Experimental infection of an African dormouse (Graphiurus kelleni) with monkeypox virus

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INTRODUCTION

Suitable animal models are needed to study monkeypox virus (MPXV) as human monkeypox clinically resembles smallpox and MPXV is a zoonotic and potential bioterroristic agent (Parker et al., 2007). The black-tailed prairie dog (p.d), Cynomys ludovicianus, and the thirteen-lined ground squirrel (g.s.), Spermophilus tridecemlineatus, have been evaluated and investigated for their suitability as animal models for MPXV research (Guarner et al., 2004; Langohr et al., 2004; Sbrana et al., 2007; Tesh et al., 2004; Xiao et al., 2005). Husbandry issues associated with these species make them challenging to propagate and maintain in a research vivarium. Female p.d. produce only one litter each year (Hoogland, 2001) while g.s. hibernate for part of the year and produce only one litter each year (Vaughan et al., 2006). Consequently, these species are obtained from their natural habitat where their health status is undefined. Another potential experimental host for MPXV is an African dormouse, Graphiurus kelleni, which is a mouse-sized rodent found in areas of Africa where human monkeypox has occurred. Graphiurus spp. were in the shipment of African rodents that introduced MPXV into the U.S. in 2003. The Centers for Disease Control and Prevention (CDC) determined that Graphiurus spp. and other rodents from this shipment tested positive for MPXV (CDC, 2003). Fortunately, this species has many characteristics similar to laboratory mice which enables their propagation and maintenance in a research setting; in a vivarium, they were determined to be non-seasonally polyestrous, litter bearing, and lacking a need for hibernation. Here we show that G. kelleni (dormice/dormouse) is highly susceptible to lethal infections with MPXV-ZAI-79 and can be used to evaluate prophylactics and therapeutics against MPXV.

RESULTS

Morbidity and mortality of dormice following intranasal infection with MPXV-ZAI-79

Initially, dormice were infected with MPXV-ZAI-79 (1.4 × 10⁴ PFU/dormouse) via a footpad injection route; mortality was 92% (12 of 13 dormice) which occurred during days 7–10 p.i. Considering that smallpox and monkeypox infections are typically initiated by the deposition of large respiratory droplets containing small quantities of virions on the respiratory epithelium, likely in the upper respiratory tract (Fenner et al., 1988; Parker et al., 2007), we infected dormice (n = 55) with a small (10 μl) volume of MPXV-ZAI-79 virus suspension at various doses by the intranasal (i.n.) route. The dormice were highly...
susceptible to morbidity and mortality over a range of virus doses (Figs. 1A,B). Virus doses of 2000 and 200 plaque forming units (PFU)/dormouse were lethal for 100% of dormice with a mean time to death of 7.9±1 and 8.7±1 days and highest mean body weight change of −21% and −14%, respectively. Infection with 20 PFU/dormouse was 63% lethal (mean time to death: 12.0±5 days) with the highest mean body weight change of −6%. Infection with 2 PFU/dormouse was 38% lethal (mean time to death: 12.3±5 days) with the highest mean body weight change of +3%. No mortality was observed at 0.2 PFU/dormouse and the highest mean body weight change was +6%. Morbidity of dormice that died due to a lethal MPXV-ZAI-79 infection was decreased activity, hunched posture, unkempt hair coat, dehydration, and conjunctivitis. There were no significant differences for day of death attributed to gender (female: 9.64; male: 9.58; p=0.973). Mortality was 100% for both genders at a virus dose of 2000 and 200 PFU/dormouse; mortality was 50% and 25% for females and 75% and 50% for males at the virus doses of 20 and 2 PFU/dormouse, respectively. The calculated LD$_{50}$ (Reed and Muench, 1938) of MPXV-ZAI-79 in dormice following i.n. infection was 12 PFU.

For comparative purposes, dormice were infected via the intranasal route with MPXV-COP-58, a MPXV strain originating from West Africa and isolated from imported cynomolgus macaques in Copenhagen, Denmark in 1958 (Von Magnus et al., 1959). This strain is less virulent to humans and primates than MPXV-ZAI-79 (Chen et al., 2005). Dormice infected with various doses of MPXV-COP-58 had similar days of death and mortality rates as dormice infected with similar doses of MPXV-ZAI-79.

