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# Two Viruses Isolated from Rodents (Clethrionomys gapperi and Microtus pennsylvanicus) Trapped in St. Lawrence County, New York<sup>\*</sup>

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#### Abstract

Four strains of C. gapperi virus were isolated from 3 Clethrionomys gapperi and 47 strains of Microtus virus from 15 Microtus pennsylvanicus and 1 Mus musculus. One of the Microtus strains was isolated from a pool of 20 mites while the others were from rodent tissues. These agents were insensitive to ether and sodium desoxycholate, withstood freezing at -70 C for 3 years and lyophilization without loss of titer, and were not killed when heated at 60 C for 1 hour. Their size as determined by filtration was less than 50 m $\mu$  and greater than 20-35 m $\mu$ . The strains within each group appear to be similar. The illness induced in suckling mice by the C. gapperi agents had a 5-day incubation period followed by prostration and death with a histologic picture of extensive encephalomalacia. The incubation period in mice for the Microtus agents was 9 to 11 days followed by convulsions and death. Histopathology showed meningeal infiltration and necrosis of the molecular layer. No antigenic similarity was detected between the C. gapperi and Microtus viruses by cross complement-fixation test.

#### Introduction

An arbovirus survey of tissues and ectoparasites from live-trapped wildlife collected from both mainland and island sites in St. Lawrence County, New York, was conducted from 30 June, 1964, to 26 August, 1965.<sup>12</sup> Infectious agents were isolated from 3 of 20 *Clethrionomys* gapperi (red-backed mouse), from 15 of 96 *Microtus pennsylvanicus* (meadow voles) and 1 of 43 *Mus musculus* (house mouse). The properties of these virus agents are the subject of this report.

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#### Materials and Methods

Five hundred and sixty-one specimens from ectoparasites (Table 1), brain, kidney, spleen, liver, and blood, or pooled suspensions of the latter 3 tissues were investigated. The methods of collection, shipment, and subsequent storage of specimens, preparation of suspensions from tissue or arthropod specimens, maintenance of virus strains, preparation of hyperimmune sera, and technic of ether and sodium desoxycholate sensitivity tests have been recently described.<sup>1,7-11</sup>

Virus isolation attempts were performed in 1-day-old mice, Nylar strain. Two groups of 8 mice were inoculated intracerebrally (ic) with 0.03 ml of the tissue or ectoparasite suspensions.

In neutralization (N) tests, equal volumes of undiluted or 2-fold serum dilutions were mixed with a constant virus dilution calculated to contain 100 LD<sub>sm</sub>. The ic route of inoculation was used. Methods and interpretation of N tests were the same as recently published.<sup>5</sup>

The complement-fixation (CF) method of Kent and Fife<sup>5</sup> using 5 units of complement was adapted to a micromethod described by Sever.<sup>6</sup> The antigens were prepared according to the sucroseacetone extraction technic of Clarke and Casals.<sup>2</sup>

In heat inactivation experiments, aliquots of 10% suckling mouse brain (smb) preparations were dispensed into glass vials and sealed before submersion in a waterbath held at stated temperatures for varying periods. They were then titrated in 2-day-old or weanling mice.

## **Isolation and Properties**

Fifty-one infectious agents were recovered from 561 specimens. Fcur strains were isolated from 3 *Clethrionomys* gapperi, 2 from the spleen and liver of one animal, the other 2 from pooled blood, and liver and spleen suspensions. Fifty-two of 64 mice inoculated with preparations from which the 4 strains were isolated showed ruffled fur, arched back, lethargy, poor balance, paralysis,

	No. animals	No. ectoparasites	No. isolates
SIPHONAPTERA			
Ctenophthalmus pseudagyrtes	2	3	U
Epitedia w. wenmanni	1	2	0
Nosopsyllus fasciatus	7	13	0
Orchopeas howardii	2	5	0
Orchopeas leucopus	9	12	0
Orchopeas sexdentatus pennsylvanicus	1	3	0
Peromyscopsylla catatina	2	3	0
Unidentified Siphonaptera	1	1	0
ACARINA			
Ixodes species	1	3	0
Unidentified Acarina	10	103+	1*

\* Pool contained 20 mites.

