

Designation: D4763 - 06 (Reapproved 2012)

# Standard Practice for Identification of Chemicals in Water by Fluorescence Spectroscopy<sup>1</sup>

This standard is issued under the fixed designation D4763; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon ( $\varepsilon$ ) indicates an editorial change since the last revision or reapproval.

# 1. Scope

1.1 This practice allows for the identification of 90 chemicals that may be found in water or in surface layers on water. This practice is based on the use of room-temperature fluorescence spectra taken from lists developed by the U.S. Environmental Protection Agency and the U.S. Coast Guard (1).<sup>2</sup> Ref (1) is the primary source for these spectra. This practice is also based on the assumption that such chemicals are either present in aqueous solution or are extracted from water into an appropriate solvent.

1.2 Although many organic chemicals containing aromatic rings, heterocyclic rings, or extended conjugated double-bond systems have appreciable quantum yields of fluorescence, this practice is designed only for the specific compounds listed. If present in complex mixtures, preseparation by highperformance liquid chromatography (HPLC), column chromatography, or thin-layer chromatography (TLC) would probably be required.

1.3 If used with HPLC, this practice could be used for the identification of fluorescence spectra generated by optical multichannel analyzers (OMA) or diode-array detectors.

1.4 For simple mixtures, or in the presence of other nonfluorescing chemicals, separatory techniques might not be required. The excitation and emission maximum wavelengths listed in this practice could be used with standard fluorescence techniques (**Refs 2-6**) to quantitate these ninety chemicals once identification had been established. For such uses, generation of a calibration curve, to determine the linear range for use of fluorescence quantitation would be required for each chemical. Examination of solvent blanks to subtract or eliminate any fluorescence background would probably be required.

1.5 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the

responsibility of the user of this standard to establish appropriate safety, health, and environmental practices and determine the applicability of regulatory limitations prior to use.

1.6 This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.

## 2. Referenced Documents

- 2.1 ASTM Standards:<sup>3</sup>
- D1129 Terminology Relating to Water
- D1193 Specification for Reagent Water
- E131 Terminology Relating to Molecular Spectroscopy
- E275 Practice for Describing and Measuring Performance of Ultraviolet and Visible Spectrophotometers

### 3. Terminology

3.1 *Definitions*—For definitions of terms used in this practice, refer to Terminology D1129, Specification D1193, and definitions under the jurisdiction of Committee E13 such as Definitions E131 and Practice E275.

# 4. Summary of Practice

4.1 This practice uses well tested fluorescence techniques to detect and identify (or determine the absence of) 90 chemicals that have relatively high fluorescence yields. Table 1 lists for each chemical an appropriate solvent (either cyclohexane, water, methyl or ethyl alcohol, depending on solubility), a suggested excitation wavelength for maximum sensitivity, a wavelength corresponding to the emission maximum, the number of fluorescence peaks and shoulders, the width (full width at half of the maximum emission intensity) of the strongest fluorescence peak and the detection limit for the experimental conditions given. Detection limits could be lowered, following identification, by using broader slit widths.

<sup>&</sup>lt;sup>1</sup> This practice is under the jurisdiction of ASTM Committee D19 on Water and is the direct responsibility of Subcommittee D19.06 on Methods for Analysis for Organic Substances in Water.

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<sup>&</sup>lt;sup>2</sup> The boldface numbers in parentheses refer to the list of references at the end of this practice.

<sup>&</sup>lt;sup>3</sup> For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

A list of corrected fluorescence spectra for the chemicals included in this practice are also available (1).

4.2 Identification of the sample is made by comparison of the obtained spectra with information in Table 1 and by direct visual comparison of appropriate spectra with positions of principal peaks in agreement to  $\pm 2$  nm and ratios of peak heights in agreement to  $\pm 10$  % if corrected spectrofluorometers are used.

4.3 Spectral distortions due to self-absorption or fluorescence quenching or dimer formation may occur at higher concentrations (for example, 100 ppm or  $\mu g/mL$ ). If this is suspected, the solution should be diluted and additional fluorescence spectra generated. If a suspected chemical is not detected on excitation at the appropriate wavelength, it usually can be assumed that it is not present above the detection limit, barring interference effects due to absorption or quenching that can usually be anticipated.

#### 5. Significance and Use

5.1 This practice is useful for detecting and identifying (or determining the absence of) 90 chemicals with relatively high fluorescence yields (see Table 1). Most commonly, this practice will be useful for distinguishing single fluorescent chemicals in solution, simple mixtures or single fluorescing chemicals in the presence of other nonfluorescing chemicals. Chemicals with high fluorescence yields tend to have aromatic rings, some heterocyclic rings or extended conjugated double-bond systems. Typical chemicals included on this list include aromatics, substituted aromatics such as phenols, polycyclic aromatic hydrocarbons (PAH's), some pesticides such as DDT, polychlorinated biphenyls (PCB's), some heterocyclics, and some esters, organic acids, and ketones.

