DAIRY WATERS

(Coliform Group and Escherichia coli) [*E. coli* verification required only on source water] IMS #24

[Unless otherwise stated all tolerances are ±5%]

1.	Lab	poratory Requirements				
	a.	Cultural Procedures (CP), items 34 & 35				
	b.	Sample volume sufficient to assure 100 mL for testing, sufficient air space for mixing, do not accept if completely filled				
	C.	Transported and maintained at 0.0-4.5°C (temperature control [TC] required)				
	d.	If samples are not refrigerated, transit not to exceed 6 hours (TC not required)				
	e.	Transit time does not exceed 30 hours				
	f.	Samples examined within 30 hours of collection or within 2 hours of receipt (item 1.d)				
		APPARATUS				
2.	CP,	, see items 1 - 33 (as necessary)				
3.	Sample Containers					
	a.	Borosilicate glass, plastic bottles or bags				
	b.	Sterile, containing 0.1 mL of 10% Sodium Thiosulfate				
	C.	Holds sufficient sample with air space for all necessary bacterial tests				
	d.	Maintains sample uncontaminated				
4.	Inc	ubator 35±0.5°C (Make/Model:)				
	a.	See CP item 15 for incubator requirements				
5.	Wa	ter Bath, 35±0.5°C (Make/Model:)				
	a.	Circulating and thermostatically controlled				
	b.	Maintain sufficient water depth				
6.		ter Bath, 44.5±0.2°C (Make/Model:) equired for EC-MUG]				
	a.	Circulating and thermostatically controlled				

	b.	Maintain sufficient water depth	
7.		ter Bath, 44.5±0.5°C (Make/Model:) ly for use with item 31f]	
	a.	Circulating and thermostatically controlled	
	b.	Maintain sufficient water depth	
8.	Feri	mentation Tubes/Bottles	
	a.	Sufficient size to conform with requirements for media, Durham tube and sample	
	b.	Tubes and bottles used for EC-MUG broth and chromogenic substrate methods do not autofluoresce	
9.	Inoc	culation Equipment	
	a.	Sterilized loops of at least 3 mm diameter, 22-24 gauge nichrome, chromel or platinum-iridium wire	
	b.	Disposable dry heat-sterilized hardwood applicator sticks, 0.2 to 0.3 cm in diameter and a minimum of 2.5 cm longer than the fermentation tubes	
	c.	Inoculating needle	
	d.	Sterile disposable plastic loops	
	e.	Commercial pre-sterilized cotton swabs on wooden sticks, 0.2 to 0.3 cm in diameter and a minimum of 2.5 cm longer than the fermentation tubes	
10.	Lon	ng Wavelength UV Light (365 – 366 nm) and PPE	
	a.	6 watts	
	b.	Keep clean	
	C.	UV protective glasses	
11.	Vac	cuum Source with Trap	
12.	Mer	mbrane Filter (MF) Funnel; Brand:	
	a.	Free from defects that may interfere with function	
	b.	Sterilizable	
	C.	Marked at 100 mL, or pre-marked checked and adjusted, using a 100 mL Class A graduated cylinder	

13.	Membrane Cellulose Filters, 47 mm, 0.45 μM (±0.02 μM), Sterilized				
	Brai	nd: Lot #:			
14.	Abs	sorbent Pads, Sterilized, Brand:			
15.	For	ceps _			
	a.	Round tipped, with smooth surface			
16.	Cult	ture (Petri) Dishes (for MF), Brand: Size:			
	a.	Sterile with plastic, tight fitting covers			
17.	Mic	roscope and Lamp, Brand: Model:			
	a.	Binocular, wide field, 10x oculars			
	b.	Fluorescent light, adjacent, above, perpendicular to filter plane			
	C.	Other optical device giving equivalent results			
		CULTURE MEDIA			
18.	Sto	rage of Media			
	a.	CP item 28-30 for media and storage requirements			
	b.	MF media			
		1. Store in dark at 0.0-4.5°C			
		2. Broth medium used within 96 hours. Date Prep.:			
		3. Plates kept no more than 1 week in a sealed container at 0.0-4.5°C. Date Prep.:			
19.	Med	dia Quality Control			
	a.	See CP item 28 for media composition			
	b.	Suitability test conducted on each new lot of commercially prepared and/or each new batch of laboratory prepared media by spiking with known coliform or <i>E. coli</i> , as applicable; records maintained			

