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LEGIONELLA IN PUERTO RICO COOLING TOWERS

by

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RECEIVED MAY 2 6 1998 OSTI

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ABSTRACT

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Water samples from air conditioning cooling towers receiving 3 4 different treatment protocols on five large municipal buildings in San ⁵ Juan, Puerto Rico were assayed for various species and serogroups of ⁶ <u>Legionella</u> spp. using direct immunofluorescence. Several water quality ⁷ parameters were also measured with each sample. Guinea pigs were ⁸ inoculated with water samples to confirm pathogenicity and recover ⁹ viable organisms. <u>Legionella pneumophila</u> (1-6), <u>L. bozemanii</u>, ¹⁰ L. micdadei, L. dumoffii, and L. gormanii were observed in at least one 11 of the cooling towers. L. pneumophila was the most abundant species, ¹² reaching 10⁵ cells/ml, within the range that is considered potentially 13 pathogenic to humans. A significantly higher density of L. pneumophila 14 was observed in the cooling tower water that was not being treated 15 with biocides. Percent respiration (INT) and total cell activity (AODC), 16 were inversely correlated with bacterial density. This study 17 demonstrates that Legionella spp. are present in tropical air-18 conditioning cooling systems, and without continuous biocide treatment 19 may reach densities that present a health risk.

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INTRODUCTION

Legionellosis accounts for almost 4% of all patients with atypical 3 4 pneumonia [7]. The disease has been reported in many parts of the 5 United States and Europe. Fliermans [3] has estimated that over 6 200,000 cases a year occur in the United States. Reports of legionellosis ⁷ from the tropics were rare until twenty four people that visited ⁸ St. Croix, U.S. Virgin Islands, acquired legionellosis [11]. Legionella ⁹ pneumophila serogroups 1 and 3, and several new species were isolated 10 from the potable water system in the resort where the patients were 11 vacationing [11]. Recently, studies in Puerto Rico have demonstrated ¹² the <u>Legionella</u> spp. are widely distributed in natural environments and 13 may reach potentially pathogenic densities [10]. Ortiz-Roque and Hazen 14 [10] also demonstrated, from autopsy analysis, that legionellosis in 15 Puerto Rico has an overall mortality of 25%, and that at least 52 cases 16 should be diagnosed every year, yet only 4 retrospective cases have 17 ever been reported. The present study was undertaken to determine 18 the incidence, density, and pathogenicity of <u>Legionella</u> spp. in cooling 19 towers for air-conditioning systems in buildings over fifteen stories 20 high in San Juan, Puerto Rico.

21 (This study was part of the M.S. thesis of A. Negrón-Alvíra at the 22 University of Puerto Rico, Río Piedras, Puerto Rico, 1987.)

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MATERIALS AND METHODS

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Sampling procedures. Samples for the detection of Legionella were taken from the air-conditioning cooling systems of buildings over 5 15 stories high in the banking area in Hato Rey (San Juan), Puerto Rico. 6 The cooling towers were examined for fecal coliforms, Legionella spp., 7 the existence of algae in the tanks, the state of maintenance of the 8 cooling units, and fill material. Samples for bacteriological analysis 9 were collected by grab sampling and placed into sterile Whirl-Pak Bags 10 (Nasco International, Fort Wilkinson, Wis) or sodium thiosulfate bags 11 (Nasco), if the water source was chlorinated. Standard fixation and 12 storage techniques were performed [1]. Time from collection to analysis 13 never exceeded 6 h.

Water quality. Conductivity, pH, temperature, and dissolved Vater quality. Conductivity, pH, temperature, and dissolved Vater quality. Conductivity, pH, temperature, and dissolved Vater quality. Conductivity, pH, temperature, and dissolved National 4041, Hydrolab Corp., Austin, Tex.). Alkalinity and hardness Vater also measured in situ by standard methods [1] using Spectrokits (Bausch and Lomb, Rochester, N.Y.). Other samples were collected in Nalgene bottles, fixed, and transported to the laboratory for further analysis. These fixed samples were tested for nitrites plus nitrates, vater sulfates, phosphates, total phosphorus, and chlorophyll *a* trichromatic using Standard Methods for Water and Waste Water Analysis [1]. In order to have an index of biological contamination, fecal coliform densities were done for every sample. Determination of fecal coliform densities was performed by membrane filtration of triplicate

1 samples, plating on m-FC media, and incubation at 44.5 \pm 0.1°C for 24 h 2 in a block type incubator [1].

Total bacteria cell counts were determined by acridine orange staining (AODC) as described by Singleton et al [12]. At the same time, total bacterial activity was measured in terms of cell ability to reduce INT to INT-formazan during respiration as described by Zimmermann et al. [14]. All methods are as described previously [10].

