ANALYSIS INSTRUCTIONS

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photoLab® 6x00 / 7x00

METHOD DATA, V 2.50





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Test kits with barcode

Available methods

Here, the method for a cell test (KT) is selected with the aid of the barcode on the cell, for a reagent test (RT) with the aid of the AutoSelector. The total measuring range is related to the shown citation form. For reagent tests, the measuring range covers all possible path length (cells from 10 to 50 mm).

Parameter	Model	Order No.	Total measuring range	Method	Type ^a	Method No.
Acid Capacity to pH 4.3 (total alkalinity)	01758	252 087	0.40 – 8.00 mmol/l	Indicator reaction	KT	208
Aluminium*	00594	252 068	0.02 – 0.50 mg/l Al	Chromazurole S	KT	196
Aluminium*	14825	250 425	0.020 - 1.20 mg/l Al	Chromazurole S	RT	043
Ammonium	A6/25	252 072	0.20 - 8.00 mg/l NH ₄ -N	Indophenol blue	KT	003
Ammonium	14739	250 495	0.010 - 2.000 mg/l NH ₄ -N	Indophenol blue	KT	104
Ammonium	14558	252 000	0.20 - 8.00 mg/l NH ₄ -N	Indophenol blue	KT	051
Ammonium	14544	250 329	0.5 - 16.0 mg/l NH ₄ -N	Indophenol blue	KT	052
Ammonium	14559	250 424	4.0 – 80.0 mg/l NH ₄ -N	Indophenol blue	KT	053
Ammonium	14752/1 14752/2	250 426 252 081	0.010 – 3.00 mg/l NH ₄ -N	Indophenol blue	RT	054
Ammonium	00683	252 027	2.0 – 75.0 mg/l NH ₄ -N	Indophenol blue	RT	155
Ammonium	00683	252 027	5 – 150 mg/l NH ₄ -N	Indophenol blue	RT	163
AOX Cell*	00675	252 023	0.05 – 2.50 mg/L AOX	Oxidation to chloride	KT	156
Arsenic*	01747	252 063	0.001 - 0.100 mg/l As	Ag-DDTC	RT	132
BOD*	00687	252 028	0.5 – 3000 mg/l BOD	Modification of Winkler method	KT	157
Boron*	00826	252 041	0.05 – 2.00 mg/l B	Azomethine H	KT	164
Boron*	14839	250 427	0.050 - 0.800 mg/l B	Rosocyanine	RT	046
Bromine*	00605	252 014	0.020 - 10.00 mg/l Br ₂	S-DPD	RT	146
Cadmium	14834	250 314	0.025 - 1.000 mg/l Cd	Cadion derivative	KT	067
Cadmium	01745	252 051	0.0020 - 0.500 mg/l Cd	Cadion derivative	KT	183
Calcium*	00858	252 047	10 – 250 mg/l Ca	Phthalein purple	KT	165
Calcium*	14815	250 428	5 – 160 mg/l Ca	Glyoxal-bis-hydroxyanil	RT	042
Calcium sensitive*	14815	250 428	1.0 - 15.0 mg/l Ca	Glyoxal-bis-hydroxyanil	RT	125
Chloride*	14730	250 353	5 – 125 mg/l Cl	Iron(III)-thiocyanat	KT	095
Chloride*	14897/1 14897/2	250 491 252 082	2.5 – 25.0 mg/l Cl	Iron(III)-thiocyanat	RT	110
Chloride*	14897/1 14897/2	250 491 252 082	10 – 250 mg/l Cl	Iron(III)-thiocyanat	RT	063
Chlorine* (free chlorine)	00595	250 419	0.03 - 6.00 mg/l Cl ₂	S-DPD	KT	141
Chlorine* (free and total chlorine)	00597	250 420	0.03 – 6.00 mg/l Cl ₂	S-DPD	KT	142
Chlorine* (free chlorine)	00598/1 00598/2	252 010 252 011	0.010 – 6.00 mg/l Cl ₂	S-DPD	RT	143
Chlorine* (total chlorine)	00602/1 00602/2	252 013 252 055	0.010 – 6.00 mg/l Cl ₂	S-DPD	RT	145
Chlorine* (free and total chlorine)	00599	252 012	0.010 – 6.00 mg/l Cl ₂	S-DPD	RT	144
Chlorine* (free and total chlorine)	00086 00087 00088	252 077 252 078 252 079	0.010 - 6.00 mg/l Cl ₂	DPD	KT	194
Chlorine dioxide*	00608	252 017	0.020 - 10.00 mg/l ClO ₂	S-DPD	RT	149
Chromate*	14552	250 341	0.05 – 2.00 mg/l Cr	Diphenylcarbazide	KT	039
Chromate* (total chromium)	14552	250 341	0.05 – 2.00 mg/l Cr	Peroxodisulfate oxidation, Diphenylcarbazide	KT	039
Chromate*	14758	250 433	0.010 - 3.00 mg/l Cr	Diphenylcarbazide	RT	040
COD*	C3/25	252 070	10 – 150 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	001

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Parameter	Model	Order No.	Total measuring range	Method	Type ^a	Method No.
COD*	14560	250 303	4.0 – 40.0 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	031
COD*	01796	252 092	5.0 – 80.0 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	211
COD*	14540	252 001	10 – 150 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	014
COD*	14895	250 359	15 – 300 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	105
COD*	14690	250 304	50 – 500 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	093
COD*	C4/25	252 071	25 – 1500 mg/l COD	Chromosulfuric acid oxidation, chromium(III) determination	KT	002
COD*	14541	252 002	25 – 1500 mg/l COD	Chromosulfuric acid oxidation, chromium(III) determination	KT	023
COD*	14691	250 351	300 – 3500 mg/l COD	Chromosulfuric acid oxidation, chromium(III) determination	KT	094
COD*	14555	250 309	500 – 10000 mg/l COD	Chromosulfuric acid oxidation, chromium(III) determination	KT	024
COD*	01797	252 093	5000 – 90000 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	209
COD (Hg free)*	09772	250 301	10 – 150 mg/l COD	Chromosulfuric acid oxidation, chromate determination	KT	137
COD (Hg free)*	09773	250 306	100 – 1500 mg/l COD	Chromosulfuric acid oxidation, chromium(III) determination	KT	138
Copper*	14553	250 408	0.05 – 8.00 mg/l Cu	Cuprizone	KT	026
Copper*	14767	250 441	0.02 – 6.00 mg/l Cu	Cuprizone	RT	027
Cyanide* (free cyanide)	14561	250 344	0.010 - 0.500 mg/l CN	Barbituric acid and pyridinecarboxylic acid	KT	075
Cyanide* (readily liberated cyanide)	14561	250 344	0.010 - 0.500 mg/l CN	Citric acid, barbituric acid and pyridinecarboxylic acid	KT	075
Cyanide* (free cyanide)	09701	250 492	0.0020 - 0.500 mg/l CN	Barbituric acid and pyridinecarboxylic acid	RT	109
Cyanide* (readily liberated cyanide)	09701	250 492	0.0020 - 0.500 mg/l CN	Citric acid, barbituric acid and pyridinecarboxylic acid	RT	109
Cyanuric Acid	19253	252 091	2 - 160 Cyan Acid	Triazine derivative	RT	210
Fluoride*	00809	252 094	0.10 – 1.80 mg/l F	Alizarin complexone	KT	215
Fluorid sensitive	00809	250 094	0.025 – 0.500 mg/l F	Alizarin complexone	KT	216
Fluorid*	14598/1 14598/2	252 048 252 083	0.10 – 2.00 mg/l F	Alizarin complexone	RT	166
Fluorid*	14598/1 14598/2	252 048 252 083	1.0 – 20.0 mg/l F	Alizarin complexone	RT	167
Formaldehyde*	14500	250 406	0.10 - 8.00 mg/l HCHO	Chromotropic acid	KT	028
Formaldehyde*	14678	250 331	0.02 - 8.00 mg/l HCHO	Chromotropic acid	RT	091
Gold	14821	250 436	0.5 – 12.0 mg/l Au	Rhodamine B	RT	045
Hardness see Total Hardnes	ss or Residu	al Hardnes	S			
Hydrazine*	09711	250 493	0.005 – 2.00 mg/l N ₂ H ₄	4-Dimethylaminobenzaldehyde	RT	044
Hydrogenperoxide*	14731	250 402	2.0 – 20.0 mg/l H ₂ O ₂	Titanyl sulfate	KT	099
Hydrogenperoxide sensitive*	14731	250 402	0.25 – 5.00 mg/l H ₂ O ₂	Titanyl sulfate	KT	128
Hydrogenperoxide	18789	252 067	0.015 – 6.00 mg/l H ₂ O ₂	Phenanthroline derivative	RT	198
Iodine*	00606	252 015	0.050 - 10.00 mg/l l ₂	S-DPD	RT	147
Iron	14549	250 349	0.05 – 4.00 mg/l Fe	Triazine	KT	037
Iron*	14896	250 361	1.0 – 50.0 mg/l Fe (Fe(II) and Fe(III))	2,2'-Dipyridyl	KT	106
Iron	14761/1 14761/2	250 435 250 439	0.005 – 5.00 mg/l Fe	Triazine	KT	038
Iron*	00796	252 042	0.010 – 5.00 mg/l Fe (Fe(II) and Fe(III))	1,10-Phenanthroline	KT	161
Lead*	14833	250 313	0.10 – 5.00 mg/l Pb	PAR	KT	066
Lead*	09717	252 034	0.010 – 5.00 mg/l Pb	PAR	RT	160