TESTS FOR PRESENCE OF MEMBERS OF THE COLIFORM GROUP AND *E. coli* BY MULTIPLE-TUBE FERMENTATION TECHNIQUE

20.	Pre	sump	otive Test	
	a.	Dou	ible Strength Lauryl Sulfate Tryptose Broth (DS-LST)	
		1.	Before inoculating arrange tubes in order and label, or otherwise identify	
		2.	Shake samples vigorously 25 times in a 1 ft movement in 7 sec before removing test portion	
		3.	Remove test portions (100 mL total) within 3 min	
		4.	Inoculate ten (10) fermentation tubes containing 10 mL DS-LST or five (5) tubes containing 20 mL DS-LST or one bottle containing 100 mL DS-LST with equal volume of sample	
		5.	Incubate tubes at 35±0.5°C for 24±2 hours	
		6.	Examine tubes for gas – any gas is considered Presumptive Positive and must be transferred to BGLB (all tests) and EC-MUG broth (if performing <i>E. coli</i> testing)	
		7.	Return negative tubes (no gas) to incubator and incubate an additional 24 hr (total of 48±3 hours)	
		8.	Re-examine tubes for gas production after 48±3 hours	
		9.	Record presence or absence of gas at each examination	
		10.	Any gas produced by 24 or 48 hours is considered positive for the Presumptive Test	
		11.	No gas after 48 hours is Not Found (NF) for the test	
		a.	Tubes showing no gas, but showing evidence of growth (turbid) can be promptly submitted to the Confirmation Test (item 21). Do not report until after completion of confirmation	
		12.	Do not report gas production after 51 hours of incubation	
		13.	Promptly submit all presumptive positive tubes showing gas production at 24 or 48 hours to the Confirmation Test	
21.	Tot	al Co	liform Confirmation Test	
	a.	Brill	iant Green Lactose Bile Broth (BGLB)	
		1.	Gently shake presumptive positive tube (item 20.a.13)	
		2	Transfer (item 9) portion of positive broth to BGLR broth	

		3.	Incubate tubes at 35±0.5°C for 24±2 hours				
		4.	Examine tubes for gas – any gas is considered positive				
		5.	Return negative tubes (no gas) to incubator and incubate an additional 24 hours (total of 48±3 hours)				
		6.	Re-examine tubes for gas production after 48 hours				
		7.	Record presence or absence of gas at each examination				
		8.	Any gas produced by 24 or 48 hours is considered positive for Total Coliform				
		9.	No gas after 48 hours is Not Found (NF) for Total Coliform				
		10.	Do not report gas production after 51 hours of incubation				
22.	E. c	oli V	erification Test				
	a.	EC-	MUG Broth				
		1.	Gently shake presumptive positive tube(s) (item 20.a.13)				
		2.	Transfer (item 9) portion of positive broth to EC-MUG broth				
			a. If using the same apparatus to transfer to both BGLB and EC-MUG, transfer to EC-MUG first then to BGLB				
		3.	Incubate tubes at 44.5±0.2°C for 24±2 hours (item 6 only)				
		4.	Place tubes in water bath within 30 min of inoculation				
		5.	Examine tubes exhibiting growth for fluorescence using Longwavelength UV light (item 10)				
		6.	Record presence or absence of fluorescence				
		7.	Bright blue fluorescence after incubation is considered positive for <i>E. coli</i>				
		8.	No fluorescence after incubation is Not Found (NF) for <i>E. coli</i>				
23.	Re	cordi	ing and Reporting				
	a.	If one or more DS-LST tubes are turbid with no gas production and confirmation in BGLB yields no gas production, invalidate the sample and request a re-sample from the same point source for heterotrophic plate count. [If history has shown a sample source to repeatedly yield growth/turbidity in LST that does not confirm, the lab may test for Heterotrophic Plate Count at the same time as testing the same sample by					