**Kinetics of virus spread and progression of disease in dormice following a lethal intranasal MPXV-ZAI-79 infection**

Dormice were infected with 2×10$^4$ PFU of MPXV-ZAI-79 by the i.n. route. At designated times following infection, dormice were sacrificed and nasal lavages, blood and selected tissues were examined for virus titers and/or histopathology (Fig. 2). MPXV-ZAI-79 was first detected in nasal lavages on day 2 p.i. in 6 of 8 dormice (75%). Detectable levels of MPXV-ZAI-79 were noted in the spleen starting day 3 p.i. for 2 of the 8 dormice (25%) sacrificed and in the liver, lung,

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**Fig. 1.** Mortality and mean body weight of dormice following i.n. infection with increasing doses of MPXV-ZAI-79 (n=55; female=28, male=27). (A) Mortality; (B) mean body weight post-inoculation of survivors.
and blood starting day 4 p.i. for 13, 13, and 25% of the 8 dormice sacrificed, respectively; all positive tissues on day 4 were from different dormice. The highest level of MPXV-ZAI-79 was found in liver tissue on day 8 p.i., with 6 of the 6 dormice demonstrating detectable levels (100%). Day 8 was the last day when surviving dormice were available for evaluation. The temporal similarity of virus detection in spleen, liver, and lung suggest that the lung was infected through lymphatic and/or hematogenous spread of the virus and not through aspiration from the upper respiratory tract.

After days 3 and 4 p.i., all dormice clinically demonstrated dehydration and conjunctivitis. At necropsy, upper gastrointestinal hemorrhage (Fig. 3), hepatomegaly, and lymphadenopathy were observed. Histopathologically, infected dormice sacrificed late in infection (starting days 4–5 p.i.) exhibited rhinitis (Figs. 4B–D), lymphoid necrosis in the submandibular lymph nodes (draining site of infection; Figs. 4F–H), spleen (Figs. 4J–L) and thymus. There was also hepatocellular necrosis in the liver (Figs. 4N–P). Hemorrhage was observed in the lungs, stomach, small intestine and elsewhere while areas of necrosis and myeloid hyperplasia were demonstrated in the bone marrow. In dormice found dead (pooled from many experiments), there was often hemorrhage from the nasal cavity as well as hemorrhage in the gall bladder (Fig. 5A) and brain (Fig. 5B) in addition to the lesions described above. In this and other studies, syncytial cell formation was present in nasal mucosal epithelial cells (Fig. 5C), and intracytoplasmic Guarnieri or basophilic (B)-type viral inclusions were noted in the liver (Fig. 5D), spleen, and nasal mucosa.

In summary, histopathological changes found in dormice (pooled from many experiments) which died due to a lethal MPXV-ZAI-79 infection are listed in Table 1 where necrosis and/or hemorrhage affecting many organs were common features.

Use of dormice for evaluating prophylactics and therapeutics against MPXV-ZAI-79

To determine if the dormouse was suitable for efficacy testing of prophylactics and therapeutics, we examined the ability of the Dryvax (smallpox) vaccine and a marketed antiviral, cidofovir, to protect against a lethal i.n. challenge with MPXV-ZAI-79 (Table 2). Dormice immunized 4 weeks previously with Dryvax vaccine were solidly protected from mortality when challenged with $2 \times 10^4$ PFU of MPXV-ZAI-79, whereas dormice in the non-vaccinated control group experienced uniform mortality. Dormice treated with a single dose of cidofovir 4 h following challenge with MPXV-ZAI-79 (pooled results from 3 independent experiments: infecting doses of 75, $4 \times 10^3$, and $5 \times 10^3$ PFU) were significantly protected from mortality, whereas dormice in the vehicle treated control group experienced uniform mortality.

Discussion

Many of the histopathological features of MPXV infected dormice are similar to those observed in smallpox infected humans (Bras, 1952;
Unfortunately, reports concerning histopathology of humans that died due to a MPXV infection do not exist. Martin (2002) reviewed accessible cases of smallpox pathology and review articles from English language journals written during the last 200 years. He classified smallpox into 3 subtypes: variola vera, modified variola, and hemorrhagic/toxic variola. Variola vera comprised >90% of cases with a case fatality rate of 30%. Modified variola represented a variable number of cases with a case fatality rate of 3%; this subtype occurred in previously vaccinated humans. Hemorrhagic/toxic variola comprised <10% of cases with a case fatality rate of nearly 100% and was characterized by lesions with bleeding diathesis and disseminated intravascular coagulation. The MPXV-dormouse model described in this paper most resembles the hemorrhagic/toxic smallpox subtype and provides a more severe disease course than other rodent models of human MPXV disease. The mechanism for severe hemorrhage in dormice infected with a lethal dose of MPXV-ZAI-79 was not determined. Extensive liver and bone marrow necrosis may have resulted in reduced coagulation ability due to the loss of clotting factors and platelets. Additionally, infection and damage of endothelium in affected tissues may have contributed to the multiorgan hemorrhage.

