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Differential Resource Allocation in Deer Mice Exposed to Sin Nombre Virus

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ABSTRACT
The resource allocation hypothesis predicts that reproductive activity suppresses immunocompetence; however, this has never been tested in an endemic disease system with free-ranging mammals. We tested the resource allocation hypothesis in wild deer mice (Peromyscus maniculatus) with natural exposure to Sin Nombre Virus (SNV). Immunocompetence was estimated from the extent of swelling elicited after deer mice were injected with phytohemagglutinin (PHA); swelling is positively correlated with immunocompetence. After livetrapping deer mice, we determined their reproductive state and SNV infection status. Males were more likely to be seropositive for SNV than females (37% vs. 25%) and exhibited 10% less swelling after PHA injection. The swelling response of females differed with both infection status and reproductive condition. There was also a significant infection status by reproductive condition interaction: nonreproductive, seropositive females experienced the least amount of swelling, whereas females in all other categories experienced significantly greater swelling. The swelling response of males differed with both SNV infection status and reproductive condition, but there was no significant infection status by reproductive condition interaction. Seronegative males elicited greater swelling than seropositive males regardless of reproductive status. In contrast to the resource allocation hypothesis, these results do not indicate that reproductive activity suppresses immunocompetence of deer mice but rather suggest that chronic SNV infection reduces immunocompetence. Sex-based differences in swelling indicate that SNV modulates the immune system of female deer mice differently than it does that of males, particularly during reproduction. We propose that differences in resource allocation between males and females could result from inherent sex-based differences in parental investment.

Introduction
The resource allocation hypothesis proposes that expenditures associated with maintenance functions (e.g., growth, locomotion, and immune system function) are dynamic and at times may be compromised to satisfy more immediate functions (Sheldon and Verhulst 1996; Sinclair and Lochmiller 2000; Nelson et al. 2002). Thus, during periods when the energetic demands of reproduction are primary, other maintenance functions, including immunocompetence, will be compromised, and animals may become more susceptible to disease (Festa-Bianchet 1989; Gustafsson et al. 1994). For example, laboratory studies have demonstrated that several species of small passerine birds expend roughly the same amount of energy when subjected to an immune challenge as during egg production (Martin et al. 2003). Similar patterns have been reported in mammals. For example, captive white-footed mice (Peromyscus leucopus) have been shown to shift their allocation of resources away from reproductive functions when mounting an immune response; their testis mass was significantly reduced after subcutaneous injection with an immune challenge agent (Derting and Compton 2003). Although the hypothesis that reproducing mammals have compromised immunocompetence is pervasive, this idea has not been empirically tested in natural populations of mammals exposed to native disease agents. We evaluated this hypothesis in wild populations of deer mice (Peromyscus maniculatus) exposed to Sin Nombre virus (SNV).

Deer mice are the primary reservoir and vector for SNV. Secondary reservoirs include other Peromyscus species (Otteson et al. 1996) and Neotoma lepida (Dearing et al. 1998). SNV infection in deer mice is chronic; the virus is maintained in the liver, lungs, kidney, and spleen for the life of the deer mouse (Netski et al. 1999). Although this infection is not fatal (Netski
et al. 1999), infected hosts experience an increase in concentrations of proinflammatory cytokines (Herbst et al. 2001), and in response to this chronic infection, deer mice continue to produce SNV-specific antibodies throughout their lifetime (Netski et al. 1999). Thus, SNV presents a constant immune challenge. Over time, persistent activation of the immune system may reduce the ability of deer mice to respond adequately to other types of infections (Klasing and Barnes 1988; Råberg et al. 1998; Borkow et al. 2000).

To measure general immunocompetence of deer mice, we used phytohemagglutinin (PHA), a rapidly acting (<12 h) immune challenge agent. PHA results in profuse cytokine infiltration of local tissue (Martin et al. 2004, 2006) and has been used in numerous ecological studies to assess lymphocyte activity in a variety of animal species (Goto et al. 1978; Williams et al. 1979; Derting and Compton 2003; Martin et al. 2004). Because lymphocytes are effectors of the cell-mediated immune response, PHA has been used to indirectly assess an animal’s cell-mediated immunocompetence. The degree of swelling that follows PHA injection is positively correlated with the level of cell-mediated immune response. Animals that are fully immunocompetent produce the greatest amount of swelling, whereas animals that are immunosuppressed produce less swelling (Bize et al. 2005; Greenman et al. 2005). In general, immunocompetence is defined as an individual’s ability to develop an immune response to infection or disease (Hecht and Shiel 2003). Thus, in this study, immunocompetence is defined as the ability of the animal to mount an immune response to PHA.