Journal of Wildlife Diseases Vol. 6, January, 1970 TABLE 2. Microtus virus isolated from

and died 9 or 10 days postinoculation. On subsequent passage of 10% smb suspension the incubation period was reduced to 5 days and the LD<sub>20</sub> titers were from  $10^{0.1}$  to  $10^{10.2}$  per gram of brain tissue. These agents are referred to in this paper as *C. gapperi* viruses. The lack of materials precluded any attempts at reisolation.

Of the 47 remaining strains, 46 were isolated from mites, kidney, brain, spleen, liver, and blood from 15 Microtus pennsylvanicus (Table 2). Another strain was isolated from the kidney of a Mus musculus. Five hundred and eighty-four of the 720 mice inoculated with these preparations developed convulsions 12 to 14 days postinoculation and died. The incubation period was reduced on further passage of smb tissue to 8 to 9 days; the  $LD_{50}$  titers ranged from  $10^{7.2}$  to  $10^{9.0}$  per gram. Successful reisolations were made from 5 original tissue suspensions frozen for from 4 to 6 weeks. In this report, these strains are called the Microtus agents.

96 Microtus pennsylva	nicus
10% tissue	No. strains
suspension	isolated
Blood	2
Spleen	3
Liver	3
Kidney	15
Brain	10
Pool (spleen and liver)	1
Pool (spleen, liver and bloc	od) 11
Ectoparasites in 1 ml volume	
One pool of 20 unidentified	1 mites 1
TOTAL	
TOTAL	46

Histologic sections of suckling mice infected with the *C. gapperi* virus showed encephalitis and severe extensive encephalomalacia. Tissue from animals infected with the *Microtus* agent demonstrated meningeal infiltration, encephalomalacia, and necrosis of the molecular

	C. gapperi 64-7855		Microtus 64-7947	
	LD∞/gram F	assage level	LD∞/gram	Passage level
Size. By filtration	<u></u>			
LD∞/gram original	9.1	smb <sub>3</sub>	8.3	smbs
50 mµ*	8.4		8.1	
20-35 mµ†	no activity		no activity	
Sensitivity				
LD <sub>30</sub> /gram control	9.9	smb <sub>2</sub>	8.1	smb₅
Sodium desoyxcholate	9.9		8.8	
LD <sub>30</sub> /gram control	10.2	smb <sub>2</sub>	9.2	smb₅
Ether	10.5		8.8	
Stability				
Original	9.8	smb	8.9	smb₄
Lyophilization	not tested		9. <b>U</b>	smb₄
Stored frozen 3 years				
—70 C	9.8	smb	9.4	smb.
Heat				
Unheated	10.1	smb <sub>3</sub>	8.9	smb。
1 hr 60 C	6.4		>8.4	

\*Millipore Filter Corporation, Bedford, Massachusetts

†Carl Schleicher and Schuell, Keene, New Hampshire

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layer of the central portion of the cerebellum. No nuclear or cytoplasmic inclusions were seen.

The incubation time, symptoms and histology suggest that the *C. gapperi* agents are different from the *Microtus* agents. There may be, however, similarity or identity among the strains within each of the two species.

The C. gapperi and Microtus agents appeared to have similar physical properties, such as size, their reaction to ether and sodium desoxycholate treatment, and the stability of storage at -70 C for 3 years without loss of titer. They differed in their resistance to heat. The C. gapperi strain showed a 4 log<sub>10</sub> loss of infectious titer after 1 hour at 60 C (Table 3).