	b.	Record results of fermentation tubes that confirm positive in BGLB as MPN Total Coliform/100 mL (≥ 1.1/100 mL if 10 tubes of 10 mL or 5 tubes of 20 mL are used) or ≥ 1 Total Coliform/100 mL if 100 mL presence/absence test used					
	C.	Record results of fermentation tubes that confirm positive in EC-MUG as MPN <i>E. coli</i> /100 mL (≥1.1/100 mL if 10 tubes of 10 mL or 5 tubes of 20 mL are used) or ≥ 1 <i>E. coli</i> /100 mL if 100 mL presence/absence test used					
	d.	Pos	rpretation: for multiple tubes, Not Found (NF) is < 1.1/100 mL and itive is ≥ 1.1/100 mL; for presence/absence, NF is < 1/100 mL and itive is ≥ 1/100 mL				
			TESTS FOR PRESENCE OF MEMBERS OF THE COLIFORM GROUP BY MEMBRANE FILTRATION TECHNIQUE				
24.	Filtr	ation	າ _				
	a.		ce (with alcohol flamed forceps, item 15) sterile membrane filter (item 13) corous plate, secure funnel				
	b.	Pou	r 100 mL test sample into funnel (item 12) and apply vacuum				
	C.		er test volume has been filtered, rinse funnel by filtering 3 volumes of 30 mL of sterile buffered water				
	d.	Turr	n off vacuum and remove filter with sterile (alcohol flamed) forceps				
	e.	M-Endo Broth					
		1.	Sterile pad (item 14) placed in culture dish				
		2.	Saturate pad with 2.0 mL of M-Endo Broth, CP item 28.u				
		3.	Allow to stand a few minutes before pouring off excess				
		4.	Prepared filter rolled (grid side up) onto pad slowly to avoid trapping air bubbles, do not drag across side of plate				
	f.	M-E	indo Agar				
		1.	Use culture dish previously prepared (CP item 28.t)				
		2.	Prepared filter placed on agar with rolling motion to avoid trapping air bubbles				
25.	Incu	ıbatio	on _				
	a.	In sa	aturated humidity, with dish inverted				
	b.	At 3	5±0.5°C for 23±1 hour				

26.	Cou	unting					
	a.	atyp		sheen colonies as typical coliforms and dark suspect colonies as coliforms, keep separate counts of each morphological type until d			
	b.			10% up to a maximum of 10 isolated colonies, with representative ns of each colony type			
27.	Tota	al Co	liforr	m Confirmation Test			
	a.			rial transfers of colonies to individual LST and then to BGLB tubes e same transfer apparatus (item 9)			
	b.	Incu	ıbate	tubes at 35±0.5°C for 24±2 hours			
	c.	Exa	mine	tubes for gas			
		1.		tubes with gas must be transferred to fresh BGLB tubes if the inal BGLB tubes show no gas			
	d.			egative tubes (no gas) to incubator and incubate an additional (total of 48±3 hours)			
	e.	Re-	exam	nine tubes for gas production after 48 hours			
	f.	Rec	ord p	resence or absence of gas at each examination			
	g.	•	-	produced in BGLB tubes by 24 or 48 hours is considered positive for mation Test			
	h.	No (gas a	fter 48 hours is Not Found (NF) for the Test			
	i.	Do r	not re	eport gas production after 51 hours of incubation			
28.	E. c	coli Verification Test					
	a.	EC-MUG Broth					
		1.	Trar	nsfer (item 9) portion of each target colony to EC-MUG broth			
			a.	If using the same apparatus to transfer to LST, BGLB and EC-MUG, transfer to LST first, then EC-MUG, then to BGLB			
		2.	Incu	ubate tubes at 44.5±0.2°C for 24±2 hours (item 6 only)			
			a.	Place tube in water bath within 30 min of inoculation			
		3.		mine tubes exhibiting growth for fluorescence using Long- velength UV light (item 10)			
		4.	Rec	ord presence or absence of fluorescence			