Manganese* 00816 252 035 0.10 – 5.00 mg/l Mn Formaldoxime KT 159 Manganese* 01846 252 097 0.05 – 2.00 mg/l Mn PAN RT 226 Manganese* 1477012 250 48 2010 – 10.00 mg/l Mn Formopyrogalidired KT 175 Molybdenum 18252 252 090 0.5 – 45.0 mg/l Mo Bromopyrogalidired KT 175 Molybdenum 18252 252 090 0.5 – 45.0 mg/l Mo Mercapteaetic acid RT 206 Mickel* 14554 250 409 0.10 – 6.00 mg/l N Dimetrylegloxime KT 117 185 Nickel* 14785 250 409 0.10 – 6.00 mg/l N Dimetrylegloxime RT 018 Nitrate* 14242 250 410 0.5 – 25.0 mg/l NO-N 2.5-Dimetryleghenol KT 059 Nitrate* 14563 252 0073 0.5 – 25.0 mg/l NO-N 2.6-Dimetrylyghenol KT 039 Nitrate* 14566 250 347 0.7 – 50.0 mg/l NO-N 2.6-Dimetrylyghenol KT<	Parameter	Model	Order No.	Total measuring range	Method	Type ^a	Method No.
Marganese* 01846 252 097 0.005 – 2.00 mg/l Mn PAN RT 226 Manganese* 147701 258 428 0.010 – 10.00 mg/l Mn Formaldoxime RT 0.19 Molybdenum 0.0860 252 040 0.02 – 1.00 mg/l Mo Bromopyrogalol red KT 173 Molybdenum 1.0862 252 050 0.05 – 4.50 mg/l Mo Mercaptocetic acid RT 1.00 Monochloramine 0.1632 252 057 0.50 – 10.00 mg/l N Dimetrylogloxime RT 0.18 Nicker* 1.4755 250 443 0.5 – 2.50 mg/l NO-N Dimetrylogloxime RT 0.18 Nikrate* 1.4542 250 410 0.5 – 2.50 mg/l NO-N 2.6-Dimetrylphenol KT 0.04 Nikrate* 1.4552 252 010 0.5 – 5.0 mg/l NO-N 2.6-Dimetrylphenol KT 0.05 Nikrate* 1.0614 250 041 0.2 – 2.50 mg/l NO-N 2.6-Dimetrylphenol KT 1.07 Nikrate* 1.0731 250 441 0.2 – 2.50 mg/l NO-N 2.6-Dimetrylphenol K	Magnesium*	00815	252 043	5.0 – 75.0 mg/l Mg	Phthalein purple	KT	158
Marganeee* 147707 250 442 0.010 - 10.00 mg/l Mn Formaldoxime RT 019 147707 252 044 0.02 - 1.00 mg/l Mn Bromopyrogallol red KT 175 17	Manganese*	00816	252 035	0.10 – 5.00 mg/l Mn	Formaldoxime	KT	159
Melybdenum	Manganese*	01846	252 097	0.005 – 2.00 mg/l Mn	PAN	RT	226
Molybedrum 19252 25 290 0.5 – 45.0 mg/l Monochloramine Morcaptoacetic acid RT 206 Monochloramine 0.1632 225 295 7.0560 – 10.00 mg/l Clz Indophenol blue RT 185 Nickel* 14584 250 493 3.02 – 5.00 mg/l Ni Dimethylgyloxime KT 018 Nikrate* N225 252 073 3.5 – 25.0 mg/l NO-N 2.6-Dimethylghonol KT 008 Nikrate* 14563 252 003 3.5 – 25.0 mg/l NO-N 2.6-Dimethylphenol KT 059 Nikrate* 14563 252 003 3.5 – 25.0 mg/l NO-N 2.6-Dimethylphenol KT 059 Nikrate* 14764 253 047 1.0 – 50.0 mg/l NO-N 2.6-Dimethylphenol KT 151 Nikrate* 14773 250 421 0.10 – 25.0 mg/l NO-N 2.6-Dimethylphenol KT 151 Nikrate* 14776/2 250 421 0.10 – 3.00 mg/l NO-N 2.6-Dimethylphenol RT 106 Nikrate* 14776/2 250 421 0.10 – 3.00 mg/l NO-N Resorcine	Manganese*			0.010 – 10.00 mg/l Mn	Formaldoxime	RT	019
Monochloramine 01632 252 057 0.050 - 10.00 mg/l Cl2 Indophenol blue RT 185 Nickola* 14554 250 409 0.10 - 6.00 mg/l Ni Dimethylglyoxime RT 018 Nickola* 14554 250 409 0.20 - 5.00 mg/l Ni Dimethylglyoxime RT 018 Nickola* 14554 250 409 0.20 - 5.00 mg/l Ni Dimethylglyoxime RT 018 Nickola* 14542 250 410 0.5 - 18.0 mg/l NO-N 2.6-Dimethylphenol KT 004 Nitrate* 14542 250 410 0.5 - 18.0 mg/l NO-N Nitrospectral KT 059 Nitrate* 14542 250 410 0.5 - 18.0 mg/l NO-N 2.6-Dimethylphenol KT 030 Nitrate* 14764 250 347 1.0 - 50.0 mg/l NO-N 2.6-Dimethylphenol KT 030 Nitrate* 00614 252 019 23 - 225 mg/l NO-N 2.6-Dimethylphenol KT 107 Nitrate* 14773 250 444 0.2 - 20.0 mg/l NO-N 2.6-Dimethylphenol KT 150 Nitrate* 03713/1 250 427 0.10 - 25.0 mg/l NO-N 2.6-Dimethylphenol KT 150 Nitrate* 03713/1 250 427 0.10 - 25.0 mg/l NO-N 2.6-Dimethylphenol RT 159 Nitrate* 03713/2 250 82 985 0.10 - 25.0 mg/l NO-N 2.6-Dimethylphenol RT 159 Nitrate* 03713/1 250 427 0.20 - 25.0 mg/l NO-N 2.6-Dimethylphenol RT 159 Nitrate* 14566 250 441 0.10 - 3.00 mg/l NO-N 2.6-Dimethylphenol RT 159 Nitrate* 14566 250 427 0.2 - 17.0 mg/l NO-N Resorcine RT 140 Nitrate* 14567 250 402 0.2 - 17.0 mg/l NO-N Resorcine RT 140 Nitrate* 14547 252 004 0.010 - 0.700 mg/l NO-N Resorcine RT 140 Nitrate* 14547 252 004 0.010 - 0.700 mg/l NO-N Nitrospectral Nitrate* 1476/1 250 450 0.000 - 0.000 mg/l NO-N Nitrospectral Nitrate* 1476/1 250 450 0.000 - 0.000 mg/l NO-N Nitrospectral Nitrate* 1476/1 250 450 0.5 - 15.0 mg/l N Peroxodisultate oxidation, KT 159 Nitrate* 1476/1 250 450 0.5 - 15.0 mg/l N Peroxodisultate oxidation, KT 168 Nitrospectral 14543 250 494 0.10 - 2.50 mg/l DEHA Peroxodisultate oxidation, KT 168 Phenol* 14551 250 450 0.5 - 15.0 mg/l PO-P Phospho	Molybdenum	00860	252 040	0.02 – 1.00 mg/l Mo	Bromopyrogallol red	KT	175
Nickel*	Molybdenum	19252	252 090	0.5 – 45.0 mg/l Mo	Mercaptoacetic acid	RT	206
Nitrate No. Nitrate No. No. Nitrate Nitrate No. Nitrate No. Nitrate Nitrate Nitrate No. Nitrate Nitrate Nitrate No. Nitrate Nitrate Nitrate Nitrate No. No. Nitrate No. No. Nitrate No.	Monochloramine	01632	252 057	0.050 – 10.00 mg/l Cl ₂	Indophenol blue	RT	185
Nitrate* N225 252 073 0.5 – 25.0 mg/l NO+N 2.6-Dimethylphenol KT 004 Nitrate* 14542 250 410 0.5 – 18.0 mg/l NO+N Nitrospectral KT 059 Nitrate* 14563 252 003 0.5 – 25.0 mg/l NO+N 2.6-Dimethylphenol KT 030 Nitrate* 14764 250 347 1.0 – 50.0 mg/l NO+N 2.6-Dimethylphenol KT 107 Nitrate* 00614 252 019 23 – 225 mg/l NO+N 2.6-Dimethylphenol KT 151 Nitrate* 00614 252 019 23 – 225 mg/l NO+N 2.6-Dimethylphenol KT 151 Nitrate* 00614 252 019 23 – 225 mg/l NO+N 2.6-Dimethylphenol KT 151 Nitrate* 006713/1 250 241 0.10 – 25.0 mg/l NO+N Nitrospectral RT 067 Nitrate* 08713/1 250 241 0.10 – 25.0 mg/l NO+N Nitrospectral RT 08713/2 252 085 Nitrate in seawater* 14942 250 422 0.2 – 17.0 mg/l NO+N Resorcine KT 072 Nitrate in seawater* 14942 250 422 0.2 – 17.0 mg/l NO+N Resorcine RT 1400 Nitrite* Ni5/25 252 074 0.010 – 0.700 mg/l NO+N Resorcine RT 005 Nitrite* 14547 252 044 0.010 – 0.700 mg/l NO+N Resorcine RT 035 Nitrite* 14776/1 250 445 0.002 – 1.00 mg/l NO+N Resorcine RT 035 Nitrite* 14776/1 250 445 0.002 – 1.00 mg/l NO+N Resorcine RT 036 Nitrogen (total) 14537 250 358 0.5 – 15.0 mg/l N Percoxodisulfate oxidation, RT 036 Nitrogen (total) 14763 250 440 0.5 – 15.0 mg/l N Percoxodisulfate oxidation, RT 036 Nitrogen (total) 14763 250 440 0.5 – 15.0 mg/l N Percoxodisulfate oxidation, RT 036 Nitrogen (total) 14764 250 043 0.5 – 15.0 mg/l N Percoxodisulfate oxidation, RT 036 Nitrogen (total) 14763 250 440 0.002 – 1.00 mg/l NO+N Percoxodisulfate oxidation, RT 036 Nitrogen (total) 14764 250 050 6.4 – 8.8 Pencol Molfication of Winkler method KT 092 Dozone* 00607/1 252 054 0.002 – 0.00 mg/l DEHA Perroxidisulfate oxidation, RT 148 Dehenol* 14551 250 080 0.002 – 0.500 mg/l DEHA Perroxidisulfate oxidation, RT 177 Phosphate Phenol* 14543 250 400 0.002 – 0.00 mg/l PO+P Phosphomolybdenum blue RT 076 Phosphate P6/25 252 075 0.5 – 5.00 mg/l PO+P Phosphomolybdenum blue RT 076 Phosphate P6/25 252 076 0.5 – 5.00 mg/l PO+P Phosphomolybdenum blue RT 006 Phosphate 14848/1 250 344 0.5 – 5.00 mg/l PO+P Phosphomolybdenum blue RT 056	Nickel*	14554	250 409	0.10 – 6.00 mg/l Ni	Dimethylglyoxime	KT	017
Nitrate* 14542 250 410 0.5 − 18.0 mg/l NO₂-N Nitrospectral KT 059 Nitrate* 14563 252 030 0.5 − 25.0 mg/l NO₂-N 2.6-Dimethylphenol KT 030 Nitrate* 14764 250 347 1.0 − 50.0 mg/l NO₂-N 2.6-Dimethylphenol KT 157 Nitrate* 00614 252 019 23 − 225 mg/l NO₂-N 2.6-Dimethylphenol KT 157 Nitrate* 071 250 444 0.2 − 20.0 mg/l NO₂-N 2.6-Dimethylphenol KT 157 Nitrate* 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 09713/2 250 685 Nitrate in seawater* 14556 250 411 0.10 − 3.00 mg/l NO₂-N Resorcine KT 072 Nitrate in seawater* 14547 250 422 0.2 − 17.0 mg/l NO₂-N Griess reaction KT 005 Nitrite* 14547 252 044 0.010 − 0.700 mg/l NO₂-N Griess reaction KT 005 Nitrite* 14547 252 045 0.000 − 0.700 mg/l NO₂-N Griess reaction KT 035 Nitrite* 14776/1 250 445 0.000 − 1.00 mg/l NO₂-N Iron(II) ethylenediammonium KT 197 Nitrogen (total) 14537 250 358 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14763 250 493 0.5 − 12.0 mg/l O₂ Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14763 250 495 0.000 − 0.500 mg/l DEHA Ferozine∞ RT 108 Doxygen* 14694 250 403 0.5 − 12.0 mg/l O₂ Modification of Winkter method KT 092 Doxygen* 252 086 0.000 − 0.500 mg/l DEHA Ferozine∞ RT 148 Doxygen* 14694 250 403 0.5 − 12.0 mg/l O₂ Modification of Winkter method KT 092 Doxygen* 252 086 0.000 − 0.500 mg/l DEHA A minoantipyrine, by extraction RT 168 Pheno1* 0.0656 252 056 0.000 − 0.500 mg/l DEHA A minoantipyrine, by extraction RT 176 Phenophate (total phospho- us) Phosphate (total phospho- us) Phosphate (total phospho- us) Phosphate (total phospho- us) Phosphate (total phospho-	Nickel*	14785	250 443	0.02 – 5.00 mg/l Ni	Dimethylglyoxime	RT	018
Nitrate* 14563 252 003 0.5 – 25.0 mg/l NO₂-N 2.6-Dimethylphenol KT 030 Nitrate* 14764 250 347 1.0 – 50.0 mg/l NO₂-N 2.6-Dimethylphenol KT 107 Nitrate* 00614 252 019 23 – 225 mg/l NO₂-N 2.6-Dimethylphenol KT 107 Nitrate* 14773 250 444 0.2 – 20.0 mg/l NO₂-N Nitrate* 09713/1 0250 440 0.2 – 20.0 mg/l NO₂-N Nitrate* 09713/2 252 052 050 0.10 – 25.0 mg/l NO₂-N Nitrate* 09713/2 252 052 050 0.10 – 25.0 mg/l NO₂-N Resorcine KT 050 Nitrate* 09713/2 252 052 050 0.10 – 25.0 mg/l NO₂-N Resorcine KT 072 Nitrate in seawater* 14556 250 411 0.10 – 3.00 mg/l NO₂-N Resorcine KT 072 Nitrate in seawater* 14942 250 422 0.2 – 17.0 mg/l NO₂-N Resorcine RT 140 Nitrite* 14547 252 004 0.010 – 0.700 mg/l NO₂-N Resorcine RT 140 Nitrite* 14547 252 004 0.010 – 0.700 mg/l NO₂-N Griess reaction KT 005 Nitrite* 14547 252 004 0.010 – 0.700 mg/l NO₂-N Griess reaction KT 035 Nitrite* 14547 252 004 0.010 – 0.700 mg/l NO₂-N Griess reaction KT 035 Nitrite* 14547 252 044 0.002 – 1.00 mg/l NO₂-N Griess reaction RT 14776/1 250 440 0.002 – 1.00 mg/l NO₂-N Griess reaction RT 036 Nitrogen (total) 14537 250 358 0.5 – 15.0 mg/l N Percoxdisulfate oxidation, KT 068 Nitrogen (total) 14537 250 358 0.5 – 15.0 mg/l N Percoxdisulfate oxidation, KT 153 Nitrogen (total) 14763 250 494 10 – 150 mg/l N Percoxdisulfate oxidation, KT 108 0.002 Nitrogen (total) 14763 250 494 10 – 150 mg/l N Percoxdisulfate oxidation, KT 108 0.000000000000000000000000000000000	Nitrate*	N2/25	252 073	0.5 – 25.0 mg/l NO ₃ -N	2,6-Dimethylphenol	KT	004
Nitrate* 14764 250 347 1.0 − 50.0 mg/l NO₂N 2.6-Dimethylphenol KT 107 Nitrate* 00614 252 019 23 − 225 mg/l NO₂N 2.6-Dimethylphenol KT 151 Nitrate* 009713/1 250 444 0.2 − 20.0 mg/l NO₂N Nitrospectral RT 060 Nitrate* 099713/1 250 2444 0.2 − 20.0 mg/l NO₂N Nitrospectral RT 060 Nitrate* 099713/2 250 285 011 0.10 − 25.0 mg/l NO₂N Resorcine RT 139 Nitrate in seawater* 14556 250 411 0.10 − 3.00 mg/l NO₂N Resorcine RT 140 Nitrate in seawater* 14942 250 422 0.2 − 17.0 mg/l NO₂N Resorcine RT 140 Nitrate* N525 252 074 0.010 − 0.700 mg/l NO₂N Resorcine RT 140 Nitrate* N525 252 074 0.010 − 0.700 mg/l NO₂N Griess reaction KT 005 Nitrite* N525 250 040 0.010 − 0.700 mg/l NO₂N Griess reaction KT 035 Nitrite* 00609 252 069 1.0 − 90.0 mg/l NO₂N Griess reaction KT 035 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.002 − 1.00 mg/l NO₂N Griess reaction RT 197 Nitrite* 1476/3 250 494 10 − 150 mg/l N 2.0000 Griess reaction RT 153 Nitrospectral Nit	Nitrate*	14542	250 410	0.5 − 18.0 mg/l NO ₃ -N	Nitrospectral	KT	059
Nitrate* 00614 252 019 23 − 225 mg/l NO3-N 2,6-Dimethylphenol KT 151 151 Nitrate* 14773 250 441 0.2 − 20.0 mg/l NO3-N Nitrate* 09713/1 250 421 0.10 − 25.0 mg/l NO3-N Nitrate* 09713/1 250 4821 0.10 − 25.0 mg/l NO3-N Resorcine RT 139 Nitrate in seawater* 14566 250 411 0.10 − 3.00 mg/l NO3-N Resorcine KT 072 Nitrate in seawater* 14942 250 422 0.2 − 17.0 mg/l NO3-N Resorcine RT 140 Nitrate in seawater* 14942 250 422 0.2 − 17.0 mg/l NO3-N Resorcine RT 140 Nitrate in seawater* 14942 250 422 0.2 − 17.0 mg/l NO3-N Resorcine RT 140 Nitrate* N5/25 252 074 0.010 − 0.700 mg/l NO3-N Griess reaction KT 005 Nitrite* 14547 252 004 0.010 − 0.700 mg/l NO3-N Griess reaction KT 035 Nitrite* 14776/2 250 440 0.020 − 0.00 mg/l NO3-N Griess reaction KT 035 Nitrite* 14776/2 250 440 0.020 − 1.00 mg/l NO3-N Griess reaction RT 197 Nitrite* 14776/2 250 440 0.020 − 1.00 mg/l NO3-N Griess reaction RT 036 Nitrogen (total) 14537 250 388 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, KT 068 Nitrogen (total) 14633 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, KT 153 Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, KT 108 20xygen* 14694 250 403 0.5 − 12.0 mg/l O2 Modification of Winkler method KT 092 Oxygen Soavengers 19251 252 089 0.020 − 0.500 mg/l DEHA FerroZinee RT 207 Dxygen* 0.0607/2 252 089 0.020 − 0.500 mg/l DEHA FerroZinee RT 148 Phenol* 14551 250 412 0.10 − 2.50 mg/l Phenole MBTH KT 073 Phenol* 0.0866 252 058 0.020 − 0.500 mg/l CeHsOH Aminoantipyrine, by extraction RT 176 Phenophate P6/25 252 075 0.5 − 5.00 mg/l PO+P Phosphate (total phospho-103) 0.5 − 2.50 mg/l PO+P Phosphate (total phospho-14543 250 440 0.5 − 5.00 mg/l PO+P Phosphate (total phospho-14543 250 344 0.5 − 5.00 mg/l PO+P Phosphate (total phospho-14543 250 344 0.5 − 5.00 mg/l PO+P Phosphate (total phospho-14543 250 344 0.5 − 5.00 mg/l PO+P Phosphonolybdenum blue KT 055 Phosphate (1454) phospho-14543 250 340 0.5 − 5.00 mg/l PO+P Phosphonolybdenum blue KT 055 Phosphate (1454) phospho-14543 250 340 0.5 − 5.00 mg/l PO+P Phosphonolybdenum blue KT 055	Nitrate*	14563	252 003	0.5 – 25.0 mg/l NO ₃ -N	2,6-Dimethylphenol	KT	030
Nitrate*	Nitrate*	14764	250 347	1.0 − 50.0 mg/l NO ₃ -N	2,6-Dimethylphenol	KT	107
Nitrogen (total) 14761 250 445 0.5 − 15.0 mg/l NO₂N 2.6-Dimethylphenol RT 139	Nitrate*	00614	252 019	23 – 225 mg/l NO ₃ -N	2,6-Dimethylphenol	KT	151
Nitrate in seawater* 14556 250 415 0.10 − 3.00 mg/l NO₂-N Resorcine RT 140	Nitrate*	14773	250 444	0.2 – 20.0 mg/l NO ₃ -N	Nitrospectral	RT	060
Nitrate in seawater* 14942 250 422 0.2 – 17.0 mg/l NO-N Resorcine RT 140	Nitrate*			0.10 – 25.0 mg/l NO ₃ -N	2,6-Dimethylphenol	RT	139
Nitrite* N5/25 252 074 0.010 − 0.700 mg/l NO₂-N Griess reaction KT 005 Nitrite* 14547 252 004 0.010 − 0.700 mg/l NO₂-N Griess reaction KT 035 Nitrite* 14547 252 069 1.0 − 90.0 mg/l NO₂-N Griess reaction KT 035 Nitrite* 14776/1 250 445 1.0 − 90.0 mg/l NO₂-N Inon(II) ethylenediammonium sufficient suffate valuation with suffate suffate valuation suffate valuation with suffate valuation in the	Nitrate in seawater*	14556	250 411	0.10 - 3.00 mg/l NO ₃ -N	Resorcine	KT	072
Nitrite* 14547 252 004 0.010 − 0.700 mg/l NO₂-N Griess reaction KT 035 Nitrite* 00609 252 069 1.0 − 90.0 mg/l NO₂-N Iron(II) ethylenediammonium KT 197 Nitrite* 14776/1 250 445 14776/2 250 440 Nitrogen (total) 14537 250 358 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, Nitrospectral Nitrogen (total) 14537 250 358 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, Nitrospectral Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, Nitrospectral Nitrogen (total) 14763 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, RT 153 Doxygen* 14694 250 403 0.5 − 12.0 mg/l N Peroxodisulfate oxidation, RT 108 Doxygen* 14694 250 403 0.5 − 12.0 mg/l O₂ Modification of Winkler method RT 092 Doxygen Scavengers 19251 252 089 0.020 − 0.500 mg/l DEHA FerroZine® RT 207 Dozone* 00607/1 252 015 0.010 − 4.00 mg/l O₂ S-DPD RT 148 DH 01744 250 050 6.4 − 8.8 Phenol red RT 148 DH 01744 250 050 6.4 − 8.8 Phenol red RT 166 Phenol* 14551 250 412 0.10 − 2.50 mg/l PO+P Phosphomolybdenum blue RT 006 Phosphate P6/25 252 075 0.05 − 5.00 mg/l CH-BOH Aminoantipyrine, by extraction RT 176 Phosphate (total phospho-18745) 252 076 0.5 − 5.00 mg/l PO+P Phosphomolybdenum blue RT 007 Phosphate (total phospho-18543 250 324 0.05 − 5.00 mg/l PO+P Phosphomolybdenum blue RT 007 Phosphate (total phospho-18543 250 324 0.05 − 5.00 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-18543 250 324 0.05 − 5.00 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-18543 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-18543 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-18543 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-1848/2 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 055 Phosphate (total phospho-1848/2 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 152 Phosphate (total phospho-1948/2 250 334 0.5 − 25.0 mg/l PO+P Phosphomolybdenum blue RT 152 Phosphate (total phospho-1948/2 250 3	Nitrate in seawater*	14942	250 422	0.2 – 17.0 mg/l NO ₃ -N	Resorcine	RT	140
Nitrite	Nitrite*	N5/25	252 074	0.010 – 0.700 mg/l NO ₂ -N	Griess reaction	KT	005
Nitrite* 14776/1 250 445 0.002 - 1.00 mg/l NO2-N Griess reaction RT 036	Nitrite*	14547	252 004	0.010 - 0.700 mg/l NO ₂ -N	Griess reaction	KT	035
Nitrogen (total) 14537 250 358 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, Nitrospectral Nitrogen (total)* 252 018 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, Nitrospectral Nitrospectral Nitrogen (total)* 00613 252 018 0.5 − 15.0 mg/l N Peroxodisulfate oxidation, 2.6-Dimethylphenol RT 153 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, 2.6-Dimethylphenol RT 108 250 494 10 − 150 mg/l N Peroxodisulfate oxidation, 2.6-Dimethylphenol RT 108 20 20 20 20 20 20 20 20 20 20 20 20 20	Nitrite*	00609	252 069	1.0 – 90.0 mg/l NO ₂ -N		KT	197
Nitrogen (total)* 00613 252 018 0.5 − 15.0 mg/l N 2,6-Dimethylphenol	Nitrite*			0.002 – 1.00 mg/l NO ₂ -N	Griess reaction	RT	036
Nitrogen (total) 14763 250 494 10 - 150 mg/l N Peroxodisulfate oxidation, 2,6-Dimethylphenol KT 108	Nitrogen (total)	14537	250 358	0.5 – 15.0 mg/l N		KT	068
2,6-Dimethylphenol 2,6-Dimethylphenol 2,6-Dimethylphenol National Principle Nation	Nitrogen (total)*	00613	252 018	0.5 – 15.0 mg/l N	Peroxodisulfate oxidation, 2,6-Dimethylphenol	KT	153
Oxygen Scavengers 19251 252 089 0.020 − 0.500 mg/l DEHA FerroZine® RT 207 Ozone* 00607/1 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 054 00607/2 252 055 00607/2 252 056 00607/2 252 056 0062 00607/2 252 058 0062 − 0.100 mg/l CeHsOH Aminoantipyrine, by extraction RT 176 Phenol* RT 176 Phenol* MBTH KT 073 Phenol* 00856 252 058 0002 − 0.100 mg/l CeHsOH Aminoantipyrine, by extraction Phenol* 00856 252 058 0.025 − 5.00 mg/l CeHsOH Aminoantipyrine RT 177 Phosphate P6/25 252 075 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 250 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 056	Nitrogen (total)	14763	250 494	10 – 150 mg/l N	Peroxodisulfate oxidation, 2,6-Dimethylphenol	KT	108
Ozone*	Oxygen*	14694	250 403	0.5 - 12.0 mg/l O ₂	Modification of Winkler method	KT	092
00607/2 252 054 DH 01744 252 050 6.4 − 8.8 Phenol red KT 186 Phenol* 14551 250 412 0.10 − 2.50 mg/l Phenole MBTH KT 073 Phenol* 00856 252 058 0.002 − 0.100 mg/l CeHsOH Aminoantipyrine, by extraction RT 176 Phenol* 00856 252 058 0.002 − 0.100 mg/l CeHsOH Aminoantipyrine, by extraction RT 177 Phosphate P6/25 252 075 0.05 − 5.00 mg/l P0₄-P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P6/25 252 075 0.05 − 5.00 mg/l P0₄-P Phosphomolybdenum blue KT 007 Phosphate P7/25 252 076 0.5 − 25.00 mg/l P0₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.00 mg/l P0₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.00 mg/l P0₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.00 mg/l P0₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l P0₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l P0₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l P0₄-P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l P0₄-P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l P0₄-P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 346 0.5 − 25.0 mg/l P0₄-P Phosphomolybdenum blue KT 152 Phosphate 00616 252 021 3.0 − 100.0 mg/l PO₄-P Phosphomolybdenum blue KT 152 Phosphate 14848/1 250 446 14848/2 250 86	Oxygen Scavengers	19251	252 089	0.020 - 0.500 mg/l DEHA	FerroZine®	RT	207
Phenol* 14551 250 412 0.10 − 2.50 mg/l Phenole MBTH KT 073 Phenol* 00856 252 058 0.002 − 0.100 mg/l C₀H₅OH Aminoantipyrine, by extraction RT 176 Phenol* 00856 252 058 0.025 − 5.00 mg/l C₀H₅OH Aminoantipyrine, by extraction RT 177 Phosphate P6/25 252 075 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P6/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 086	Ozone*			0.010 – 4.00 mg/l O ₃	S-DPD	RT	148
Phenol* 00856 252 058 0.002 − 0.100 mg/l C₀H₅OH Aminoantipyrine, by extraction RT 176 Phenol* 00856 252 058 0.025 − 5.00 mg/l C₀H₅OH Aminoantipyrine RT 177 Phosphate P6/25 252 075 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P6/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) 14543 250 324 0.05 − 5.00 mg/l PO₄-P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 − 25.0 mg/l PO₄-P Phosphomolybdenum blue KT 086 <td>рН</td> <td>01744</td> <td>252 050</td> <td>6.4 – 8.8</td> <td>Phenol red</td> <td>KT</td> <td>186</td>	рН	01744	252 050	6.4 – 8.8	Phenol red	KT	186
Phenol* 00856 252 058 0.025 – 5.00 mg/l C ₆ H ₅ OH Aminoantipyrine RT 177 Phosphate P6/25 252 075 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P6/25 252 075 0.05 – 5.00 mg/l PO ₄ -P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 006 Phosphate P7/25 252 076 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) 14543 250 324 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate 0616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT <t< td=""><td>Phenol*</td><td>14551</td><td>250 412</td><td>0.10 - 2.50 mg/l Phenole</td><td>МВТН</td><td>KT</td><td>073</td></t<>	Phenol*	14551	250 412	0.10 - 2.50 mg/l Phenole	МВТН	KT	073
Phosphate P6/25 252 075 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P6/25 252 075 0.05 – 5.00 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 006 Phosphate (total phosphorus) P7/25 252 076 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) 14543 250 324 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate 14848/1 250 446 0.01 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/2 252 086 0.010 – 5.00 mg/l PO ₄ -P <td>Phenol*</td> <td>00856</td> <td>252 058</td> <td>0.002 - 0.100 mg/l C₆H₅OH</td> <td>Aminoantipyrine, by extraction</td> <td>RT</td> <td>176</td>	Phenol*	00856	252 058	0.002 - 0.100 mg/l C ₆ H ₅ OH	Aminoantipyrine, by extraction	RT	176
Phosphate (total phosphorus)	Phenol*	00856	252 058	0.025 – 5.00 mg/l C ₆ H ₅ OH	Aminoantipyrine	RT	177
Phosphate P7/25 252 076 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 252 076 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 007 Phosphate (total phosphorus) P7/25 250 324 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) P7/25 250 324 0.05 – 5.00 mg/l PO ₄ -P Proxodisulfate oxidation, Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) P7/25 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) P7/25 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) P7/25 250 334 0.5 – 25.0 mg/l P Proxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate P14729 250 334 0.5 – 25.0 mg/l P Phosphomolybdenum blue KT 086 Phosphate P1488/1 250 446 0.010 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue RT 056	Phosphate	P6/25	252 075	0.05 - 5.00 mg/l PO ₄ -P	Phosphomolybdenum blue	KT	006
Phosphate (total phosphorus)	Phosphate (total phosphorus)	P6/25	252 075	0.05 – 5.00 mg/l P		KT	006
Phosphate 14543 250 324 0.05 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue KT 055 Phosphate (total phosphorus) 14543 250 324 0.05 – 5.00 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 055 Phosphate 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate 00616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/1	Phosphate	P7/25	252 076	0.5 – 25.0 mg/l PO ₄ -P	Phosphomolybdenum blue	KT	007
Phosphate (total phosphorus) 14543 250 324 0.05 – 5.00 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 055 Phosphate 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate 00616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/1 14848/2 250 446 252 086 0.010 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue RT 056	Phosphate (total phosphorus)	P7/25	252 076	0.5 – 25.0 mg/l P		KT	007
Phosphate 14729 250 334 0.5 – 25.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 250 334 0.5 – 25.0 mg/l P Phosphomolybdenum blue KT 086 Phosphate (total phosphorus) 250 334 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate 00616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/1 250 446 252 086 0.010 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue RT 056	Phosphate	14543	250 324	0.05 – 5.00 mg/l PO ₄ -P	Phosphomolybdenum blue	KT	055
Phosphate (total phosphorus) 14729 250 334 0.5 – 25.0 mg/l P Peroxodisulfate oxidation, Phosphomolybdenum blue KT 086 Phosphate 00616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/1 14848/2 250 446 252 086 0.010 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue RT 056	Phosphate (total phosphorus)	14543	250 324	0.05 – 5.00 mg/l P		KT	055
Phosphate 00616 252 021 3.0 – 100.0 mg/l PO ₄ -P Phosphomolybdenum blue KT 152 Phosphate 14848/1 250 446 14848/2 252 086 0.010 – 5.00 mg/l PO ₄ -P Phosphomolybdenum blue RT 056	Phosphate	14729	250 334	0.5 - 25.0 mg/l PO ₄ -P	Phosphomolybdenum blue	KT	086
Phosphate 14848/1 250 446 252 086 0.010 − 5.00 mg/l PO₄-P Phosphomolybdenum blue RT 056	Phosphate (total phosphorus)	14729	250 334	0.5 – 25.0 mg/l P		KT	086
14848/2 252 086	Phosphate	00616	252 021	3.0 – 100.0 mg/l PO ₄ -P	Phosphomolybdenum blue	KT	152
Phosphate 00798 252 045 1.0 − 100.0 mg/l PO₄-P Phosphomolybdenum blue RT 162	Phosphate	14848/1 14848/2		0.010 – 5.00 mg/l PO ₄ -P	Phosphomolybdenum blue	RT	056
	Phosphate	00798	252 045	1.0 – 100.0 mg/l PO ₄ -P	Phosphomolybdenum blue	RT	162

