REPORT DOC	IMENTATION PA	GE	12
1a. REPORT SECURITY CLASSIFICATION	16. RESTRICTIVE MARKINGS		
25. AD-A268 307 ———————————————————————————————————	3. DISTRIBUTION/AVAILABILITY OF REPORT Approved for public release; distribution is unlimited		
4. P NMR I 93-54	S. MONITORING ORGA	ANIZATION REPORT NUME	SER(S)
6a. NAME OF PERFORMING ORGANIZATION Naval Medical Research Institute 6b OFF CE SYMBOL (If applicable)	7a. NAME OF MONITORING ORGANIZATION Naval Medical Command		
6c. ADDRESS (City, State, and ZIP Code) 8901 Wisconsin Avenue Bethesda, MD 20889-5607	7b. ADDRESS (City, State, and ZIP Code) Department of the Navy Washington, DC 20372-5120		
8a. NAME OF FUNDING/SPONSORING ORGANIZATION Naval Medical (If applicable) Research & Development Comman	9. PROCUREMENT INST	RUMENT IDENTIFICATION	NUMBER
8c. ADDRESS (City, State, and ZIP Code)	10. SOURCE OF FUNDING NUMBERS		
8901 Wisconsin Avenue Bethesda, MD 20889-5606	PROGRAM PRO SLEMENT NO. N.A.	JECT TASK NO.	WORK UNIT ACCESSION NO.
12. PERSONAL AUTHOR(S) Olson PE, Richards AL, Dasch CA and Kennedy CA 13a. TYPE OF REPORT Technical report 13b. TIME COVERED FROM TO 16. SUPPLEMENTARY NOTATION Reprinted from The Journal of I	14. DATE OF REPORT (Y 1993 nfectious Diseases 19		GE COUNT 2 257-258
FIELD GROUP SUB-GROUP rickettsial disc epidemiology, to	ases, Lyme borrelios ck-borne diseases, l		reaction,
Accession For Accession For Accession For Anti-School String Codes Availability Codes Special Special or Speci		S ELEC AUG19 E 93-1921	1993
20. DISTRIBUTION/AVAILABILITY OF ABSTRACT WUNCLASSIFIEDUNUMITED SAME AS RET DIC USERS 22a. NAME OF RESPONSIBLE INDIVIDUAL	21. ABSTRACT SECURITY Unclassific		SYMAOL
Phyllis Blum, Librarian DD FORM 1473 R4 MAR 83 APR edition may be used un	(301) 295-2188	MRL/NMP	(I

All other editions are obsolete.

SECURITY CLASSIFICATION OF THIS PAGE UNCLASSIFIED

References

- Ziegler EJ. Projective antibody to endotoxin core: the emperor's new clothes? J Infect Dis 1988:158:286-90.
- Silva AT, Appeimelk B, Buurman WA, Bayston KF, Chien J, Monoclonal antibody to endotoba core protects might from Escherichia coli sepsis by a mechanism independent of trailor necrosis factor and interleukin-6. J Infect Dis 1990;162—3—9.
- Appelmelk BJ, Verweij-Van Vught AM, Maaskant JJ, Schouten WF, Thijs LG, MacLauri DM, Monoclonal artibodies detecting novel structures in the core region of Szimonella minimuma lipopolysaccharide. FEMS Microbiol Lett 1987;40:71-4.
- 4. Galanus C. Lüdentz O. Westphal O. Preparation and properties of anti-

- sera against the hpid-A component of bacterial hpopolysacchande Eur J Biochem 1971.24 116~22
- Tene NN, Kaplan HS, Hebert JM, et al. Protection against gram-negative bacteria and endotoxemia with human membelonal IgM antibodies. Proc Natl Acad Sci USA 1985;82:1734-4
- Baumgartner, JD. Heumann, D. Germ, J. Weinbreck, P. Grau, Gf., Glauser MP. Association between protective efficacy of anti-lipopolysaccharide (LPS) antibodies and suppression of LPS-induced tumor necrosis factor α and interleukin 6. Compansion of O side chain-specific antibodies and core LSS antibodies. J Exp. Med 1990;171:889-96
- Gearing AJ, Leung H, Bird CR. Thorpe R. Presence of the inflammatory cytokings IL-1, TNF, and IL-6 in preparations of monoclonal antibodies. Psychologia 1989:8:361-7

Failure to Identify Borrelia burgdorferi in Southern California Ticks by DNA Amplification

Colleagues—In the absence of erythema migrans or late systemic sequelae, the diagnosis of Lyme borreliosis after tick exposure remains difficult. Problems include the nonspecific nature of patient complaints, lack of serologic standardization, and difficulty propagating the spirochete in artificial culture medium. Magid et al. [1] recently suggested that prophylactic antibiotics given after tick exposure may be cost-effective in areas where Borrelia burgdorferi is hyperendemic. Knowledge of the prevalence patterns of B. burgdorferi infection in potential tick vectors would be useful in assessing the risk of Lyme borreliosis in exposed patients and the utility of antimicrobial prophylaxis. We used polymerase chain reaction (PCR) to screen Southern California ticks for B. burgdorferi.