		5. Bright blue fluorescence after incubation is considered positive for <i>E. coli</i>	
		6. No fluorescence after incubation is Not Found (NF) for <i>E. coli</i>	
29.	Rep	porting	
	a.	Report confirmed colony count/100 mL	
	b.	Invalidate all samples with confluent growth or TNTC, and request a re-sample from the same point source for heterotrophic plate count	
	C.	Interpretation: Not Found (NF) is < 1/100 mL and Positive is ≥ 1/100 mL	
		HETEROTROPHIC BACTERIA STANDARD PLATE COUNT METHOD	
30.	Het	erotrophic Plate Count Method	
	a.	Plate samples as in SPC, items 2 - 10, 13 and 14. a and b	
	b.	Incubate at 35±0.5°C for 48±3 hours	
	C.	Count as in SPC items 15 and 16	
	d.	Report counts as in SPC item 19.a	
	e.	Record as "Heterotrophic Plate Count/mL at 35°C"	
	f.	Interpretation: Not Found (NF) if < 500 CFU/mL and Positive if ≥ 500 CFU/mL	
		CHROMOGENIC SUBSTRATE (MMO-MUG) PRESENCE – ABSENCE TEST FOR DAIRY WATERS (SOURCE WATER SUPPLIES <u>ONLY</u>)	
31.	Mat	terials	
	a.	Sterile non-fluorescent borosilicate glass or clear plastic bottles to contain 100 mL sample with sufficient air space for mixing	
	b.	Color comparator (required for Colilert® and Colilert®-18)	
	C.	Commercially prepared substrate used	
		1. Colilert® (CP item 28.v)	
		2. Colilert®-18 (CP item 28.w) (see 32.d)	
		3. Colisure® (CP item 28.x)	
		4. Modified Colitag™ (CP item 28.z)	
	d.	Suitability test conducted on each lot of substrate received, by spiking with known coliform; maintain records	

	e.	water Bath, circulating, maintains 35±0.5°C or; maintain records during periods of use (required for Colilert-18, optional for Modified Colitag; see item 32.d or e)
	f.	Water Bath, circulating, maintains 44.5±0.2°C (item 6 or 7); maintain records during periods of use (optional for Colilert-18 and Modified Colitag; see item 32.d or e)
32.	Pro	cedure
	a.	Aseptically add pre-weighed substrate to 100 mL of the water sample
	b.	Optionally, add 100 mL of sample to the substrate in a sterile container provided by the manufacturer
	C.	Aseptically cap and mix thoroughly by shaking 25 times to dissolve reagent (does not completely dissolve)
	d.	For Colilert-18 thermally equilibrate test solution for 20 min in a 35±0.5°C circulating water bath or alternatively 7-10 min (not to exceed 10 min) in a 44.5±0.5°C circulating water bath (item 6 or 7), and then continue incubation in water bath or dry incubator (35±0.5°C) for a total of 18 hours (minimum), not to exceed 22 hours.
	e.	For Modified Colitag that could be read prior to 24 hours, thermally equilibrate test solution for 20 min in a 35±0.5°C circulating water bath or alternatively 7-10 min (not to exceed 10 min) in a 44.5±0.2°C circulating water bath (item 6 or 7), and then continue incubation in a water bath or dry incubator (35±0.5°C) for a minimum of 18 to a maximum of 48 hours.
	f.	For Colilert and Colisure, incubate at 35±0.5°C in water bath or dry incubator for a minimum of 24 hours, not to exceed 28 hours for Colilert, 48 hours for Colisure
	g.	For Modified Colitag not pre-warmed (item 32.e), incubate at 35±0.5°C in water bath or dry incubator for a minimum of 24 to a maximum of 48 hours
	h.	Examine containers for the production of color change
	i.	Examine containers that exhibit color change for fluorescence
33.	Inte	rpretation and Reporting
	a.	Colilert and Colilert-18
		1. If no yellow color is observed
		a. Record test result as Not Found (NF) for Total Coliform
		b. Report as Total Coliform Not Found (NF) in 100 mL sample:< 1/100 mL

