Hantaviruses cause 2 zoonotic diseases, hemorrhagic fever with renal syndrome (HFRS) and hantavirus cardiopulmonary syndrome. Infection is usually initiated after inhalation of virus-contaminated rodent excreta. In addition to the zoonotic infection route, growing evidence suggests person-to-person transmission of Andes virus. For this reason, we studied whether saliva from HFRS patients contained hantavirus. During an outbreak in northern Sweden of nephropathia epidemica (NE), a milder form of hemorrhagic fever with renal syndrome, we collected saliva and plasma from 14 hospitalized NE patients with verified Puumala virus (PUUV) infection. PUUV RNA was detected in saliva from 10 patients (range 1,530–121,323 PUUV RNA copies/mL) by quantitative reverse transcription–PCR. The PUUV S-segment sequences from saliva and plasma of the same patients were identical. Our data show that hantavirus RNA could be detected in human saliva several days after onset of disease symptoms and raise the question whether interhuman transmission of hantavirus may occur through saliva.

Members of the family Bunyaviridae cause severe and often fatal human diseases in a large and increasing number of persons worldwide each year (1). This family contains 5 genera, and the genus Hantavirus causes 2 febrile illnesses: hemorrhagic fever with renal syndrome (HFRS) in Europe and Asia and hantavirus cardiopulmonary syndrome (HCPS) in North and South America. Hantaviruses are rodent-borne pathogens. In Sweden, Finland, Norway, Russia, and parts of central Europe, Puumala virus (PUUV) is endemic in bank voles (Myodes glareolus). PUUV causes nephropathia epidemica (NE), a milder form of HFRS. The most common symptoms of NE are fever, headache, nausea, vomiting, myalgia, abdominal pain, back pain, and visual disturbances (2). One third of the patients have hemorrhagic manifestations, 10%–20% have respiratory tract symptoms, and most have signs of renal failure (2). There is no effective treatment or available vaccine.

The most common route of hantavirus infection is infectious aerosols originating from saliva, urine, and feces of infected rodents (1). Because rodent bites have been demonstrated to cause human hantavirus infections (3,4), saliva of hantavirus-infected rodents must contain infectious virus. Infectious PUUV has been found in oropharyngeal secretions from M. glareolus (5) and Andes virus (ANDV) RNA is present in saliva from infected pigmy rice rats (Oligoryzomys longicaudatus) (6). Furthermore, Sin Nombre virus RNA is present in saliva but not in urine and feces from deer mice (Peromyscus maniculatus) (7), confirming that hantaviruses are transmitted to humans through rodent saliva. However, person-to-person transmission has recently been documented for the ANDV that causes HCPS (8–13), and contaminated human saliva suggests a possible route of infection (13). For herpesviruses, such as Epstein-Barr virus and herpesviruses-7 and -8, person-to-person transmission can occur from saliva (14–16), but human salivary secretions could also negatively modulate herpesvirus infectivity as shown for herpesvirus-1 (17).

Because the saliva of hantavirus-infected rodents contains infectious virus that is transmitted to humans (3,4,7) and person-to-person transmission is strongly suspected for ANDV (13), we studied whether the virus could be detected in saliva of hantavirus-infected patients. For this reason, we collected saliva from patients during an NE outbreak in northern Sweden and analyzed the samples for the presence and levels of PUUV RNA by using a real-time reverse transcription–PCR (RT-PCR) assay.
Material and Methods

Collection of Human Saliva

Saliva was collected from PUUV-infected patients who were hospitalized from January through May 2007 at the Department of Infectious Diseases at Umeå University Hospital (Umeå, Sweden). Samples were obtained from patients with typical clinical symptoms of acute NE, which were verified by detection of PUUV-specific immunoglobulin (Ig) M in serum, a PUUV-specific real-time RT-PCR, or both. The collection was random, with no consideration of time of the day or food intake. The samples were only collected on 1 occasion during each patient’s hospitalization by asking the patient to spit in a single-use plastic mug. The saliva was then removed with a syringe (5 mL; Becton Dickinson, Franklin Lakes, NJ, USA) and put in sterile plastic test tubes (Nunc, Roskilde, Denmark). The samples were immediately stored at –80°C until they were further analyzed. Plastic gloves and protective clothing were used during the procedure. A total of 9 samples were stored without additives, and 5 were diluted 1:3 in virus transport medium (2% HEPES, 10 μg/mL bovine serum albumin, 50 μg/mL sucrose, 0.016 μg/mL fungizone, and 2.5 μg/mL garamycin) before storage. The project was approved by the Research Ethics Committee of Umeå University and all patients gave written and informed consent.