Our primary objective was to test the resource allocation hypothesis in wild deer mice in different reproductive states that had been exposed to SNV, using PHA to gauge cell-mediated immunity. We first validated the efficacy of PHA in generating immune responses in laboratory-bred deer mice and then applied this technique to wild populations. We compared differences in immune response between SNV-seropositive and seronegative wild deer mice that were either reproductive or nonreproductive. The resource allocation hypothesis predicts that reproductive animals should be less immunocompetent, as evidenced by smaller immune responses after PHA injection, than nonreproductive animals. Furthermore, because chronic infection is thought to depress the immune system, we predicted that SNV-seropositive deer mice would mount a smaller immune response, as indicated by reduced swelling after PHA injection, than seronegative deer mice. Finally, we predicted that nonreproductive deer mice who were also SNV seronegative would mount the greatest immune response after PHA challenge, whereas deer mice who were both reproductive and SNV seropositive were expected to mount the smallest immune response.

Methods

Laboratory Validation of PHA Technique

Adult (>3 mo of age), nonreproductive, laboratory-bred deer mice were obtained from the Peromyscus Genetic Stock Center (Columbia, SC) and were held under standard conditions at the University of Utah for the duration of the laboratory study. Because a range of PHA doses have been used on a variety of species, we first determined the appropriate dose for deer mice. To this end, we randomly assigned individuals to low-dosage or high-dosage PHA treatment groups. Each treatment group contained nine males and nine females. Before injection with PHA, we measured diameters of left and right hind limbs to the nearest 0.01 mm using a handheld Starrett micrometer (Athol, MA). The low-dosage treatment group was injected in the musculature of the ventral side of the left hind limb with 0.1 mL of a 0.04 mg/mL solution of crystalline PHA (Sigma L9017) diluted in phosphate-buffered saline (PBS). The high-dosage treatment group was injected with 0.1 mL of a 0.08 mg/mL PHA solution. As an internal control, the right hind limbs of all deer mice were injected with an identical volume (0.1 mL) of PBS. Hind limb diameters were measured 3, 6, 12, and 24 h after injection.

Field-Based Immunocompetence Study

In May 2004, deer mice were livetrapped (Sherman Traps) at four study sites near the West Tintic Mountains in the Great Basin Desert of central Utah (Juab County). Each site consisted of a web design of 148 traps (Mills et al. 1999) distributed over 3.14 ha. Vegetative communities of each site were dominated by big sagebrush (Artemisia tridentata) and Utah juniper (Juniperus osteosperma).

After capture, deer mice were weighed, sexed, and uniquely marked with numbered ear tags. Deer mice weighing less than 14 g were eliminated from the study because these animals are considered to be juveniles (Borucki et al. 2000; Calisher et al. 2001). Reproductive condition of individual deer mice was determined by physical examination; females were considered reproductive if they were visibly perforate, pregnant, or lactating, whereas males were considered reproductive if they were visibly scrotal. To determine the SNV infection status of animals, we collected ∼0.2 mL of blood from the retro-orbital sinus of all deer mice. Blood was immediately stored on dry ice and later in a −80°C freezer until being tested for SNV antibodies. After blood collection, diameters of the left and right hind limbs of all deer mice were measured to the nearest 0.01 mm. Left hind limbs of wild deer mice were injected with 0.1 mL of a 0.08 mg/mL PHA solution, whereas their right hind limbs were injected with 0.1 mL of PBS. After PHA injections, deer mice were held in live traps in a shaded, isolated outdoor area for at least 6 h before remeasurement of hind limbs. Hind limbs were remeasured between 6 and 7 h after capture. Once final
hind limb diameters had been determined, deer mice were released at their locations of capture. In total, deer mice remained in captivity for less than 8 h and were directly handled for less than 10 min each. All personnel involved in trapping and handling rodents took precautions for working with animals potentially infected with hantavirus (CDC 1995), and all techniques used in the handling, capturing, and processing of deer mice were approved by the Institutional Animal Care and Use Committee at the University of Utah (05-03011).

