Airborne transmission of lyssaviruses

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In 2002, a Scottish bat conservationist developed a rabies-like disease and subsequently died. This was caused by infection with European bat lyssavirus 2 (EBLV-2), a virus closely related to Rabies virus (RABV). The source of this infection and the means of transmission have not yet been confirmed. In this study, the hypothesis that lyssaviruses, particularly RABV and the bat variant EBLV-2, might be transmitted via the airborne route was tested. Mice were challenged via direct introduction of lyssavirus into the nasal passages. Two hours after intranasal challenge with a mouse-adapted strain of RABV (Challenge Virus Standard), viral RNA was detectable in the tongue, lungs and stomach. All of the mice challenged by direct intranasal inoculation developed disease signs by 7 days post-infection. Two out of five mice challenged by direct intranasal inoculation of EBLV-2 developed disease between 16 and 19 days post-infection. In addition, a simple apparatus was evaluated in which mice could be exposed experimentally to infectious doses of lyssavirus from an aerosol. Using this approach, mice challenged with RABV, but not those challenged with EBLV-2, were highly susceptible to infection by inhalation. These data support the hypothesis that lyssaviruses, and RABV in particular, can be spread by airborne transmission in a dose-dependent manner. This could present a particular hazard to personnel exposed to aerosols of infectious RABV following accidental release in a laboratory environment.

INTRODUCTION

The genus Lyssavirus within the family Rhabdoviridae comprises classical Rabies virus (RABV; genotype 1) and at least six additional genetically divergent lineages that discriminate the genotypes of virus within the genus. The viral genome is composed of a single, negative strand of RNA encoding five proteins: the nucleoprotein, phosphoprotein, matrix protein, glycoprotein and an RNA polymerase (Tordo, 1996). The genus Lyssavirus includes: Lagos bat virus (genotype 2), Mokola virus (genotype 3), Duvenhage virus (genotype 4), European bat lyssavirus 1 and 2 (EBLV-1 and -2, genotypes 5 and 6, respectively) and Australian bat lyssavirus (genotype 7) (Bourhy et al., 1993). Members of this genus can be placed into two broad groups. Genotypes 1, 4, 5, 6 and 7 belong to phylogroup I, whilst genotypes 2 and 3 have been placed in phylogroup II on the basis of glycoprotein sequence, pathogenicity and immunogenicity (Badrane et al., 2001). Additionally, there are at least four unclassified lyssaviruses – Aravan, Khujand, Irkut and West Caucasian bat virus – that have been isolated recently from bats in Eurasia and proposed as new members of the genus Lyssavirus, indicating that these are emerging zoonotic viruses (Botvinkin et al., 1996, 2003; Arai et al., 2003). Lyssaviruses are all capable of causing clinical ‘rabies’ disease in mammals. All genotypes, with the exception of Lagos bat virus, have been responsible for human fatalities (Fooks, 2004).

