EFFICACY OF BIOCIDES IN CONTROLLING MICROBIAL POPULATIONS, INCLUDING LEGIONELLA, IN COOLING SYSTEMS

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ABSTRACT

Ten cooling towers were tested to determine possible correlations between Legionella numbers and physical, chemical, biological, and operational parameters. Significant correlations were found between total bacteria and Legionella levels as measured by direct fluorescent antibody techniques. Five biocides were tested for efficacy in controlling Legionella and other microbial populations in four towers. The results showed that most biocides were ineffective at manufacturers' suggested dosages. Only frequent, high doses of organic biocides or low-level continuous treatment with an oxidizing biocide showed promise for long-term control of microorganisms in the towers tested.

INTRODUCTION

Cooling systems, including open recirculating cooling towers and evaporative condensers, are known to harbor microorganisms both in the bulk water and on surfaces. These microorganisms cause a variety of problems, including loss of heat transfer efficiency and deterioration of system components. These systems have also been implicated in the growth and dissemination of pathogenic microorganisms, particularly those belonging to the genus Legionella (Howland and Pope 1983). Early work implied that aerosols from various heat rejection units (cooling towers, evaporative condensers) were responsible for many of the confirmed cases of legionellosis. In one instance it was felt that the role of the aerosol was demonstrated in an outbreak of legionellosis in a hospital in Tennessee (Dondero et al. 1980). The plume from an adjacent tower could have reached the air intakes of the patients' rooms. However, in another case that implicated a cooling tower in an outbreak in Rochester, New York, it was found that the water distribution system of the facility was contaminated with L. pneumophila (Nolte et al. 1984). Many hospital outbreaks of legionellosis have been traced back to the water distribution system (see Muraca et al. [1988] for details), and the role of cooling towers in outbreaks has become much less clear. Other cases have further implicated water distribution systems (States et al. 1987; Stout et al. 1985) and their role in contaminating shower heads (Bollin et al. 1985), hot water tanks (Fields et al. 1989), and potable water supplies (Tison and Seidler 1983).

Several investigators have studied the efficacy of biocides in controlling microorganisms, including Legionella, in cooling tower environments. Soracco et al. (1983) demonstrated that many commercially available biocides kill Legionella and many of them kill the algae capable of supporting Legionella growth (Tison et al. 1980) when tested in a laboratory environment. Braun (1982) and Soracco and Pope (1983) demonstrated, however, that some of these same biocides are not as effective in "real world" cooling systems.

Fliermans and Harvey (1984) suggested that brominecontaining biocides might not be as effective as chlorine in controlling Legionella in open cooling systems. Later work by McCoy and Wireman (1989) found that bromochlorodimethylhydantoin (BCDMH as 1.0 ppm chlorine) was 99.9% effective against L. pneumophila in industrial cooling water. Muraca et al. (1988) state that biocides have been found to be ineffective in "eradicating" Legionella from cooling towers and are "only marginally effective in reducing organism numbers." The literature is divided regarding the case for chlorine, especially in potable water systems. In certain instances, continuous chlorination (1.0 to 1.5 mg/L) and shock chlorination (\geq 50 mg/L) were found to be ineffective (Muraca 1988). Studies by Baird et al. (1984) and Massanari et al. (1984) found that hyperchlorination (i.e., greater than 2.0 mg/L) successfully controlled Legionella. Although chlorine has been found to be effective in some cases, the above literature suggests that chlorine may be contraindicated in some industrial environments.

Pope et al. (1984) demonstrated that ozone, if properly applied, could effect control of microbial populations, including *Legionella*, in a variety of cooling systems. The degree to which general microbial communities in the cooling system can, or should be, controlled is an open question. Most investigators would contend that maintaining a cooling system in a sterile condition is impractical and uneconomical. Instead, the target is to keep the microorganisms under control and to limit their effects on operation of the system. The situation with respect to

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Legionella control is little different. Some investigators (Fliermans and Nygren 1987) have used fluorescent antibody methods to measure Legionella populations in such devices and have proposed a practical "working level" of 10,000 Legionella pneumophila per milliliter. above which serious consideration should be given to cleaning and chemically disinfecting the facility. Other investigators (Morris and Shelton 1990) have used culture methods and have suggested that levels of viable (culturable) Legionella in the range of 10 to 100 per milliliter might be cause for enhanced treatment.

The objectives of the two-phase study reported here were to 21219 .

- 1. determine whether the levels of Legionella in cooling systems could be statistically correlated with other cooling system parameters and
- 2. test the ability of several commercially available biocides to control microorganisms, including Legionella, in cooling systems. E

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MATERIALS AND METHODS

Cooling Systems Studied

The objective of phase I was to determine if significant correlations between Legionella levels and other biological and operational parameters existed. In this phase, 10 sites were sampled. These are listed in Table 1, along with system characteristics. As can be seen, the sites included open cooling towers and evaporative condensers, some of which operated as pairs (two identical units operating in the same facility). All operated on the same make-up water (city of Troy, New York) and all had been on similar chemical treatments (using the same biocides and corrosion inhibitors) for more than 10 years. A wide range of system? parameters (e.g., volume, temperatures) were represented.

Make-up volumes were recorded from meters on the individual units. Event recorders were installed on all towers used in phase II in which biocides were tested. These were to be used with blowdown rates to determine blowdown volumes over time. Unfortunately, these rarely gave accurate readings due to faulty operation of the towers (e.g., dirt and leaves can clog blowdown valves).