Sin Nombre Antibody Detection

Enzyme-linked immunosorbent assays (ELISAs) were used to screen deer mouse blood for immunoglobulin G (IgG) antibodies to SNV. Because deer mice produce virus-specific IgG antibodies continuously after infection with SNV, presence of antibodies is a reliable indicator of SNV infection (Borucki et al. 2000; Botten et al. 2003; Safronetz et al. 2006). In this process, wells of polyvinyl chloride microtiter plates (Dynatech) were coated overnight at 4°C with recombinant nucleocapsid antigen diluted 1:2,000 in PBS. A nonhantavirus recombinant antigen was used as a negative control. After incubation, unbound antigen was removed from wells by washing three times with wash buffer. Deer mouse sera were heat inactivated by placing in a 55°C water bath for 30 min. Heat-inactivated sera were diluted 1:100 in serum-dilution buffer containing powdered nonfat milk, Tween 20, and 10× PBS in a 1:1:20 ratio. The diluted sera solution was added to the antigen-coated wells, and plates were then incubated at 37°C for 60 min. Plates were then washed three times with wash buffer (1:20 Tween and 10× PBS) and incubated at 37°C for 30 min with 100 µL of ABTS Microwell Peroxidase Substrate solution (Kirkegaard and Perry Laboratories; Borucki et al. 2000). Absorbance (405 nm) was recorded with a Versa Max Tunable Microplate Reader (VWR International), and values >3 standard deviations from those of the negative control wells contained on each plate were considered positive for anti-SNV antibodies (Borucki et al. 2000). All steps in the heat activation of sera were performed in a laminar flow hood in a BSL-3 facility at the University of Nevada. The presence of SNV-specific antibodies in adults is strongly correlated with active SNV infections; ELISA results for SNV antibodies have a concordance of about 70% with the presence of viral RNA in blood as determined by reverse-transcriptase polymerase chain reaction (Rowe et al. 1995; Otteson et al. 1996). The remaining 30% of seropositive animals includes adults with antibody titers too low for detection and/or uninfected juveniles with maternal antibodies from SNV-positive dams.

Statistical Analyses

To evaluate the possibility that left and right hind limbs of deer mice were naturally different in size, we used paired $t$-tests to compare differences in diameter between left and right hind limbs before initial injections. We used repeated-measures ANOVA to determine whether injection with PBS alone resulted in hind limb swelling and whether male and female deer mice differed in their response to PBS injection. In this model, hind limb diameter was the dependent, repeatedly measured variable and sex was the independent variable.

For the laboratory validation study, we used repeated-measures ANOVA to determine whether PHA-induced swelling differed between the high and low dosages of PHA or between male and female deer mice across time intervals. In this model, hind limb diameter was the dependent, repeatedly measured variable, whereas sex, dosage, and interaction terms were the independent variables. Differences between individual time intervals were determined using least squares means comparisons, with Bonferroni adjustments for multiple comparisons. Results of these post hoc tests were used to determine the most appropriate time interval to measure postinjection swelling in deer mice held under field conditions.

To determine whether wild deer mice respond differently to PHA injection than laboratory-bred deer mice, we used one-way ANOVA for both males and females. PHA-induced swelling was the dependent variable, and the habitat (laboratory or field) was the independent variable.

To determine differences in the proportion of SNV-seropositive male and female deer mice, $\chi^2$ analysis was used. To determine whether PHA-induced swelling differed between males and females, a one-way ANOVA was used, with sex as the independent categorical factor. Then, for both males and females, independent two-way ANOVAs were used to determine whether PHA-induced swelling differed with reproductive condition or SNV infection status. Differences between individual categories were measured using least squares means comparisons with Tukey-Kramer adjustments for multiple comparisons. Differences in all statistical analyses were considered to be statistically significant if $\alpha \leq 0.05$.

Results

Laboratory Validation of PHA Technique

We found no differences between the left and right hind limb diameters of captive deer mice before injection (right = 3.05 mm vs. left = 3.13 mm; $t_{19} = 1.46, P = 0.17$). Injection with PBS as an intra-animal control did not elicit a swelling response in captive deer mice; their hind limb diameters did not change over consecutive time intervals ($F_{1,19} = 0.32, P = 0.87$). Likewise, there was no difference in response to PBS injection between male and female deer mice ($F_{1,19} = 0.05, P = 0.90$). Therefore, we estimated PHA swelling response by subtracting the diameter of the treatment limb before injection with PHA from the treatment limb diameter after injection.