ba75728e11 03/2021 **11**

Parameter	Model	Order No.	Total measuring range	Method	Type ^a	Method No.
Phosphate*	14546	250 413	0.5 – 25.0 mg/l PO ₄ -P	Vanadatomolybdate	KT	069
Phosphate*	14842	250 447	0.5 – 30.0 mg/l PO ₄ -P	Vanadatomolybdate	RT	070
Potassium	14562	250 407	5.0 – 50.0 mg/l K	Kalignost, turbidimetric	KT	103
Potassium	00615	252 020	30 – 300 mg/l K	Kalignost, turbidimetric	KT	150
Residual Hardness*	14683	250 404	0.50 - 5.00 mg/l Ca	Phthalein purple	KT	098
Silicate (Silicic acid)	14794	250 438	0.11 - 10.70 mg/l SiO ₂	Silicomolybdenum blue	RT	079
Silicate (Silicic acid)	14794	250 438	0.011 - 1.600 mg/l SiO ₂	Silicomolybdenum blue	RT	081
Silicate (Silicic acid)*	00857	252 046	1.1 – 107.0 mg/l SiO ₂	Molybdatosilicate	RT	169
Silicate (Silicic acid)*	00857	252 046	11 - 1070 mg/l SiO ₂	Molybdatosilicate	RT	171
Silver*	14831	250 448	0.25 – 3.00 mg/l Ag	Eosine / 1,10-Phenanthroline	RT	047
Sodium in nutrient solutions*	00885	252 044	10 – 300 mg/l Na	indirectly as chloride	KT	168
Sulfate	14548	250 414	5 – 250 mg/l SO ₄	Bariumsulfate, turbidimetric	KT	064
Sulfate	00617	252 022	50 – 500 mg/l SO ₄	Bariumsulfate, turbidimetric	KT	154
Sulfate	14564	250 415	100 – 1000 mg/l SO ₄	Bariumsulfate, turbidimetric	KT	082
Sulfate*	14791	250 449	25 – 300 mg/l SO ₄	Tannin	RT	065
Sulfate	02537	252 103	5 – 300 mg/l SO ₄	Bariumsulfate, turbidimetric	RT	236
Sulfide*	14779	250 450	0.020 – 1.50 mg/l S	Dimethyl-p-phenylendiamine	RT	080
Sulfite*	14394	250 416	1.0 − 20.0 mg/l SO ₃	Ellman's reagens	KT	127
Sulfite sensitive*	14394	250 416	0.05 - 3.00 mg/l SO ₃	Ellman's reagens	KT	127
Sulfite*	01746	252 053	1.0 − 60.0 mg/l SO ₃	Ellman's reagent	RT	187
Surfactants (anionic)	14697	250 333	0.05 – 2.00 mg/I MBAS (methylen blue active substances)	Methylene blue	KT	087
Surfactants (anionic)	02552	250 333	0.05 – 2.00 mg/l MBAS (methylen blue active substances)	Methylene blue	KT	231
Surfactants (cationic)*	01764	252 062	0.05 – 1.50 mg/l k-Ten	Disulfine blue	KT	192
Surfactants (nonionic)*	01787	252 061	0.10 – 7.50 mg/l n-Ten	TBPE	KT	193
Tin*	14622	250 401	0.10 – 2.50 mg/l Sn	Pyrocatechol violet	KT	100
TOC	14878	252 036	5.0 - 80.0 mg/l TOC	Peroxodisulfate oxidation, indicator	KT	172
TOC	14879	252 037	50 – 800 mg/l TOC	Peroxodisulfate oxidation, indicator	KT	173
Total Hardness*	00961	252 039	5 – 215 mg/l Ca	Phthalein purple	KT	178
Water hardness see Total Ha	rdness or F	Residual Ha	ardness	,	'	
Volatile Organic Acids*	01749	252 096	50 – 3000 mg/l CH₃COOH	Esterification	KT	222
Volatile Organic Acids*	01809	252 095	50 – 3000 mg/l CH₃COOH	Esterification	KT	223
Zinc	00861	252 049	0.025 – 1.000 mg/l Zn	PAR	KT	174
Zinc	14566	250 417	0.20 – 5.00 mg/l Zn	PAR	KT	074
	i	1	The state of the s			

a. Turbidity correction possible

^{**} KT = reaction cell test (16 mm round cell); RT = reagent test



01758 · Acid capacity to pH 4.3 (total alkalinity)

a xylem brand

Measuring range: 0.40 – 8.00 mmol/l

20 - 400 mg/l CaCO₃



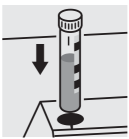
Pipette 4.0 ml of **AC-1** into a round cell.



Add 1.0 ml of the sample with pipette, close the cell with the screw cap, and mix.



Add 0.50 ml of **AC-2** with pipette, close the cell with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a sodium hydroxide solution 0.1 mol/l can be used after diluting accordingly (see section "Standard solutions").



Measuring 0.02-0.50 mg/l Al

range: Expression of results also possible in mmol/l.



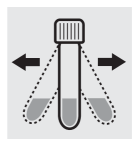
Check the pH of the sample, specified range: pH 3-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 6.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 level blue microspoon of **Al-1K**, close with the screw cap.



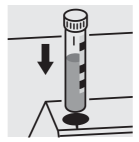
Shake the cell vigorously to dissolve the solid substance.



Add 0.25 ml of **Al-2K** with pipette, close with the screw cap, and mix.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

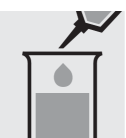
Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use aluminium standard solution, Cat.No. 250460, concentration 1000 mg/l Al can be used after diluting accordingly.

Measuring	0.10 -1.20 mg/I AI	10-mm cell
range:	0.05 -0.60 mg/I AI	20-mm cell
	0.020-0.200 mg/I AI	50-mm cell
	Expression of results also	possible in mmol/l.



Check the pH of the sample, specified range: pH 3-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1 level blue microspoon of Al-1 to the test tube and dissolve the solid substance.



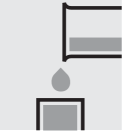
Add 1.2 ml of Al-2 with pipette and mix.



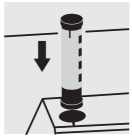
Add 0.25 ml of **Al-3** with pipette and mix.



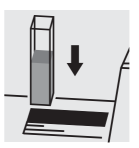
Reaction time: 2 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

Ready-for-use aluminium standard solution, Cat.No. 250460, concentration 1000 mg/l Al, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.

Measuring	0.20- 8.00 mg/I NH ₄ -N
range:	0.26-10.30 mg/I NH ₄
	0.20 - 8.00 mg/l NH ₃ -N
	0.24- 9.73 mg/I NH ₃
	Expression of results also possible in mmol/l.



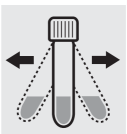
Check the pH of the sample, specified range: pH 4–13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



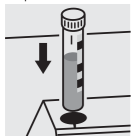
Add 1 dose of **NH**₄-1**K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄⁺, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

Measuring	0.010-2.000 mg/l NH ₄ -N		
range:	0.01 -2.58 mg/l NH ₄		
	0.010 -2.000 mg/l NH ₃ -N		
	0.01 −2.43 mg/l NH ₃		
	Expression of results also possible in mmol/l.		



Check the pH of the sample, specified range: pH 4–13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell close with the screw cap, and mix.



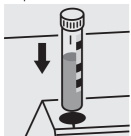
Add 1 dose of **NH**₄-**1K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄⁺, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.

Measuring	0.20- 8.00 mg/I NH ₄ -N
range:	0.26-10.30 mg/l NH ₄
	0.20 - 8.00 mg/l NH ₃ -N
	0.24− 9.73 mg/l NH ₃
	Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4–13 If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell close with the screw cap, and mix.



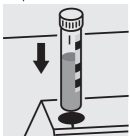
Add 1 dose of **NH**₄-**1K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄⁺, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

Measuring	0.5-16.0 mg/l NH ₄ -N
range:	0.6-20.6 mg/l NH ₄
	0.5 – 16.0 mg/l NH ₃ -N
	0.6-19.5 mg/l NH ₃
	Expression of results also possible in mmol/l.



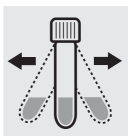
Check the pH of the sample, specified range: pH 4–13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 0.50 ml of the sample into a reaction cell close with the screw cap, and mix.



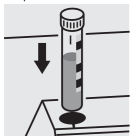
Add 1 dose of **NH₄-1K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄⁺, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.

Measuring	4.0- 80.0 mg/l NH ₄ -N
range:	5.2-103.0 mg/I NH ₄
	4.0- 80.0 mg/I NH ₃ -N
	4.9 - 97.3 mg/l NH ₃
	Expression of results also possible in mmol/l.



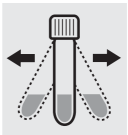
Check the pH of the sample, specified range: pH 4-13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 0.10 ml of the sample into a reaction cell close with the screw cap, and mix.



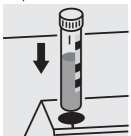
Add 1 dose of NH₄-1K using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 70, Cat.No. 250488.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 70) is highly recommended.



Measuring	0.05 -3.00 mg/I NH ₄ -N	0.06 -3.86 mg/I NH ₄	10-mm cell
range:	$0.05 - 3.00 \text{ mg/l NH}_3\text{-N}$	0.06 -3.65 mg/I NH ₃	10-mm cell
	0.03 -1.50 mg/l NH ₄ -N	0.04 -1.93 mg/l NH ₄	20-mm cell
	0.03 -1.50 mg/l NH ₃ -N	0.04 -1.82 mg/I NH ₃	20-mm cell
	0.010-0.500 mg/I NH ₄ -N	0.013-0.644 mg/I NH ₄	50-mm cell
	0.010-0.500 mg/I NH ₃ -N	0.016-0.608 mg/I NH ₃	50-mm cell
Expression of results also possible in mmol/l.			



Check the pH of the sample, specified range: pH 4-13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



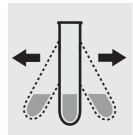
Pipette 5.0 ml of the sample into a test tube.



Add 0.60 ml of NH₄-1 with pipette and mix.



Add 1 level blue microspoon of NH₄-2.



Shake vigorously to dissolve the solid substance.



Reaction time: 5 minutes



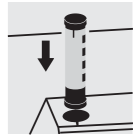
Add 4 drops of NH₄-3 and mix.



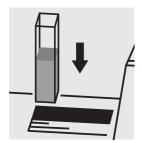
Reaction time: 5 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.



Measuring range:	2.0-75.0 mg/I NH ₄ -N	2.6- 96.6 mg/l NH ₄	10-mm cell
	5 -150 mg/I NH ₄ -N	6 -193 mg/l NH ₄	10-mm cell
	2.0-75.0 mg/I NH ₃ -N	2.4 - 91.2 mg/I NH ₃	10-mm cell
	5 -150 mg/I NH ₃ -N	6 -182 mg/I NH ₃	10-mm cell
	Expression of results also pos	ssible in mmol/l.	

Measuring range: 2.0−75.0 mg/l NH₄-N



Check the pH of the sample, specified range: pH 4–13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



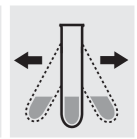
Pipette 5.0 ml of **NH₄-1** into a test tube.



Add 0.20 ml of the sample with pipette.



Add 1 level blue microspoon of **NH₄-2**.



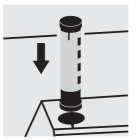
Shake vigorously to dissolve the solid substance.



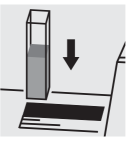
Reaction time: 15 minutes



Transfer the solution into a cell.



Select method with AutoSelector measuring range 2.0–75.0 mg/l NH₄-N.



Place the cell into the cell compartment.

Measuring range: 5-150 mg/l NH₄-N



Check the pH of the sample, specified range: pH 4–13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.

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Pipette 5.0 ml of **NH₄-1** into a test tube.



Add 0.10 ml of the sample with pipette.

Continue as mentioned above; starting from the addition of NH_4 -2 (Fig. 4). Select method with AutoSelector measuring range 5–150 mg/l NH_4 -N.

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 70, Cat.No. 250488.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄, can also be used after diluting accordingly.

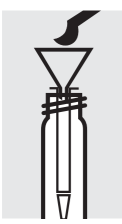
To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 70) is highly recommended.



Adsorbable organic halogens (x)

Measuring range: 0.05-2.50 mg/I AOX

Preparation of the adsorption column:



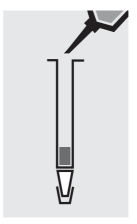
Place the column in an empty cell (Empty cells, Cat.No. 250621). Fill 1 level blue microspoon of AOX-1 into the column using the glass funnel.



Run 3 separate 1-ml portions of AOX-2 through the column. Discard the wash solution.



Run 3 separate 1-ml portions of AOX-3 through the column. Discard the wash solution.

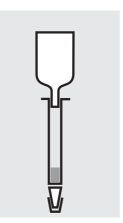


Close the bottom end of the column with the stopper. Apply to the column 1 ml of AOX-3. Close the top end of the column with the stopper and swirl to eliminate air bubbles. Remove the stopper on the top end and fill the column to the brim with AOX-3.

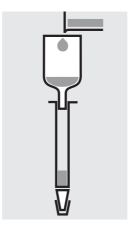
Sample enrichment:



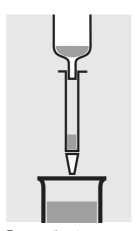
Check the pH of the sample, specified range: pH6-7.If required, add dilute sodium hydroxide solution or nitric acid drop by drop to adjust the pH.



Attach the glass reservoir to the prepared column (closed at the bottom end).



Fill 100 ml of the sample and 6 drops of AOX-4 into the reservoir.



Remove the stopper from the column outlet and run the sample through completely.



Detach the column from the reservoir. Apply 3 separate 1-ml portions of AOX-3. Discard the wash solution.



Adsorbable organic halogens (x)

Digestion:



Fill the 10-ml syringe with Add 2 level green micro- Heat the cell at 120 °C 10 ml of reagent AOX-5 and attach the syringe with the column outlet using the connector. Place the top end of the column on an empty cell (Empty cells, Cat.No. 250621) and rinse the charcoal filling of the column into an empty 16-mm cell.



spoons of AOX-6, close the cell with the screw cap, and mix.



in the thermoreactor for 30 minutes.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Add 5 drops of AOX-4, close the cell and mix; clear supernatant: pretreated sample.

Determination:



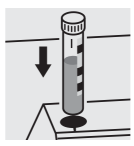
Pipette 0.20 ml of AOX-1K into a reaction cell, and mix.



Add 7.0 ml of **pretreated** Reaction time: sample with glass pipette, close the cell with the screw cap, and mix.



15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) AOX Standard, Cat. No. 250026, concentration 0.2-2.0 mg/l AOX, can be used.

Measuring0.005 – 0.100 mg/l As10-mm cellrange:0.001 – 0.020 mg/l As20-mm cellExpression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-13.



Place 350 ml of the sample into an Erlenmeyer flask with ground joint.



Add 5 drops of **As-1** and mix.



Add 20 ml of **As-2** with pipette and mix.



Add 1 level green dosing spoon of **As-3** and dissolve.



Add 1.0 ml of **As-4** with pipette and mix.



Pipette 5.0 ml of **As-5** into the absorption tube.



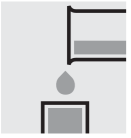
Add 1.0 ml of **As-6** with pipette to the solution in the Erlenmeyer flask and mix.



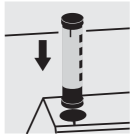
Add 3 level red dosing spoons of **As-7. Immediately** attach the absorption tube to the Erlenmeyer flask.



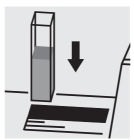
Leave to stand for 2 hours. During this time carefully swirl the flask several times or stir slowly with a magnetic stirrer.



Transfer the solution from the absorption tube into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use arsenic standard solution, concentration 1000 mg/l As, can be used after diluting accordingly.

Biochemical oxygen demand

Measuring 0.5-3000 mg/l BOD

range: $0.5-3000 \text{ mg/l O}_2$

Expression of results also possible in mmol/l.

Preparation and incubation:



Check the pH of the sample, specified range: pH 6-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Fill 2 oxygen reaction bottles each with **pretreated sample** and 2 glass beads to overflowing. Close bubble-free with the slanted ground-glass stoppers.



Fill 2 oxygen reaction bottles each with inoculated nutrient-salt solution and 2 glass beads to overflowing. Close bubble-free with the slanted ground-glass stoppers.

Measurement of inital oxygen concentration

= Result 1 (measurement sample) = Result 1 (blank)

Use one bottle of pretreated sample and one of inoculated nutrient-salt solution for the measurement of the initial oxygen concentration.



Incubate one bottle of pretreated sample and one of inoculated nutrient-salt solution closed in a thermostatic incubation cabinet at 20 ± 1°C for 5 days.

Determination:

Measurement of final oxygen concentration

= Result 2 (measurement sample) = Result 2 (blank)

After incubation, use one bottle of **pretreated sample** and one of **inoculated nutrient-salt solution** for the measurement of the final oxygen concentration.



Add 5 drops of **BSB-1K** and then 10 drops of **BSB-2K**, close bubble-free, and mix for approx. 10 seconds.



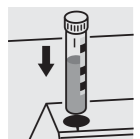
Reaction time: 1 minute



Add 10 drops of **BSB-3K**, reclose, and mix.



Fill the solution into a round cell.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

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Calculation:

BOD of measurement sample:

Result 1 - Result 2 (measurement sample) = A in mg/l

BOD of blank:

Result 1 – Result 2 (blank) = B in mg/l

BOD of original sample in mg/l = A · dilution factor – B

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) BOD Standard (acc. to EN 1899), Cat.No. 252030, can be used.



Measuring 0,05-2,00 mg/l B

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 2–12. If required, add dilute sodium hydroxide solution or nitric acid drop by drop to adjust the pH.



Pipette 1.0 ml of **B-1K** into a reaction cell, close with the screw cap, and mix.



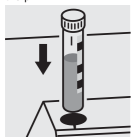
Add 4.0 ml of the sample with pipette into a reaction cell, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 60 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

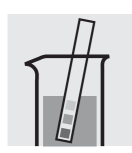
Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use boron standard solution, Cat.No. 250463, concentration 1000 mg/l B can also be used after diluting accordingly.





Measuring 0.050 – 0.800 mg/l B 10-mm cell range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 1–13.



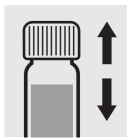
Pipette 5.0 ml of the sample into a test tube with screw cap. (Important: Do not use test tubes made of glass containing boron!)



Add 1.0 ml of **B-1** with pipette, close with the screw cap, and mix.



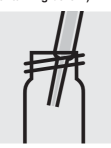
Add 1.5 ml of **B-2** with pipette and close with the screw cap.



Shake the tube vigorously for 1 minute.



Aspirate 0.5 ml of the clear lower phase from the tube with pipette.



Transfer the extract to a separate fresh tube.



Add 0.80 ml of **B-3** with pipette, close with the screw cap, and mix.



Add 4 drops of **B-4**, close with the screw cap, and mix.



Add 15 drops of **B-5**, close with the screw cap, and mix.



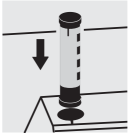
Reaction time: 12 minutes



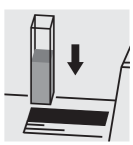
Add 6.0 ml of **B-6** with pipette, close with the screw cap, and mix.