It should be emphasized that all other routine chemicaltreatments (corrosion/scale inhibitors) continued in the towers during all phases of the study. Biocides that were in use by the host facility during the study were replaced only. for the duration of the testing. There were approximately three cycles of concentration (i.e., the number of times the solids in a particular volume of water are concentrated [Drew 1979]) in systems tested." 51 1232 · · ?*

Microbiological Analyses

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2. 8 9 3 C maximize Water samples were obtained from the towers, and temperature, pH, and total dissolved solids (i.e., by

Site Number and Tower Designation	Manufacturer 9 and Model	і на Тура	Ľócation	Volume (gal)
1 - CIIE	BAC CFT 2427 FCR	Open	Outside shelter, roof	7000
2 - CIIW	BAC CFT 2427 FCR	Open i	Outside shelter,	27000
3 - Cogs	Marley 222-621	Open	Outside, root	
4 - CC	BAC VCT 375 CR	Open	Outside, ground	7000
5 - Union, .	Marley 7128	Орел	Outside, roof	4. A
6 - Wvls IC	Marley 8202	Open.	Outside, ground	
7 - Lib1 🕄 🗍	BAC VS1-100-1	Closed	Inside, top floor	1100
8 - Lib 2 ¹¹ 1	BAC VS1-100-1	Closed	Inside, top floor	1100
9 - JEC1	BAĆ [®] VS1-200-3 W .1911	Closed /	Inside, ground floor	i1:100
10 - JEC2	BAC VS1-200-3	Closed	Inside, ground floor	5110Q

TABLE 1

- 1 88 V dar o 51 (21.) 1 conductivity) were determined on site using portable probes. Aliquots of each sample were processed on site using serial decimal dilution to an endpoint to determine the numbers of viable bacteria. Acid-producing (APB; facultative anaerobes and anaerobes that produce organic acids) and sulfate-reducing bacteria (SRB) were enumerated in this fashion using commercially available, prepared fluid

media (proprietary media formulations; see Table 3). The remainder of the sample was returned to the laboratory within 30 minutes of collection. Dilutions of the samples were performed in autoclaved and filter-sterilized deionized water, and these were plated onto standard plate count agar to obtain the total viable count. Plating was also done to obtain total viable Pseudomonas spp: counts.

Legionella analysis was done on both unconcentrated and concentrated samples. Concentration was achieved by filtering aliquots (up to 100 mL) of the samples onto polycarbonate membrane filters with a pore size of 0.2 µm. Aliquots of these concentrates were treated with acid and/or heat according to the procedures outlined by Dennis (1988). These samples were plated onto selective and : nonselective Legionella agar plates (i.e., buffered charcoal yeast extract (BCYE) with and without antibiotics [APHA] 1989]). Media from several sources were used and compared. Tavist.

Aliquots of each sample were incubated with INT dye (2-[4-iodophenyl]-3-[4-nitrophenyl]-5-phenyltetrazolium chloride) and respiratory substrates for one hour at room temperature in an attempt to determine the numbers of viable cells in the total microbial and presumptive Legionella populations, Details of these procedures have been described by Fliermans et al. (1981). Total microscopic counts were done by spotting concentrated, unconcentrated, and diluted aliquots of the samples incubated with INT onto

toxoplasmosis.slides and then air drying, heat fixing, and staining them with fluorescein isothiocyanate (FITC). The microbial cells were enumerated using an epifluorescence microscope at 1000× magnification. The number of INTpositive cells was determined by counting the percentage of cells having internal formazan deposits characteristic of cells having active respiratory activity (sufficient to reduce the INT to formazan in the cell) at the time of sampling. Separate aliquots of each sample were prepared as above and stained using FITC-labeled polyvalent antisera (i.e., direct fluorescent antibody method or DFA) to Legionella. Positive and negative controls were included with each set of samples. These slides were viewed (using the epifluorescence microscope) to enumerate the numbers of DFApositive cells with morphological characteristics consistent with designation as Legionella. The numbers of DFApositive cells having a positive INT reaction were also determined. 415

It should be noted that indigenous populations of *Legionella* were used since inoculating the cooling towers with *Legionella* was forbidden by the contract. The disadvantage of this approach was realized prior to initiation of the work: If phase I work showed only low levels of *Legionella* in the systems under consideration, documentation of blocide kill of *Legionella* (specifically) would be unlikely.

Surface' samples for the enumeration of surfageassociated microorganisms (SAM) were obtained by swabbing 4 cm² of the interior wetted surface of the cooling tower. This was processed into a slurry in sterile diluting solution and treated as for water samples.

Algal components were enumerated by direct microscopic observation using both brightfield and epifluorescence (to observe red fluorescence of chlorophyll-con_T taining organisms). Very few algae were observed in the samples. Those found were identified as blue-green bacteria (cyanobacteria) using morphological and -color characteristics.

Enumeration of protozoans was attempted using several methods. Direct microscopic observation, enrichment in tubes containing samples plus nutrient agar blocks, and enrichment on plates inoculated with bacteria as the food source were all tried with little success. Protozoans in a control sample from a local pond were easily observed by all three methods.

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Chemical Analyses

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Aliquots of each sample were provided to a water chemistry laboratory. All samples were preserved and analyzed for the chemical parameters given in Table 2, according to methods outlined in APHA (1989).

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Methods for measuring biocide residuals were available only for bromochlorodimethylhydantoin (BCDMH). This was measured as free chlorine residual, immediately after sampling, using the N,N-diethyl-p-phenylenediamine method (DPD). A list of the biocides used and the treatment levels, as ppm product, is given in Table 4.

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RESULTS

Phase I Correlation of *Legionella* Populations with Cooling Tower Parameters

The phase I effort attempted to correlate the chemical, physical, and biological parameters for the various towers to the levels of *Legionella* in these towers. Table 2 gives the raw data for all chemical and operational parameters measured throughout the first eight months of the study. Table 3 presents the biological data. It should be noted that 10 towers were sampled. This is six more than were to be studied in phase II (biocide testing). The other towers were included in order to broaden the data base and assist in attempts to find correlations between operational parameters and *Legionella* levels, even though it was known that the additional towers would be shut down in the fall and consequently that only a few samples would be obtained from these towers.