Deer mice produced considerable swelling in response to PHA injection (within subjects: $F_{5,96} = 360.95, P < 0.01$). There
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was no significant difference in the amount of hind limb swelling between the low- and high-dose injections of PHA ($F_{1,32} = 0.69, P = 0.41$) across time intervals. Female deer mice produced a more than 28% greater swelling response than males (females = 0.23 mm vs. males = 0.18 mm; $F_{1,32} = 12.04, P < 0.01$), and there were no sex by dose interactions ($F_{1,32} = 0.40, P = 0.53$), indicating that male and female deer mice had similar responses to PHA concentrations. Differences in PHA-induced swelling between male and female deer mice prompted us to consider sex as a factor in the remainder of our statistical analyses for the laboratory validation study.

The magnitude of swelling differed across time intervals in both male ($F_{1,32} = 12.5, P < 0.01$) and female ($F_{1,32} = 19.9, P < 0.01$; Fig. 1) deer mice. Hind limb diameters peaked 6 h after PHA injection and at that time were significantly greater than at the time of injection (male: $t_{17} = 29.07, P < 0.01$; female: $t_{17} = 36.20, P < 0.01$).

Field-Based Immunocompetence Study

Over 6 d (888 trap nights), we captured 112 unique adult deer mice—51 females and 61 males. There was no difference between the left and right hind limb diameters of wild deer mice (left = 2.98 mm vs. right = 2.96 mm; $t_{111} = 0.64, P = 0.53$). Injection with PBS as an intra-animal control did not elicit a marked swelling response in wild deer mice; their hind limb diameters were unchanged from the initial to the final measurement ($t_{111} = 0.39, P = 0.70$). Of 112 deer mice captured, IgG antibodies against SNV were detected in 36 individuals, indicating that overall SNV seroprevalence was about 32%. SNV seroprevalence was higher among male deer mice than among females (males = 37% vs. females = 25%; $\chi^2 = 60.02, P < 0.01$). The SNV prevalence we observed is similar to that reported in other deer mouse populations in the same geographic region (Otteson et al. 1996; Douglass et al. 2001; Kuenzi et al. 2001; Mackelprang et al. 2001). Nearly 71% of deer mice sampled were reproductive, including 36 females (71%) and 43 males (70%).

As in the laboratory trials, there were differences in swelling between male and female deer after following injection with PHA. Females produced a 12% greater swelling response than males (females = 0.28 mm, males = 0.25 mm; $F_{1,109} = 4.29, P = 0.04$). Because of these sex-based differences, we considered male and female deer mice independently in the remainder of our statistical analyses.

The PHA swelling response of female deer mice differed with both infection status ($F_{1,66} = 11.99, P < 0.01$) and reproductive condition ($F_{1,66} = 8.20, P < 0.01$). There was also a significant infection status by reproductive condition interaction ($F_{1,66} = 23.56, P < 0.01$). Nonreproductive, seropositive females experienced the least amount of swelling, whereas females in all other categories (reproductive, seropositive; nonreproductive, seronegative; reproductive, seronegative) had significantly greater ($P < 0.05$) and similar amounts of swelling ($P > 0.05$; Fig. 2). The PHA swelling response of male deer mice differed with both SNV infection status ($F_{1,56} = 306.77, P < 0.01$) and reproductive condition ($F_{1,56} = 10.08, P < 0.01$), but there was no significant infection status by reproductive condition interaction ($F_{1,56} = 0.09, P = 0.77$). Seronegative males elicited greater swelling than seropositive males regardless of reproductive status ($P < 0.01$; Fig. 3).

Comparison of Laboratory and Wild Deer Mice

Comparisons of swelling between captive and wild deer mice indicated that deer mice in the field elicited greater swelling than laboratory deer mice. Wild females had 22% greater swelling than laboratory-bred females (0.28 ± 0.01 vs. 0.23 ± 0.01 mm; $F_{1,66} = 9.56, P < 0.01$), whereas the swelling of wild male deer mice was 39% greater than that of laboratory-bred male deer mice (0.25 ± 0.01 vs. 0.18 ± 0.01 mm; $F_{1,66} = 15.95, P < 0.01$).