Host spectrum. Suckling mice were susceptible to the C. gapperi agents by both ic and intraperitoneal (ip) routes of inoculation but to Microtus agents only by the ic route. Both agents infected weanling mice by ic inoculation. With the Microtus strains deaths were irregu-

TABL	E 4. N	eutrali	TABLE 4. Neutralization tests in suckling mice with C. gapperi and Microtus strains	ts in st	ickling	: mice w	ith C.	gappe	ri and M	licrotus	strain	S
					Ĥ	Hyperimmune mouse sera	une mo	ouse a	era			
Virus			C. gapperi	eri					W	Microtus		
strains	Vo. Vo. Virus	64-7855 64-7855 Ser. dil Re	855 Result	LD. Vo.	Pool 452 Ser. dil Re-	452 Result	LD. Vo.	64-7947 Ser. dil Re	047 Result	LD. Virus	Pool 593 Ser. dil Re	593 Result
C. gapperi 64-7855	100	1:2 1:8 8	16/16* 10/16 6/16	199	6	6/8						
Pool 452	316	1:2 1:4 1:8 1:16		316	1:2 1:4 1:8 1:16							
Pool 497	398	θ	13/16									
Microtus 64-7947 18 other strains							199 100- 1000	$\theta$ 1:4 -1:2 1	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			
Pool 593 House mouse strain	200	ø	0/15	200	θ	0/10	166 200	0 0	5/8 16/16	166 200	θ 1:2	9/16
* Number of mice surviving/number of mice inoculated $\theta =$ undiluted	of mice	surviv	ing/numb	er of n	nice in	oculated						

lar at the higher dilutions. Convulsions began 8 to 9 days postinoculation and frequentiy persisted for 10 to 12 additional days.

Guinea pigs infected ic with 0.1 ml of a 10% smb suspension of *Microtus* strain 64-7947 showed no rise in temperature or other signs of infection during a 28-day observation period. Ten-percent chorioallantoic membrane (cam) suspensions from 3 serial passages of the *Microtus* strain 64-7947 on cam of 12day-old embryonated hens' eggs failed to elicit any infection in suckling mice.

Cell cultures. Ten-per-cent smb suspensions of C. gapperi strain 64-7855 and Microtus strain 64-7947 failed to propagate in cell cultures of continuous lines of FL<sup>+</sup>, HEL<sup>+</sup>, or BHK21<sup>+</sup>. The intermediate and terminal passages were checked for virus by ic inoculation of suckling mice.

To rule out ectromelia, footpads of weanling mice were inoculated with 0.03 ml of a 10% smb suspension of *Microtus* strain 64-7947. There were 3 control animal groups: one received 0.03 ml of 10% normal smb suspension; the 2nd, 0.03 ml of diluent, 0.75% bovine plasma albumin: and the 3rd was uninoculated. In 2 experiments no swelling or redness was detected; in the third trial the test

of three Microtus strains					
Sucrose-	Мо	Mouse sera			
acetone	64-7947	Normal			
antigens	4/13/65	5/18/62			
64-7947	1024/16*	<16			
64-8906	1024/16	<16			
64-8912	1024/16	<16			
Normal	<16	<16			

\*Reciprocal highest dilution of serum giving 50% hemolysis/reciprocal highest dilution of antigen giving 50% hemolysis.

group showed slight transient swelling and redness on days 7 to 11.

Antigenic relationships between C. gapperi and Microtus virus strains were determined by N and CF tests. In N tests with hyperimmune sera prepared against 2 C. gapperi strains (64-7855 and pool 452), the 3 C. gapperi viruses appeared similar (Table 4).

The hyperimmune serum prepared with the *Microtus* strain 64-7947 in a 1:2 dilution neutralized not only 199 LD<sub>50</sub> of the homologous strain but 100 to 1000 LD<sub>50</sub> of 18 other *Microtus* 

- +FL = human amnion cell line originally established by Dr. J<sub>φ</sub>rgen Fogh and Rosemary O. Lund (Proc. Soc. Exper. Biol. & Med., 1957, 94, 532-537).
- #HEL = human embryonic lung received August 9, 1962 from Dr. E. V. Davis, Communicable Disease Center Field Station, Phoenix, Arizona.

IBHK21 = clone 13 baby hamster kidney cells received from Dr. Sonja Buckley, Rockefeller Foundation Laboratory, N. Y. C. on April 17, 1964.