The ranges for most parameters were quite broad and therefore gave a data base that should yield reasonable correlations, if such exist, between Legionella levels and other chemical or biological parameters. The exception was pH, which ranged only from 8.0 to 8.9. Many other parameters had reasonably wide ranges but only at quite low absolute values. This was true for nitrate, nitrite, ammonia, and iron. Algae (n up to 45) only had one positive sample and protozoans were not recovered (n' =37). Since there was a reasonable sample size in each case $(n \ge 27)$, low correlations with these parameters indicate that their influence on the Legionella populations was minimal. Statistical analyses were performed using two statistical packages. Pearson pairwise correlation coefficients and stepwise regression analysis using Legionella-DFA (Legionella by DFA method) numbers as the dependent variable all revealed good correlations between total bacterial levels (FITC) and Legionella-DFA levels (r =0.770). This was true for both water and SAM samples. The level of iron in the water correlated positively with total bacterial levels in the water (r = 0.630). Excellent correlations were found between those chemical parameters that are expected to correlate well, e.g., calcium, magnesium, hardness, and alkalinity (r = 0.860 through 0.995). A positive correlation was seen between nitrite levels and Legionella-DFA levels (r = 0.510; with FITC, r = 0.720). However, as discussed above, the values for nitrite were all quite low and thus the interrelationship must be considered with caution. Interestingly, there seemed to, be little effect of temperature on the Legionella-DFA levels (r = -0.1980).1 61.51

When a comparison of surface and water (or planktonic) samples was made, the correlations among the

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Date	Month	Site	Label	Temp	рн	TDS mg/L	Ortho Phosphate ug/L	Total - Phosphate ug/L	Nitrate mg/L	Nitrite mg/L	Ammonia mg/L	Cl mg/L	Ca == mg/L
10/16/89		2	CIIE	20	8.8	280	360	120	0.65	0.01	0.01	46	-74
11/21/89	1	3	CITE	23	1 8.6	e 220	155	1250	0.32	0.02	0.01	22.5	. 38
12/18/89	1	4	CI LE	18	8.9	270	28	1900	1.36	. 0.02	- 0.01	39	53
1/29/90	1	5	CITE	22	8.4	250	265	3120	1.2	0	0.01	35	- 59.7
3/12/90		6 1.	CI IE	20	1 8.2	290	215	2100	2.39	0.01	0.2		67
4/6/90		7	CHIE	40		300	380	190	(1) m (1) mm				
5/11/90		8	CIIE	32	8.1	280							
10/16/89		2 #1	1110	20	8.8	280	360	3370	···. 1.1	-0:01	0.01	- 46	69
11/21/89		3 11	1110	23	8.6	220	155	1250	0.32	- 0.02	0.01	22.5	- 38
12/18/80?			cin cin	18	8 0	270	28	1000		0.02	0.01	30	- 53
1/20/00	1			1 22	.: 0.7	250	245	71700	11.50	10.02	0-01	75	-50.7
7/12/00				22	0.4	200	203	3100	1.2	0.01	0.01	33	59.1
5/12/90				20	0.0	290	213	2100	1 6.34	0.01	0.2	and in the second	01
4/6/90	1	1	CIII	40	1	300	380			+			
5/11/90		8 -1	CIN	32	8.1	280	102	111 11					-
10/16/89		2 14	S Cógs	23	8.7	210	105	120	0.65	0.01	0.01	45	74
10/16/89	1	2	Cogs Sedim	ent	1. 10	20.	2 ^{k - 1} *			F			
10/16/89		2	Cogs Scale	1	11	21					4.5	1.10	44
9/22/89		1	4 . · · C(28	8.6	290	180	200	0.32	0.03	0.01	*****	63.2
10/16/89	1	2	6 CC	26	8.8	e' 390	280	1890	2.2	0.01	0.01	75	126
10/16/89		2	5 Unior	n 1 17,	8.8	400	400	465	2.36	0.02	0.01	77.5	111
9/22/89	1	1	6 WVI	27	8.6	260	190	200	- 1.24	0.01	0.01		87.8
10/16/89		2	6 . WVI	14	8.5	300	320	745	0.38	0.08	0.01	22.5	106
9/22/89	i i	1	7 Lid	1 26.5	8.6	110	1 - 1	105	., 0.21		0.03	10	34.3
10/16/89	-	2	7 Lib	27	8.6	100		40	0.23	0.01	0.1	22.5	51
11/21/89		3	7 Lib	1, 28	8.8	270	21	600	0:2	0:01	0.02	40	52
12/18/89	.1	4	7 Lib	1 19	8.4	280	1	, 20	1 0:2	0.01	~ 0.01	8	- 41
9/22/89		1	B Libi	2 26.5	8.5	180	89	200	0.26	0.08	0.02		49.8
10/16/89		2 '	B Lib	2 23	8.2	240	420	4500	1.13	0.5	0.01	87.5	96.5
11/21/89		3	8 Lib	2 16	8.6	220	130	2750	0.9	- 0.31	A	37	43.5
12/18/89	1	4 1	8 Lib	2 17	8.4	320	19	2900	0.9	0.4	2.08	36	21
9/22/89		1	9 JEC	1 28	8.4	240	1 . 47	200	- 0.67	0.05	- 0.01		96.7
10/16/89:	100	2	9 JEC	1 27	8.7	400	99	4500	2.02	0.01		77.5	107.5
11/21/89		3	9 JEC	1 9	8.8	230	134	2700	0.5	0.08	0:23	- 42.5	64.5
12/18/89	1	4	9 JEC	1 20	21 8:5	150	0	14550	0.5	- 0.08	0.01	16	- 25
1/29/90		5	9 JEC	1 23	8.4	430	48	2430	2.2	0.05	0.02	56	96.7
3/12/90		6	9 JEC	1 29	8.7	110	20	300	4- 0.7	0	0.03		20
4/6/90		7	9	25	8 9	300				week to			20
5/11/90		8	O JEC	1 28	- 8/	150	1.1.00	1.0	- (1 W		12.1	1.0	
9/22/80		1 1	0 160	2 20	0	100		27		- 0-0/	0.01	Printoph	74 5
10/16/90		31. 1		37	0	190	ie d			0.04	0,01	11	20.3
11/21/80		7	JEC.	2/	8.0	110	1	140	0.21	0.01		22,5	- 37
12/19/09			JEC	2 23	8.9	260		8 0	0.38	0.02	- 0:01	-15	26.5
1/20/09		4 1	JEC	22	.8.8	110	28	10	0.38	0.03	0.07	- 15	17
1/29/90		21 1	U : nto JEC	14	8.	917 33 80		10	0.5	D.02	0.02	13	21
5/12/90	-	01 1	JEC:	18	8.3	100	1. 11	20	0.6	0.13	0.02		22
4/6/90		1 1	JEC	2 22	8.4	75		Cert State State	+				

TABLE 2 Chemical and Operational Parameters for Cooling Towers during Phase I Tests