The prairie dog and ground squirrel have been examined as potential rodent models of human MPXV disease. The prairie dog has been experimentally infected with approximately $1 \times 10^5$ to $2 \times 10^5$ PFU of MPXV via intraperitoneal (i.p.) and i.n. routes (Xiao et al., 2005). The MPXV strain utilized (MPX-2003) was isolated from a skin lesion of a human infected with MPXV during the 2003 outbreak in the United States. Neither route of infection with MPX-2003 produced histopathologic abnormalities in the heart, pancreas, kidneys, adrenal glands, or gastrointestinal tract. The i.p. infection route was uniformly lethal and produced necrosis of the spleen, liver, and abdominal adipose tissue in 3 of 3 prairie dogs. The i.n. infection route did not produce necrosis in the spleen or liver in 5 of 5 prairie dogs, but it did produce necrosis of the thymus and mediastinal lymph nodes in some.

Fig. 4. Histopathologic changes in selected tissues of dormice following a lethal i.n. infection with MPXV-ZAI-79; same study as in Fig. 2. (A–D) Nasal mucosa, bar = 20 μm. Submucosal inflammation, mucosal necrosis (arrow) and erosion/ulceration were observed with increasing severity over time; (E–H) submandibular lymph node, bar = 100 μm. Necrosis (arrow) increased in severity and distribution over time; (I–L) Spleen, bar = 20 μm. Increased death of lymphocytes were followed by the development of patchy areas of necrosis (arrow) over time; (M–P) Liver, bar = 100 μm. Large necrotic foci (arrows) of hepatic parenchyma were observed at 8 days p.i., which corresponded to maximal viral load in this tissue.
prairie dogs while necrosis, edema, and hemorrhage were observed in the lungs of 3 of 5 prairie dogs. Death occurred in 3 of 5 prairie dogs infected via the i.n. route; the 2 surviving prairie dogs did not have significant histopathological lesions when their tissues were examined following euthanasia on day 25 p.i.

The ground squirrel was also experimentally infected with approximately $1 \times 10^5$ to $2 \times 10^5$ PFU of MPX-2003 via i.p. and i.n. routes (Tesh et al., 2004). Both routes were uniformly lethal for the 10 ground squirrels tested. Regardless of the route of infection, all ground squirrels exhibited necrosis of the liver and spleen. Additionally, some i.n. infected ground squirrels demonstrated consolidation and interstitial inflammation of pulmonary tissue and necrosis of peribronchial lymphoid tissue. Necrosis in other lymph nodes was also noted. Another study (Sbrana et al., 2007) utilizing ground squirrels compared the pathology following the subcutaneous injection with 100 PFU of either MPX Z79 (same strain as MPXV-ZAI-79) or MPX US03 (same strain as MPX-2003). Both strains were 100% lethal at the same virus dose for the 40 ground squirrels tested, but the MPXV-ZAI-

### Table 1

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Morphological diagnosis</th>
<th>Distribution</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal mucosa</td>
<td>Necrosis</td>
<td>Diffuse</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Liver</td>
<td>Necrosis</td>
<td>Multifocal to coalescing</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Spleen</td>
<td>Necrosis</td>
<td>Diffuse</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Spleen</td>
<td>Hemorrhage</td>
<td>Diffuse</td>
<td></td>
</tr>
<tr>
<td>Liver</td>
<td>Hemorrhage</td>
<td>Diffuse</td>
<td></td>
</tr>
<tr>
<td>Lung</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Stomach</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Lymph node (submandibular)</td>
<td>Necrosis</td>
<td>Diffuse</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Adrenal gland</td>
<td>Hemorrhage</td>
<td>Multifocal to coalescing</td>
<td>100% (16/16)</td>
</tr>
<tr>
<td>Adrenal gland</td>
<td>Necrosis</td>
<td>Diffuse</td>
<td>56% (9/16)</td>
</tr>
<tr>
<td>Pancreas</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>94% (15/16)</td>
</tr>
<tr>
<td>Thymus</td>
<td>Necrosis</td>
<td>Multifocal to coalescing</td>
<td>93% (14/15)</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>Necrosis</td>
<td>Multifocal</td>
<td>91% (10/11)</td>
</tr>
<tr>
<td>Heart</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>87% (13/15)</td>
</tr>
<tr>
<td>Small intestine</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>75% (12/16)</td>
</tr>
<tr>
<td>Gall bladder</td>
<td>Hemorrhage</td>
<td>Diffuse</td>
<td>75% (9/12)</td>
</tr>
<tr>
<td>Kidney</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>63% (10/16)</td>
</tr>
<tr>
<td>Brain</td>
<td>Hemorrhage</td>
<td>Multifocal</td>
<td>56% (9/16)</td>
</tr>
</tbody>
</table>

* All dormice ($n=16$) were found dead; mean time to death was 9.3 ± 1.5 days.