Real-Time RT-PCR

RNA from 140 μL of patient saliva and plasma was extracted by using a QIAamp Viral RNA kit (QIAGEN, Valencia, CA, USA) according to the manufacturer’s instructions. The real-time RT-PCR was performed as previously described (18). Briefly, the RNA was reverse-transcribed followed by a real-time PCR TaqMan assay in triplets with PUUV-specific primers and probe from the S segment (18). The real-time PCR was performed with an ABI Prism 7900HT Sequence Detection System 2.0 (Applied Biosystems). The PCR and sequencing reactions were performed with primers from the S-segment. For patients 1 and 4, the primers for PCR was Puu1s (5′-CAAGAGGATATGACGGCA-3′) and Puu6as (5′-GCCATCCTCCTGCAATAGAT-3′) followed by Puu3s (5′-AAGTGGGATTGAGCCAGAT-3′) and Puu5as (5′-TTGGCACATTCTTTTTCCATAA-3′) for sequencing. For patient 2, the primers for PCR was Puu1s and Puu6as followed by Puu2s (5′-ACCCGCAAGGACAAACT-3′) and Puu4as (5′-TAGGGCTTTTCATAATAGGTA-3′) for sequencing. Before the sequencing reaction, the PCR fragment was purified by using a QIAquick PCR Purification kit (QIAGEN) and precipitated in ethanol. Assembly, analysis, and alignment of sequences were performed by using Geneious Basic 3.0.6 (www.geneious.com) and BLAST (www.ncbi.nlm.nih.gov/blast).

Results

Hantavirus RNA in Saliva from NE Patients

We collected saliva 2–9 days after onset of symptoms from 14 hospitalized NE patients to determine whether hantaviruses were present in human saliva. All patients were positive for PUUV RNA in plasma by a real-time RT-PCR, and 13 patients had PUUV-specific IgM antibodies in serum (Table). We then analyzed the saliva by real-time RT-PCR for the presence and levels of viral RNA. Saliva samples from 10 of 14 patients were positive for PUUV RNA (range 1,530–121,323 PUUV RNA copies/mL), and IgG levels were determined by using fluorescein-conjugated rabbit anti-human IgG (F202, DAKO A/S, Glostrup, Denmark) diluted in phosphate-buffered saline (PBS) with Evans blue. For IgM analysis, patient serum was pretreated with rheumatoid factor absorbent (Virion Serion GmbH, Würzburg, Germany) to eliminate possible interference of rheumatoid factor and PUUV-specific IgG. Slides were incubated overnight at 37°C, and antibody was detected by using fluorescein-conjugated rabbit F(ab′)2 anti-human IgM antibodies (F0317, DAKO A/S) diluted in PBS with Evans blue.
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demonstrating that viral RNA could be detected in saliva
during disease (Table). Furthermore, we detected no inhi-
bition of real-time RT-PCR in saliva or plasma when we
analyzed our patient samples by using an internal positive
control (data not shown). All patients were also IgG posi-
tive in serum with varying titers.

Sequencing of PUUV RNA in Saliva
To study in more detail whether the RNA we detected
in saliva was truly speci-
cfic, we sequenced regions of RT-
PCR products. Sequencing of ampli-
fication products from
saliva and plasma from 3 NE patients—patient 1 (GenBank
accession nos. EU337014 and EU337015), patient 2 (Gen-
Bank accession nos. EU177629 and EU177630), and patient
4 (GenBank accession nos. EU337016 and EU337017)—
demonstrated a PUUV S-segment sequence that was iden-
tical for each saliva/plasma pair (data not shown). When
we compared the sequences for patients 1 and 4, we found
24 mismatches in the 292-nt S-segment sequence, which
clearly demonstrated that they were derived from different
strains (Figure). The observed nucleotide mismatches did
not result in different amino acid sequences. The sequence
from patient 2 was from another S-segment region. All
PUUV sequences from the NE patients were most closely
related to PUUV strains from the disease-endemic region
in northern Sweden and grouped with the northern branch
of PUUV strains (19,20).

Respiratory Symptoms in NE patients
All 14 NE patients required hospital care and dis-
played the symptoms of NE. All survived and none had
to be treated by dialysis. Seven patients had respiratory
symptoms including cough, dyspnea, or both. In 2 of the
patients with respiratory symptoms, a chest radiograph re-
ealed infiltrates (data not shown). The 2 patients with the
highest levels of viral RNA in saliva had dyspnea (patient
1) and cough and lung infiltrates (patient 2) (Table). All
patients with respiratory symptoms, except 1 (patient 12),
had PUUV RNA in their saliva.