	۷.	II ye	silow color present.	
		a.	Gently invert container several times until color is uniformly dispersed through the sample	
		b.	Compare yellow color to color comparator dispersed into the SAME type of sample container	
		C.	If color is equal to or greater than that of the color comparator, record test result as Positive (POS) for Total Coliform	
		d.	Report as total coliform Present in 100 mL sample: ≥ 1/100 mL	
		e.	If yellow color is obvious but less than the comparator, record test result as Not Found (NF) for Total Coliform; report as for no yellow color above (33.a.1.b)	
	3.	Pla	ce yellow containers under Long-Wavelength UV light (item 10)	
		a.	If the container fluoresces, record test result as Positive (POS) for <i>E. coli</i>	
		b.	Report as <i>E. coli</i> Present in 100 mL sample: ≥ 1/100 mL	
		C.	If container does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i>	
		d.	Report as E. coli Not Found (NF) in 100 mL sample: <1/100 mL	
Ο.	Coli	sure		
	1.	If no	o red or magenta color is observed	
		a.	Record test result as Not Found (NF) for Total Coliform	
		b.	Report as Total Coliform Not Found (NF) in 100 mL sample: < 1/100 mL	
	2.	If re	ed or magenta color present	
		a.	Gently invert container several times until color is uniformly dispersed through the sample	
		b.	If red or magenta color is present, record test result as Positive for Total Coliform	
		c.	Report as Total Coliform Present in 100 mL sample: ≥ 1/100 mL	
	3.	Pla	ce red or magenta containers under Long-Wavelength UV light	
		a.	If the container fluoresces, record test result as Positive (POS) for	

		b.	Report as <i>E. coli</i> Present in 100 mL sample: ≥ 1/100 mL	
		C.	If container does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i> . Report as < 1/100 mL for <i>E. coli</i>	
		d.	Report as <i>E. coli</i> Not Found (NF) in 100 mL sample: < 1/100 mL	
C.	Mod	dified	Colitag	
	1.	If no	o yellow color is observed	
		a.	Record test result as Not Found (NF) for Total Coliform	
		b.	Report as Total Coliform Not Found (NF) in 100 mL sample: < 1/100 mL	
	2.	If ye	ellow color present	
		a.	Gently invert container several times until color is uniformly dispersed through the sample	
		b.	Optionally, compare yellow color to one or more of the following controls that have been dispersed into the SAME type of sample container:	
			Color comparator	
			Positive and negative control strain	
			3. Blank	
		C.	If yellow color is present, record test result as Positive for Total Coliform	
		d.	Report as Total Coliform Present in 100 mL sample: ≥ 1/100 mL	
	3.	Pla	ce yellow containers under Long-Wavelength UV light (Item 10)	
		a.	If the sample fluoresces, record test result as Positive (POS) for E. coli	
		b.	Report as <i>E. coli</i> Present in 100 mL sample: ≥ 1/100 mL	
		C.	If container does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i> .	
		d.	Report as <i>E. coli</i> Not Found (NF) in 100 mL sample: < 1/100 mL	

CHROMOGENIC SUBSTRATE (MMO-MUG) MULTIPLE TUBE PROCEDURE FOR THE PRESENCE OF TOTAL COLIFORM (SOURCE WATER SUPPLIES ONLY)

34.	Materials						
	a.	Sterile non-fluorescent borosilicate glass or clear plastic tubes 10 mL or 20 mL capacity					
	b.	Color comparator: See item 31.b (comparator solution must be in container of same size and type (34.a.)					
	c.	Water Bath: See item 31.e					
35.	Pro	cedure					
	a.	Before transferring sample portions arrange tubes in order and identify					
	b.	Shake sample vigorously 25 times in 7 sec with a 1 ft movement prior to adjusting to test volume					
	C.	Aseptically add pre-weighed substrate to 100 mL sample					
	d.	Optionally, add 100 mL of sample to container with substrate provided by manufacturer					
	e.	Aseptically cap and mix thoroughly by shaking 25 times to dissolve reagent (does not completely dissolve)					
	f.	Remove test portions (100 mL total) within 3 min					
	g.	Transfer 20 mL of sample/reagent mixture to five (5) tubes, or 10 mL to ten (10) tubes					
	h.	Optionally, transfer 100 mL of mixed (see item 35.b) sample to 10 tubes containing pre-dispensed substrate provided by manufacturer					
	i.	For Colilert-18, thermally equilibrate test solution for 20 min in a 35±0.5°C circulating water bath and then continue incubation in water bath or dry incubator for a total of 18 hours (minimum), not to exceed 22 hours					
	j.	For Modified Colitag that could be read prior to 24 hours, thermally equilibrate test solution for 20 min in a 35.0±0.5°C circulating water bath, and then continue incubation in a waterbath or dry incubator (35.0±0.5°C) for a minimum of 18 to a maximum of 48 hours.					
	k.	For Colilert and Colisure, incubate at 35±0.5°C in water bath or dry incubator for a minimum of 24 hours, not to exceed 28 hours for Colilert, 48 hours for Colisure					
	l.	For Modified Colitag not pre-warmed, incubate at 35.0±0.5°C in water bath or dry incubator for a minimum of 24 to a maximum of 48 hours					