Ixodes pacificus is the predominant vector for B. burgdorferi transmission to humans on the West Coast of the United States and is found throughout the coastal mountain ranges of Washington, Oregon, and California. Despite the wide distribution of this vector, well-documented cases of Lyme borreliosis remain rare in southern California. Ecologic factors including enzootic cycles, tick feeding preferences, and nonuniform distribution of infected ticks may be responsible for the low incidence of documented human infection [2-4]. To better define the epidemiology and risk of clinical infection with B. burgdorferi in this area, we examined ticks from multiple locations in rural San Diego County and attempted to amplify spirochetal DNA with PCR primers known to be highly sensitive and specific.

We obtained 1046 adult *I. pacificus* by dragging appropriate habitat on 32 widely dispersed rural sins during 1990–1991. From this sample, 160 specimens (5 from each site) were randomly chosen for DNA amplification studies; assays were also done on a single engorged *I. pacificus* adult that was removed from the skin of a clinic patient. *B. burgdorferi* cultures were obtained from R. A. Wirtz (Department of Entomology, Walter Reed Army Institute of Research, Washington DC). Oligonucle-

otides were synthesized and purified by Synthecell Corp. (Rock-ville, MD). PCR reagents (Perkin-Elmer Cetus, Norwalk, CT) were used as prescribed by the manufacturer with an automated thermocycler device. Nucleotide sequences of PCR primers were defined according to Barbour [5, 5a, 5b].

On initial PCR analysis, two pooled tick homogenates proved negative for *B. burgdorferi*. When the same homogenates were amplified in the presence of exogenous *B. burgdorferi*, bands of appropriate molecular weights for primers specific for *B. burgdorferi* were identified. Subsequent PCR analysis of all tick samples without exogenous *B. burgdorferi* added failed to detect DNA sequences of *B. burgdorferi* (as a positive control, *B. burgdorferi* alone was successfully amplified in each test).

We believe this represents the first use of PCR in screening significant numbers of competent vector ticks in southern California. Our test group (n = 161) represents a reasonably random sample. In adjacent Orange County, researchers tested 359 *I. pacificus* ticks by culture on BSK II medium and found only 1 to be positive (subsequently confirmed by immunofluorescent microscopy using specific monoclonal antibodies) [6]. According to the revised Centers for Disease Control and Prevention case definition [7], there have been no locally acquired cases of human Lyme borreliosis reported by San Diego County (E. Haas Department of Health Services, personal communication). Adjacent Orange and San Bernardino Counties have reported a total of 3 patients with characteristic lesions of erythema migrans; however, the causative organism was not isolated [6].

Patients bitten by ticks are frequently concerned about developing Lyme disease. Clinicians can counsel their patients with reasonable certainty that the risk of acquiring the disease after tick exposure in this area remains low. Our data show a low prevalence of *B. burgdorferi* infection of competent tick vectors in southern California and argue strongly against routine antibiotic prophylaxis of tick-exposed patients. Future studies may be warranted as changing land use and development bring more people into contact with vectors of *B. burgdorferi*.

Patrick E. Olson, Allen L. Richards, Gregory A. Dasch, and Charles A. Kennedy

Department of Internal Metacoc (Infections Discuse), Naval Hospital, San Diego, California Department of Imminology, Naval Medical Research Unit Two, Jazarra, Indonesia, Naval Medical Research Inscreee Hoteltons Discusses, Bethesda, Maryland

Reprints or correspondence: Commander Patrick E. Olson, c/o Infectious Disease Division, Naval Hospital, San Diego, CA 92134-5000

References

- Magid D. S., Skartz B. Craft J. Schwartz JS. Prevention of Lyme disease after the present a cost-effectiveness analysis. N. Engl. J. Med. 1992;327:534-41.
- Brown RN. Lane RS. Lyme disease in California a novel enzootic transmission cycle of Borrelia burgdorferi. Science 1992;256:1439-42.
- Lane RS 1. SelfE. Lyme disease in California: interrelationship of Ixodes
 page 2. Acarr. Ixodidae), the western fence lizard (Sceloporus occidents).
 and Borrelia burgdorfers. J Med Entomol 1989;26:272–8.
- Telford SR & moste SS, Spielman A, Clustering of host-seeking nymphal deer ticks (Trodes dammini) infected by Lyme disease spirochetes (Bosec and regdorferi). Am J Trop Med Hvg 1992;47:55-60.