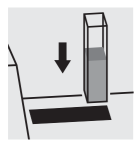
Reaction time: 2 minutes



Transfer the solution into a cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

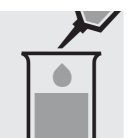
Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use boron standard solution, Cat.No. 250463, concentration 1000 mg/l B, can also be used after diluting accordingly.

Measuring	0.10 -10.00 mg/l Br ₂	10-mm cell
range:	$0.05 - 5.00 \text{ mg/l Br}_2$	20-mm cell
	0.020 - 2.000 mg/l Br ₂	50-mm cell



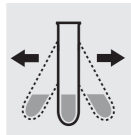
Check the pH of the sample, specified range: $pH \dot{4} - \dot{8}$. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



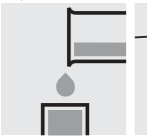
Add 1 level blue microspoon of Br₂-1.



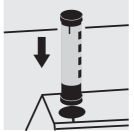
Shake vigorously to dissolve the solid substance.



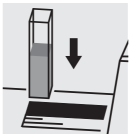
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high bromine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").

Measuring 0.025 – 1.000 mg/l Cd

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-11. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



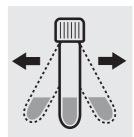
Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 0.20 ml of **Cd-1K** with pipette, close the cell with the screw cap, and mix.



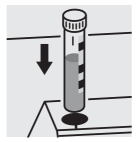
Add 1 level green microspoon of **Cd-2K**, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 2 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total cadmium** a pretreatment with Crack Set 10C, Cat.No. 252033 or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of cadmium (Σ Cd).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

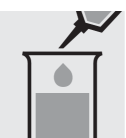
Ready-for-use cadmium standard solution, Cat.No. 250464, concentration 1000 mg/l Cd, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.

Measuring	0.010 -0.500 mg/l Cd 10-mm cell
range:	0.005 -0.250 mg/l Cd 20-mm cell
	0.0020 – 0.1000 mg/l Cd 50-mm cell
	Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-11. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 1.0 ml of Cd-1 into a test tube.



Add 10 ml of the sample with pipette and mix.



Add 0.20 ml of Cd-2 with Add 1 level green pipette and mix.



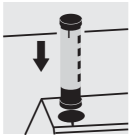
microspoon of Cd-3 and dissolve the solid substance.



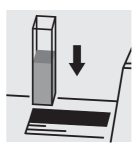
Reaction time: 2 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

For the determination of total cadmium a pretreatment with Crack Set 10C, Cat.No. 252033 or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of cadmium (Σ Cd).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use cadmium standard solution, Cat.No. 250464, concentration 1000 mg/l Cd, can be used after diluting accordingly.



 Measuring
 10-250 mg/l Ca

 range:
 14-350 mg/l CaO

 25-624 mg/l CaCO₃

Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-9. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



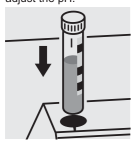
Add 1.0 ml of **Ca-1K** with pipette, close the cell with the screw cap, and mix.



Reaction time: exactly 3 minutes



Add 0.50 ml of **Ca-2K** with pipette, close the cell with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use calcium standard solution, Cat.No. 250465, concentration 1000 mg/l Ca, can be used after diluting accordingly.





Measuring	10 -160 mg/l Ca	14 -224 mg/l CaO	25 -400 mg/l CaCO ₃	10-mm cell
range:	5 - 80 mg/l Ca	7 - 112 mg/l CaO	12 -200 mg/l CaCO ₃	20-mm cell
	1.0- 15.0 mg/l Ca	1.4- 21.0 mg/l CaO	2.5- 37.5 mg/l CaCO ₃	10-mm cell (see "sensi-
				tive" preparation procedure)
				avo proparation procedure

Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 0.10 ml of the sample into a test tube.



Add 5.0 ml of **Ca-1** with pipette and mix.



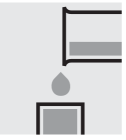
Add 4 drops of **Ca-2** and mix.



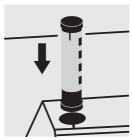
Add 4 drops of **Ca-3** and mix.



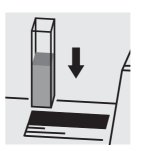
Reaction time: 8 minutes, **measure immediately.**



Transfer the solution into a corresponding cell



Select method with AutoSelector.



Place the cell into the cell compartment.

Calcium sensitive

Use the same preparation procedure as above, but add 0.50 ml of sample instead of 0.10 ml. For measurement transfer the solution into a 10-mm cell and select method **Ca sens.** in the menu (method no. 125).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use calcium standard solution, Cat.No. 250465, concentration 1000 mg/l Ca, can be used after diluting accordingly.





Measuring 5-125 mg/l Cl

range: Expression of results also possible in mmol/l.



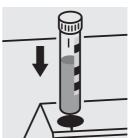
Check the pH of the sample, specified range: pH 1–12. If required, add dilute ammonia solution or nitric acid drop by drop to adjust the pH.



Pipette 0.50 ml of **CI-1K** into a reaction cell, close with the screw cap, and mix.



Add 1.0 ml of the sample with pipette, close with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10 and 20, Cat.Nos. 250482 and 250483.

Ready-for-use chloride standard solution, Cat.No. 250466, concentration 1000 mg/l Cl¯, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.

Measuring	10 -250 mg/l Cl	10-mm cell
range:	2.5- 25.0 mg/l Cl	10-mm cell
	Expression of results als	so possible in mmol/l.

Measuring range: 10-250 mg/l Cl



Check the pH of the sample, specified range: pH 1-12. If required, add dilute ammonia solution or nitric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a test tube.



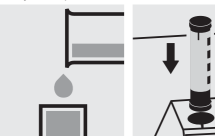
Add 2.5 ml of CI-1 with pipette and mix.



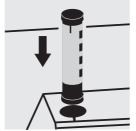
Add 0.50 ml of CI-2 with pipette and mix.



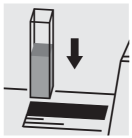
Reaction time: 1 minute



Transfer the solution into a cell.



Select method with AutoSelector measuring range 10-250 mg/l Cl.



Place the cell into the cell compartment.

Measuring range: 2.5-25.0 mg/l Cl



Check the pH of the sample, specified range: pH 1-12. If required, add dilute ammonia solution or nitric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.

Continue as mentioned above; starting from the addition of CI-1 (Fig. 3). Select method with AutoSelector measuring range 2.5-25.0 mg/l Cl.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 60, Cat.No. 250487.

Ready-for-use chloride standard solution, Cat.No. 250466, concentration 1000 mg/l Cl-, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 60) is highly recommended.

35

Determination of free chlorine

Measuring $0.03-6.00 \text{ mg/I Cl}_2$

range: Expression of results also possible in mmol/l.



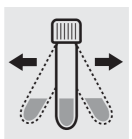
Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a round cell.



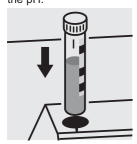
Add 1 level blue microspoon of **Cl₂-1**, close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").



Determination of free chlorine and total chlorine

Measuring 0.03 – 6.00 mg/l Cl₂
range: Expression of results also possible in mmol/l and also in free Cl₂ [Cl₂(f)], combined Cl₂
[Cl₂(b)], total Cl₂ [Cl₂(t)].

Determination of free chlorine



Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a round cell.



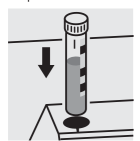
Add 1 level blue microspoon of Cl_2 -1, close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 3 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Determination of total chlorine

Same preparation as described above, add 2 drops of Cl₂-2, close the cell with the screw cap, and mix after dissolving solid.

A differentiation between free and combined chlorine $[\operatorname{Cl}_2(f)]$ and $\operatorname{Cl}_2(b)]$ can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the free chlorine, press enter, remove the cell, add 2 drops of Cl_2 -2, close with the screw cap, mix, and measure the total chlorine. After pressing enter, the individual measuring values for free and combined chlorine are shown on the display.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check). After each determination of total chlorine rinse the cell with sulfuric acid 25 % and subsequently several times with distilled water.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").

Determination of free chlorine

Measuring	$0.05 - 6.00 \text{ mg/l Cl}_2$	10-mm cell
range:	0.02 -3.00 mg/l Cl ₂	20-mm cell
	0.010 – 1.000 mg/l Cl ₂	50-mm cell



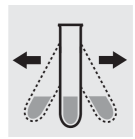
Check the pH of the sample, specified range: $pH \dot{4} - \dot{8}$. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



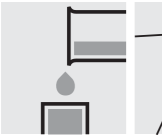
Add 1 level blue microspoon of Cl₂-1.



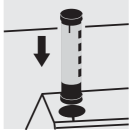
Shake vigorously to dissolve the solid substance.



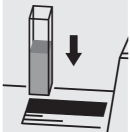
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").

Determination of total chlorine

Measuring	0.05 -6.00 mg/l Cl ₂	10-mm cell
range:	0.02 -3.00 mg/l Cl ₂	20-mm cell
	0.010 – 1.000 mg/l Cl ₂	50-mm cell



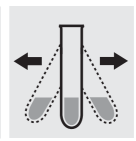
Check the pH of the sample, specified range: pH4-8.If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



Add 1 level blue microspoon of Cl₂-1.



Shake vigorously to dissolve the solid substance.



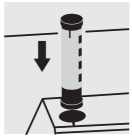
Add 2 drops of Cl₂-2 and mix.



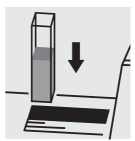
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check). After each determination of total chlorine rinse the cell with sulfuric acid 25 % and subsequently several times with distilled water.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard from Chloramine T GR can be used (see section "Standard solutions").



00599 · Chlorine

a xylem brand

Determination of free chlorine and total chlorine

Measuring	0.05 -6.00 mg/l Cl ₂	10-mm cell
range:	$0.02 - 3.00 \text{ mg/l Cl}_2$	20-mm cell
	0.010 - 1.000 mg/l Cl ₂	50-mm cell

Measuring	Expression of results also possible in mmol/l
range:	and also in free Cl ₂ [Cl ₂ (f)], combined Cl ₂
	$[Cl_2(b)]$, total $Cl_2[Cl_2(t)]$.

Determination of free chlorine



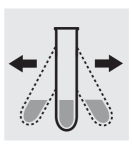
Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



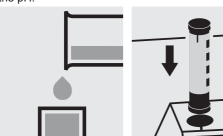
Add 1 level blue microspoon of Cl₂-1.



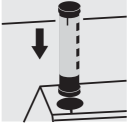
Shake vigorously to dissolve the solid substance.



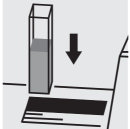
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Determination of total chlorine

Same preparation as described above, add 2 drops of Cl₂-2 and mix after dissolving solid.

A differentiation between free and combined chlorine $[Cl_2(f) \text{ and } Cl_2(b)]$ can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the free chlorine, press enter and measure the total chlorine. After pressing enter, the individual measuring values for free and combined chlorine are shown on the display.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check). After each determination of total chlorine rinse the cell with sulfuric acid 25 % and subsequently several times with distilled water.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").



00086/00087/00088 · Chlorine (with liquid reagents)

a xylem brand

Determination of free chlorine and total chlorine

 $\begin{array}{ll} \textbf{Measuring} & 0.03-6.00 \text{ mg/l Cl}_2 \\ \textbf{range:} & \text{Expression of results also possible in mmol/l} \\ & \text{and also in free Cl}_2 \left[\text{Cl}_2(f) \right] \text{, combined Cl}_2 \\ & \left[\text{Cl}_2(b) \right] \text{, total Cl}_2 \left[\text{Cl}_2(t) \right]. \\ \end{array}$

Determination of free chlorine



Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Place 6 drops of Cl₂-1 into a round cell.



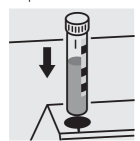
Add 3 drops of Cl₂-2, close with the screw cap, and mix.



Add 10 ml of the sample with pipette, close with the screw cap, and mix.



Reaction time: 3 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Determination of total chlorine

Same preparation as described above, add 2 drops of ${\it Cl}_2$ -3, close with the screw cap, and mix after the end of the reaction time.

A differentiation between free and combined chlorine $[Cl_2(f)]$ and $Cl_2(b)$ can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the free chlorine, press enter, remove the cell, add 2 drops of Cl_2 -3, close with the screw cap, mix, and measure the total chlorine. After pressing enter, the individual measuring values for free and combined chlorine are shown on the display.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check). After each determination of total chlorine rinse the cell with sulfuric acid 25 % and subsequently several times with distilled water.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").



00086/00087/00088 · Chlorine (with liquid reagents)

a xylem brand

Determination of free chlorine and total chlorine

Measuring	0.10-1.00 mg/I Cl ₂	50-mm cell
range:	Expression of results also	possible in mmol/l
	and also in free Cl ₂ [Cl ₂ (f)], combined Cl ₂
	[Cl ₂ (b)], total Cl ₂ [Cl ₂ (t)].	

Determination of free chlorine



Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Place 6 drops of Cl₂-1 into a test tube.



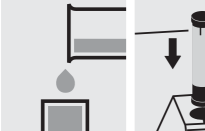
Add 3 drops of Cl₂-2, close with the screw cap, and mix.



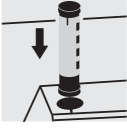
Add 10 ml of the sample with pipette, close with the screw cap, and mix.



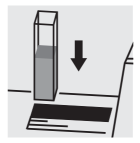
Reaction time: 3 minutes, measure immediately.



Transfer the solution into Select method with a cell.



AutoSelector.



Place the cell into the cell compartment.

Determination of total chlorine

Same preparation as described above, add 2 drops of Cl₂-3 and mix after the end of the reaction time.

A differentiation between free and combined chlorine $[Cl_2(f) \text{ and } Cl_2(b)]$ can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the free chlorine, press enter, remove the cell, add 2 drops of Cl₂-3, mix using the microspatula, and measure the total chlorine. After pressing enter, the individual measuring values for free and combined chlorine are shown on the display.

Important:

Very high chlorine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check). After each determination of total chlorine rinse the cell with sulfuric acid 25 % and subsequently several times with distilled water.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").





Measuring	0.10 -10.00 mg/l ClO ₂	10-mm cell
range:	0.05 - 5.00 mg/l ClO ₂	20-mm cell
	0.020- 2.000 mg/l ClO ₂	50-mm cell
	Expression of results also p	oossible in mmol/l.



Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



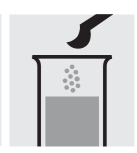
Pipette 10 ml of the sample into a test tube.



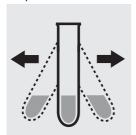
Add 2 drops of CIO₂-1 and mix.



Reaction time: 2 minutes



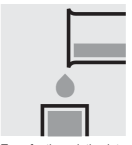
Add 1 level blue microspoon of CIO₂-2.



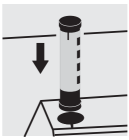
Shake vigorously to dissolve the solid substance.



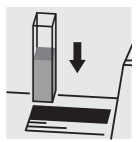
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").





Determination of chromium(VI)

Measuring 0.05-2.00 mg/l Cr

range: 0.11 – 4.46 mg/l CrO₄

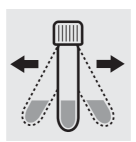
Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 1–9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Add 6 drops of **Cr-3K** into a reaction cell, close with the screw cap.



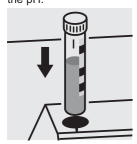
Shake the cell vigorously to dissolve the solid substance and leave to stand for 1 minute.



Add 5.0 ml of the sample with pipette, close the cell with the screw cap, and mix.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use chromate standard solution, Cat.No. 250468, concentration 1000 mg/l CrO₄²⁻, can be used after diluting accordingly.



Determination of total chromium = sum of chromium(VI) and chromium(III)

Measuring 0.05 – 2.00 mg/l Cr

range: 0.11 – 4.46 mg/I CrO₄

Expression of results also possible in mmol/l and also in Cr total (Σ Cr), Cr(III), and Cr(VI).



Check the pH of the sample, specified range: pH 1–9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1 drop of **Cr-1K**, close with the screw cap, and mix.



Add 1 dose of **Cr-2K** using the blue dosemetering cap, close the reaction cell with the screw cap.



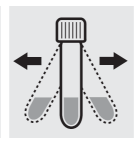
Heat the cell in the thermoreactor at 120 °C (100 °C) for 1 hour.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature: **pretreated sample.**



Add 6 drops of **Cr-3K** into a reaction cell, close the cell with the screw cap.



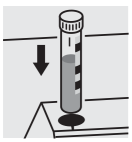
Shake the cell vigorously to dissolve the solid substance and leave to stand for **1 minute**.



Add 5.0 ml of the **pre-treated sample** with pipette, close with the screw cap, and mix.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between chromium(VI) and chromium(III) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the total chromium, press enter and measure the chromium(VI) (see analytical procedure for chromium(VI)). After pressing enter, the individual measuring values for Cr VI and Cr III are shown on the display.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use chromate standard solution, Cat.No. 250468, concentration 1000 mg/l CrO₄²⁻, can be used after diluting accordingly.

Determination of chromium(VI)

Measuring	0.05 -3.00 mg/l Cr	0.11 - 6.69 mg/I CrO ₄	10-mm cell
range:	0.03 -1.50 mg/l Cr	0.07-3.35 mg/I CrO ₄	20-mm cell
	0.010-0.600 mg/l Cr	0.02-1.34 mg/l CrO ₄	50-mm cell
	Expression of results also	oossible in mmol/l.	



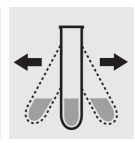
Check the pH of the sample, specified range: pH 1-9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Place 1 level grey microspoon of Cr-1 into a dry test tube.



Add 6 drops of Cr-2.



Shake the test tube vigorously to dissolve the solid substance.



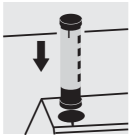
Add 5.0 ml of the sample with pipette and mix.



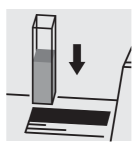
Reaction time: 1 minute



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

For the determination of total chromium = sum of chromium(VI) and chromium(III) a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of chromium (Σ Cr).

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use chromate standard solution, Cat.No. 250468, concentration 1000 mg/l CrO₄²⁻, can be used after diluting accordingly.

Chemical oxygen demand

10-150 mg/I COD or O₂ Measuring range:



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 3.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.





Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



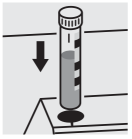
Remove the cell from the thermoreactor and place in a test-tube rack to cool.



Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature. Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.



Chemical oxygen demand

4.0-40.0 mg/I COD or O₂ Measuring range:

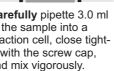


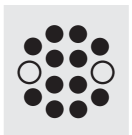
Suspend the bottom sediment in the cell by swirling.



Carefully pipette 3.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously. Caution, the cell

becomes hot!





Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



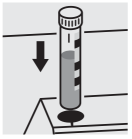
Remove the cell from the thermoreactor and place in a test-tube rack to cool.



Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature. Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.

Chemical oxygen demand

Measuring range: 5.0-80.0 mg/I COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

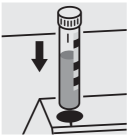


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.

Chemical oxygen demand

Measuring range: 10-150 mg/l COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 3.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.





Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

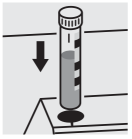


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.



Chemical oxygen demand

Measuring range: 15-300 mg/l COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

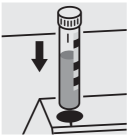


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 60, Cat.No. 250487.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 60) is highly recommended.



Chemical oxygen demand

Measuring range: 50-500 mg/l COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



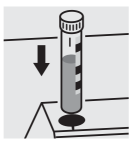
Remove the cell from the thermoreactor and place in a test-tube rack to cool.



Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature. **Very important!**



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 60, Cat.No. 250487.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 60) is highly recommended.

Chemical oxygen demand

Measuring range: 25-1500 mg/I COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 3.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

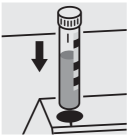


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 20) is highly recommended.

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Chemical oxygen demand

Measuring range: 25-1500 mg/I COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 3.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

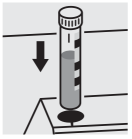


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 20) is highly recommended.

Chemical oxygen demand

Measuring range: 300-3500 mg/l COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.
Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

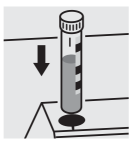


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 80, Cat.No. 250489.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 80) is highly recommended.



Chemical oxygen demand

Measuring range: 500-10000 mg/I COD or O₂



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 1.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

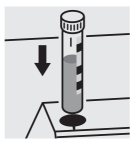


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 70, Cat.No. 250488.

To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 70) is highly recommended.



Chemical oxygen demand

5000-90000 mg/I COD or O₂ Measuring range:



Suspend the bottom sediment in the cell by swirling.



Carefully pipette 0.10 ml of the sample into a reaction cell, close 148 °C for 2 hours. tightly with the screw cap, and mix vigorously. Caution, the cell becomes hot!



Heat the reaction cell in the thermoreactor at



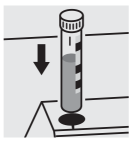
Remove the cell from the thermoreactor and place in a test-tube rack to cool.



Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature. Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.



09772 · COD (Hg-free)

a xylem brand

Chemical oxygen demand

Measuring range: 10-150 mg/l COD or O₂



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell

becomes hot!



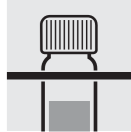
Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

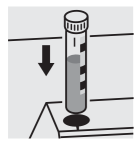


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

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09773 · COD (Hg-free)

a xylem brand

Chemical oxygen demand

Measuring range: 100-1500 mg/I COD or O₂



Carefully pipette 2.0 ml of the sample into a reaction cell, close tightly with the screw cap, and mix vigorously.

Caution, the cell becomes hot!



Heat the reaction cell in the thermoreactor at 148 °C for 2 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool.

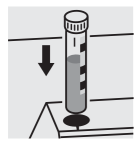


Swirl the cell after 10 minutes.



Replace the cell in the rack for complete cooling to room temperature.

Very important!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 20) is highly recommended.





Measuring 0.05 – 8.00 mg/l Cu

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



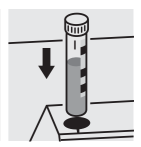
Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 5 drops of **Cu-1K**, close the cell with the screw cap, and mix.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high copper concentrations in the sample produce turquoise-coloured solutions (measurement solution should be blue) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

For the determination of **total copper** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of copper (Σ Cu).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use copper standard solution, Cat.No. 250473, concentration 1000 mg/l Cu, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.



Measuring	0.10-6.00 mg/l Cu	10-mm cell
range:	0.05-3.00 mg/l Cu	20-mm cell
	0.02-1.20 mg/l Cu	50-mm cell
	Expression of results also possible in mmol/l.	



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1 green dosing spoon of Cu-1 and dissolve the solid substance.



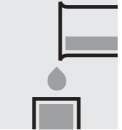
Check the pH, specified range: pH 7.0-9.5. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



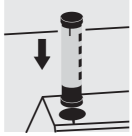
Add 5 drops of Cu-2 and mix.



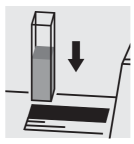
Reaction time: 5 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high copper concentrations in the sample produce turquoise-coloured solutions (measurement solution should be blue) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

For the determination of total copper a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of copper (Σ Cu).

To measure in the 50-mm cell, only the sample volume has to be doubled.

Alternatively, the semi-microcell can be used.

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Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use copper standard solution, Cat.No. 250473, concentration 1000 mg/l Cu, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.



14561 · Cyanide

a xylem brand

Determination of free cyanide

Measuring 0.010 – 0.500 mg/I CN

range: Expression of results also possible in mmol/l

and cyanide free [CN(f)].



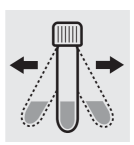
Check the pH of the sample, specified range: pH 4.5–8.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and dissolve the solid substance.



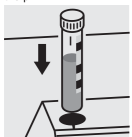
Add 1 level blue microspoon of **CN-3K**, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) cyanide standard solution can be used.

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14561 · Cyanide

a **xylem** brand

Determination of readily liberated cyanide

Measuring 0.010 – 0.500 mg/I CN

range: Expression of results also possible in mmol/l

and cyanide readily liberated [CN(v)].



Check the pH of the sample, specified range: pH 4.5–8.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1 dose of **CN-1K** using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Swirl the cell before opening.



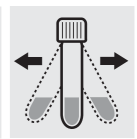
Add 3 drops of **CN-2K**, close with the screw cap, and mix: **pretreated** sample.



Pipette 5.0 ml of the **pretreated sample** into a reaction cell, close with the screw cap, and dissolve the solid substance.



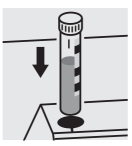
Add 1 level blue microspoon of **CN-3K**, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) cyanide standard solution can be used.

Determination of free cyanide

Measuring	0.010 -0.500 mg/I CN	10-mm cell
range:	0.005 -0.250 mg/l CN	20-mm cell
	0,0020-0,1000 mg/l CN	50-mm cell
	Expression of results also possible in mmol/l and cyanide free [CN(f)].	



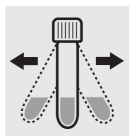
Check the pH of the sample, specified range: pH 4.5-8.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



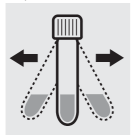
Add 1 level green microspoon of CN-3, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Add 1 level blue microspoon of CN-4, close the cell with the screw



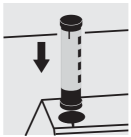
Shake the cell vigorously to dissolve the solid substance.



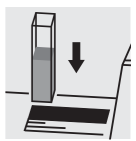
Reaction time: 10 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus preventing any gas losses.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents CN-3 and CN-4 have to be doubled for each.

Alternatively, the semi-microcell can be used.

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Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) cyanide standard solution can be used.

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Determination of readily liberated cyanide

Measuring	0.010 -0.500 mg/I CN	10-mm cell
range:	0.005 -0.250 mg/I CN	20-mm cell
	0,0020-0,1000 mg/l CN	50-mm cell
	Expression of results also possib	le in mmol/l and cyanide readily liberated [CN(v)].



Check the pH of the sample, specified range: pH 4.5-8.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



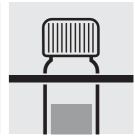
Add 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1 dose of CN-1 using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Swirl the cell before opening.



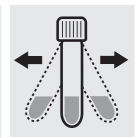
Add 3 drops of CN-2, close with the scew cap, and mix: pretreated sample.



Pipette 5.0 ml of the pretreated sample into an empty round cell (Empty cells, Cat.No. 250621).



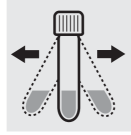
Add 1 level green microspoon of CN-3, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Add 1 level blue microspoon of CN-4, close the cell with the screw сар.



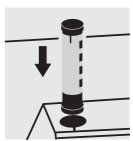
Shake the cell vigorously to dissolve the solid substance.



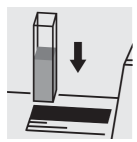
Reaction time: 10 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

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Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus preventing any gas losses.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents CN-3 and CN-4 have to be doubled for each.

Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) cyanide standard solution can be used.

Measuring range: 2 – 160 mg/l cyanuric acid 20-mm cell



Filter turbid samples.



Pipette 5.0 ml of the sample into into a empty round cell (Empty cells, Cat.No. 250621).



Add **5.0 ml of distilled water** with pipette, close with the screw cap, and mix



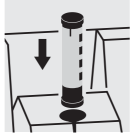
Add 1 tablet **Cyanuric Acid**, crush with stirring rod, and close with the screw cap.



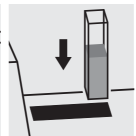
Swirl the cell to dissolve the solid substance.



Transfer the solution into a cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a cyanuric acid standard solution must be prepared (see section "Standard solutions").



Measuring	0.10 -1.80 mg/l F	Round cell
range:	0.025-0.500 mg/l F	50-mm cell
	Expression of results also possible in mmol/l.	

Measuring range: 0.10 – 1.80 mg/l F



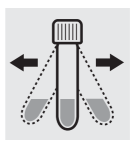
Check the pH of the sample, specified range: pH 3 - 8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 level blue microspoon of **F-1K**, close the cell with the screw cap.



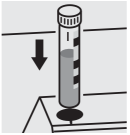
Shake the cell vigorously to dissolve the solid substance.



Reaction time: 15 minutes



Swirl the cell before measurement.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high fluoride concentrations in the sample produce brown-coloured solutions (measurement solution should be violet) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use fluoride standard solution, Cat.No. 250470, concentration 1000 mg/l F⁻, can be used after diluting accordingly.



Measuring range: 0.025 – 0.500 mg/l F



Check the pH of the sample, specified range: pH 3 - 8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



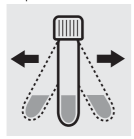
Pipette 10 ml of the sample into a reaction cell, close with the screw cap, and mix.



Pipette 10 ml of distilled water into a second reaction cell, close with the screw cap, and mix. (Blank)



Add 1 level blue microspoon of **F-1K** to each cell, close with the screw cap.



Shake both cells vigorously to dissolve the solid substance.



Select method F sens

in the menu (method

no. 216).

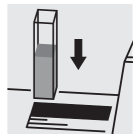
Reaction time: 15 minutes



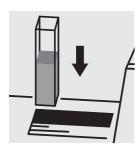
Swirl the cells.



Transfer both solutions into two separate 50-mm-cells.



Place the blank cell into the cell compartment.



Place the cell containing the sample into the cell compartment.

Important:

Very high fluoride concentrations in the sample produce brown-coloured solutions (measurement solution should be violet) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use fluoride standard solution, Cat.No. 250470, concentration 1000 mg/l F⁻, can be used after diluting accordingly.

Measuring range: 0.10 - 2.00 mg/l F 10-mm cell

1.0 -20.0 mg/l F 10-mm cell

Expression of results also possible in mmol/l.

Measuring range: 0.10-2.00 mg/l F



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



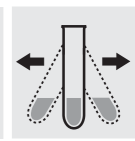
Pipette 2.0 ml of **F-1** into a test tube.



Add 5.0 ml of the sample with pipette and mix.



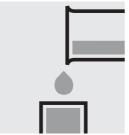
Add 1 level blue microspoon of **F-2** and mix.



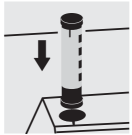
Shake the test tube vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Transfer the solution into a cell.



Select method with AutoSelector measuring range 0.10 – 2.00 mg/l F.



Place the cell into the cell compartment.

Measuring range: 1.0-20.0 mg/l F



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 2.0 ml of **F-1** into a test tube.



Add 5.0 ml of water and 0.5 ml of the sample with pipette and mix.

Continue as mentioned above; starting from the addition of **F-2** (Fig. 4). Select method with AutoSelector measuring range 1.0–20.0 mg/l F.

Important:

Very high fluoride concentrations in the sample produce brown-coloured solutions (measurement solution should be violet) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use fluoride standard solution, Cat.No. 250470, concentration 1000 mg/l F⁻, can be used after diluting accordingly.

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Measuring 0.10-8.00 mg/I HCHO

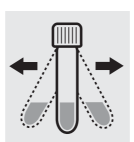
range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-13.



Add 1 level green microspoon of **HCHO-1K** into a reaction cell, close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.

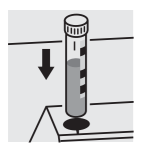


Add 2.0 ml of the sample with pipette, close the cell with the screw cap, and mix.

Caution, cell becomes hot!



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a formaldehyde standard solution must be prepared from Formaldehyde solution 37% (see section "Standard solutions").





Measuring	0.10-8.00 mg/I HCHO	10-mm cell
range:	0.05-4.00 mg/I HCHO	20-mm cell
	0.02-1.50 mg/I HCHO	50-mm cell
	Expression of results also possible in mmol/l.	



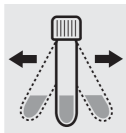
Check the pH of the sample, specified range: pH 0-13.



Pipette 4.5 ml of **HCHO-1** into an empty round cell (Empty cells, Cat.No. 250621).



Add 1 level green microspoon of HCHO-2, close the cell with the screw cap.



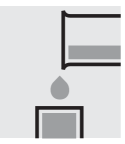
Shake the cell vigorously to dissolve the solid substance.



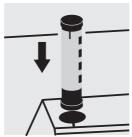
Add 3.0 ml of the sample with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



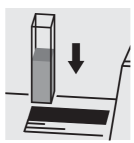
Reaction time: 5 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a formaldehyde standard solution must be prepared from Formaldehyde solution 37% (see section "Standard solutions").



0.5-12.0 mg/l Au Measuring 10-mm cell range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 1-9. If required, add dilute hydrochloric acid drop by drop to adjust the pH.



Pipette 2.0 ml of the sample into a test tube with screw cap.



Add 2 drops of Au-1 and Add 4 drops of Au-2 and Add 6 drops of Au-3 and mix.



mix.



mix.



Add 6.0 ml of Au-4 with pipette, close with the screw cap.



Shake the tube vigorously for 1 minute.



Add 6 drops of Au-5, close with the screw сар.



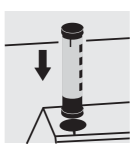
Shake the tube vigorously for 1 minute.



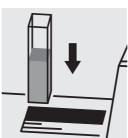
Aspirate the clear upper phase from the tube with pipette.



Transfer the solution into Select method with a cell.



AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) gold standard solution can be used.





Measuring	$0.02 - 2.00 \text{ mg/l N}_2\text{H}_4$	10-mm cell
range:	$0.01 - 1.00 \text{ mg/l N}_2\text{H}_4$	20-mm cell
	$0.005 - 0.400 \text{ mg/l N}_2\text{H}_4$	50-mm cell
	Expression of results also	possible in mmol/l.



Check the pH of the sample, specified range: pH 2–10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



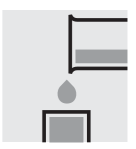
Pipette 5.0 ml of the sample into a test tube.



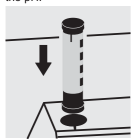
Add 2.0 ml of **Hy-1** with pipette and mix.



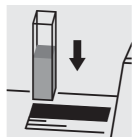
Reaction time: 5 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a hydrazine standard solution must be prepared (see section "Standard solutions").





Measuring range: 2.0 – 20.0 mg/l H₂O₂



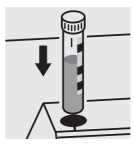
Check the pH of the sample, specified range: pH 0 – 10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a reaction cell, close with the screw cap, and mix.



Reaction time: 2 minutes

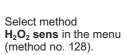


Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Measuring range: 0.25 – 5.00 mg/l H₂O₂



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.





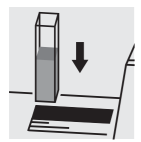
Pipette 10 ml of the sample into a reaction cell, close with the screw cap, and mix.



Reaction time: 2 minutes



Transfer the solution into a 50-mm cell.



Place the cell into the cell compartment.

Important:

The contents of the reaction cells may be slightly yellow. However, this does not influence the measurement result.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a hydrogenperoxide standard solution must be prepared from Perhydrol $^{\otimes}$ 30 % H_2O_2 GR (see section "Standard solutions").





Measuring 0.03 -6.00 mg/I H₂O₂ 10-mm cell range: 0.015 - 3.000 mg/I H₂O₂ 20-mm cell Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 0.50 ml of H_2O_2 -1 into a test tube.



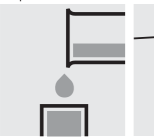
Add 8.0 ml of the sample with pipette and mix.



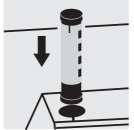
Add 0.50 ml of H_2O_2 -2 with pipette and mix.



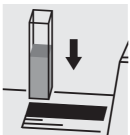
Reaction time: 10 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a hydrogenperoxide standard solution must be prepared from Perhydrol® 30% H₂O₂ GR (see section "Standard solutions").



Measuring	$0.20 - 10.00 \text{ mg/l l}_2$	10-mm cell
range:	$0.10 - 5.00 \text{ mg/l l}_2$	20-mm cell
	0.050- 2.000 mg/l l ₂	50-mm cell



Check the pH of the sample, specified range: $pH \dot{4} - \dot{8}$. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



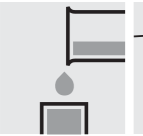
Add 1 level blue microspoon of I_2 -1.



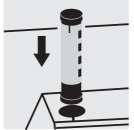
Shake vigorously to dissolve the solid substance.



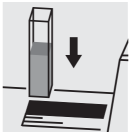
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high iodine concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").



Measuring 0.05-4.00 mg/l Fe

range: Expression of results also possible in mmol/l.



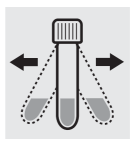
Check the pH of the sample, specified range: pH 1–10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



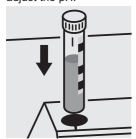
Add 1 level blue microspoon of **Fe-1K**, close the cell with the screw



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 3 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total iron** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of iron (Σ Fe).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use iron standard solution, Cat.No. 250469, concentration 1000 mg/l Fe, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.



Determination of iron(II) and iron(III)

Measuring 1.0-50.0 mg/l Fe

range: Expression of results also possible in mmol/l

and also in Fe(II), Fe(III).

Determination of iron (II)



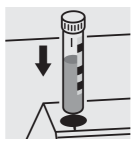
Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Determination of iron (II + III)



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



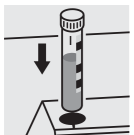
Add 1 dose of **Fe-1K** using the blue dosemetering cap, close the reaction cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between iron(II) and iron(III) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form.

Then measure the iron(II + III), press enter and measure the iron(II). After pressing enter, the individual measuring values for Fe II and Fe III are shown on the display.

Important:

For the determination of **total iron** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of iron (Σ Fe).

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Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use iron standard solution, Cat.No. 250469, concentration 1000 mg/l Fe(III), can be used after diluting accordingly.



Measuring	0.05 -5.00 mg/l Fe	10-mm cell
range:	0.03 -2.50 mg/l Fe	20-mm cell
	0.005-1.000 mg/l Fe	50-mm cell
	Expression of results also	possible in mmol/l.



Check the pH of the sample, specified range: pH 1–10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



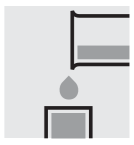
Pipette 5.0 ml of the sample into a test tube.



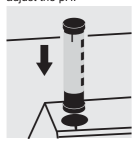
Add 3 drops of **Fe-1** and mix.



Reaction time: 3 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

For the determination of **total iron** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of iron (Σ Fe).

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use iron standard solution, Cat.No. 250469, concentration 1000 mg/l Fe, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.



Determination of iron(II) and iron(III)

Measuring	0.10 -5.00 mg/l Fe	10-mm cell	
range:	0.05 -2.50 mg/l Fe	20-mm cell	
	0.010-1.000 mg/l Fe	50-mm cell	
	Expression of results also possible in mmol/l.		

Determination of iron(II)



Check the pH of the sample, specified range: pH 2-8. If required, add dilute sodium hydroxide solution or nitric acid drop by drop to adjust the pH.



Pipette 8.0 ml of the sample into a test tube.



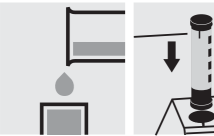
Add 1 drop of Fe-1 and mix.



Add 0.50 ml of Fe-2 with Reaction time: pipette and mix.

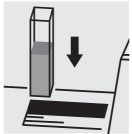


5 minutes



Transfer the solution into Select method with a corresponding cell.

AutoSelector.



Place the cell into the cell compartment.

Determination of iron(II + III)

Same preparation as discribed above. After adding of Fe-2 continue with Fe-3.



Add 1 dose of Fe-3 using the blue dosemetering cap and dissolve the solid substance.



Reaction time: 10 minutes, then measure.

A differentiation between iron(II) and iron(III) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form.

Then measure the iron(II), press enter and measure the iron(II + III). After pressing enter, the individual measuring values for Fe II and Fe III are shown on the display.

Important:

For the determination of total iron a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use iron standard solution, Cat.No. 250469, concentration 1000 mg/l Fe(III), can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.

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Measuring 0.10-5.00 mg/l Pb

range: Expression of results also possible in mmol/l.

Samples of total hardness 0-14 °d



Check the total hardness of the sample.



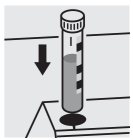
Check the pH of the sample, specified range: pH 3-6. If required, add dilute ammonia solution or nitric acid drop by drop to adjust the pH.



Add 5 drops of **Pb-1K** into a reaction cell and mix.



Add 5.0 ml of the sample with pipette, close the cell with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer = Result A

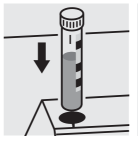
Samples of total hardness > 14 °d



Add 1 level grey microspoon of **Pb-2K** to the already measured cell, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer

= Result B

Result A

- Result B

= mg/l Pb

Important:

For the determination of **total lead** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of lead (Σ Pb).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

Ready-for-use lead standard solution, Cat.No. 250462, concentration 1000 mg/l Pb, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.

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Measuring	0.10 -5.00 mg/l Pb	10-mm cell
range:	0.05 -2.50 mg/l Pb	20-mm cell
	0.010-1.000 mg/l Pb	50-mm cell
	Expression of results also	possible in mmol/l.



Check the pH of the sample, specified range: pH 3-6. If required, add dilute ammonia solution or nitric acid drop by drop to adjust the pH.



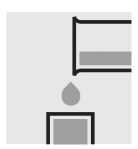
Pipette 0.50 ml of **Pb-1** into a test tube.



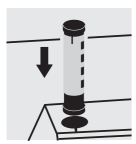
Add 0.50 ml of **Pb-2** with pipette and mix.



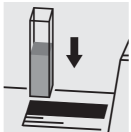
Add 8.0 ml of the sample with pipette and mix.



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

For the determination of **total lead** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of lead (Σ Pb).

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Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

Ready-for-use lead standard solution, Cat.No. 250462, concentration 1000 mg/l Pb, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.