Lyssaviruses in bats present a public health risk and are classified as ‘category C emerging infectious disease threats’. In Europe, classical rabies is being eliminated from much of the continent; however, two related viruses, EBLV-1 and -2, are widely distributed (Bourhy et al., 1992). Between 1985 and 2002, 673 cases of EBLV infection were detected; the majority (> 95%) of virus isolates were EBLV-1 and were predominantly associated with the Serotine bat (Eptesicus serotinus), whereas EBLV-2 appeared to be associated with Myotis species (Fooks et al., 2003a). In the UK, a recent fatality due to infection with EBLV-2 occurred in Scotland (Fooks et al., 2003b). A bat conservationist was diagnosed with rabies infection following exposure to bats; the route of transmission has never been accurately ascertained, although he admitted to numerous exposures involving close handling of different bat species and previously had been bitten by bats on some occasions whilst not wearing gloves. This is the first recorded case of indigenous human rabies in the UK in 100 years. In the majority of cases, animal-to-animal transmission of RABV occurs following saliva exposure after a bite from an infected animal (Warrell & Warrell, 2004). In the Americas, however, ‘cryptic
transmission’ of RABV bat variants to man is reported annually (Messenger et al., 2002). The ‘small vector hypothesis’ (Messener et al., 2003) suggests a failure to recognize the seriousness of a bat bite, especially from a small bat, which accounts for numerous cryptic cases of human rabies as a result of being bitten. The bite is often unrecognized, largely due to the perceived negligible size of the lesion. As frequent urination and defecation is commonplace among bats, and a large amount of guano is present, resulting in the possibility of aerosolized virus, it is possible that vertical transmission of virus among bats in enclosed areas (caves, roof spaces and cellars) occurs as a result of airborne virus (Constantine, 1962, 1967; Winkler, 1968). Bats naturally infected with RABV have been shown to harbour virus in the nasal mucosa, leading to the supposition that airborne infected with RABV have been shown to harbour virus in the nasal mucosa, leading to the supposition that airborne virus transmission via the airborne route and the viral load required for dispersal are not known (Winkler, 1975). It is assumed that environmental conditions for airborne transmission (Winkler et al., 1972) and the virus titre will dictate the effectiveness of transmission. The possibility exists for exposure via the respiratory tract in susceptible hosts that enter an infected area (Atanasiu, 1965). As a consequence of exposure via the airborne route, it is suggested that this route provides an enhanced capability for RABV to invade the brain via the olfactory neuroepithelium and olfactory nerve in susceptible individuals (Jenson et al., 1969; Lafay et al., 1991). Human cases of rabies from accidental airborne transmission have been reported previously (Irons et al., 1957; Kent & Finegold, 1960; Winkler et al., 1973; Tillotson et al., 1977; summarized in Table 1). It is possible that EBLV-2 can be transmitted by a number of routes, principally through bites, but also by the generation of local aerosols. To assess this, we compared EBLV-2 transmission via the airborne route with classical RABV.

**METHODS**

**Viruses.** The Challenge Virus Standard (CVS) strain of RABV was used as a representative of genotype 1. EBLV-2 strain RV1332 was isolated from a Daubenton’s bat (Myotis daubentonii) in 2002 (Johnson et al., 2003). Both viruses were grown in BHK-21 cells (<10 passages) using Leibovitz L-15 media (Invitrogen) with 1 % fetal calf serum (FCS). At this concentration of FCS, both viruses grew suboptimally and, as a result, virus suspensions were prepared by combining the supernatant and trypsinized cell monolayer in an attempt to maximize the level of virus. TCID₅₀ (fluorescent foci of infected cells) was calculated using a tenfold serial dilution of BHK-21 cells as described previously (King, 1996). Suspensions were stored at −70°C until required.

**Mouse strain.** Male RIIt mice (k/k haplotype) were used throughout the study (VLA stock, inbred, albino). Mice were provided with food and water ad libitum and all experiments were conducted under Home Office Project Licence 70/5327. Mice challenged with lyssa-virus were monitored using a scoring system. This rated the mice as 0 (no effects), 1 (ruffled fur, hunched back), 2 (slow or circular movements, gait affected), 3 (trembling, shaking and lameness) and 4 (paralysis). Animals found with scores of 3 or above were killed humanely.

**Virus challenge.** Groups of mice (n=5–7) were challenged by direct intracranial (i.c.) inoculation, intranasal (i.n.) inoculation or subdermally through the footpad. Aerosol exposure of groups of mice (n=10) was achieved using a bespoke portable apparatus as described previously (Phillpotts et al., 1997). Briefly, compressed air (BOC) was used to create an aerosol from a Collison spray. Groups were exposed to the aerosol, unrestrained, in an 80 l box, closed at one end, with a HEPA filter. Subsequently, the exposure cabinet was flushed out with air (30 l min⁻¹) for 10 min before the animals