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Date	Month	Site	Label	Mg 1	Fe ma/L	Total Alkalinity mg/L	"Hardness mg/L	TSS	Residue	Volatile Solids	Water Heter gal	Blow- down
10/16/89	2	1	CITE	11.5	0.61	127	202.2	8	370	14	7749180	
11/21/89	3	1	CITE	. 6	.0.61	1 75	119.6	- 4	- 390	190	8038800	1-
12/18/89	4	121	CITE	9.6	0,68	112	171.9	13	800	360	8174200	1.
1/29/90	5	1	CITE	10	0.95	106	- 190.3	4	220	140	8412100	- 22.75
3/12/90	6	-1	CLIE		0.81	118	· · · · · · · · ·	2	600	280	8673000	39.4
4/6/90		1	CLIE	- Colo	1,000	96		1		70	8830900	52.6
5/11/90	- 8	2.1	CIIE	1.11	etter -		10 m m m m m m m m m m m m m m m m m m m				1. Same a car	
10/16/89	/2	2	CIIV	11.5	0.61	126	219.7	25	430	11	7749100	
11/21/89	3	2	CITY	6	0.61	75	119.6	4	390	190	8038800	-
12/18/89	4	-2	CITY	9.6	0.68	112	171.9	13	800	360	8174200	
1/29/90	5	2	CILV	10	0.95	106	190.3	4	220	140	8412100	22.75
3/12/90	6	2	CHIV		0.81	118		2	- 600	280	8673000	39.4
4/6/90	.7	2	CI IV	1.		96			310	70	8830900	52.6
5/11/90	8	. 2	CIIV	- (***	10			the states	a	7. 9 m and	14. IA.	77.2
10/16/89	2	3	Cogs	. 11.5	0.05	141	232.1	10	- 460		in a Tari	
10/16/89	2	3	Cogs Sedimen	nt	r: 4	1 A 1	4 1.5					· · · · ·
10/16/89	2	13	Cogs Scale							and the second	a la cya i la	- 446
9/22/89	1	4	V. CÇ	17.2	0.27	192	228.6	15	- 1844	5. m.	4160270	
10/16/89	2	4	CC	18	0.05	218	388.7	8	690	25	556960	are
10/16/89	2	-5	Union	17	0.41	218	347.2	54	· · Lanerary	.24	** 1 *	
9/22/89	1	6	Wyl	16.4	0.17	246	, 286.8	16	508	v	7184540	
10/16/89	2	0.6	WVL	22.5	0.04	218	357.3	14	570	14		
9/22/89		÷.7	Lidt	6.8	0.04	84	113.6	4	- 92		3396690	5 M
10/16/89	. 5	7	Libl	. 6	0.03	64	152.1	18	160	4	4304350	
11/21/89			Lib1	8.5	0.06	s 98	164.8	-3	580	320	4451420	
12/18/89	514	27	- Libl	8.4	0.96	42	137	- 6	1360	360	4473460	145
9/22/89	1	8	: Lib2	8.6	0.08	168	159.8	- 2	504	Pier I	5162280	м н
10/16/89	. 2	8	Lib2	/15	1.17	196	302.7	16	600	10		
11/21/89	3	- 8	Lib2	7.5	0,147	102	139.5	2	500	330	7188367	11-17-16-1
12/18/89	. 4	8	Lib2	.3.6	0.07	98	, 67.3	. 8	520	240	7188100	-
9/22/89	-11	9 اد	JEC1	14.6	0.07	172	301.6	8	592	1.44	53785	
10/16/89	2	-; 9	JEC1	:17.5	0.05	198	. 340.5	. 4	160	6	3419060	
11/21/89	3.3	9	JEC1	10	0.08	134	197.2	0	500	. 140	3465445	
12/18/89	- 4	. 9	JE01	4.8	0.08	66	82.2	Fry 3	360	240	3472870	
1/29/90	5	119	i JEC1	15.8	10.1	. 95	306.5	; 9	, 360	240	3532159	10.75
3/12/90	6	1.9	JEC1		0.04	6		2	220	120	3639281	72.8
4/6/90	7	. 9	JEC1			84	1	Para .	370	110	3965149	86.7
5/11/90	- 8	1.9	JEC1	1.19.1		1 A 1		2171			3789690	110.
9/22/89	21	: 10	JEGS	5	0.04	60	86:8	2.5	188		627264	and the
10/16/89	2	. 10	JEC2	5.5	0.01	· - · 60	115	560	10	10	5291110	1
1.1/21/89	. 3	10	JEC2	. 4	0.03	44	:on 82.6	-0	340	. 190	5454568	
12/18/89	4	10	JEC2	c 3.6	0.07	42	57.3	1	560	480	5471055	المشدوي
1/29/90	n - 5	. 10	JEC2	3.4	5 0 .11	40	66.4	. 2	. 60	. 60	5505768	1.1
3/12/90	6	10	JEC2		0.03	.42		: 4	260	20	5560476	3.0
4/6/90	7	10	JEC2		A	58		A 11-4-1	150	30	5604106	7 .

TABLE 2 (continued) Chemical and Operational Parameters for Cooling Towers during Phase I Tests