Discussion
This investigation showed that hantavirus RNA could
be detected in saliva from HFRS patients. We found PUUV
RNA with varying PUUV genome copy numbers in the sa-
liva of most of the 14 hospitalized NE patients. Previously,
we have shown that PUUV RNA in plasma decreases with
time in the individual patient (18), and although we did
not study this finding, PUUV in saliva likely would dis-
play similar kinetics. PUUV viremia is thought to persist
for 5–7 days, but in our study, 1 patient had PUUV in both
saliva and plasma 9 days after onset of symptoms. Fur-
thermore, we recently detected PUUV RNA in plasma 16
days after first appearance of disease (18). It would also be
interesting to study whether nonhospitalized patients with
milder symptoms also had PUUV RNA in their saliva. We
do not know if PUUV RNA in saliva could originate from
PUUV viremia, but the PUUV RNA detected in saliva was
likely produced in salivary glands or through coughing. Re-
cently, ANDV antigen was found in secretory cells of the
salivary glands of human patients (21). In our study, 7 of
the patients had respiratory symptoms, and virus shedding
in saliva is known to occur for several respiratory virus-
es, such as severe acute respiratory syndrome–associated
coronavirus, respiratory syncytial virus, and human meta-
pneumovirus (22,23). Symptoms in the upper respiratory
tract are also often present in NE (2,24), which makes our

<table>
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<tr>
<th>Patient no.</th>
<th>Sex</th>
<th>Age, y</th>
<th>Date saliva collected (d of disease)</th>
<th>PUUV RNA† in saliva, copies/mL</th>
<th>PUUV RNA† in plasma,‡ copies/mL</th>
<th>PUUV IgM§ in serum‡</th>
<th>PUUV IgG titer in serum‡</th>
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*PUUV, Puumala virus. Ig, immunoglobulin.
†Detected by real-time reverse transcription–PCR.
‡Most plasma and serum samples were collected the same day or within 1 day before or after the saliva sample and antibody was detected by
immunofluorescence assay.
§Serum was diluted 1:16 and scored as negative (–), positive (+), or weakly positive (±).
finding of PUUV in saliva as credible. One may argue that NE, the HFRS endemic in northern and central Europe, is a different entity than HCPS, caused by Sin Nombre hantavirus and ANDV, and that the finding of viral RNA in saliva from NE patients is a pure coincidence. However, NE in Scandinavia shows several common characteristics with HCPS in the Americas. For example, pulmonary involvement and respiratory symptoms are common in NE. In addition, radiologic examination–detectable infiltrates, decreased pulmonary function, and a local inflammatory response in the lungs have previously been demonstrated (25–28).

Could hantaviruses in human saliva be infectious and initiate infection? Today, the only hantavirus suggested to be transmitted between humans is ANDV (8–13). By using sophisticated epidemiologic data, Ferres et al. showed that the risk for HCPS caused by ANDV was higher among sex partners of the index case than among other household contacts (13). These investigators suggested that ANDV needs close person-to-person contact, such as sexual relations or deep kissing, to be transmitted between humans (13). Our finding of hantavirus in saliva support the conclusion that hantavirus infection could be transmitted within humans through saliva, such as during kissing or coughing. However, we do not know at this stage whether the hantavirus RNA in saliva detected in our study is infectious. When we tried to infect bank voles with human PUUV RNA–positive saliva, no seroconversion was found after 21 days (J. Hardestam et al., unpub. data). Apparently, hantaviruses in rodent saliva are infectious, but whether the composition of rodent and human saliva differs is not known. Further studies are needed to elucidate the mechanisms behind hantavirus transmission between the natural rodent hosts and humans.

After finding PUUV RNA in saliva from NE patients, we searched for evidence of possible person-to-person transmission of PUUV. When the NE patients were followed-up 1–2 months later, we discovered several clusters of household members with NE symptoms. However, none had severe enough disease to require physician care, and no samples were collected for laboratory diagnosis. Furthermore, during the large NE outbreak in early 2007, there was a very high incidence (313/100,000) of NE in the region where our university hospital is situated (36). Since all patients lived in this area, we cannot exclude inhalation of rodent excreta as the main or only transmission route. Although it remains to be clearly shown, our results sup-

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**Figure.** Comparison of nucleotide sequences between reverse transcription–PCR products isolated from nephropathia epidemica patients 1 and 4. Mismatches are underlined.
port the hypothesis that person-to-person transmission may occur by this route.

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References


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