<table>
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<th>Year</th>
<th>Comments</th>
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<td>1956</td>
<td>Rabies was confirmed in an entomologist who had worked in a number of caves in Texas containing large colonies of Mexican Free-tailed bats. The patient had no recollection of being bitten.</td>
<td>Irons et al. (1957)</td>
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<td>1959</td>
<td>A consultant mining engineer was admitted to hospital in Los Angeles, California, on 1 June, complaining of shortness of breath and retching. Rabies was confirmed following death. Previously, he had been working in caves in both Mexico and Texas. No evidence of a biting incident could be confirmed, although there was a report that he emerged from one cave with a bleeding wound on his face.</td>
<td>Kent &amp; Finegold (1960)</td>
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<td>1972</td>
<td>A 56-year-old veterinarian who had been working in a laboratory preparing rabies vaccine developed headaches, vomiting, diarrhoea and general weakness. The patient had been involved in the preparation of rabies vaccine from brain obtained from rabies-infected goats. This process involved the use of a kitchen blender, which, on subsequent investigation, was observed to produce a visible aerosol.</td>
<td>Winkler et al. (1973)</td>
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<td>1977</td>
<td>A further case of laboratory-acquired rabies was reported in 1977 in a 32-year-old technician, who had been participating in experiments to prepare an oral vaccine by coating small pellets with aerosolized virus. He felt weak and developed a fever. On admission to hospital, he became lethargic and entered a comatose state. Unlike previous cases, the patient began to recover from 3 May, although with neurological impairment.</td>
<td>Tillotson et al. (1977)</td>
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were removed. All exposures were undertaken within a microbiological safety cabinet operating at class III mode under guidelines covering Advisory Committee on Dangerous Pathogens Category 3/Specified Animal Pathogens Order Category 4 containment. Immediately following exposure, two mice from the control group exposed to medium and the highest virus titre group were sacrificed, and brain, tongue, lungs and stomach were removed and stored at −70°C. The remaining mice were transferred to a flexible-film isolator (Bell Isolation Systems) and monitored twice daily for 28 days unless otherwise stated.

Following challenge, subjects were killed humanely when clinical signs of disease developed or at the termination of the study. The brain was removed from all animals exposed to either media or virus aerosols and tested for rabies using a fluorescent antibody test. Serum was obtained from all animals that survived to the end of each experiment and tested for neutralizing antibody by a fluorescent antibody virus neutralization test.

Fluorescent antibody test (FAT). Viruses were initially detected by a FAT using standard protocols (Dean & Abebe, 1973). Briefly, impression slides of brain tissue were fixed in acetone, washed and stained with anti-nucleoprotein fluorescein isothiocyanate-labelled monoclonal antibody (Centocor).

Antibody detection. RABV-specific antibody was detected using a fluorescent antibody virus neutralization test (Cluquet et al., 1998).

Extraction of RNA. Detection of virus within total organ samples was performed using sensitive hemi-nested RT-PCR as described previously (Johnson & Fooks, 2005). Briefly, RNA was extracted from whole organs using TRIzol (Invitrogen). RNA was stored at −80°C.

RT-PCR and sequencing. RT-PCR was performed as described previously using primers JW12 and JW6(DPL) (Heaton et al., 1997). Hemi-nested PCR was carried out using primer combinations JW12 and JW10(P) or JW12 and GT1/GT6 (Johnson & Fooks, 2005). PCR products were separated on a 1% agarose gel and visualized by ethidium bromide (1 μg ml⁻¹) staining and UV illumination.

RESULTS AND DISCUSSION

Virus distribution in mice challenged directly with RABV

The survival of mice inoculated with RABV was assessed by the i.c., subdermal (footpad) and i.n. routes. The challenge virus induced rabies in all of the inoculated mice (Table 2).

The only apparent difference among the routes of challenge was the time taken to reach late-stage disease, with the two peripheral routes of inoculation (footpad and i.n.) taking a further 2 days to induce disease compared with the more direct i.c. route.

Two mice challenged by the i.n. route or control animals were sacrificed immediately after challenge to assess the distribution of CVS following introduction of virus (Fig. 1). Total RNA was extracted from the tongue, lungs and stomach (including stomach contents) to detect virus using hemi-nested PCR. All first-round amplifications were negative. However, nested PCR detected RABV in the stomach (Fig. 1, lanes 10 and 13), lung (Fig. 1, lane 11) and tongue (Fig. 1, lane 12). This suggested that virus introduced through the i.n. route became widely distributed and could find multiple routes of entry. The consistent finding of virus in the stomach (Fig. 1, lanes 10 and 13) indicated that most virus introduced by the i.n. route was swallowed and presumably rendered non-infectious.