Direct enumeration of Legionella spp. Ten liters of water 8 ⁹ were collected in sterile polycarbonate containers at each sampling site ¹⁰ incubated with INT for 30 min [14], fixed with formalin, and 11 transported on ice to the laboratory. These samples were centrifuged at 12 5,000 x g for 15 min at 4°C. The pellet and residual water was filtered 13 onto a 0.2 µm pore size, 47-mm diameter membrane (Nuclepore Corp., 14 Pleasanton, Calif.). The filter was eluted by shaking with sample water 15 and 10 μ l aliquots placed into the 8 wells of a toxoplasmosis slide (Cell 16 Line Associates, Newfield, N.J.). The the aliquots were fixed with 17 formalin and the slide subsequently stained with fluorescent antibody 18 to <u>L. pneumophila</u> (serogroup 1-6), <u>L. gormanii</u> (serogroup 1), 19 L. dumoffii (serogroup 1), L. bozemanii, L. micdadei, L. longbeachae, 20 and L. oakridgensis. All sera and antigens were supplied by the U.S. 21 Dept. of Health and Human Services, Center for Disease Control, Atlanta, 22 Georgia. Stained slides were examined with an epifluorescence 23 microscope (Model 16 + IV FL Vertical illuminator, Carl Ziess Inc., N.Y.). 24 The percentage of respiring Legionella spp. was determined using the 25 FAINT technique as described by Fliermans et al. [6]

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Inoculation of guinea pigs. Sample processing and inoculum 2 dosages varied with the total number of organisms (DFA) found. 3 Unfixed water samples were prepared for inoculation into guinea pigs 4 as follows: if the sample contained more than $1 \ge 10^3$ Legionella-like 5 cells/ml, 2 ml was inoculated intraperitoneally; if at least $1 \ge 10^2$ 6 cells/ml but less than $1 \ge 10^3$ cells/ml were present, 3 ml were 7 inoculated intraperitoneally; if less than $1 \ge 10^2$ cells/ml were present, 8 the sample was concentrated by centrifuging the sample at 2,900 x g for 9 30 min, discarding the supernatant, resuspending the sediments in 6 ml 10 sucrose phosphate glutamate buffer, and inoculating 3 ml 11 intraperitoneally, as described by Morris et al. [8].

Five guinea pigs were used in each sampling. One guinea pig was 12 13 used as a positive control, inoculating it directly with Legionella 14 pneumophila (ATCC 33152), and another guinea pig as a negative is control, inoculating it with sample water filtered through a 0.2 µm-pore 16 size membrane filter. Before inoculation, each animal's mean baseline 17 temperature was established from 5 daily measurements. After 18 inoculation the guinea pig's temperature was measured at a 19 predetermined time each day for 7 days. A rise of 0.6°C over the 20 baseline temperature for 2 consecutive days was considered a fever, 21 and febrile animals were sacrificed immediately as well as the other 22 animals with other signs of illness (ruffled fur, watery eyes, prostration, 23 and hypothermia). All guinea pigs were sacrificed at 7 days. The tissue 24 homogenates were examined by fluorescent antibody and inoculated 25 onto media as described below [2].

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Legionella viable counts and isolation. Four liter samples were collected in sterile polycarbonate containers and transported on ice to the laboratory. All samples were than pretreated with acid to reduce background organisms as described by Cherry et al. [2]. Treated samples were than plated on Legionella Agar Base and Legionella Agar Enrichment (Difco Laboratories, Detroit, Mich.). After 2 to 5 days of rincubation in an aerobic and humid chamber containing 2.5% carbon dioxide at 35°C, colonies that appeared light blue to blue-gray in color were considered positive [5]. Isolates were then gram-stained and subcultured to a fresh agar plate and to a blood agar plate that did not contain L-cysteine. Typical isolates were than subjected to iz immunofluorescent staining for confirmation.

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Data analysis. Statistical analysis were done with programs developed for Apple IIe and Macintosh computers. Heteroscedastic data were made more homoscedastic using the appropriate formation prior to analysis. Any statistical probability equal or l7 less than 0.05 were considered significant [13].