Chemical	Code	Concentra- tions, ppm	Solvent	$\lambda_{\text{exc}}$ , nm	$\lambda_{\max}^{em}$ , nm	Number of Peaks	WHM, nm	Shoulder Number	Detection Limit (DL), ppm	$\lambda_{DL},  \text{nm}$	Comments
Acenaphthene	ACN	1.03	СН	290	323	4		3	0.001	290	
Acetone	ACT	227	СН	290	410	1			212	290	
Acridine	ACR	96	CH	285/355	386/422	4/2		2/0			
	ACR	9.6	ETOH	290/355	357/415	2/2		1/1	0.02/0.04	290/355	
Aniline	ANL	15.5	CH	280	316	10 91	2 A.Y		0.037	280	
Anthracene	ATH	1.03	СН	355	378	4		1	0.001	355	
	ATH	1.55	ETOH	355	380	4		1	0.001	355	
roclor 1242	PC4	131	CH	270	317	2	35	<b>h</b>	0.3	270	
254	PC5	129	CH	270	317	2	36	4 <b>1</b> • <b>41</b> /	2	270	
trazine	ATZ	369	СН	290	350	1			300	290	
zinphosmethyl	AZP	112	CH	350	410	2	60		10	350	
P 7	AZP	122	ETOH	340	420	2	80	V	4	340	
enz(a)anthracene	BAT	1.1	CH	280	386	4		1	0.003	280	
enzene	BNZ	79	CH	250	279	3	24	1	2/4	250/265	
enzonitrile	BZN	9.9	CH	260	287	2	28	1	0.1/0.1	260/270	
enzo(a)pyrene	BAP	0.088	CH	370	405763	-66(201)	2.1.	2	0.002	370	
enzyl alcohol	BAL	99	CH	250	284	2	27	1	0.1/0.1	250/260	
enzyl amine <sup>Catalog/s</sup>	BZM	ls118 tm/86	CH 93	250	283 76-2	a <b>6</b> 33-90	c <sub>27</sub> 270d	e208/astm	-3/2 /63-06	250/260	
enzyl triethylam- monium chloride	BMA	210	H <sub>2</sub> O	250	280	1	28		59	250	
isphenol A	BPA	10.5	ETOH	270	304	1	30	1	0.04/0.02	270/285	
rucine	BRU	13.5	ETOH	280	327	1	56		2/2	280/295	
-tert-Butylphenol	BOP	21	CH	265	295	1	30	1	0.1/0.1	265/275	
-tert-Butylphenol	BTP	17.5	CH	260	295	1	31	1	0.6/0.4	260/280	
arbaryl	CBY	1.0	СН	285	335	2	36	2	0.01	285	
arnauba wax	WCA	63.5	СН	260	310	1	64		42	260	
astor oil	OCA	390	ETOH	290	328	1	43	2	20	290	
	OCA	286	CH	280/320		1			180/300	280/320	
atechol	CTC	8.7	H <sub>2</sub> O	265	310	1	46		0.4/0.2	265/280	
-Chloroaniline	CAP	17.2	CH	290	328	1	36	1	0.2	290	
-Chloronaphthalene	CNA	11.3	СН	290	328	3	34	4	0.1	290	
-Chlorophenol	CPN	101	CH	260	305	1	30		1/0.1	260/285	
hlorpyrifos (Duraban)	DUR	25.3	CH	280	326	1	52		1/0.5	280/295	
-Chlorotoluene	CTN	23.8	CH	265	288	1	29	3	1/0.8	265/275	
-Chloro-o-toluidine	COT	25	CH	290	328	1	39	1	0.09	300	
hrysene	CRY	1.0	CH	270	383	5			0.002	270	
oconut oil	OCC	286	CH	290	330				100	290	
od liver oil	OCL	323	CH	260/280	320/320	1/1	150		260,140	260,280	
	001	020	011	330	500	1			65	330	
opper naphthenate	CNN	98	СН	260	326	1	60	3	3/1	260/280	
ottonseed oil	OCS	305	CH	280/320	320/380				165,300	280,320	
oumaphos	COU	11.4	CH	320	320/380	1	 74		0.3	320	
Cresol	CRO	12.0	CH	265	293	1	30	 1	0.04	280	
-Cresol	CRP	12.0	CH	265	293	1	30		0.04	280	
umene	CUM	10.3	CH	205 250	299 283	2	28	 1	3	250	
Cymene	CMP	11.8	CH	260	285	2	20 28	2	0.4/0.2	260/270	
DD		61.0	CH	200 240	205 294	1	20 30	2	4	240	
DT	DDD DDT	87	СН	240 245	294 291	2	30 28	2	4 7	240 245	
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