Aerosol challenge of mice with RABV

Having established the consequences of i.n. challenge, we proceeded to assess the ability of RABV to infect mice using aerosolized virus. Tenfold dilutions of challenge virus were prepared and used to expose groups of mice to increasing concentrations of CVS. Two mice were sacrificed from the control group and the group exposed to aerosol generated from a suspension containing 10⁵ TCID₅₀ CVS ml⁻¹. Virus distribution was assessed as described above. Hemi-nested PCR demonstrated the presence of RABV in the lungs (Fig. 2, lanes 7 and 10) and stomach (lanes 9 and 12) in both mice. This observation suggested that aerosolized virus was inhaled and ingested in a similar manner to direct i.n. challenge.

The detection of virus in lung vacuoles suggested that the aerosol particles were of the optimum size for uptake. The remaining mice were monitored for 28 days using a scoring system developed for i.c. challenge for RABV (see Methods). Mice challenged with 10⁵ TCID₅₀ ml⁻¹ uniformly developed signs of disease by day 7 following challenge and were

Table 2. Comparison of survival following challenge with RABV through different inoculation routes

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<th>Route of challenge</th>
<th>Survival at 28 days</th>
<th>Mean days to late-stage disease (± SD)</th>
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<tr>
<td>Intranasal (media)</td>
<td>5/5</td>
<td>–</td>
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<tr>
<td>Intracranial</td>
<td>0/5</td>
<td>4.8 (±0.45)</td>
</tr>
<tr>
<td>Subdermal (footpad)</td>
<td>0/5</td>
<td>6.8 (±0.45)</td>
</tr>
<tr>
<td>Intranasal</td>
<td>0/5</td>
<td>7 (±0)</td>
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Fig. 1. RT-PCR detection of RABV in the lungs, stomach and tongue of mice challenged by intranasal inoculation. Organ samples were removed immediately after i.n. challenge with CVS (30 μl, 10⁵ TCID₅₀). Lanes: 1, 30 μl inocula (CVS) (positive control); 2, no-template (negative) control; 3–8, control mice: tongue (3 and 6), stomach (4 and 7), lungs (5 and 8); 9–14, CVS-challenged mice: tongue (9 and 12), stomach (10 and 13), lungs (11 and 14); M, φX174 HaeIII digest DNA marker (Invitrogen).
groups (reciprocal titres of 3). titre derived from the medium- and low-titre challenge virus neutralization test. No difference was observed in the rabies neutralizing antibodies by a fluorescent antibody four of the survivors from each group were assessed for at day 28 were all FAT negative. Serum samples taken from disease from each group were FAT positive and the survivors of disease worsened. All of the mice that developed signs of day 10 (score of 1) and was sacrificed on day 12 when signs developing aerosol exposure. Groups of 8–10 mice were challenged with a dilution series of RABV by aerosol exposure. Samples were obtained from duplicate animals exposed to media (lanes 1–6) or CVS (lanes 7–12) and RNA was extracted from lungs (lanes 1, 4, 7 and 10), tongue (lanes 2, 5, 8 and 11) and stomach (lanes 3, 6, 9 and 12). The positive (+) and negative (−) controls were the same as those described in the legend to Fig. 1. M, φX174 HaeIII digest DNA marker (Invitrogen).

These data confirmed previous studies of experimental oral and nasal transmission of RABV in mice, which demonstrated that CVS was able to kill mice via the i.n. route (Charlton & Casey, 1979). Similiar i.n. challenge studies with CVS have shown that the virus can be detected shortly after exposure within the olfactory bulb, suggesting that this is the most probable route from the nasal mucosa to the brain (Jenson et al., 1969; Lafay et al., 1991). However, early infection of the trigeminal nerve has also been observed following i.n. challenge (Lafay et al., 1991) and could indicate an alternative route of spread.

**EBLV-2 aerosol challenge of mice**

EBLV-2 is a lyssavirus closely related to RABV and causes a rabies-like disease in bats of the Myotis species, the probable host reservoir for this virus, and in man (Fooks, 2004). Current knowledge of the virulence of this virus is limited and is based mainly on analogous data from RABV (Constantine, 1966; Baer & Bales, 1967; Selimov et al., 1969; Baer et al., 1977).