	Titers wi	th mouse sera	
Sucrose-acetone antigens	C. gapperi 64-7855	Microtus 64-7947	Normal
C. gapperi — 64-7855	256/16*	<4	<4
Microtus — 64-7947	<4	512/16	<4
Normal	<4	<4	<4

\*Reciprocal highest dilution of serum giving 50% hemolysis/reciprocal highest dilution of antigen giving 50% hemolysis

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agents. Similarity of the strains is also indicated by cross-neutralization tests of *Microtus* Pool 593 with the two *Microtus* sera (Table 4).

The results of the CF test with 3 antigen preparations and serum 64-7947 indicate that the *Microtus* agents are at least similar if not identical (Table 5).

The infectious agent isolated from the kidney of the house mouse was shown to belong to the *Microtus* group of agents and not to the *C. gapperi* (Table 4).

Major antigenic differences between the C. gapperi and Microtus viruses were noted in cross CF tests and are shown in Table 6.

Sera prepared against the following infectious agents failed to neutralize C. gapperi and Microtus strains: eastern and western equine encephalomyelitis, Powassan, St. Louis encephalitis, Maguari, Cache Valley, Flanders, epizootic hemorrhagic disease (EHD) of deer, MM, herpes simplex, Theilers TO, lymphocytic choriomeningitis (LCM), and psittacosis. Immune sera to Colorado tick fever and Q fever also failed to neutralize the *Microtus* agents; sera against Modoc, California encephalitis virus, trivittatus, and reovirus type 3 failed to neutralize the *C. gapperi* strains.

Mouse hyperimmune sera prepared in September, 1965, against *C. gapperi* strain 64-7855 and in April, 1965, against *Microtus* strain 64-7947 were checked for antigenic similarity with several mouse agents by Microbiological Associates, Inc.\*\* No reactivity was noted with antigens of reovirus type 3, Theilers GDVII, and K virus. Both sera reacted in hemagglutination-inhibition (HI) tests with antigen of minute virus of mice (MVM) with titers of 1:80 and 1:40, respectively. In the CF test with mouse hepatitis virus (MHV) antigen, both sera had a titer of 1:80. Two pools of

\*\*Microbiological Associates, Inc., 4733 Bethesda Avenue, Bethesda, Maryland 20014.

Species	Number animals trapped	Site	Trapped Date
Clethrionomys			
rapperi	1	Mainland, Louisville, Wilson Hill Rd.	8/ 8/64
	1	Mainland, Norfolk, O'Brien Rd.	10/29/64
Microtus	1	Mainland, Norfolk, O'Brien Rd.	11/17/64
pennsylvanicus	1	Barnhart Island, Pole 37A	8/ 4/64
	1	Barnhart Island, Pole 37A	8/ 6/64
	1	Barnhart Island, Pole 42C	8/20/64
	2	Barnhart Island, Pole 44B	8/31/64
	1	Barnhart Island, Pole 44B	9/ 1/64
	1	Barnhart Island, Power Dam	8/19/64
	1	Barnhart Island, South Picnic	9/10/64
	1	Barnhart Island, South Picnic	11/17/64
	1	Barnhart Island, Pole 121C	3/ 8/65
	1	Mainland, Louisville, Swamp	10/17/64
	1	Mainland, Louisville, Swamp	10/23/64
	2	Mainland, Wonderland, Route 37	3/ 3/65
	1	Mainland, Wonderland, Route 37	3/ 5/65
Mus musculus	1	Mainland, Brasher, Keenan Farm	6/30/64

# TARIE 7 Transites of animals vielding C conneri and Microtus agents

weanling Nylar strain mouse sera collected in September, 1965, and in July, 1967, from animals of approximately the same age as those used for immunization were submitted to Microbiological Associates for murine virus antibody determination. Again, no reactivity was noted with reovirus type 3, Theilers GDVII, and K virus, but both pools of sera reacted in HI tests with antigens of MVM with titers of 1:160 and 1:1280, respectively. In the CF test with MHV antigen, the titers were 1:10 and 1:20, respectively.

Neutralizing antibodies against agents isolated from either *Microtus* or *C. gapperi* were not detected among human residents of St. Lawrence County in 53 and 32 sera, respectively.