Discussion

Our primary objective was to test the resource allocation hypothesis in wild deer mice in different reproductive states with natural exposure to SNV. We predicted that reproductive deer mice or those infected with SNV would mount a smaller immune response than nonreproductive or uninfected individuals, as gauged by swelling response after PHA injection. Below, we
Efficacy of PHA in Deer Mice

Our results demonstrate that PHA is effective in generating a measurable swelling response in deer mice, a species in which this technique has not been previously used. We found that maximal swelling occurred in a relatively short period of time (6–12 h), which is in contrast to several published studies in birds that typically measure swelling 24 h after PHA injection (Smits et al. 1999; Granbom et al. 2004; Haussmann et al. 2005). That the immune response generated by PHA injection occurs within a short time frame underscores the utility of this technique and makes the use of PHA particularly attractive for studies where maintaining captive animals infected with biohazardous agents such as SNV is not possible.

Differential Resource Allocation

In general, our results do not support the idea that reproductive activity suppresses immunocompetence of deer mice but rather indicate that a chronic SNV infection can reduce the ability of the immune system to respond to additional challenges. Our results show that there are no appreciable differences in PHA-induced swelling between reproductive and nonreproductive

Sex-Based Differences in Immunocompetence

The results of our study are consistent with the hypothesis that males have lower immunocompetence and therefore may be more susceptible to infection with SNV. Male deer mice were generally less immunocompetent than females; they experienced less swelling in response to PHA challenge in both laboratory and field settings. Furthermore, SNV seroprevalence of males was nearly 1.5 times greater than that of females. In general, disease prevalence is often higher in males than in females, including parasitic, bacterial, and viral infections (Grossman 1985; Möller et al. 1998; Moreno et al. 2001). A higher incidence of infection among males has been observed in several strains of Hantavirus, including Sin Nombre, El Moro Canyon, Puumala, and Seoul viruses (Weigler et al. 1996; Mills et al. 1997; Bernstein et al. 1999). Laboratory studies report that SNV is shed and transmitted less efficiently than other hantaviruses, which suggests that infection with SNV may depend on reduced immunocompetence of the host (Dohm ae et al. 1993; Botten et al. 2002). Our results support this prediction, indicating that in the natural environment, reduced immunocompetence of male deer mice may confer greater susceptibility to SNV infection.
males with the same infection status, and we found similar patterns for SNV-seronegative females. Although laboratory-based studies have shown that hantaviruses do have an immediate impact on deer mouse survival (Hjelle and Yates 2001; Yee et al. 2003), our results in wild deer mice counter this interpretation: reduced immunocompetence associated with SNV infection could render hosts more susceptible to other pathogenic infections, which may directly limit their longevity or fertility. It is noteworthy that the incidence of new hantavirus infections among hosts varies across seasons and is generally highest during periods coinciding with peak reproductive activity (Klein et al. 2002). Higher incidences of SNV infection may result from behavioral changes associated with reproduction, such as increased contact and aggression among conspecifics (Childs et al. 1987; Glass et al. 1998; Hinson et al. 2004; Klein et al. 2004), rather than from differences in immunocompetence.

Sex-based differences in the PHA-induced immune response of deer mice indicate that SNV modulates the immune system of females differently from that of males, particularly during reproduction. Reproductive female deer mice that were also SNV seropositive appeared to have elevated cell-mediated immunocompetence; they experienced significantly greater swelling in response to PHA injection compared to females that were SNV seropositive but nonreproductive. It is important to emphasize that PHA measures only one facet of an animal’s immune system function (i.e., cell-mediated immunity) and also that energetic costs of reproduction could differ between females in different reproductive states (e.g., perforate, pregnant, lactating), which we did not identify. However, our results are in stark contrast to our original prediction that reproductive activity leads to a reduction in general immunity and suggest that interpretation of the resource allocation hypothesis may not be as straightforward as originally expected.

In its most basic interpretation, the resource allocation hypothesis predicts that animals will shift energetic resources to support functions that promote reproductive success; however, the mechanisms that males and females use to promote reproductive success are inherently different. For example, male deer mice have very little parental investment and thus enhance reproductive success by increasing mating frequency (Armitage 1986, 1998). As such, after conception, immunocompetence of the father has little or no influence on the fitness of their offspring. In contrast, female deer mice have much higher parental investment because they must care for offspring from the time of conception until weaning. It is arguable that the investment associated with intestinal helminth infections results in energy conservation and differences between the sexes. J Mammal 79:385–393.


Martin L.B., M. Pless, J. Svoboda, and M. Wikelski. 2004. Im-
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