To investigate the ability of this virus to infect mice by the i.n. and aerosol routes, we challenged two groups of mice with EBLV-2. The titre for this virus was over tenfold less than that obtained for RABV due to a failure of this virus to adapt to low-serum conditions in BHK-21 cells. However, this was equivalent to an effective aerosol dose for RABV. Intranasal challenge with 30 μl inoculum (10^5 TCID<sub>50</sub>) resulted in the development of disease in two mice on days 16 and 18. Both were FAT positive for lyssavirus. Following aerosol challenge, two mice were sacrificed and viral genomic RNA was measured within the tongue, lungs and stomach. No virus could be detected during the first round of RT-PCR (data not shown). However, EBLV-2 could be detected in the lungs of the two challenged mice using a hemi-nested pan-lyssavirus PCR (Fig. 4a) and a hemi-nested PCR designed to differentiate genotype 1 from genotype 6 (Fig. 4b). No virus could be detected associated with the tongue or stomach. Despite evidence of low-level exposure, no mice from the virus-challenged group developed signs of rabies (Fig. 5). The aerosol-challenged group was maintained for 90 days without development of disease (data not shown). In the surviving mice, there was no evidence of seroconversion, suggesting that the mice had been incompletely challenged. Although there was no evidence that aerosolization was effective in the EBLV-2 challenge model, we cannot exclude that this observation was as a result of the low dose used to challenge the mice.

In conclusion, we found that mice (strain RIII) were highly susceptible to RABV (CVS) via all routes of challenge. There was an extended incubation period via the i.n. route compared with direct stereotactic injection of virus into the brain of susceptible mice. We were unable to calculate the precise dose of virus delivered, as the airborne stability of RABV is unknown. However, in this apparatus, an aerosol generated from a suspension containing 10<sup>5</sup> p.f.u. ml<sup>−1</sup> of a stable virus such as *Venezuelan equine encephalitis virus* (VEEV) would be expected to deliver a mouse-received dose of <10 p.f.u. (R. Phillpotts, unpublished data). These data suggest that RABV is highly infectious for mice by the aerosol route. Similar results have been found in a range of
rodent models (Hronovsky & Benda, 1969a, b). RABV was detectable in the lungs and stomach of the airborne-challenged mice, suggesting that the virus was both inhaled and ingested during exposure. This has implications for the site at which the virus enters the host and suggests that RABV entry is likely to occur at the nasal mucosa or possibly at sites within the lung. The greater innervation found within the nasal mucosa favours this site. In a separate aerosol-challenge experiment, aerosols generated from a suspension of EBLV-2 at a concentration $10^{3.5}$ TCID$_{50}$ ml$^{-1}$ failed to infect mice, suggesting that EBLV-2 has a lower virulence in direct comparison with ‘fixed’ RABV strains. In previous studies using VEEV, $10^{3.0}$ TCID$_{50}$ ml$^{-1}$ in the spray fluid delivered approximately 3 TCID$_{50}$ to each mouse (Elvin et al., 2002). In this study, the fixed RABV strain used was relatively pathogenic by the airborne route, as this dose killed 10% of the airborne-challenged mice. In contrast, none of the mice died from the airborne EBLV-2 challenge at $10^{3.5}$ TCID$_{50}$ ml$^{-1}$, with each mouse receiving an estimated 6 TCID$_{50}$. Our data suggest that EBLV-2 is less pathogenic by the airborne route than RABV or is less stable following aerosolization.

In this experiment, there was no evidence of seroconversion in the surviving mice, indicating that EBLV-2 failed to cause even a limited infection. There was evidence of viral RNA within the exposed lung (Fig. 4a, b); however, we must conclude that this was not sufficient to cause infection or induce an immune response.

This study confirms the ability of two lyssaviruses to infect a mammalian model through i.n. exposure. RABV was highly effective at infecting mice by aerosol exposure, emphasizing the need to minimize risk of generating lyssavirus aerosols within the laboratory environment. Moreover, it is essential that all laboratory staff receive pre-immunization against RABV to reduce substantially the risk following exposure to the virus. This study also suggests a risk, albeit low, to individuals in confined spaces that have contact with infected bats, which may be excreting virus in airborne droplets. Our results do not exclude the possibility of transmission of RABV by routes other than a direct bite.

**ACKNOWLEDGEMENTS**

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**REFERENCES**