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		1	4.5	3.04		-1	WATER		SAMPLES			ent
Date	Month	Site	Label	APB	SRB	Plate Count	Pseudocel	- FITC	LDB/FA	INT	Algae	Protozoa
1.11		+- +-		per ml	. per ml	per mi	per ml	per ml	· per ml	per ml	per ml	per m
0/16/89	2	1	1 CIIE	1.0E+02	1.0E+01	1.1E+04	1.8E+03	1.9E+06	1.3E+05		0	
1/21/89	3		1 CIIE	1.0E+02	1.0E+01	4.8E+03	4.3E+02	3.2E+04	3.0E+04	!	0	1
2/18/89	4		1 CITE	1.0E+03	1.0E+02	9.3E+03	2.0E+00		1.5E+02		. 0	
/29/90	. 5		1 CIIE	1.0E+02	1.0E+01	5.0E+03	0.0E+00		2.5E+03		- 0	-
/12/90	. 6		1 CIIE	1.0E+01	0.0E+00	1.1E+05	2.0E+00		1:0E+02		0	
/6/90	7		1 CITE	1.0E+02	0.0E+00	1.3E+05	0.0E+00	tain-	2:5E+02	1-	0	:
/11/90.	8	5'	1 CITE	1.0E+02	0.0E+00	6.3E+04	1.0E+01		-	- 14.80 0 1 mag.	0	
0/16/89	. 2		2	1.0E+02	10.0E+00	2.1E+04	2-4E+01	2.4E+06	4-8E+04	11.1 C	0	
1/21/89	3		2 0119	1.0E+02	1.0E+01	3.8E+D3	4.3E+02	3.2E+04	3.0E+04	4	0	1. L
2/18/80	4	31 0	2 0114	1.0E+03	1.0E+02	9.3E+03	2.0E+00		1.5E+02		-0	* Aur
/20/00 : *	5	1.4	2	1.05+02	1 05+01	5 0E+03	0.05+00		2 5E+03	1	0	*Mar > A share
(12/00)		1.	2 0110	1 05+01	0.05+00	1 16405	2.05100		1 05+02		0	
16100	. 0		2 0110	1 05+02	0.05+00	1 36405	0.05+00		2 55+02		0	
(11/00	1 0			1.05+02	0.05+00	A 3640/	1 05+00		2.JETUZ		0	-
0/16/90	. 0		Z CODE	1.00002	1.05+01	1.05+04	7.05+00	1 45204			0	-
0/16/09	2		3 Coop Codi	1.02402	1.00+01	1.02404	1.0E+00	1.00700	4.02+04		0	
0/16/29	2	-V1	3 Cogs Sealing	ant .	11 . 1- 1		1 1 1 1 1 1 1				- 0	
0/22/80	1		/ co	1 05+02	1 05+02	1 25+03	1 25403	1 25+03	/ 25+03	/ 205+03		
0/16/80	1 2		4 60	1.00+02	0.05+00	1 05+04	4.2E+03	0 45+05	-2 05+04	4.202703	0	
0/16/09/	2		SUL Union	1.05+02	0.05+00	0 75+03	5 (5402	1 35+06	1 65+04		0	
1/22/80	1		A INI	1.00+02	1 05+05	7 75402	2 25401	T. 5E+00	1.00+04	1 005+01		
0/16/80		-	6 WVL	1 05+03	0.05+00	5.05402	1 05402	1 65+06	-8-05403	1.002+01	0	
10/10/09	1 4		7 1	1.00+02	1 05+01	1 35+04	1.02402	5 05+0/	6.0E+03		-0	
0116/00				1.05+01	0.05+00	1.52+04	6 75+01	9-05-05	4.02+03		0	
11/21/80			7 LIDI	1.00+01	0.00+00	2 /5407	2.55+01	2 45+04	4.0E+04		0	
11/21/09	3	1	7 LIDI	0.05.00	0.00+00	2.4E+03	2.5E+01	2.02704	1.00.02		0	
12/10/09		11		0.02+00	0.0E+00	0.0E+01	0.02+00		1.00402	1 005-04	0	-
1/22/89		1	8 + L102	1.02+02	1.0E+02	1.0E+00	1.1E+01	1.42+00	1.000	1.002+01	- 0	
10/16/89	2	1	8 L102	1.0E+02	1 0.0E+00	· 5.3E+04	2.22+03	9.02+00	1.22+05		0	<u> </u>
11/21/89	3	31	8 L102	1.02+01	1.02+01	8.8E+05	7.02701	4.02+04	4.02+04		0	
12/18/89	4	11	8 J LIDZ	1.0E+02	1.02+01	1.02+04	2.0E+01	F 05.05	- 1 05.07	4 005 -04	0	
122/89	1		Y JECT	1.02+03	1.0E+03	0.2E+U3	1.02+00	5.9E+05	1.0E+05	1.00E+01	0.1	
11/21/00	1	1	JECT	1.00+01	1.05+00	1.1E+04	1.02+00	1.52+06	2.40+04		0	
11/21/89		1	JEC1	1.0E+02	1.0E+02	8.3E+02	1.0E+01	2.0E+04	5.9E+04		0	
1/20/00	4		JEC3	1.0E+02	1.02+01	3.4E+03	0.00+00		3.0E+02		0	Transaction of the
1/29/90		1	JEC1	0.02+00	0.02+00	3.22+04	0.0E+00		1.02+02			
12/901	0		y JECI	1.06+02	1.05+00	2.52+03	0.0E+00		2.25+03		- 0	
10/90			JEC1	1.0E+02	1.02+02	1.3E+04	0.0E+00	1440	2.22+04		.0	
0/11/90	. 8		JEC1	1.0E+01	1.0E+01	1.7E+04	10.0E+00	0 10.00	1 00.00	2 705.04	0	-
122/89	+		JEC2	1.0E+01	1.0E+04	1.2E#03	2.0E+00	9.6E+05	1.0E+03	2.30E+01	0	
10/16/89	2		JEC2	1.0E+03	0.0E+00	1.6E+03	3.0E+03	1.6E+06	2.0E+04		- 0	100
11/21/89	3		JEC2	1.0E+01	1.0E+01	1.2E+04	1.0E+01		-4.8E+03		0	
12/18/89	4	1	IO ; JEC2	1.0E+01	1.0E+01	5.6E+03	0.0E+00	î	5.0E+01	1. S. S. S.	0	
1/29/90	40. 5	1 1 3	IO JEC2	1.0E+01	0.0E+00	1.5E+04	0.0E+00	· · · · · ·	0.0E+00		0	1.00
3/12/90	6	1	IO JEC2	. 1.0E+05	1.0E+02	5.7E+05	0.0E+00		5.0E+01	1.	0	
4/6/90	7		10 JEC2	: 1.0E+03	1.0E+03	2.0E+03	0.0E+00		1:0E+02	A	- 0	