Measuring 5.0-75.0 mg/l Mg

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-9. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



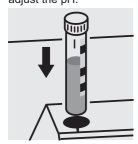
Add 1.0 ml of **Mg-1K** with pipette, close the cell with the screw cap, and mix.



Reaction time: Exactly 3 minutes.



Add 3 drops of **Mg-2K**, close the cell with the screw cap and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").



Measuring 0.10-5.00 mg/l Mn

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 2-7. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 7.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 2 drops of **Mn-1K**, close the cell with the screw cap, and mix.



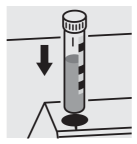
Reaction time: 2 minutes



Add 3 drops of **Mn-2K**, close the cell with the screw cap, and mix.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use manganese standard solution, Cat.No. 250474, concentration 1000 mg/l Mn, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.

Measuring	0.05 -2.00 mg/l Mn	10-mm cell
range:	0.03 -1.00 mg/l Mn	20-mm cell
	0.005 - 0.400 mg/l Mn	50-mm cell
	Expression of results also possible in mmol/l.	



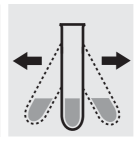
Check the pH of the sample, specified range: pH 3 – 10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



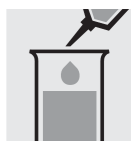
Pipette 8.0 ml of the sample into a test tube.



Add 1 level grey microspoon of **Mn-1**.



Shake the tube vigorously to dissolve the solid substance.



Add 2.0 ml of **Mn-2** with pipette and mix.



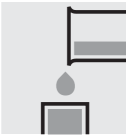
Add **carefully** 3 drops of **Mn-3** and mix.



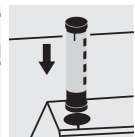
Add carefully 0.25 ml of Mn-4 with pipette and mix carefully (Foams! Wear eye protection!).



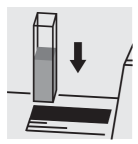
Reaction time: 10 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

When using the 50-mm cell, perform the measurement against a separately prepared blank (preparation as per measurement sample, but with distilled water instead of sample).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use manganese standard solution, Cat.No. 250474, concentration 1000 mg/l Mn, can be used after diluting accordingly.

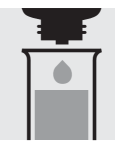
Measuring	0.50 -10.00 mg/l Mn	10-mm cell
range:	0.25 - 5.00 mg/l Mn	20-mm cell
	0.010- 2.000 mg/l Mn	50-mm cell
	Expression of results also possible in mmol/l.	



Check the pH of the sample, specified range: pH2-7.If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 4 drops of Mn-1 and mix.



Add 2 drops of Mn-2 and mix. Check the pH, specified pH: approx. 11.5.



Reaction time: 2 minutes



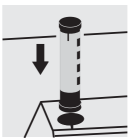
Add 2 drops of Mn-3 and mix.



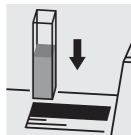
Reaction time: 2 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 30, Cat.No. 250484.

Ready-for-use manganese standard solution, Cat.No. 250474, concentration 1000 mg/l Mn, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 30) is highly recommended.



 $\begin{tabular}{lll} \textbf{Measuring} & 0.02-1.00 \ mg/l \ Mo \\ \hline \textbf{range:} & 0.02-1.67 \ mg/l \ MoO_4 \\ & 0.04-2.15 \ mg/l \ Na_2MoO_4 \\ \hline & Expression \ of \ results \ also \ possible \ in \ mmol/l. \\ \hline \end{tabular}$



Check the pH of the sample, specified range: pH 1–10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Place 2 drops of **Mo-1K** into a reaction cell and mix.



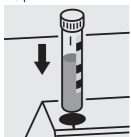
Add 10 ml of the sample with pipette, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 2 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) molybdenum standard solution can be used.

Measuring range: 0.5 - 45.0 mg/l Mo	20-mm cell
0.8 - 75.0 mg/I MoO ₄	20-mm cell
1.1 – 96.6 mg/l Na ₂ MoO ₄	20-mm cell



Pipette 10 ml of the sample into into a empty round cell (Empty cells, Cat.No. 250621).



Add 1 powder pack of Molybdenum HR1, close with the screw cap, and dissolve the solid substance.



Add 1 powder pack of Molybdenum HR2, close with the screw cap, and dissolve the solid substance.



Add 1 powder pack of Molybdenum HR3 and close with the screw



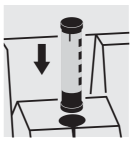
Swirl the cell to dissolve the solid substance.



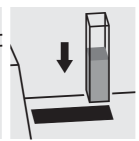
Reaction time: 5 minutes, measure immediately



Transfer the solution into Select method with a cell.



AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) molybdenum standard solution can be used.





Measuring	0.25 -10.00 mg/l Cl ₂	0.18 -7.25 mg/I NH ₂ CI	0.05 -1.96 mg/l NH ₂ Cl-N	10-mm-cell
range:	$0.13 - 5.00 \text{ mg/l Cl}_2$	0.09 -3.63 mg/I NH ₂ CI	$0.03 - 0.98 \text{ mg/l NH}_2\text{Cl-N}$	20-mm cell
	0.050- 2.000 mg/l Cl ₂	0.036-1.450 mg/I NH ₂ CI	0.010-0.392 mg/l NH ₂ Cl-N	50-mm cell
Expression of results also possible in mmol/l.				



Check the pH of the sample, specified range: pH 4-13. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 10 ml of the sample into a test tube.



Add 0.60 ml of **MCA-1** with pipette and mix.



Reaction time: 5 minutes



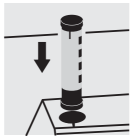
Add 4 drops of MCA-2 and mix.



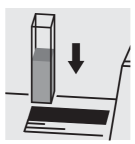
Reaction time: 5 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high monochloramine concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a standard solution must be prepared (see section "Standard solutions").

Measuring 0.10-6.00 mg/l Ni

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Reaction time: 1 minute



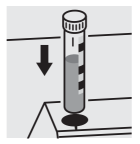
Add 2 drops of **Ni-1K**, close with the screw cap, and mix.



Add 2 drops of **Ni-2K**, close the cell with the screw cap, and mix.



Reaction time: 2 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total nickel** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of nickel (Σ Ni).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

A nickel standard solution, Cat.No. 250475, concentration 1000 mg/l Ni, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.

90



Measuring	0.10-5.00 mg/l Ni	10-mm cell
range:	0.05-2.50 mg/l Ni	20-mm cell
	0.02-1.00 mg/l Ni	50-mm cell
	Expression of results also possible in mmol/l.	



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1 drop of Ni-1 and mix. If the colour disappears, continue adding drop by drop until a slight yellow colouration persists.



Reaction time: 1 minute



Add 2 drops of Ni-2 and mix.



Check the pH, specified range: pH 10-12. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



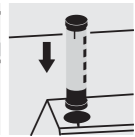
Add 2 drops of Ni-3 and mix.



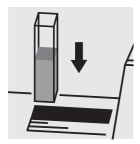
Reaction time: 2 minutes



a corresponding cell.



Transfer the solution into Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

For the determination of total nickel a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496 and thermoreactor is necessary.

Result can be expressed as sum of nickel (Σ Ni).

03/2021

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

A nickel standard solution, Cat.No. 250475, concentration 1000 mg/l Ni, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.

 $\label{eq:measuring} \textbf{Measuring} \qquad 0.5 - \ 25.0 \ \text{mg/l NO}_3\text{-N}$

range: 2.2-110.7 mg/I NO₃

Expression of results also possible in mmol/l.



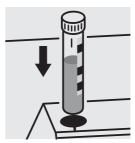
Pipette 1.0 ml of the sample into a reaction cell, **do not mix**.



Add 1.0 ml of NO₃-1K with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.



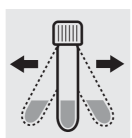
 $\textbf{Measuring} \qquad 0.5-18.0 \text{ mg/l NO}_3\text{-N}$

range: 2.2-79.7 mg/l NO₃

Expression of results also possible in mmol/l.



Add 1 level yellow micro-spoon of NO₃-1K into a reaction cell and close with the screw cap.



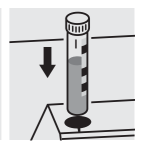
Shake the cell vigorously for 1 minute to dissolve the solid substance.



Add very slowly 1.5 ml of the sample with pipette, close with the screw cap, and mix briefly. Caution, cell becomes hot!



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.



 $\label{eq:measuring} \textbf{Measuring} \qquad 0.5 - \ 25.0 \ \text{mg/l NO}_3\text{-N}$

range: 2.2-110.7 mg/I NO₃

Expression of results also possible in mmol/l.



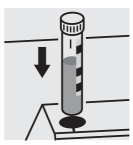
Pipette 1.0 ml of the sample into a reaction cell, **do not mix.**



Add 1.0 ml of NO₃-1K with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 20) is highly recommended.

 $\textbf{Measuring} \qquad 1.0 - \;\; 50.0 \; \text{mg/I NO}_3\text{-N}$

range: $4 - 221 \text{ mg/l NO}_3$

Expression of results also possible in mmol/l.



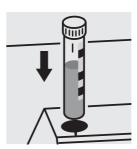
Pipette 0.50 ml of the sample into a reaction cell, **do not mix.**



Add 1.0 ml of NO₃-1K with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 80, Cat.No. 250489.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 80) is highly recommended.





Measuring $23-225 \text{ mg/I NO}_3-\text{N}$ range: $102-996 \text{ mg/I NO}_3$

Expression of results also possible in mmol/l.



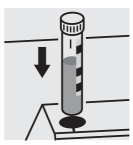
Pipette 1.0 ml of NO₃-1K into a reaction cell, do not mix.



Add 0.10 ml of the sample with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



Reaction time: 5 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO $_{\overline{3}}$, can also be used after diluting accordingly.



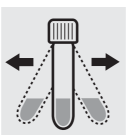
Measuring	0.5-20.0 mg/I NO ₃ -N	2.2-88.5 mg/l NO ₃	10-mm cell
range:	0.2-10.0 mg/I NO ₃ -N	0.9-44.3 mg/I NO ₃	20-mm cell
	Expression of results also	possible in mmol/l.	



Place 1 blue microspoon of NO₃-1 into a dry empty round cell (Empty cells, Cat.No. 250621).



Add 5.0 ml of NO₃-2 with Shake vigorously for pipette into the cell. Close the cell with the screw cap.



1 minute to dissolve the solid substance.



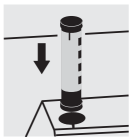
Add very slowly 1.5 ml of the sample with pipette, close the cell with the screw cap, and mix briefly. Caution, cell becomes hot!



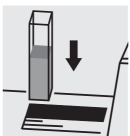
Reaction time: 10 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10 and 20, Cat.Nos. 250482 and 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO₃, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.



Measuring	1.0 – 25.0 mg/I NO ₃ -N	4.4-110.7 mg/I NO ₃	10-mm cell
range:	0.5 - 12.5 mg/l NO ₃ -N	2.2- 55.3 mg/I NO ₃	20-mm cell
	0.10- 5.00 mg/I NO ₃ -N	0.4- 22.1 mg/I NO ₃	50-mm cell
	Expression of results also poss	sible in mmol/l.	



Pipette 4.0 ml of NO₃-1 into a dry empty round cell (Empty cells, Cat. No. 250621).



Add 0.50 ml of the sample with pipette, do not mix.



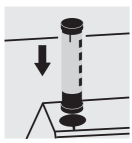
Add 0.50 ml of NO₃-2 with pipette, close the cell with the screw cap, and mix. Caution, cell becomes hot!



Reaction time: 10 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO₃, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.





in seawater

 $\label{eq:measuring} \textbf{Measuring} \qquad 0.10 - \ 3.00 \ \text{mg/I NO}_3\text{-N}$

range: $0.4 - 13.3 \text{ mg/l NO}_3$

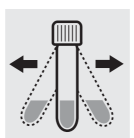
Expression of results also possible in mmol/l.



Pipette 2.0 ml of the sample into a reaction cell, **do not mix.**



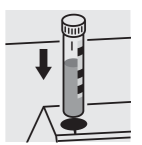
Add 1 level blue microspoon of NO₃-1K, immediately close the cell tightly with the screw cap. Caution, foams strongly (eye protection, protective gloves)!



Shake the cell **vigorously for 5 seconds** to dissolve the solid substance.



Reaction time: 30 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

in seawater

Measuring $0.2-17.0 \text{ mg/l NO}_3-\text{N}$ $0.9-75.3 \text{ mg/l NO}_3$ 10-mm cell

range: Expression of results also possible in mmol/l.



Pipette 5.0 ml of **NO₃-1** into a dry empty round cell (Empty cells, Cat. No. 250621).

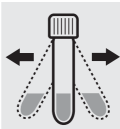


Add 1.0 ml of the sample with pipette.

Caution, cell becomes hot!



Immediately add 1.5 ml of NO₃-2 with pipette.



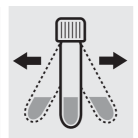
Close cell tightly and shake **vigorously**.



Reaction time: 15 minutes



Add 2 level grey microspoons of NO₃-3.



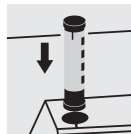
Close cell tightly and shake **vigorously** until the reagent is completely dissolved.



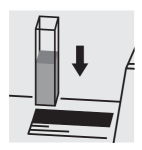
Reaction time: 60 minutes



Transfer the solution into a cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use nitrate standard solution, Cat.No. 250476, concentration 1000 mg/l NO_3^- , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.





 $\textbf{Measuring} \qquad 0,010-0,700 \text{ mg/l NO}_2\text{-N}$

range: 0,03 -2,30 mg/l NO₂

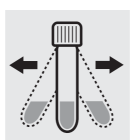
Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



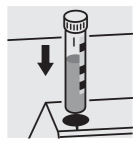
Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use nitrite standard solution, Cat.No. 250477, concentration 1000 mg/l NO_2^- , can be used after diluting accordingly.



Measuring 0.010 – 0.700 mg/l NO₂-N

range: 0.03 -2.30 mg/l NO₂

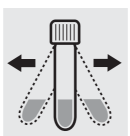
Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



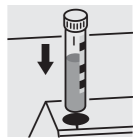
Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use nitrite standard solution, Cat.No. 250477, concentration 1000 mg/l NO_2^- , can be used after diluting accordingly.

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Measuring 1.0 - 90.0 mg/l NO₂-N

range: 3.3 – 295.2 mg/l NO₂

Expression of results also possible in mmol/l.



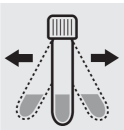
Check the pH of the sample, specified range: pH 1–12. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Add 2 level blue microspoons of **NO₂-1K** into a reaction cell.



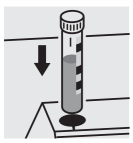
Add 8.0 ml of the sample with pipette and close with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 20 minutes, measure immediately. Do not shake or swirl the cell before the measurement.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use nitrite standard solution, Cat.No. 250477, concentration 1000 mg/l NO_2^- , can be used after diluting accordingly.



Measuring	0.02 -1.00 mg/I NO ₂ -N	0.07 -3.28 mg/I NO ₂	10-mm cell
range:	0.010-0.500 mg/I NO ₂ -N	0.03 -1.64 mg/I NO ₂	20-mm cell
	0.002-0.200 mg/I NO ₂ -N	0.007-0.657 mg/I NO ₂	50-mm cell
	Expression of results also possible in mmol/l.		



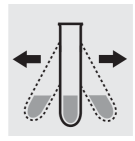
Check the pH of the sample, specified range: pH 2-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1 level blue microspoon of NO₂-1.



Shake vigorously to dissolve the solid substance.



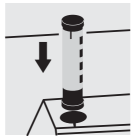
Check the pH, specified range: pH 2.0-2.5. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



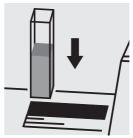
Reaction time: 10 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use nitrite standard solution, Cat.No. 250477, concentration 1000 mg/l NO₂, can be used after diluting accordingly.





Measuring 0.5-15.0 mg/l N

range: Expression of results also possible in mmol/l.



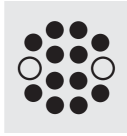
Pipette 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1 level blue microspoon of **N-1K**.



Add 6 drops of **N-2K**, close the cell with the screw cap, and mix.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 1 hour.



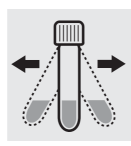
Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature: **pretreated sample.**



Swirl the cell after 10 minutes.



Add 1 level yellow micro-spoon of **N-3K** into a reaction cell, close the cell with the screw cap.



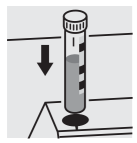
Shake the cell vigorously for 1 minute to dissolve the solid substance.



Add very slowly 1.5 ml of the **pretreated sample** with pipette, close the cell tightly with the screw cap, and mix briefly. **Caution, cell becomes hot!**



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.





Measuring 0.5-15.0 mg/l N

range: Expression of results also possible in mmol/l.



Pipette 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



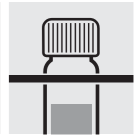
Add 1 level blue microspoon of **N-1K**.



Add 6 drops of **N-2K**, close the cell with the screw cap, and mix.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 1 hour.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature: **pretreated sample.**



Swirl the cell after 10 minutes.



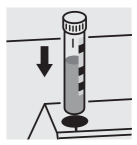
Pipette 1.0 ml of the **pretreated sample** into a reaction cell, **do not mix!**



Add 1.0 ml of **N-3K** with pipette, close the cell with the screw cap, and mix. Caution, cell be comes hot!



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 50, Cat.No. 250486.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 50) is highly recommended.





Measuring 10-150 mg/l N

range: Expression of results also possible in mmol/l.



Pipette 1.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 9.0 ml of distilled water with pipette.



Add 1 level blue microspoon of **N-1K**.



Add 6 drops of **N-2K**, close the cell with the screw cap, and mix.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 1 hour.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature: **pretreated sample**.



Swirl the cell after 10 minutes.



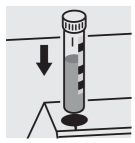
Pipette 1.0 ml of the **pretreated sample** into a reaction cell, **do not mix!**



Add 1.0 ml of **N-3K** with pipette, close the cell with the screw cap, and mix. **Caution, cell becomes hot!**



Reaction time: 10 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 70, Cat.No. 250488.

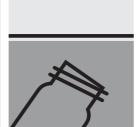
To check for sample-dependent effects the use of addition solutions (e. g. in CombiCheck 70) is highly recommended.



Measuring range: 0.5-12.0 mg/I O₂



Check the pH of the sample, specified range: pH 6-8. If required, add dilute sodium hydroxide solution or nitric acid drop by drop to adjust the pH.



Fill watersample into a reaction cell to overflowing and make sure, that no air bubbles are present.



Place the filled cell in a test-tube rack.



Add with microspoon 1 glass bead.



Add 5 drops of O₂-1K.



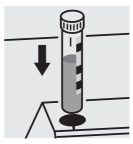
Add 5 drops of O_2 -2K, close the cell with the screw cap, and shake for 10 seconds.



Reaction time: 1 minute



Add 10 drops of O_2 -3K, close the cell with the screw cap, mix, and clean from outside.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Measuring range:	: 0.020 – 0.500 mg/l DEHA*	20-mm cell
	* N,N-diethylenhydroxylamine	
	0.027 - 0.667 mg/l Carbohy*	20-mm cell
	* carbohydrazide	
	0.053 - 1.315 mg/l Hydro*	20-mm cell
	* hydroquinone	
	0.078 - 1.950 mg/I ISA*	20-mm cell
	* isoascorbic acid	
	0.087 - 2.170 mg/l MEKO*	20-mm cell
	* methylethylketoxime	



Pipette 10 ml of the sample into into a empty round cell (Empty cells, Cat.No. 250621).



Add 1 powder pack of Oxyscav 1 and close with the screw cap.



Swirl the cell to dissolve the solid substance.



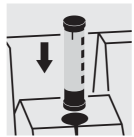
Add 0.20 ml of Oxyscav 2 with pipette, close with the screw cap, and mix.



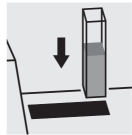
Reaction time: 10 minutes, **protect** from light in the process, measure immediately.



Transfer the solution into Select method with a cell.



AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a oxygen scavengers standard solution must be prepared (see section "Standard solutions").



Measuring	0.05 -4.00 mg/I O ₃	10-mm cell
range:	$0.02 - 2.00 \text{ mg/l O}_3$	20-mm cell
	0.010 - 0.800 mg/I O ₃	50-mm cell
	Expression of results also possible in mmol/l.	



Check the pH of the sample, specified range: pH 4-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



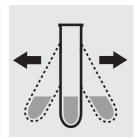
Pipette 10 ml of the sample into a test tube.



Add 2 drops of O₃-1 and mix.



Add 1 level blue microspoon of O_3 -2.



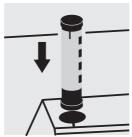
Shake vigorously to dissolve the solid substance.