	m.	Exa	mine	tubes for the development of color change	
	n.	Exa	mine	tubes that exhibit color change for fluorescence	
36.	Inte	rpret	atior	n	
	a.	Coli	lert a	and Colilert-18	
		1.	Mix	tubes to uniformly distribute yellow color	
		2.	Con	mpare tubes to color comparator tube (SAME size and type)	
		3.		cord test result of tubes without color or obvious yellow color but less n comparator as Not Found (NF) for Total Coliform	
		4.		cord test result of tubes with yellow color equal to or greater than or comparator tube as Positive (POS) for Total Coliform	
		5.	Plac	ce yellow tubes under Long-Wavelength UV light (item 10)	
			a.	If the tube fluoresces, record test result as Positive (POS) for E. coli	
			b.	If tube does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i>	
	b.	Coli	sure		
		1.	Mix	tubes to uniformly distribute red or magenta color	
		2.		cord test result of tubes without red or magenta color as Not Found ———————————————————————————————————	
		3.		cord test result of tubes with red or magenta color as Positive (POS) Total Coliform	
		4.		ce red or magenta tubes under Long-Wavelength UV light	
			a.	If the tube fluoresces, record test result as Positive (POS) for E. coli	
			b.	If tube does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i>	
	C.	Mod	dified	Colitag	
		1	Miv	tubes to uniformly distribute vellow color	

		2.	•	ionally, compare yellow color to one or more of the following trols that have been dispersed into the SAME type of tube:	
			a.	Color comparator	
			b.	Positive and negative control strain	
			C.	Blank _	
		2.		ord test result of tubes without color as Not Found (NF) for Total form	
		3.		ord test result of tubes with yellow color as Positive (POS) for Total form	
		4.	Plac	ce yellow tubes under Long-Wavelength UV light (item 10)	
			a.	If the tube fluoresces, record test result as Positive (POS) for E. coli	
			b.	If tube does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i>	
37.	Rep	ortin	g		
	a.			es exhibit no color change (36.a.3, 36.b.2, or 36.c.2), report as and (NF): < 1.1/100 mL for Total Coliform and <i>E. coli</i>	
	b.			more tubes exhibit color change (36.a.4, 36.b.3, or 36.c.3), report as (POS): ≥ 1.1/100 mL for Total Coliform	
	C.			more tubes exhibit fluorescence, report as Positive (POS): mL for <i>E. coli</i>	
	F			MOGENIC SUBSTRATE (MMO-MUG) QUANTI-TRAY PROCEDURE PRESENCE OF TOTAL COLIFORM (SOURCE WATER SUPPLIES ONL)	<u>Y</u>)
38.	Mat	erials	5	_	
	a.	Qua	nti-T	ray or Quanti-Tray 2000	
	b.	See	item	31.b (comparator solution must be in same type of tray (34.a.)	
	c.	Lon	g Wa	velength UV Light (Item 10)	
39.	Pro	cedu	re	-	
	a.	Turr	n on (Quanti-Tray Sealer and allow to pre-heat	
	b.	Befo	ore tra	ansferring sample portion identify tray	