Clarification of Dietary Risk Factors and Religion in a Botulism Outbreak

Colleagues—We would like to explain why, in an outbreak of botulism in Egypt caused by eating faseikh (salted fish), the majority of patients were Coptic Christians [1]. Faseikh is traditionally eaten on a national holiday, Sham-el-Nessim, but in 1991, Sham-el-Nessim and Ramadan coincided. During the month of Ramadan, most Muslims eat and drink only after sundown and avoid salty food because it makes them thirsty the following day.

Reprints or correspondence: Dr. J. Todd Weber, Enteric Diseases Branch, National Center for Infectious Diseases, Centers for Disease Control and Prevention/MS C-09, Atlanta, GA 30333.

The Journal of Infectious Diseases 1993;168:258. This article is in the public domain.

Therapy with Atovaquone for Cryptosporidium parvum Infection in Neonatal Severe Combined Immunodeficiency Mice

Colleagues—Cryptosporidium parvum is a coccidial protozoan that causes protracted and severe diarrhea in immunocompromised patients, especially in patients with AIDS and in malnourished children in developing countries [1]. Currently, there is no available effective therapy for cryptosporidiosis.

The hydroxynaphthoquinone 2-[trans-4-(4-chlorophenyl)cyclohexyl]-3-hydroxy-1,4-naphthoquinone (atovaquone, 566C80; Wellcome Research Laboratories, Beckenham, UK) has activity against several protozoal pathogens, suggesting its potential usefulness for therapy of cryptosporidiosis. In vitro studies have demonstrated the efficacy of atovaquone against the coccidial

- Barbour AG 1st 2005 and cultivation of I state disease spirochetes. Yale. J Biol Med 1984;27:521–5.
- 5a. Rosa PA, Hegi e D. Senwan TG, Polymerase chain feaction analyses identify two emitter classes of *Boure in Internation*, J. Clin Microbiol 1991;29:524-52.
- 56 Goodman JL, Jusse vich P, Kramber JM, Johnson RC. Molecular detection of personent Borrelia burgdorleri in the urine of patients with active Lyme e-sease. Infect Immun 1991;59:269–78.
- Meyers HB, Morre DE, Gellert G, Euler GL, Prendergast LJ, Barr M, Isolation of Exercise purgeletter from ticks in southern California. West J Med 1992;157

 455-6.
- Centers for Disease Control. Case definitions for public health surveillance. MMWR 1990.39(RR-13):19-21

Coptic Christians are under no such constraint and ate the faseikh, causing this group to make up 79% of the patients, even though they represent only 15% of the population. One Muslim patient in the family case-control study ate fascikh because she was menstruating, which exempted her from the Ramadan fast.

J. Todd Weber, Charles L. Hatheway, Paul A. Blake, and Robert V. Tauxe

Enteric Diseases Branch, Division of Bacterial and Mycotic Diseases, National Center for Infectious Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia

Reference

 Weber JT, Hibbs RG, Darwish A, et al. A massive outbreak of type E botulism associated with traditional salted fish in Cairo, Egypt. J Infect Dis 1993;167-451-4

protozoans Eimeria species and Toxoplasma gondu [2, 3]. In a murine model, 100% of mice that received 100 mg/kg atovaquone survived infection with 5 different strains of T. gondii [3]. Early animal model experiments of atovaquone therapy for cryptosporidiosis, largely unpublished, were reported in summary as equivocal [4].

The severe combined immunodeficiency (SCID) strain results from an autosomal recessive mutation in the C.B-17 inbred strain of BALB/c mice, resulting in the absence of functional B and T lymphocytes [5]. The course of cryptosporidial infection in SCID mice has been reported [6, 7]. Oral infection of 5-dayold SCID mice with 10° C. parvium occysts resulted in cryptosporidial infection that was uniformly fatal within 7 weeks [7]. We chose this model to assess the efficacy of atovaquone for treatment of cryptosporidiosis.

C. parvium oocysts were obtained and prepared for inoculation as previously described [7]. The SCID mice were raised and housed as previously described [7]. Mice had food and water available ad libitum.

Five-day-old SCID mice were inoculated orally with 10^6 C. parvium oocysts in $10^{\circ}\mu$ l of PBS. Two weeks after inoculation, these mice were weaned, and littermates were randomly assigned to either the treatment or control group. The treatment group consisted of 19 mice given 0.1-mL daily intragastric doses of ≥ 100 mg/kg atovaquone (gift of S. Lefon, Burroughs Well-

Presented in part: 32nd Interscience Conference on Antimicrobial Agents and Chemotherapy, Anaheim, California, October 1992 (abstract 1368).

Reprints or correspondence: Dr. Ronald A. Greenfield, Infectious Diseases Section (1111/c), VA Medical Center, 921 NE 13th St., Oklahoma City, OK, 7316-2.

The Journal of Infectious Diseases 1993;168:258-60 © 1993 by The University of Chicago. All rights reserved. 0022-1899/93/€€01-0055\$01.00