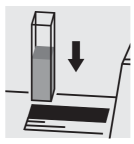
Reaction time: 3 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

Very high ozone concentrations in the sample produce yellow-coloured solutions (measurement solution should be red) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").

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Measuring range: pH 6.4-8.8



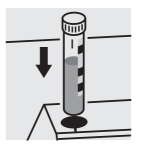
Pipette 10 ml of the sample into a round cell.



Add 4 drops of pH-1, close the cell with the screw cap, and mix.

Attention!

The reagent bottle must be held vertically by all means!



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (technical test reagents, measurement device, and handling) buffer solution pH 7.00, e.g. Cat.No. 108708, can be used.

Measuring $0.10 - 2.50 \text{ mg/I C}_6\text{H}_5\text{OH}$

range: Expression of results also possible in mmol/l.



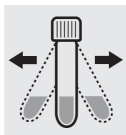
Check the pH of the sample, specified range: pH 2-11. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust



Pipette 10 ml of the sample into a reaction cell, close with the screw the cell with the screw cap, and mix.



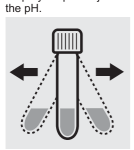
Add 1 level grey microspoon of Ph-1K, close



Shake the cell vigorously to dissolve the solid substance.



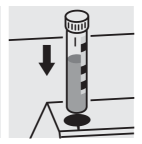
Add 1 level green microspoon of Ph-2K, close the cell with the screw



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

Very high phenol concentrations in the sample result in a weakening of the colour and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a phenol standard solution must be prepared from Phenol GR (see section "Standard solutions").



 $0.002 - 0.100 \text{ mg/I C}_6\text{H}_5\text{OH}$ Measuring 20-mm cell range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 2-11. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 200 ml of sample Add 5.0 ml of Ph-1 with into a separation funnel.



pipette and mix.



Add 1 level green microspoon of Ph-2 and shake to dissolve the solid substance.



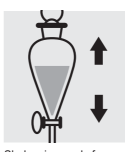
Add 1 level green microspoon of Ph-3 and shake to dissolve the solid substance.



Reaction time: 30 minutes (protected from light)



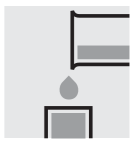
Add 10 ml of chloroform with pipette, close separation funnel.



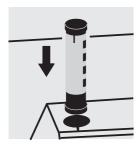
Shake vigorously for 1 minute.



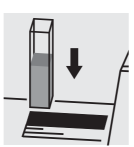
Leave to stand for 5-10 minutes to allow the phases to separate.



Transfer the clear lower phase into a cell.



Select method with AutoSelector measuring range 0.002-0.100 mg/l.



Place the cell into the cell compartment.

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Measuring	0.10 -5.00 mg/l C ₆ H ₅ OH	10-mm cell
range:	0.05 -2.50 mg/l C ₆ H ₅ OH	20-mm cell
	0.025 – 1.000 mg/l C ₆ H ₅ OH	50-mm cell
	Expression of results also possi	ble in mmol/l.



Check the pH of the sample, specified range: pH 2–11.
If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



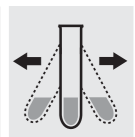
Pipette 10 ml of the sample into a test tube.



Add 1.0 ml of **Ph-1** with pipette and mix.



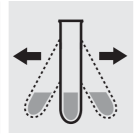
Add 1 level grey microspoon of **Ph-2**.



Shake vigorously to dissolve the solid substance.



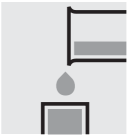
Add 1 level grey microspoon of **Ph-3**.



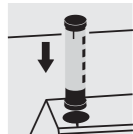
Shake vigorously to dissolve the solid substance.



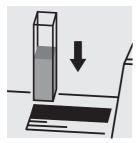
Reaction time: 10 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector measuring range 0.025-5.00 mg/l.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a phenole standard solution must be prepared from Phenol GR (see section "Standard solutions").

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P6/25 · Phosphate

a xylem brand

Determination of orthophosphate

0.05- 5.00 mg/I PO₄-P Measuring range: 0.2 -15.3 mg/I PO₄ 0.11 - 11.46 mg/I P₂O₅ Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



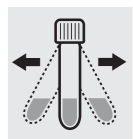
Pipette 5.0 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 5 drops of P-2K, close the cell with the



Add 1 dose of P-3K using the blue dosemetering cap, close the cell with the screw cap.



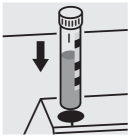
Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes

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Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.



P6/25 · Phosphate

a xylem brand

Determination of total phosphorus = sum of orthophosphate, polyphosphate, and organophosphate

Measuring 0.05 – 5.00 mg/l P range: 0.2 –15.3 mg/l PO₄ 0.11 –11.46 mg/l P₂O₅ Expression of results also possible in mmol/l and also in P total (Σ P), and P org* [P(o)].



Check the pH of the sample, specified range: pH 0-10.
If required, add dilute sulfuric acid drop by

drop to adjust the pH.



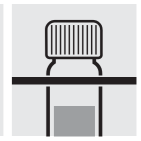
Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 dose of **P-1K** using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



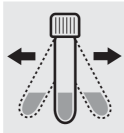
Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Add 5 drops of **P-2K**, close the cell with the screw cap, and mix.



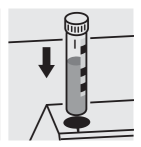
Add 1 dose of **P-3K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between orthophosphate (PO_4-P) and P org* (P(o)) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the P total, press enter and measure the orthophosphate (see analytical procedure for orthophosphate). After pressing enter, the individual measuring values for PO_4 -P and P(o) are shown on the display.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO_4^{3-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

^{*}Porg is the sum of polyphosphate and organophosphate.

P7/25 · Phosphate

a xylem brand

Determination of orthophosphate

0.5-25.0 mg/I PO₄-P Measuring range: 1.5-76.7 mg/I PO₄ 1.1-57.3 mg/I P₂O₅



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Expression of results also possible in mmol/l.

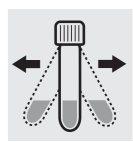
Pipette 1.0 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 5 drops of P-2K, close the cell with the



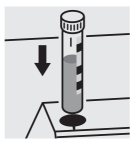
Add 1 dose of P-3K using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20 and 80, Cat. Nos. 250483 and 250489.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.

03/2021





Determination of total phosphorus = sum of orthophosphate, polyphosphate, and organophosphate

 $\begin{array}{ll} \textbf{Measuring} & 0.5-25.0 \text{ mg/I P} \\ \textbf{range:} & 1.5-76.7 \text{ mg/I PO}_4 \\ & 1.1-57.3 \text{ mg/I P}_2\text{O}_5 \\ & \text{Expression of results also possible in mmol/I and also in} \\ & \text{P total } (\Sigma \text{ P}), \text{ and P org* } [\text{P(o)}]. \end{array}$



Check the pH of the sample, specified range: pH 0-10.
If required, add dilute sulfuric acid drop by

drop to adjust the pH.



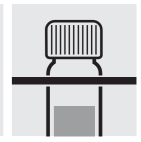
Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 dose of **P-1K** using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



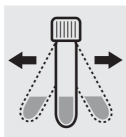
Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Add 5 drops of **P-2K**, close the cell with the screw cap, and mix.



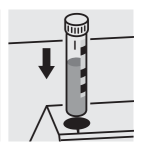
Add 1 dose of **P-3K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between orthophosphate (PO_4-P) and P org* (P(o)) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the P total, press enter and measure the orthophosphate (see analytical procedure for orthophosphate). After pressing enter, the individual measuring values for PO_4 -P and P(o) are shown on the display.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20 and 80, Cat. Nos. 250483 and 250489.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO_4^{3-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.

^{*}Porg is the sum of polyphosphate and organophosphate.



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Determination of orthophosphate

0.05- 5.00 mg/I PO₄-P Measuring range: 0.2 -15.3 mg/I PO₄ 0.11 - 11.46 mg/I P₂O₅ Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



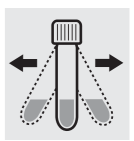
Pipette 5.0 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 5 drops of P-2K, close the cell with the



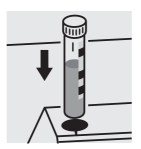
Add 1 dose of P-3K using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.



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Determination of total phosphorus

= sum of orthophosphate, polyphosphate and organophosphate

 $\begin{array}{lll} \textbf{Measuring} & 0.05 - \; 5.00 \; \text{mg/l P} \\ \textbf{range:} & 0.2 \; -15.3 \; \; \text{mg/l PO}_4 \\ & 0.11 - 11.46 \; \text{mg/l P}_2\text{O}_5 \\ & \text{Expression of results also possible in mmol/l and also in} \\ & \text{P total } (\Sigma \; \text{P}), \; \text{and P org*} \; [\text{P(o)}]. \end{array}$



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 dose of **P-1K** using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Add 5 drops of **P-2K**, close the cell with the screw cap, and mix.



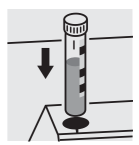
Add 1 dose of **P-3K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between orthophosphate (PO_4 -P) and P org* (P(o)) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the P total, press enter and measure the orthophosphate (see analytical procedure for orthophosphate). After pressing enter, the individual measuring values for PO_4 -P and P(o) are shown on the display.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO_4^{3-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

^{*}Porg is the sum of polyphosphate and organophosphate.



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Determination of orthophosphate

0.5-25.0 mg/I PO₄-P Measuring range: 1.5-76.7 mg/I PO₄ 1.1-57.3 mg/I P₂O₅ Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



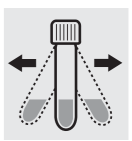
Pipette 1.0 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 5 drops of P-2K, close the cell with the



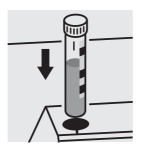
Add 1 dose of P-3K using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20 and 80, Cat.Nos. 250483 and 250489.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.



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Determination of total phosphorus

= sum of orthophosphate, polyphosphate and organophosphate

 $\begin{array}{lll} \textbf{Measuring} & 0.5-25.0 \text{ mg/l P} \\ \textbf{range:} & 1.5-76.7 \text{ mg/l PO}_4 \\ & 1.1-57.3 \text{ mg/l P}_2\text{O}_5 \\ & \text{Expression of results also possible in mmol/l and also in} \\ & \text{P total } (\Sigma \text{ P}), \text{ and P org* } [\text{P(o)}]. \end{array}$



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



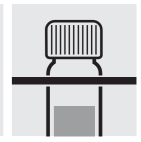
Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1 dose of **P-1K** using the green dosemetering cap, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100 °C) for 30 minutes.



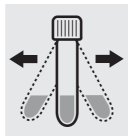
Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Add 5 drops of **P-2K**, close the cell with the screw cap, and mix.



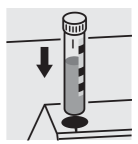
Add 1 dose of **P-3K** using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

A differentiation between orthophosphate (PO_4 -P) and P org* (P(o)) can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form. Then measure the P total, press enter and measure the orthophosphate (see analytical procedure for orthophosphate). After pressing enter, the individual measuring values for PO_4 -P and P(o) are shown on the display.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20 and 80, Cat.Nos. 250483 and 250489.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO_4^{3-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck) is highly recommended.

^{*}Porg is the sum of polyphosphate and organophosphate.



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Determination of orthophosphate

Measuring	3.0-100.0 mg/I PO ₄ -P
range:	9 −307 mg/l PO ₄
	7 –229 mg/l P ₂ O ₅
	Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



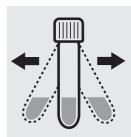
Pipette 0.20 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 5 drops of PO₄-1K, close the cell with the



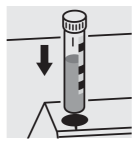
Add 1 dose of PO₄-2K using the blue dosemetering cap, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can be used after diluting accordingly.



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Determination of orthophosphate

Measuring	0.05 -5.00 mg/I PO ₄ -P	0.2 -15.3 mg/I PO ₄	0.11 – 11.46 mg/I P ₂ O ₅	10-mm cell
range:	0.03 -2.50 mg/I PO ₄ -P	0.09- 7.67 mg/l PO ₄	$0.07 - 5.73 \text{ mg/l P}_2\text{O}_5$	20-mm cell
	0.010-1.000 mg/I PO ₄ -P	0.03- 3.07 mg/I PO ₄	$0.02 - 2.29 \text{ mg/l P}_2\text{O}_5$	50-mm cell
	Expression of results also possible in mmol/l.			



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



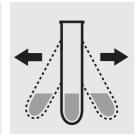
Pipette 5.0 ml of the sample into a test tube.



Add 5 drops of PO₄-1 and mix.



Add 1 level blue microspoon of PO₄-2.



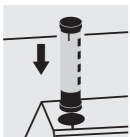
Shake vigorously to dissolve the solid substance.



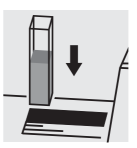
Reaction time: 5 minutes



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

For measurement in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each.

Alternatively, the semi-microcell can be used.

For the determination of total phosphorus = sum of orthophosphate, polyphosphate, and organophos**phate** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of phosphorus (ΣP).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

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Determination of orthophosphate

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 8.0 ml of distilled water into a test tube.



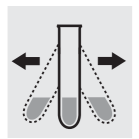
Add 0.50 ml of the sample with pipette and mix.



Add 0.50 ml of **PO₄-1** with pipette and mix.



Add 1 dose of PO_4 -2 using the blue dosemetering cap.



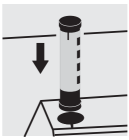
Shake vigorously to dissolve the solid substance.



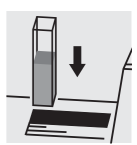
Reaction time: 5 minutes



Transfer the solution into a cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₃³⁻, can be used after diluting accordingly.



a xylem brand

Determination of orthophosphate

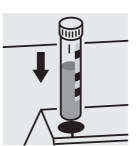
 $\begin{tabular}{lll} \textbf{Measuring} & 0.5-25.0 \text{ mg/I PO}_4\text{-P} \\ \textbf{range:} & 1.5-76.7 \text{ mg/I PO}_4 \\ & 1.1-57.3 \text{ mg/I P}_2\text{O}_5 \\ & \text{Expression of results also possible in mmol/I.} \\ \end{tabular}$



Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total phosphorus = sum of orthophosphate**, **polyphosphate**, **and organophosphate** use Phosphate Cell Test, Cat.Nos. 250324 and 252076, or Phosphate Test, Cat.No. 250446, with the Crack Set 10 or 10C, Cat.Nos. 250496 or 252033.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₃⁴⁻, can be used after diluting accordingly.



a xylem brand

Determination of orthophosphate

Measuring 1.0-30.0 mg/I PO₄-P 3.1-92.0 mg/I PO₄ 2.3-68.7 mg/l P₂O₅ 10-mm cell range: 0.5-15.0 mg/I PO₄-P 1.5-46.0 mg/I PO₄ 1.1-34.4 mg/I P₂O₅ 20-mm cell Expression of results also possible in mmol/l.



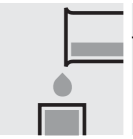
Check the pH of the sample, specified range: pH 0-10. If required, add dilute sulfuric acid drop by drop to adjust the pH.



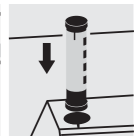
Pipette 5.0 ml of the sample into a test tube.



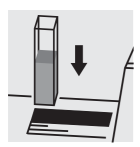
Add 1.2 ml of PO₄-1 with Transfer the solution into Select method with piette and mix.



a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

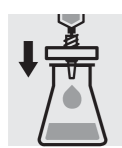
For the determination of total phosphorus = sum of orthophosphate, polyphosphate, and organophos**phate** use Phosphate Cell Test, Cat.Nos. 250324 and 252076, or Phosphate Test, Cat.No. 250446, with the Crack Set 10 or 10C, Cat.Nos. 250496 or 252033.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use phosphate standard solution, Cat.No. 250478, concentration 1000 mg/l PO₄³⁻, can be used after diluting accordingly.

Measuring 5.0-50.0 mg/l K

range: Expression of results also possible in mmol/l.



Filter turbid samples.



Check the pH of the sample, specified range: pH 3–12. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 2.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



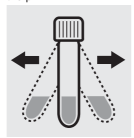
Check the pH, specified range: pH 10.0-11.5.



Add 6 drops of **K-1K**, close the cell with the screw cap, and mix.



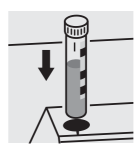
Add 1 level blue microspoon of **K-2K**, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

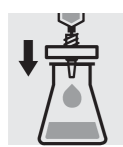
Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use potassium standard solution, Cat.No. 252471, concentration 1000 mg/l K, can be used after diluting accordingly.



Measuring 30-300 mg/l K

range: Expression of results also possible in mmol/l.



Filter turbid samples.



Check the pH of the sample, specified range: pH 3–12. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust



Pipette 0.50 ml of the sample into a reaction cell, close with the screw cap, and mix.



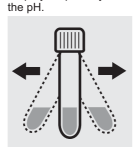
Check the pH, specified range: pH 10.0-11.5.



Add 6 drops of **K-1K**, close the cell with the screw cap, and mix.



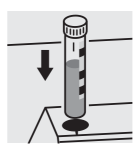
Add 1 level blue microspoon of **K-2K**, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use potassium standard solution, Cat.No. 252471, concentration 1000 mg/l K, can be used after diluting accordingly.



14683 · Residual Hardness

a xylem brand

Measuring	0.50 -5.00 mg/l Ca
range:	0.070-0.700 °d
	0.087-0.874 °e
	0.12 -1.25 °f

0.70- 7.00 mg/I CaO Measuring 1.2 -12.5 mg/l CaCO₃ range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 5-8. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



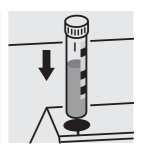
Pipette 4.0 ml of the sample into a reaction cell, close with the screw screw cap, and mix. cap, and mix.



Add 0.20 ml of RH-1K, close the cell with the



Reaction time: 10 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use calcium standard solution, Cat.No. 250465, concentration 1000 mg/l Ca, can be used after diluting accordingly. (Pay attention to pH value!)

Measuring	0.21 -10.70 mg/I SiO ₂	0.10 - 5.00 mg/l Si	10-mm cell
	0.10 - 5.35 mg/l SiO ₂	0.05 - 2.50 mg/l Si	20-mm cell
	0.011-1.600 mg/I SiO ₂	0.005-0.750 mg/l Si	50-mm cell
	Expression of results also possible in mmol/l.		



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust



Pipette 5.0 ml of the sample into a test tube.



Add 3 drops of Si-1 and mix.



Check the pH, specified range: pH 1.2-1.6.



Reaction time: 3 minutes



Add 3 drops of Si-2 and mix.



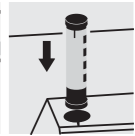
Add 0.50 ml of Si-3 with pipette and mix.



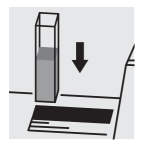
Reaction time: 10 minutes



a corresponding cell.



Transfer the solution into Select method with AutoSelector. (Method 079 for 10 mmand 20 mm-cells, and method 081 for the 50 mm-cell.)



Place the cell into the cell compartment.

Important:

The test kit contains two AutoSelectors that are to be chosen according to the measuring range or rectangular cell used (see label).

To measure in the 50 mm-cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use silicon standard solution, Cat.No. 252472, concentration 1000 mg/l Si, can be used after diluting accordingly. (Attention! Do not store standard solutions in glass vessels!)



Measuring	1.1 - 107.0 mg/I SiO ₂	0.5- 50.0 mg/l Si	10-mm cell
range:	11 -1070 mg/l SiO ₂	5 -500 mg/l Si	10-mm cell
	Expression of results also p	oossible in mmol/l.	

Measuring range: 1,1–107,0 mg/l SiO₂



Check the pH of the sample, specified range: pH 2–10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 4.0 ml of the sample into a test tube.



Add 4 drops of **Si-1** and mix.



Add 2.0 ml of **Si-2** with pipette and mix.



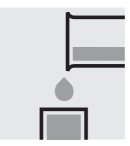
Reaction time: 2 minutes



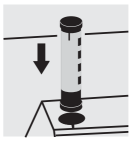
Add 4 drops of **Si-3** and mix.



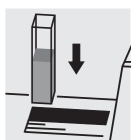
Reaction time: 2 minutes



Transfer the solution into Select method with a cell. Selector measurement and selector measurements.



Select method with AutoSelector measuring range 0.5 – 50.0 mg/l Si.



Place the cell into the cell compartment.

Measuring range: 11–1070 mg/l SiO₂



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust



Pipette 5.0 ml of distilled water into a test tube.



Add 0.50 ml of the sample with pipette and mix.

Continue as mentioned above; starting from the addition of **Si-1** (Fig. 3). Select method with AutoSelector measuring range 5–500 mg/l Si.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use silicon standard solution, Cat.No. 252472, concentration 1000 mg/l Si, can be used after diluting accordingly. (Attention! Do **not** store standard solutions in glass vessels!)

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the pH.