	C.			ample vigorously 25 times in 7 sec with a 1 ft movement prior to to test volume	
	d.	Ase	ptical	lly add pre-weighed substrate to 100 mL sample	
	e.		•	lly cap and mix thoroughly by shaking 25 times to dissolve reagent t completely dissolve)	
	f.	Trar	nsfer	test portions (100 mL total) into tray within 3 min	
	g.	Plac	ce tra	y in appropriate support and seal tray	
	h.			ert-18, incubate at 35±0.5°C in a dry incubator for a total of 18 hours n), not to exceed 22 hours	
	i.			ert, incubate at 35±0.5°C in dry incubator for a minimum urs, not to exceed 28 hours	
	j.	Exa	mine	wells in tray for the development of color change	
	k.	Exa	mine	wells in tray that exhibit color change for fluorescence	
40.	Inte	erpretation			
	a.	Coli	lert a	nd Colilert-18	
		1.	Mix	tray to uniformly distribute yellow color in wells	
		2.	Con	npare wells to color comparator tray (SAME size and type)	
		3.		ord test result of wells without color or obvious yellow color but less not comparator as Not Found (NF) for Total Coliform	
		4.		ord test result of wells with yellow color equal to or greater than or comparator tray as Positive (POS) for Total Coliform	
		5.		ce Quanti trays with yellow wells under Long-Wavelength UV light n 10)	
			a.	If any well fluoresces, record test result as Positive (POS) for <i>E. coli</i>	
			b.	If all wells do not fluoresce, record test result as Not Found (NF) for <i>E. coli</i>	
41.	Rep	ortin	g		
	a.			s exhibit no color change, report as Not Found (NF): < 1.0/100 mL for iform	
	b.			more wells exhibit color change, report as Positive (POS):) mL for Total Coliform	

c. If one or more wells exhibit fluorescence, report as Positive (POS):
 ≥ 1.0/100 mL for *E. coli*

CHROMOGENIC SUBSTRATE PRESENCE (XGAL – MUG) PRESENCE – ABSENCE TEST FOR DAIRY WATERS (SOURCE WATER SUPPLIES <u>ONLY</u>)

42.	2. Materials				
	a.	E*C	colite substrate, see CP item 28.y		
	b.	Quality control procedures conducted on each lot of substrate received, as recommended by manufacturer, test by spiking with known coliform, records maintained			
43.	Pro	cedu	re		
	a.	Add	water sample to the E*Colite substrate		
		1.	Tear perforated strip		
		2.	Open bag by pulling the white tabs		
		3.	Aseptically pour 100 mL of water sample into bag (do not touch inside of bag)		
		4.	Flatten bag to remove air		
		5.	Twirl bag 2-3 times around twister wires to form a leak proof seal		
		6.	Fold twisters around back of bag		
		7.	Shake bag 25 times in 7 sec to dissolve sodium thiosulfate tablet, if present		
		8.	Continue rolling to build pressure in water compartment		
		9.	Maintain pressure on rolled area and push water through first seal into powder section of bag ONLY		
		10.	Shake bag 25 times in 7 sec to completely dissolve powder in water (push mixture against bag sides to pull apart any remaining seal)		
	b.	Plac	ce sealed bag in 35°C water bath for 10 minutes		
	C.	Trar	nsfer to 35±0.5°C incubator for 28 hours		
	d.		mine bags for the production of blue or blue/green color or blue color in ers of bag		

44.	Interpretation and Reporting			
	a.	If yellow color is observed:		
		1.	Record sample as Not Found (NF) for Total Coliform	
		2.	Report as Total Coliform Not Found (NF) in 100 mL sample: < 1/100 mL	
	b.	If blo	ue or blue/green (or blue in corners) color observed:	
		1.	The sample is Positive for Total Coliform	
		2.	Report as Total Coliform present in 100 mL sample: ≥ 1/100 mL	
	C.	Plac	ce blue or blue/green containers under Long-Wavelength UV light (item 10)	
		1.	If the container fluoresces, record test result as Positive (POS) for <i>E. coli</i>	
		2.	Report as <i>E. coli</i> Present in 100 mL sample: ≥ 1/100 mL	
		3.	If container does not fluoresce, record test result as Not Found (NF) for <i>E. coli</i> . Report as < 1/100mL for <i>E. coli</i>	
			MISCELLANEOUS	
45.	Copy of current in-use edition of <u>Standard Methods for the</u> <u>Examination of Water and Wastewater</u> in laboratory			