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RESULTS AND DISCUSSION

³ Previous studies by our laboratory [10] demonstrated that ⁴ <u>Legionella bozemanii, L. dumofii, L. micdadei, L. gormanii,</u>

5 L. longbeachae, and L. pneumophila were found widely distributed in 6 natural waters of Puerto Rico. The present study has shown that air-⁷ conditioning cooling towers in the tropics can also harbour <u>Legionella</u> ⁸ spp. <u>Legionella</u> spp. were found at all five sites with densities from 10⁴ ⁹ to 10⁵ cells/ml (Table 1). Densities of 10⁵ to 10⁶ cells/ml, are believed 10 to be potentially pathogenic [4]. The densities reported in this study 11 were similar, though slightly lower than those reported for cooling 12 tower waters in temperate areas [5, 9]. L. bozemanii, L. micdadei, 13 L. pneumophila, L. gormanii, and L. dumoffii were isolated from the 14 cooling towers (Table 2). L. longbeachae and L. oakridgesis were not 15 detected in the cooling towers, but were observed in natural waters of 16 Puerto Rico [10]. Only L. pneumophila was found in all 5 cooling towers. 17 L. dumofii and L. gormanii were found in four of the 5 cooling towers, 18 whereas L. micdadei was only found in 2 cooling towers, and 19 L. bozemanii was only found at 1 site (Table 2). The most abundant 20 species was L. pneumophila (40.75%). L. pneumophila serogroups 1 and 21 3 were the most abundant serogroups found, each accounting for 39.4% 22 and 29.6%, respectively (Table 3). The most abundant species found in 23 the potable water system linked to an outbreak of legionellosis on the 24 adjacent island of St. Croix were also L. pneumophila serogroups 1 and 3 25 [11]. Natural waters of Puerto Rico were also shown to be dominated by

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serogroups 1 - 3 [10] as were cisterns on the adjacent island of St.
Thomas [Hazen, unpublished data). This suggests that in the Caribbean
and perhaps in other tropical areas <u>L</u>. pneumophila is the dominant
species of <u>Legionella</u> spp. and that serogroups 1-3 are the dominant
serotypes.

The pathogenicity of the Legionella spp. from each cooling tower was established through guinea pig inoculation and recovery from homogenized tissues of moribund animals. Though all animals that became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation, had isolatable Legionella spp. in their became ill after inoculation is could indicate that the became ill after inoculation for Legionella recovery. Isolation is using media is very difficult due to high levels of contamination [9]. If Indeed, in the present study Legionella spp. could not be isolated be directly from cooling tower water using media due to over-growth by seasts, similar results were obtained previously for natural waters in puerto Rico [10].

The cooling tower which was not being treated by antimicrobial 19 compounds (site B), had the highest densities of Legionella (Table 2). 20 Biocidal treatment of sites A, C, D, and E helped control to some extent 21 Legionella, even though the organisms in site C had a high level of 22 activity (Table 2). Fliermans et al. [4] reported 5-36% respiration for 23 Legionella spp. in water samples taken from freshwater lakes and 24 ponds. In this study the percentage of respiration ranged from 10 to 25 35% for the total bacterial community and from 5 to 30% for

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1 L. pneumophila. At site B, where the highest cell densities were 2 observed, the lowest percentage of respiring cells was observed; 3 conversely, at site C where the lowest cell densities were observed, the 4 highest proportion of respiring cells was observed. The total bacterial 5 population was also more active, as indicated by AODC, in the cooling ⁶ towers which were receiving biocides (Table 2). This suggests that ⁷ biocides reduce the density of <u>Legionella</u> spp. and other bacteria in the ⁸ cooling tower water, but that the remaining population is more active, ⁹ since there is less competition and more resources. It remains to be 10 seen if a more active population of <u>Legionella</u> is also more pathogenic. The presence of pathogenic Legionella spp. in air-conditioning 11 12 cooling towers in the tropics at concentrations high enough to cause 13 disease, especially in the immunocompromised or the elderly, suggests 14 that legionellosis may be under-diagnosed in the tropics. Considering 15 the constant year-round use that these cooling tower receive and the 16 large proportion of the population that may be exposed, monitoring and 17 treatment of these systems is essential for prevention of legionellosis.

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ACKNOWLEDGMENTS

We are especially grateful to Carl B. Fliermans who made many helpful suggestions to the manuscript. This work was supported by the Water Resources Research Institute of the University of Puerto Rico at Mayagüez and in part by Sea Grant R/LR-08-87-THA1 and by Public Health Service grants RR-2657 and RR-8102 from the National Institutes of Health. In addition, portions of the information contained in this article was developed during the course of work under Contract No. DE-AC09-76SR00001 with the U. S. Department of Energy.

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Table 1. Cooling tower water quality by site.