### Table 2

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Mortality</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dryvaca²</td>
<td>0/6 (0%)</td>
<td>$&lt;0.0001$</td>
</tr>
<tr>
<td>Cidofovir²</td>
<td>7/36 (19%)</td>
<td>$&lt;0.0001$</td>
</tr>
<tr>
<td>Vehicle</td>
<td>41/41 (100%)</td>
<td>N/A</td>
</tr>
</tbody>
</table>

* Dormice immunized 4 weeks previously with Dryvax vaccine were solidly protected from mortality when challenged with $2 \times 10^4$ PFU of MPXV-ZAI-79 by the i.n. route.

* Pooled results from 3 independent experiments using an infectious dose of 75, $4 \times 10^3$, or $5 \times 10^3$ PFU of MPXV-ZAI-79. Dormice were treated with a single dose of cidofovir 4 h following i.n. challenge with MPXV-ZAI-79. Cidofovir was administered as described in the experimental protocol.
was considered more virulent. Additionally, frequent nosebleeds/bleeding diathesis were observed in ground squirrels infected with MPX Z79 but infrequently observed in ground squirrels infected with MPX US03. However, Tesh et al. (2004) did not report nosebleeds/bleeding diathesis in ground squirrels infected with a higher dose of the MPX-2003 strain administered via i.p. or i.n. routes.

Other rodents evaluated for their susceptibility to MPXV were the cotton rat and the multimammate mouse/natal mastomys (Wilson, 1993, 2005). Marenkikova et al. (1998) reported the cotton rat (Signmodon hispidus) as very susceptible to MPXV; MPXV strain and dose was not reported. The cotton rat was described as having acute generalized illness with difficulty breathing, coughing, rhinitis, conjunctivitis, and progressive emaciation following the i.n. administration of MPXV; this route of administration caused 50% mortality. Moreover, a more severe disease course and 100% mortality was noted with this rodent following an intranasal inoculation of MPXV.

Takashi Kitamura (personal comm.) described to the World Health Organization in 1978 the necropsy findings of MPXV (reported as strain MP2; 1 × 10⁵ PFU) infected multimammate mice (Mastomys natalenis) as hemorrhagic pleuritis or peritonitis and inflammatory hyperemia in multiple organs following the i.p. inoculation of MPXV.

A non-human primate model for human MPXV and possibly smallpox diseases is the cynomolgus monkey (Macaca fascicularis; Zaucha et al., 2001). Histopathology of this species following lethal doses of aerosolized MPXV-ZAI-79 (inhhaled doses of 1 × 10⁴ to 1.41 × 10⁵ PFU) demonstrated necrotizing lesions at all affected sites which included: lungs, lymph nodes, thymus, spleen, skin, oral mucosa, gastrointestinal tract, and reproductive systems. Hepatic involvement was uncommon. The monkeys died naturally or were killed 9 to 17 days p.i.; cases of natural death were attributed to fibrinonecrotic bronchopneumonia.

When compared to other rodent models which are presently being explored for MPXV research, dormice have many traits similar to those of laboratory mice. This similarity enables the successful propagation and maintenance of dormice within a research vivarium and allows for an accessible supply of dormice with a defined health status. Dormice can easily utilize customary rodent caging and accessories. In our studies, cidofovir significantly protected against a lethal MPXV-ZAI-79 infection and the Dryvax vaccine when administered 28 days prior to a lethal MPXV-ZAI-79 infection was solidly protective. These findings support the applicability of the dormouse in pathogenesis, vaccine, and therapeutic studies of MPXV.