			TABLE	3			
Indianal	Daramatare	for	Cooling	Toware	during	Phace	I T

1.4		1.1	12	11	(4)		SURFACE	· /**	SAMPLES	(A) 4	 ***/* 1 	
Date	Month	Site	Label	APB	SRB per cm2	Plate Count ; per cm2	Pseudocel per cm2	FITC per cm2	LBD/FA	INT Der cm2	Algae per cm2	Protozoa per cm2
10/16/89	2	1	CITE	1.0E+05	1.0E+04	1.8E+04	- 2.5E+01	1.9E+07	2.2E+05		0	0
11/21/89	3	1	CITE		1	and the second s						
12/18/89	4	1	CITE	1.0E+03	1.0E+02	4.4E+04	0.0E+00		1.5E+02			. 0
1/29/90	5	- 11	CLIE	1.0E+03	1.0E+03		0.0E+00		5.5E+02	det 1		0
3/12/90 ;	6	.1	CITE	1							0	- 0
4/6/90	. 7	10	CIJE	1.0E+06	1.0E+02	7.3E+05	0.0E+00		3.0E+02		0	- 0
5/11/90	. 8	1	CITE				The second second				0	
10/16/89	2	2	CHIN	1_0E+04	1.05+03	1.4E+04	1.0E+01	1.35+07	4 DE+04		0	
11/21/89	-3	.2	CITY					1			0	
12/18/89	5.4	2	CITY	1.0F+04	1.0E+04	1.4F+84	0.0E+00	1 5 6 2	5 0E+01			. 0
1/29/90	5	2	CIT	1.0E+03	1.0E+03	1-5E+04	0.0E+00	- Carl	7-6E+03			- 0
3/12/90	716	12	CLIV	1102.05	1102.05	1152.04	0.02.00			5 A 11	0	0
4/6/90	7	:2		1.0E+03	1.0E+03	9:3E+04	0.0E+00	177.0	3.5E+02		0	0
5/11/90	8	2	VII3		1102 05	7152.001		· · · · · · · · · · · · · · · · · · ·	3132.02		-0	- 0
10/16/89	2	3	· Cogs	1.0E+04	0.0E+00	7.3E+03	1.0E+01	2.7E+06			0	0
10/16/89	2	3	Cogs Sedim	1.0E+05	1.0E+05	1.0E+05	1012	1.6E+07	2.4E+06		0	0
10/16/89	.2	3	Cogs Scale	1:0E+05	1.0E+05	2.0E+04	1.0E+01	6.7E+06	-1.5E+05		- 0	0
9/22/89	1	4	· 17 1CC	0.0E+00		· 1.2E+05	1.0E+00	4.2E+06	1:3E+05		0	0
10/16/89	2	. 4	CC	1.0E+03	0.0E+00	5.2E+03	1.0E+01	9.6E+05	- 2:0E+04		0	0
10/16/89:	'2	- 5	Union	- 1 - 155	4174 P	- 12. Aut -	3 fat				0	0
9/22/89	:+1	516	i WVI		Terest	3.3E+03	0.0E+00	2.3E+06	2.7E+04	9	0	0
10/16/89	. 2	6	WVI	1.0E+02	D. 0E+00	1.8E+04	1.0E+01	2.1E+06	1.5E+05		0	0
9/22/89	1	7	Lid1	1.0E+02	1.0E+03	2.5E+04	1.0E+00	2.7E+06	1.3E+05		0	- 0
10/16/89	. 2	7	Libl	1.0E+02	1.0E+02	6.7E+04	4.7E+03	3.3E+07	8.7E+05	11. A	0	- 0
11/21/89	: 3	7	Lib1	0.0E+00	0.0E+00	1.2E+05	1.0E+02	1.4E+05	8.0E+03		0	0
12/18/89	14	: 7	Libl	0.0E+00	0.0E+D0	7.6E+04	0.0E+00		1.5E+02	111 a. + 111-	0	0
9/22/89	1	8	Lib2) 0.0E+00	0.0E+00	7.DE+02	0.0E+00	3.2E+05	7.2E+03		0	0
10/16/89	2	. 8	Lib2	1.0E+02	1.0E+02	2.4E+04	1.0E+01	6.7E+07	2.7E+06	-1	0	0
11/21/89	3	8	Lib2	1.0E+02	1.0E+02	1.4E+04	1.0E+02	2.0E+05	7.2E+04		0	0
12/18/89	: 4	8	Lib2	1.0E+02	0.0E+00	2.0E+04	0.0E+00	1		Ser. 25	0	0
9/22/89	11	9	JEC1	14			Sec. Same			* m.m. 1	0	0
10/16/89	2	9	JEC1	0.0E+00	0.0E+00	8.4E+02	-1.0E+01	1.3E+07	6.0E+05		0	- 0
11/21/89	3	9	JEC1	1.0E+02	1.0E+02	1.7E+03	1.0E+02	6.0E+04	2.4E+04	2.44	0	0
12/18/89	14	9	JEC1	1.0E+03	n 1.0E+02	5.4E+04	0.0E+00		6:0E+02		0	0
1/29/90	5	: 9	JEC1	1.0E+02	1.0E+02	3.8E+05	0.0E+00		0.0E+00	-	0	C
3/12/90.	6	9	JEC1	1.0E+03	1.0E+02	1.0E+05	0.0E+00		- 6.3E+03	÷.	- 0	C
4/6/90	7	9	JEC1	1.06+02	0.0E+00	6.2E+04	0.0E+00	1.1	1.2E+04		0	0
5/11/90	: 8	. 9	JEC1	1.00		-				Sec. 1.		1
9/22/89	1	10	JEC2		1 1 1	1.1E+04	0.0E+00	1.9E+05	7.2E+03	100		~
10/16/89	2	10	JEC2	1.0E+02	1.0E+02	2.5E+03	.3.2E+03	9.3E+11	3.0E+05	1.1.1	ing .	
11/21/89	- 3	10	JEC2	1.0E+02	1.0E+02	1.0E+02	1.0E+02		4.8E+04			
12/18/89	4	10	JEC2	1.0E+04	1.0E+04	1.0E+04	0.0E+00	A.	5.0E+01			
1/29/90	5	1 10	JEC2	1.0E+04	1.0E+03	1-0E+04	0.0E+00		0.0E+00			
3/12/90	6	10	JEC2	1.0E+03	1.0E+02	1.0E+03	0.0E+00	4 9	5.0E+03			140 3
4/6/90	7	10	JEC2	1.0E+05	1 0E+0Z	1.05+05	0:0E+00	1.1.1.2.4	- 9.0E+03			
	the second s	10	arce!	1.02.05	1.00.04	1.02.00	0.00.001	the second se	1.02.05			

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TABLE 3 (continued) Biological Parameters for Cooling Towers during Phase I Tests

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Biocide	Concentration of Product (ppm)
Disodium cyanodithioimidocarbonate + potassium N-methyldithiocarbamate	35-454
Polymeric quaternary amine	80
Isothiazolin	150
Glutaraldehyde	150
Bromochlorodimethylhydantoin	0.1-10

TABLE 4 Biocides Tested in Phase II

various measured populations (FITC total count, plate count, pseudomonads) were generally low (r < 0.4). The surface and waterborne populations of *Legionella* by DFA showed a reasonably good correlation (r = 0.798). There was no clear trend regarding the size of the surface and planktonic populations. For example, surface-associated and planktonic *Legionella*-DFA values were dominant an equal amount of the time.