SITES	WTEMP	DO	рН	HARD	NO ₂₊₃	PO ₄	TP	CHLA	%R	%A	FC
A	27 ± 0.5	6.6 ± 0.2	7.1 ± 0.2	76 ± 6.3	1.7 ± 0.2	1.7 ± 0.2	0.6 ± 0.1	8.1 ± 0.2	16.7 ± 1.1	38.9 ± 2.0	11 ± 1.0
В	29 ± 0.3	8.8 ± 0.2	7.9 ± 0.3	40 ± 6.0	1.4 ± 0.1	3.4 ± 0.1	1.7 ± 0.2	6.8 ± 0.2	16.6 ± 1.5	32.4 ± 1.3	66 ± 4.0
с	28 ± 0.6	5.0 ± 0.3	7.1 ± 0.1	43 ± 3.3	5.2 ± 0.1	4.8 ± 0.1	4.1 ± 0.1	8.4 ± 0.2	30.1 ± 3.7	39.7 ± 1.1	10 ± 4.8
D	27 ± 0.1	4.0 ± 0.2	7.2 ± 0.2	30 ± 5.8	4.4 ± 0.1	4.4 ± 0.1	0.5 ± 0.1	8.1 ± 0.2	14.1 ± 2.6	58.9 ± 4.0	7.4 ± 1.0
E	28 ± 0.3	4.3 ± 0.6	7.2 ± 0.6	37 ± 5.8	6.9 ± 0.1	3.9 ± 0.1	0.4 ± 0.1	7.5 ± 0.3	14.0 ± 2.0	37.7 ± 7.0	7.4 ± 1.5

*All values are mean \pm one standard error, WTEMP = water temperature (°C), D O = dissolved oxygen (mg/L), HARD = Hardness (mg/L CaCO₃), NO_{2±3} = nitrites plus nitrates (mg/L), PO₄ = orthophosphate (mg/L), TP = total phosphorus (mg/L), CHLA=chlorophyll *a* (mg/L), %A = percenta₄ of total bacteria active (AODC), %R = percentage of total bacteria respiring (INT), FC = fecal coliforms (CFU/ml).

SITES	TL	LG	IJ	LB	LM	LL	ю	LP	FAINT	GP
A	25 ± 5.1	29 ± 9.0	25 ± 7.0	ND	ND	ND	ND	13 ± 3.5	15 ± 1.5	19/20(2)
В	290 ± 37	ND	27 ± 5.3	ND	11 ± 4.1	ND	ND	110 ± 37	14 ± 2.4	20/20(1)
С	20 ± 3.9	25 ± 6.8	ND	8.2 ± 5.3	ND	ND	ND	13 ± 3.2	22 ± 5.5	15/16(0)
D	22 ± 6.4	28 ± 8.0	20 ± 5.8	ND	ND	ND	ND	14 ± 8.8	11 ± 1.9	15/16(0)
E	19 ± 1.8	15 ± 3.1	12 ± 4.5	ND	12 ± 4.2	ND	ND	1.4 ± 0.9	9.7 ± 1.8	16/16(0)
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Table 2. Density, activity and pathogenicity of Legionella by site.

*All densities are mean \pm one standard error x 10³ cells/ml (n = 4), TL = total Legionella, LG = <u>L</u>. <u>gormanii</u>, LD = <u>L</u>. <u>dumoffii</u>, LB = <u>L</u>. <u>bozemanii</u>, LM = <u>L</u>. <u>micdadei</u>, LL = <u>L</u>. <u>longbeachae</u>, LO = <u>L</u>. <u>oakridgensis</u>, LP = <u>L</u>. <u>pneumophila</u> (serogroup 1-6), FAINT = percentage of LP that were respiring as measured by INT reduction, GP = guinea pig recovery of <u>Legionella</u> spp. number of positive recoveries / number tested (number of fatal infections).

SITE	SEROTYPES								
	1	2	3	4	5	6			
A	1.6 x 10 ⁴	2.6 x 10 ³	1.1 x 10 ⁴	3.0 x 10 ³	1.1 x 10 ³	2.4×10^3			
В	2.3 x 10 ⁴	8.5 x 10 ³	3.7 x 10 ⁴	2.4 x 10 ³	4.5 x 10 ³	2.2 x 10 ³			
C	9.4 x 10 ³	2.0 x 10 ³	7.9 x 10 ³	2.8 x 10 ³	2.7 x 10 ³	3.4 x 10 ³			
D	6.3 x 10 ³	2.4 x 10 ³	6.4 x 10 ³	3.9 x 10 ³	0	7.0×10^2			
E	9.0 x 10 ³	9.2 x 10 ³	0	8.1 x 10 ²	8.9 x 10 ²	0			
Percent									
of total	39.4	10.4	29.6	7.6	6.5	5.9			

Table 3. Densities of Legionella pneumophila serotypes by site.

*All densities in cells/ml by DFA