# Discussion

Two different rodent viruses were discovered, one from Clethrionomys gapperi, the other from Microtus pennsylvanicus and Mus musculus. Tissues from 20 other mammalian species totalling 438 animals, also from 201 amphibians and 95 reptiles failed to yield any infectious agents. The C. gapperi viruses were isolated from animals trapped over a 4-month period at 2 different mainland sites. Two animals in the same area were caught 19 days apart (Table 7). The Microtus agents were from animals collected over an 8-month interval; three of the *Microtus* were trapped at Pole 44B on 8/31 and 9/1/64 (Table 7). The persons trapping reported no observations of sick animals.

Hamilton in 1937<sup>1</sup> described among the *Microtus* of New York State an illness with signs of twitching about neck and shoulders, thrusting of hind legs straight backward, followed shortly by death. Neither gross nor microscopic examination with particular attention to the brain revealed any evidence of toxoplasma or other contributing disease organisms. Mice inoculated with our *Microtus* viruses developed convulsions similar to those described by Hamilton. It is not yet possible to classify the agents isolated from *Clethrionomys gapperi* and *Microtus pennsylvanicus*. Their small size and their apparent lack of envelope may place them in either the picorna or the parvovirus group.<sup>a</sup> Determination of the nucleic acid type, DNA or RNA, is essential to gain further information for their classification. Such studies are at present, however, not possible due to the failure of the viruses to propagate in tissue cultures.

The findings by Microbiological Associates of HI titers with MVM antigen and CF titers with MHV antigen not only in the hyperimmune mouse sera prepared with viruses of *C. gapperi* and *Microtus* but also in two pools of sera from uninoculated mice of the same age and strain collected in 1965 and 1967. indicate that these reactions are the result of latent infection of the Nylar mice with MVM and MHV viruses, rather than cross reactions with the *C. gapperi* and *Microtus* agents.

We failed to find serologic evidence of past infections with C. gapperi or Microtus viruses in a survey of a small group of residents in St. Lawrence County.

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#### Literature Cited

- 1. ANDREWES C., and HORSTMANN, D. M. 1949. The susceptibility of viruses to ethyl ether. J. Gen. Microbiol. 3: 290-297.
- CLARKE, D. H., and CASALS, J. 1958. Techniques for hemagglutination and hemagglutination-inhibition with arthropod-borne viruses. Amer. J. Trop. Med. & Hyg. 7: 561-573.
- 3. FENNER, F. 1968. The Biology of Animal Viruses. Academic Press, N.Y. and London. 845 p.
- 4. HAMILTON, W. J., Jr. 1937. The biology of the microtine cycles. J. Agricultural Res. 54: 779-790.
- 5. KENT, J. F., and FIFE, E. H., Jr. 1963. Precise standardization of reagents for complement-fixation. Amer. J. Trop. Med. & Hyg. 12: 103-116.
- SEVER, J. L. 1962. Application of a microtechnique to viral serological investigations. J. Immunol. 88: 320-329.
- 7. THEILER, M. 1957. Action of sodium desoxycholate on arthropod-borne viruses. Proc. Soc. Exper. Biol. and Med. 96: 380-382.
- 8. WHITNEY, E. 1963. Serologic evidence of group A and B arthropod-borne virus activity in New York State. Amer. J. Trop. Med. & Hyg. 12: 417-424.
- 9. WHITNEY, E. 1964. Flanders strain, an arbovirus newly isolated from mosquitoes and birds of New York State. Amer. J. Trop. Med. & Hyg. 13: 123-131.
- WHITNEY, E. 1965. Arthropod-borne viruses in New York State: Serologic evidence of Groups A, B and Bunyamwera viruses in dairy herds. Amer. J. Vet. Res. 26: 914-919.
- 11. WHITNEY, E., and JAMNBACK, H. 1965. The first isolations of Powassan virus in New York State. Proc. Soc. Exper. Biol. and Med. 119: 432-435.
- WHITNEY, E., JAMNBACK, H., MEANS, R. G., and WATTHEWS, T. H. 1968. Arthropod-borne virus survey in St. Lawrence County, New York. Arbovirus reactivity in serum from amphibians, reptiles, birds, and mammals. Amer. J. Trop. Med. & Hyg. 17: 645-650.