One important conclusion from the study was that each tower is essentially unique. This is even true for the two pairs of towers that were included in this study. It should also be remembered that all of the towers in this study had been using the same water source and chemical treatments for at least the last 10 years. The consequence of this is that "lumping" the data from all the towers together, although useful in trying to find general correlations between Legionella and other parameters, can obscure the relationships seen when individual towers are thoroughly examined and tested over fairly long time periods (weeks to months). Examples of this are shown in Figures 1 and 2. As can be seen in Figure 1, the JEC 1 tower, which operates all year, had fairly constant levels of viable (plate count) bacteria throughout the eight months, including the winter. The levels of Legionella-DFA in this tower decreased in the colder months and the level of iron was fairly constant at 0.1 ppm. Figure 2 demonstrates that a tower that operates all year, but in which the fans are not operated during cooler months and there are high levels of cyanobacteria (blue-green bacteria) only in the warmer months, had a quite different picture. The levels of platecount (viable) bacteria were fairly constant throughout the first five months and then increased in the spring, whereas the Legionella-DFA lagged behind this increase in total viable cells. This is probably due to the fact that this was the only tower having significant growth of blue-green bacteria. The organisms in these towers grow principally. in the form of "mats" on the sides of the tower. Only when the weather gets warm enough for the fans to be 100 operated do the blue-green bacteria grow well. As the authors and colleagues have demonstrated (Tison et al. 1980), cyanobacteria often support the growth of Legionella. Therefore, the summer increase in Legionella in this tower (the tail end of which was seen in September and October) is probably related to the accumulation of bluegreen bacteria and not to the total viable bacteria counts. It should also be noted that the levels of iron in this tower are essentially constant but are almost 10 times the concentration seen in the tower shown in Figure 1. In spite of this, the levels of Legionella-DFA are similar in the two towers.

Other than the significant accumulation of cyanobacteria in the CII tower, algae were actually detected only one other time in the JEC tower. It must be remembered that these towers were on chemical biocide treatments and receiving city make-up water (treated with chlorine) throughout this test period.

Attempts to culture viable Legionella met with little success, despite the fact that positive controls behaved as predicted. One set of samples was collected according to specifications provided by a laboratory and split, with one portion of the sample sent to the laboratory for analysis and the other being analyzed according to the methods normally employed by the university's team. The same lot number of environmental BCYE agar was used by both laboratories. The university's tests gave no culturable Legionella, while the laboratory recovered 30 Legionella per mL from the CC tower; 1 isolate from a JEC tower, and 2 Legionella isolates each from the CII and MRC towers. The Legionella belonged to four different serogroups (1, 4, 6, and CDC #SH2156), and each tower had only one type of Legionella. Possible explanations for these results are that the laboratory's personnel are better at recovering Legionella than the university's personnel; or the differences arose from the fact that the university's personnel processed samples within 1 hour of collection while the laboratory's personnel processed samples up to 48 hours after sample collection (according to their normal procedures). The low recovery rates for culturable Legionella presented a problem in the sense that it was not possible to perform classic "kill curves" for Legionella using viable Legionella counts. (It should be pointed out; however, that contract guidelines would have made such tests impossible, since towers having numbers of Legionella high enough for kill tests would exceed the number-at which the towers must be immediately treated.) Accordingly, kill tests done in phase II of this study relied on total viable microbial numbers to determine the potential of biocides to control microorganisms. Use of microscopic counts to determine kill kinetics is possible, as will be seen, only after longterm treatment, with continued measurements of viable bacteria, total microbial cells, and Legionella-DFA counts. This is due to the fact that incoming make-up water is very low in Legionella and other bacteria. Therefore, if a biocide is capable of controlling microbes in the tower, the level of total cells present (living and dead) will, in the absence of growth in the tower and significant scrubbing of bacteria from the air, approach the level of bacterial cells



Example of data as in Figure 1, except that this cooling tower operates all year but without fans during the cool Figure 2 periods of the year (CIIW). This tower supports considerable algal growth in the warm months. Note that the Legionella-DFA levels were high during September and October 1989 (late summer and fall) but lagged behind increases in total bacterial levels in the late winter (due to lack of algal regrowth during this period). (Fe = ppm; LDB [Legionnaires' disease bacteria] = mL for water, cm² for surface, and mL for plate count.) S STIC .C. A BOY A CONCERNMENTED AND AN ANTIMATION

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in the make-up water. Consideration must be given to the cycles of concentration for the tower in question. Failure to at least approach these make-up concentrations of bacteria may indicate that the biocide is not controlling microbial growth in the tower.

Phase II-Tests of Biocide Efficacy

A list of the biocides and dosages used is shown in Table 4. All tests were started by turning off the blowdown, taking a zero time sample, dosing with biocide, and restarting the tower. The blowdown was left off for two hours and restarted after the two-hour sampling. Additional samples were generally taken at 6, 24, 48, and 72 hours after the addition of biocide. In a few cases, samples could not be obtained due to tower malfunction. Make-up water (Troy, New York, city water) was sampled at 0, 24, 48, and 72 hours. The results for isothiazolin will be used to illustrate several points.