Materials and methods

Animals

The FDA-Center for Veterinary Medicine granted permission for the procurement of dormice from Ohio and Illinois animal vendors (see Acknowledgements) and the use of this species in MPXV research. The Institutional Animal Care and Use Committee at the Saint Louis School of Medicine approved all experimental protocols. All experimental animal procedures were completed in an ABSL-3 facility where mental animal procedures were completed in an ABSL-3 facility where

Cells and virus

The strain of MPXV used in the majority of studies was MPXV-ZAI-79 which was isolated from a fatal human case in Zaire during 1979 (Democratic Republic of the Congo; Zaucha et al., 2001). In one study, MPXV-COP-58, a West African strain of MPXV isolated from cynomolgus macaques in 1958 was utilized (Von Magnus et al., 1959). The virus strains were propagated on BSC-1 cell monolayers which were grown in Eagle's minimum essential medium (EMEM; Bio-Whittaker, Walkersville, MD) as previously described (Chen et al., 1992).

Antiviral compounds and vaccine

Cidofovir was provided as a gift from Mitch Hitchcock of Gilead Sciences Inc., (lot: A201A1, 75 mg/ml). Dryvax vaccine was provided by the CDC.

Pathology examination

After external and internal gross examination of each dormouse carcass, selected tissues were immersion-fixed in 10% neutral-buffered formalin and then routinely processed to generate paraffin sections which were stained with hematoxylin-and-eosin. Tissues selected for microscopic examination: brain, submandibular lymph node, salivary glands, heart, thymus, lung, kidney, adrenal gland, liver, gall bladder, spleen, bone marrow, stomach, small intestine, colon, and pancreas. Sections of nasal tissue were also routinely processed and stained after fixation and decalcification.

Virus assay

Lung, liver, and spleen were ground with tissue homogenizers which contained a measured amount of phosphate buffered saline (PBS; without Ca²⁺ and Mg²⁺) with 1% fetal calf serum. Nasal lavages and blood were serially diluted in the same state as they were collected. Virus infectivity of all tissue, nasal lavages, and blood was measured by plating serial 10-fold dilutions on monolayers of BSC-1 cells, which were overlayed with carboxymethyl cellulose as previously described (Chen et al., 1992).

Experimental infection

For all studies involving the i.n. administration of MPXV, dormice were anesthetized with intraperitoneal injections of a ketamine (9 mg/ml) and xylazine (1 mg/ml) mixture at 0.1 ml/10 g of body weight. Once anesthetized, each dormouse was securely positioned in a dorsal recumbent position on a 60 degree incline and one or both nostrils were inoculated with PBS w/o Ca²⁺ and Mg²⁺ alone or containing the specified MPXV strain via a micropipette. The dormice were maintained in this position for 5 min and then returned to their cages for recovery. Dormice were weighed every-other-day and observed daily for morbidity and mortality over the duration of each study. For the study examining virus spread and disease course, 104 dormice were infected with a lethal dose of MPXV-ZAI-79 (2 × 10⁴ PFU). Groups of 6–8 infected dormice were humanely euthanized every 24 h with carbon dioxide; this was continued through day 8 p.i., a point at which there were no remaining dormice that survived infection. Spleen, liver, blood, lung, and nasal lavages were collected for virus infectivity and tissues from 2 of these dormice (both genders) at each time point were collected for histopathology. Each nasal lavage utilized 0.5 ml of PBS containing 1% fetal calf serum. In order to prevent blood contamination of nasal lavage fluid, the lavage was administered retrograde with a syringe and 18 gauge needle via the nasopharyngeal meatus.

During one study, dormice (n = 13) were infected with 1.4 × 10⁴ PFU of MPXV-ZAI-79 via a left footpad injection. These dormice were made temporarily unconscious due to a very brief exposure to a CO₂/⁰₂ gas mixture immediately before the footpad injection.
Efficacy testing of Dryvax and cidofovir

Dormice in the study testing the efficacy of Dryvax were initially vaccinated with Dryvax (10 μl of a 1:10 dilution containing approximately 1×10^5 PFU) or the vaccine diluent (10 μl) via an injection into the left hind limb footpad. Four weeks later, each dormouse was challenged i.n. with a lethal dose of MPXV-ZAI-79. All dormice were observed daily for morbidity and mortality for the following 21 days.

Dormice in studies involving cidofovir were initially infected i.n. with a lethal dose of MPXV-ZAI-79 and 4 h later they were injected i.p. with 2 mg (0.1 ml of 20 mg/ml solution; 100 mg/kg) of cidofovir or 0.1 ml of sterile saline alone. All dormice were observed daily for morbidity and mortality for the following 21 days.

Statistical methods

Statistical significance of difference was determined by the t test. p<0.05 was considered significant.

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References


CDC (Centers for Disease Control and Prevention), 2003. Update: multistate outbreak of monkeypox — Illinois, Indiana, Kansas, Missouri, Ohio, and Wisconsin. MMWR 52, 616–618.


