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</tr>
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<td>S4-XX</td>
<td>≈15/F</td>
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<td>I</td>
<td>+</td>
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<td>1st</td>
<td>I</td>
<td>+</td>
<td>+</td>
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</table>

†Extraction controls and PCR controls had no amplicons. +, amplicon; (+), weak amplicon; –, no amplicon; NT, not tested.
‡Independent PCRs with the primer pair.
of 54°C, and 45 amplification cycles. PCR blanks containing all reagents except for DNA and extraction blanks were included in every PCR set.

Results of the amplification reactions are listed in the Table. All accompanied extraction and PCR controls remained free of amplification products. All amplicons resulting from suicide PCRs were sequenced. Amplicons resulting from the use of primer pairs YP14F/YP13R and pst-F/pst-R matched the reference sequence to 100% (GenBank accession no. AL109969.1). Amplicons resulting from the use of primer pair PCP-F/PCP-R matched this reference sequence to only 97.78%. This deviation is because of a 2-bp insertion (2 Ts, positions 8531 and 8532, GenBank accession no. AL109969.1) at Y. pestis strain CO92 plasmid pPCP1. The sequences obtained from 3 persons’ remains showed in the pPCP1 sequence section between nucleotide positions 8528–8532 only 3 Ts instead of 5 Ts described for Y. pestis strain CO92 plasmid pPCP1 (GenBank accession no. AL109969.1). The sequences found in this study were deposited in GenBank under accession nos. HQ290521–HQ290523.

To conclude, the successful recovery of several Y. pestis plasmid pPCP1 DNA sequences in skeletal finds from the mass burial site excavated in Manching-Pichl suggests that these persons died of plague. Moreover, our findings constitute a molecularly supported confirmation for the presence of Y. pestis, the etiologic agent of plague, in late medieval (1250–1500 CE) southern Germany. In future studies, we will attempt to recover chromosomal Y. pestis DNA from the mass grave skeletal remains to obtain clues as to the specific Y. pestis strain and the microbiology of past plague in Europe.

Ingrid Wiechmann, Michaela Harbeck, and Gisela Grupe

Author affiliations: Ludwig Maximilian University of Munich, Munich, Germany (I. Wiechmann, G. Grupe); and Bavarian State Collection of Anthropology and Palaeoanatomy, Munich (M. Harbeck, G. Grupe)

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References


Address for correspondence: Ingrid Wiechmann, Ludwig Maximilian University of Munich, Department Biology I, Biodiversity research/Antropology, Grosshaderner Str. 2, 82152 Planegg-Martinsried, Germany; email: i.wiechmann@lrz.uni-muenchen.de

Two Clusters of HIV-1 Infection, Rural Idaho, USA, 2008

To the Editor: Prevalence of HIV-1 infection in rural areas of the United States has been increasing (1). During 2003–2007, an average of 30 (range 24–42) cases of new HIV-1 infection diagnoses per year among Idaho residents were reported. Of the 152 reported cases during this period, 54 (36%) were related to a person living in a rural area of ≤75,000 residents and a 60-minute drive from an urban area (2). Of these 54 cases, 19 (35%) were in men who have sex with men (MSM), 5 (9%) were in injection drug users (IDU), and 2 (4%) were in those in both categories.

In March 2008, a cluster of newly identified HIV-1 infections that included 5 cases (cluster A) in a rural southeastern Idaho city (city A) was reported to the Idaho Department of Health and Welfare. Two patients were men and the median age was 26 years (range 18–32 years). One patient was an IDU (Table). Through epidemiologic investigation, 3 additional patients were suspected to be IDUs, but confirmation was not practicable. All reported methamphetamine use. One man and 2 women reported both male and female sex partners.

During September–December of that year, another increase in newly identified HIV-1 infections in southeastern Idaho (cluster B) was reported