The results for the other biocides will be given in an abbreviated form. The data in Figure 3a show the results for isothiazolin treatment in the four test towers. It is obvious that viable bacteria declined over a 24- to 48-hour period and then began to rise to pretreatment levels. Makeup water levels were consistently very low, indicating that most of the viable bacteria in the tower came from air washing and/or growth in the tower. Figure 3b gives the data for total microscopic counts (i.e., total microscopic count = live + dead + culturable + nonculturable bacterial cells). It is clear that these results do not follow those for the viable counts. (The drop in the MRC system was due to a blowdown valve stuck in the open position, which resulted in washout of the microorganisms to near make-up water levels for a short period.) The reason for the difference in viable and total counts is explained by the fact that the viable counts represent only that portion of the total counts that was culturable with the methods used at the time of sampling. The total population not only takes into account the contribution of bacteria made by the water but also will include the scrubbing of bacteria from the airpassing through the tower. A similar explanation applies to the Legionella-DFA results shown in Figure 3c. Note that the MRC tower was again in the washout mode. Figure 3d shows the results for Pseudomonas spp. While the data are relatively uniform for most of the towers, a spike in CII at 72 hours may have occurred. Since blooms of Pseudomonas would create safety, slime accumulation, and corrosion concerns, it is important to avoid treatments that kill the "normal flora" in the towers while allowing pseudomonads to bloom.

Figure 4 shows the viable count data for the cyanocarbonate/carbamate biocide, which has been used in the towers for the last 10 or more years. It is obvious that it has little effect on the viable counts.

The data in Figure 5 are for a polymeric quaternary amine biocide treatment. It appears generally ineffective in CC and CII towers. The MRC tower was on continuous blowdown due to a stuck valve. After treatment, the viable bacteria in the JEC tower appeared to decline for the duration of the sampling period. This may have been partially due to operational problems with the tower, but the drop in this tower, relative to the other test towers, remains unexplained. In general, however, the data suggest that the polymeric quaternary amine did not work very well at the dose levels employed.

Figure 6 shows the data for glutaraldehyde. The viable bacteria levels fell off in most towers over a 6- to 24-hour period with rapid recovery and return to pretreatment levels in 24 to 72 hours. Viable bacteria in the MRC tower did not return to pretreatment levels due to the faulty control device being in the continuous blowdown mode between 24 and 48 hours.

The data for the bromochlorodimethylhydantoin (BCDMH) test in the CII tower are presented in Figure 7. The biocide residuals are also plotted. It is clear that once the biocide residual rose above about 0.2 ppm free chlorine, the levels of total viable microorganisms rapidly decreased. As the concentration of biocide dropped, the total population rapidly recovered. The MRC tower was also tested, with similar results (data not shown). It should be noted that these towers were chosen for BCDMH testing since they have sumps that allowed buckets containing biocide tablets to be placed in the tower system. It is believed that the good kill over relatively long time periods was due to maintenance of biocide residuals in the tower over these same periods. The fact that levels of viable bacteria rapidly recovered as biocide decreased supports this view.

CONCLUSIONS

The results of the phase I testing to determine correlations between Legionella-DFA levels and other parameters can be summarized as follows:

- Good correlation was found between Legionella-DFA levels and total population count, which was performed by the FITC method.
- 2. Aqueous iron levels correlated well with the total population count as determined by FITC.
- Nitrite correlated well with the surface Legionella-DFA values, but the level of correlation was less with the planktonic Legionella-DFA populations.
- As expected, water quality parameters such as calcium, magnesium, hardness, and alkalinity showed a high degree of correlation.
- 5. There was little relationship between Legionella-DFA levels and temperature.
- 6. Other parameters tested in the survey gave poor correlations and their effect on the levels of total viable microorganisms and the level of *Legionella*-DFA may be minimal at these sites.







Figure 3b Effect of isothiazolin treatment on total microscopic count. It should be noted that this procedure measures live + dead + culturable + nonculturable microorganisms. The decrease in the MRC tower is due in part to a malfunctioning blowdown valve. Note that, in general, the levels of total bacteria do not change very much through the study.

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	1992 B. 1993 A.		the attended	1000	120 6	
1. A. A.	200 - 1 - 12 ¹⁰ - 10 ¹⁰ - 10 ¹⁰ - 10 ¹⁰		57 142 2 14	22	194	



Effect of isothiazolin treatment on Legionella-DFA, that is, the level of cells, whether live or dead, reacting Figure 3c specifically with fluorescent antibodies to Legionella sp. See "Materials and Methods" section for details of analytical procedures. - 76-Auto r 500

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Figure 3d Effect of isothiazolin treatment on Pseudomonas populations. Note that the levels of pseudomonads spiked in one tower at two points after treatment. 5... the store-12 $\{\xi_{i,k}^{\prime}\}_{i=1}^{\prime}$ 19.1 14.0 1.98 COL A CALMARK 5 36 300 . 50 24 CONTRACTOR AND AND A 1 3









e 5 Effect of polymeric quat treatment on viable bacterial populations in the test towers. This biocide was generally not effective at the dosage used. Note that the MRC tower was on continuous blowdown during most of this test. Biocide was added immediately after collecting the t = 0 samples.







Figure 7 Effect of bromochlorodimethylhydantoin treatment on viable bacterial populations in the CII test tower. Note

that the levels of viable bacteria decreased rapidly as the free residual biocide increased and then rapidly increases as the biocide residual diminished. Biocide was added immediately after collecting the t = 0 samples.

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The results of the phase II testing to determine the efficacy of commercially available biocides to control microbial populations, including *Legionella*, can be summarized as follows:

- The data suggest that several commercially available biocides are capable of killing microorganisms in cooling systems. In many cases, however, the kill is very short lived and regrowth results in a tower operating a large percentage of the time with high microbial populations.
- Use of these biocides at large doses and on a very frequent basis could, undoubtedly, control the microorganisms, but the economics of such treatment may often be unfavorable.

It is possible that continuous treatment of towers with biocides that are effective at low levels is a viable treatment option. This agrees with the findings of one of the authors in a previous study (Pope et al. 1984) in which treatment with ozone was effective as long as the treatment was continuously applied. The economics of the continuous treatment approach are not yet clear and would certainly depend in part on the biocide used and the conditions of the particular tower. In the case of the BCDMH used in the present study, economic analysis is not yet possible, since the minimum long-term dose levels required for control has not been established. If low (< 1.0 ppm) free chlorine residuals with the use of BCDMH are demonstrated to be effective, then there is a good possibility of economical, effective, and reliable continuous treatment. However, the uniqueness of each site must be considered when developing any treatment regime.

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