Measuring 0.50-3.00 mg/l Ag 10-mm cell range: $0.25 - 1.50 \, \text{mg/l Ag}$ 20-mm cell Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



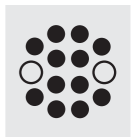
Pipette 10 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 2 drops of Ag-1.



Add 1 level green microspoon of Ag-2, close the cell with the screw cap.



Heat the cell in the thermoreactor at 120 °C (100°C) for 1 hours.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



Swirl the cell before opening.



Add 3 drops of Aq-3, close with the screw cap, and mix.



Check the pH, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Add 1 drop of Aq-4, close with the screw cap, and mix.



Add 5 drops of Ag-5, close with the screw cap, and mix.



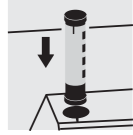
Add 1.0 ml of Ag-6, close with the screw cap, and mix.



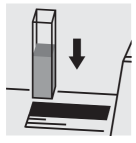
Reaction time: 5 minutes



Transfer the solution into a corresponding cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Important:

Very high silver concentrations in the sample produce turbid solutions (measurement solution should be clear). In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) readyfor-use silver standard solution, Cat.No. 250479, concentration 1000 mg/l Ag, can be used after diluting accordingly.

00885 · Sodium

a **xylem** brand

in nutrient solutions

Measuring range: 10-300 mg/l Na



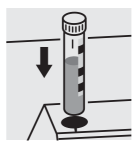
Pipette 0.50 ml of Na-1K Add 0.50 ml of the into a reaction cell and mix.



sample with pipette, close the cell with the screw cap, and mix.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

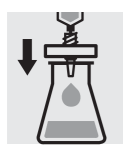
Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use chloride standard solution, Cat.No. 250466, concentration 1000 mg/l Cl⁻ (corresponds to 649 mg/l Na), can be used after diluting accordingly (see section "Standard solutions").



Measuring 5-250 mg/I SO₄

range: Expression of results also possible in mmol/l.



Filter turbid samples.



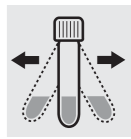
Check the pH of the sample, specified range: pH 2–10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



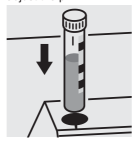
Add 1 level green microspoon of **SO₄-1K**, close the cell with the screw



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 2 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

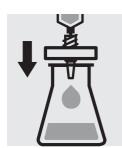
Ready-for-use sulfate standard solution, Cat.No. 250480, concentration 1000 mg/l SO_4^{2-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

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Measuring 50-500 mg/I SO₄

range: Expression of results also possible in mmol/l.



Filter turbid samples.



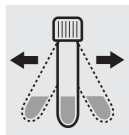
Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 2.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



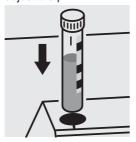
Add 1 level green microspoon of **SO₄-1K**, close the cell with the screw



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 2 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

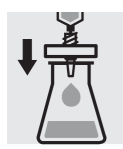
Ready-for-use sulfate standard solution, Cat.No. 250480, concentration 1000 mg/l SO_4^{2-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.



Measuring 100-1000 mg/I SO₄

range: Expression of results also possible in mmol/l.



Filter turbid samples.



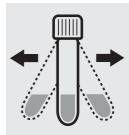
Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



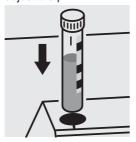
Add 1 level green microspoon of **SO₄-1K**, close the cell with the screw



Shake the cell vigorously to dissolve the solid substance.



Reaction time: 2 minutes, measure immediately.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use sulfate standard solution, Cat.No. 250480, concentration 1000 mg/l SO_4^{2-} , can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.



Measuring 25-300 mg/I SO₄ 10-mm cell

Expression of results also possible in mmol/l. range:



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to



Pipette 2.5 ml of the sample into a test tube with screw cap.



Add 2 drops of SO₄-1 and mix.



Add 1 level green microspoon of SO₄-2, close the test tube with the screw cap, and mix.



Temper the test tube in a water bath at 40 °C for 5 minutes.



pipette and mix.



Add 2.5 ml of SO₄-3 with Filter the content of the test tube with a round filter into another test tube with screw cap.



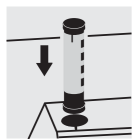
Add 4 drops of SO₄-4 to the filtrate, close the test tube with the screw cap, and mix.



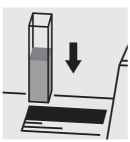
Temper the test tube again in the water bath for 7 minutes.



Transfer the solution into a cell.



Select method with AutoSelector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 10, Cat.No. 250482.

Ready-for-use sulfate standard solution, Cat.No. 250480, concentration 1000 mg/l SO₄²⁻, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 10) is highly recommended.

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02537 · Sulfate

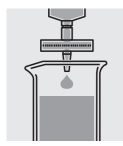
a xylem brand

Category: RT (reagent test)

Cell: 16 mm

Measuring range: 5 - 300 mg/l SO₄

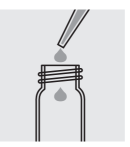
Display in mmol/l possible



Filter turbid sample solutions.



Check the pH value of the sample. Desired range: pH 2-10. Correct with diluted sodium hydroxide solution or hydrochloric acid as necessary.



Pipette 0.50 ml of **SO₄-1** into the empty cell.



Add 5.0 ml sample with a pipette and mix.



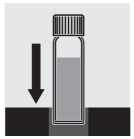
Add 1 level blue microspoon of SO_4 -2 and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



Allow to react for 2 minutes. **Then measure immediately.**



Insert the cell in the photometer cell shaft and start measurement.

Notes:

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- For further notes please refer to the package insert of the test.



Measuring	0.10 -1.50 mg/IS	0.10 -1.55 mg/I HS	10-mm cell
range:	0.050 - 0.750 mg/I S	0.052-0.774 mg/I HS	20-mm cell
	0.020 - 0.500 mg/I S	0.021-0.516 mg/I HS	50-mm cell
	Expression of results also possible in mmol/l.		



Check the pH of the sample, specified range: pH 2-10. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1 drop of S-1 and mix.



Add 5 drops of S-2 and mix.



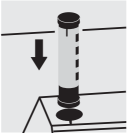
Add 5 drops of S-3 and mix.



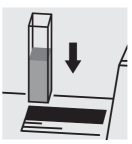
Reaction time: 1 minute.



Transfer the solution into Select method with a corresponding cell.



AutoSelector.



Place the cell into the cell compartment.

Important:

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, the semi-microcell can be used.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a sulfide standard solution must be prepared from sodium sulfide GR (see section "Standard solutions").

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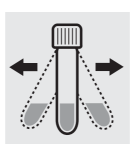
Measuring	1.0 -20.0 mg/I SO ₃	Round cell
range:	0.8 - 16.0 mg/I SO ₂	Round cell
	0.05-3.00 mg/I SO ₃	50-mm cell (see "sensitive" preparation procedure)
	0.04-2.40 mg/I SO ₃	50-mm cell (see "sensitive" preparation procedure)
	Expression of results also possible in mmol/l.	



Check the pH of the sample, specified range: pH 4-9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Add 1 level grey microspoon of **SO₃-1K** into a reaction cell, close with the screw cap.



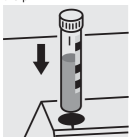
Shake the cell vigorously to dissolve the solid substance.



Add 3.0 ml of the sample with pipette, close the cell with the screw cap, and mix.



Reaction time: 2 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Sulfite sensitive

Use the same preparation procedure as above, but add 7.0 ml of the sample instead of 3.0 ml. Prepare an own blank by using 7.0 ml of distilled water and all reagents. For measurement transfer the solution into a 50-mm cell. Configure the photometer prior for blank-measurement. Select method **SO**₃ **sens** in the menu (method no. 127).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a sulfite standard solution must be prepared from sodium sulfite GR (see section "Standard solutions").

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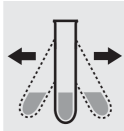
Check the pH of the sample, specified range: pH 4-9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



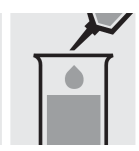
Place 1 level grey microspoon of **SO**₃-1 into a dry test tube.



Add 3.0 ml of **SO₃-2** with pipette.



Shake vigorously to dissolve the solid substance.



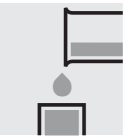
Add 5.0 ml of distilled water with pipette and mix.



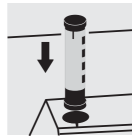
Add 2.0 ml of the sample with pipette and mix.



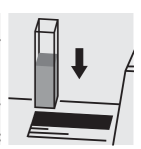
Reaction time: 2 minutes



Transfer the solution into a cell.



Select method with Auto-Selector.



Place the cell into the cell compartment.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a sulfite standard solution must be prepared from sodium sulfite GR (see section "Standard solutions").



Measuring range: 0,05 - 2,00 mg/l MBAS*

* Methylene-blue-active substances

Ergebnisangabe auch in mmol/l möglich



Check the pH value of the sample. Required range: pH 5-10. Correct with diluted sodium hydroxide solution or hydrochloric acid as necessary.

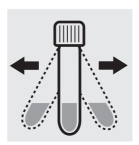


Pipette 5.0 ml of sample into a reaction cell.

Do not mix the contents!



Add 2 drops of **T-1K**, close the cell with the screw cap.



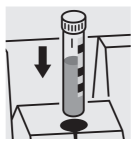
Shake the cell vigorously for 30 seconds.



Reaction time: 10 minutes



Swirl the cell before the measurement.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a surfactants standard solution must be prepared from dodecane-1-sulfonic acid sodium salt GR (see section "Standard solutions").



02552 · Surfactants (anionic)

a xylem brand

Measuring 0.05-2.00 mg/I MBAS*

range: * Methylene-blue-active substances

Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 5-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, **do not mix!**



Add 3 drops of **T-1K**, **do not mix!**



Add 2 drops of **T-2K**, close the cell with the screw cap.



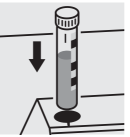
Shake the cell for 30 seconds.



Reaction time: 10 minutes



Swirl the cell before the measurement.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a surfactants standard solution must be prepared from dodecane-1-sulfonic acid sodium salt GR (see section "Standard solutions").



01764 · Surfactants (cationic)

a xylem brand

0.05-1.50 mg/l k-Ten Measuring

range: (calculated as

N-cetyl-N,N,N-trimethylammonium bromide)



Check the pH of the sample, specified range: pH 3-8. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a reaction cell, do not mix!



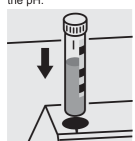
Add 0.50 ml of T-1K with Swirl the cell for pipette and close with the screw cap.



30 seconds.



Reaction time: 5 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a surfactants standard solution must be prepared from N-cetyl-N,N,N-trimethylammonium bromide (see section "Standard solutions").



01787 · Surfactants (nonionic)

a xylem brand

Measuring 0.010 – 7.50 mg/l n-Ten

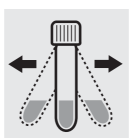
range: (calculated as Triton® X-100)



Check the pH of the sample, specified range: pH 3-9. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 4.0 ml of the sample into a reaction cell. Close with the screw cap.



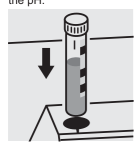
Shake the cell for 1 minute vigorously.



Reaction time: 2 minutes



Swirl the cell before measurement.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a surfactants standard solution must be prepared from Triton® X-100 (see section "Standard solutions").



0.10-2.50 mg/l Sn Measuring

Expression of results also possible in mmol/l. range:



Check the pH of the sample, specified range: pH < 3.

If required, add dilute sulfuric acid drop by drop to adjust the pH.



Add 6 drops of Sn-1K into a reaction cell, close sample with pipette, with the screw cap, and



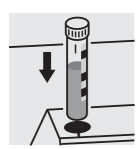
Add 5.0 ml of the close the cell with the screw cap, and mix.



Check the pH, specified range: pH 1.5-3.5. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use tin standard solution, Cat.No. 250501, concentration 1000 mg/l Sn, can be used after diluting accordingly in diluted hydrochloric acid.



Total Organic Carbon

Measuring range: 5.0-80.0 mg/I TOC

Removal of inorganic bound carbon (TIC):



Check the pH of the sample, specified range: pH 2-12. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Place 25 ml of the sample into a suitable glass vessel.



Add 3 drops of TOC-1K and mix.



Check the pH, specified range pH < 2.5.



Stir for 10 minutes.

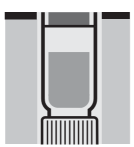
Preparation of measurement sample:



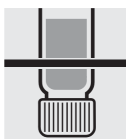
Pipette 3.0 ml of stirred sample into a reaction cell.



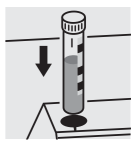
Add 1 level grey microspoon of TOC-2K. **Immediately** close the cell tightly with an aluminium cap (Cat.No. 73500).



Heat the cell, standing on its head, at 120 °C in the thermoreactor for 2 hours.



Remove the cell from the thermoreactor and let it, standing on its head, to cool for 1 hour.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a TOC standard solution, Cat.No. 250499, concentration 1000 mg/l TOC, can be used after diluting accordingly.



Total Organic Carbon

Measuring range: 50-800 mg/I TOC

Removal of inorganic bound carbon (TIC):



Check the pH of the sample, specified range: pH 2-12. If required, add dilute sulfuric acid drop by drop to adjust the pH.



Pipette 1.0 ml of the sample and 9.0 ml of distilled water into a suitable glass vessel.



Add 2 drops of TOC-1K and mix.



Check the pH, specified range pH < 2.5



Stir for 10 minutes.

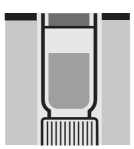
Preparation of measurement sample:



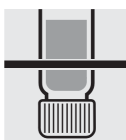
Pipette 3.0 ml of stirred sample into a reaction cell.



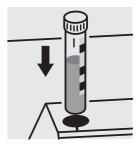
Add 1 level grey microspoon of TOC-2K. **Immediately** close the cell tightly with an aluminium cap (Cat.No. 252038).



Heat the cell, standing on its head, at 120 °C in the thermoreactor for 2 hours.



Remove the cell from the thermoreactor and let it, standing on its head, to cool for 1 hour.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a TOC standard solution, Cat.No. 250499, concentration 1000 mg/l TOC, can be used after diluting accordingly.



00961 · Total Hardness

a **xylem** brand

Determination of total hardness

Measuring	5 –215 mg/l Ca
range:	0.7- 30.1 °d
	0.9- 37.6 °e
	1.2- 53.7 °f

Measuring 7-301 mg/l CaO
range: 12-537 mg/l CaCO₃

Expression of results also possible in mmol/l
and also in mg/l Mg.



Check the pH of the sample, specified range: pH 3-9. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



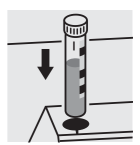
Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1.0 ml of **H-1K** with pipette, close the cell with the screw cap, and mix



Reaction time: 3 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a freshly prepared standard solution can be used (see section "Standard solutions").

00961 · Total Hardness

a xylem brand

Differentiation between Ca- and Mg-hardness

Measuring	0.12 - 5.36 mmol/l
range:	0.7 -30.1 °d
	0.9 -37.6 °e
	1.2 -53.7 °f

Differentiation possible only in mmol/l.

A differentiation between calcium- and magnesium-hardness can be performed on the photometer. Prior to measuring, select the differentiation measurement and choose the corresponding citation form.



Check the pH of the sample, specified range: pH 3-9. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



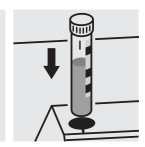
Pipette 1.0 ml of the sample into a reaction cell, close with the screw cap, and mix.



Add 1.0 ml of **H-1K** with pipette, close the cell with the screw cap, and



Reaction time: 3 minutes

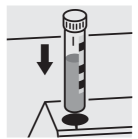


Place the cell into the cell compartment. Align the mark on the cell with that on the photometer = Result total hardness

Press enter, remove the cell.



Add 3 drops of **H-2K** to the already measured cell, close the cell with the screw cap, and mix.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer = Result magnesium

After pressing enter, the individual measuring values for Ca- and Mg-hardness are shown on the display.



01749 · Volatile Organic Acids

a xylem brand

Measuring50 – 3000 mg/l volatile organic acid(calculated as acetic acid)range:71 – 4401 mg/l volatile organic acid(calculated as butyric acid)



Check the pH of the sample, specified range: pH 2– 12.



Pipette 0.50 ml of **OA-1** into a round cell.



Add 0.50 ml of the sample with pipette, close with the screw cap, and



Heat the cell in the thermoreactor at 100 °C for 15 minutes. Then cool to room temperature under running water.



Add 1.0 ml of **OA-2** with pipette.



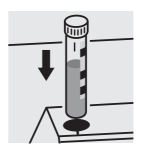
Add 1.0 ml of **OA-3** with pipette, close the cell with the screw cap, and mix.



Add 1.0 ml of **OA-4** with pipette, close the cell with the screw cap, and shake vigorously.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a standard solution must be prepared from sodium acetate anhydrous (see section "Standard solutions").



01809 · Volatile Organic Acids

a xylem brand

Measuring 50 – 3000 mg/l volatile organic acid (calculated as acetic acid)

range: 71 – 4401 mg/l volatile organic acid (calculated as butyric acid)



Check the pH of the sample, specified range: pH 2– 12.



Pipette 0.75 ml of **OA-1** into a round cell.



Add 0.50 ml of **OA-2** with pipette.



Add 0.50 ml of the sample with pipette, close with the screw cap, and mix.



Heat the cell in the thermoreactor at 100 °C for 15 minutes. Then cool to room temperature under running water.



Add 1.0 ml of **OA-3** with pipette.



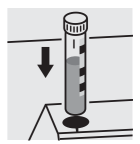
Add 1.0 ml of **OA-4** with pipette, close the cell with the screw cap, and mix.



Add 1.0 ml of **OA-5** with pipette, close the cell with the screw cap, and shake vigorously.



Reaction time: 1 minute



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a standard solution must be prepared from sodium acetate anhydrous (see section "Standard solutions").



Measuring 0.025 – 1.000 mg/l Zn

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 1–7. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 10 ml of sample into a glass vessel.



Add 1 level green microspoon of **Zn-1K** and shake to dissolve the solid substance: **sample-reagent mixture.**



Pipette 0.50 ml of **Zn-2K** into a reaction cell, close with the screw cap, and mix



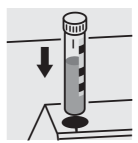
Add 2.0 ml of the sample-reagent mixture with pipette, close the cell with the screw cap, and mix.



Add 5 drops of **Zn-3K**, close the cell with the screw cap, and mix.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total zinc** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of zinc (Σ Zn).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use zinc standard solution, Cat.No. 250481, concentration 1000 mg/l Zn, can be used after diluting accordingly.



Measuring 0.20-5.00 mg/l Zn

range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 3-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Add 5 drops of **Zn-1K** into a reaction cell, close with the screw cap, and mix.



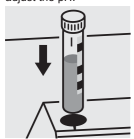
Add 0.50 ml of the sample with pipette, close the cell with the screw cap, and mix.



Add 5 drops of **Zn-2K**, close the cell with the screw cap, and mix.



Reaction time: 15 minutes



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Important:

For the determination of **total zinc** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of zinc (Σ Zn).

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Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 40, Cat.No. 250485.

Ready-for-use zinc standard solution, Cat.No. 250481, concentration 1000 mg/l Zn, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 40) is highly recommended.



Measuring 0.05–2.50 mg/l Zn 10-mm cell range: Expression of results also possible in mmol/l.



Check the pH of the sample, specified range: pH 4-10. If required, add dilute sodium hydroxide solution or hydrochloric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube with screw cap.



Add 5 drops of **Zn-1**, close the test tube with the screw cap, and mix.



Check the pH, specified range: pH 12–13. If required, add dilute sodium hydroxide solution drop by drop to adjust the pH.



Add 2 drops of **Zn-2**, close the test tube with the screw cap, and mix.



Add 5 drops of **Zn-3**, close the test tube with the screw cap, and mix.



Add 3 drops of **Zn-4**, close the test tube with the screw cap, and mix.



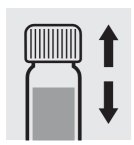
Reaction time: 3 minutes



Add 1 level grey microspoon of **Zn-5**, close the test tube with the screw cap, and dissolve the solid substance.



Add 5.0 ml of **Zn-6** (Cat. No. 06146, Isobutylmethy ketone) with pipette and close the test tube with the screw cap.



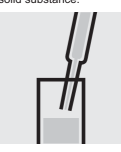
Shake the tube vigorously for 30 seconds.



Leave to stand for 2 minutes.



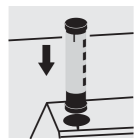
Aspirate the clear upper phase from the tube with a cell. pipette.



Transfer the solution into a cell.

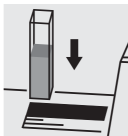


Leave to stand for 3 minutes.



Select method with AutoSelector.

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Place the cell into the cell compartment.

Important:

For the determination of **total zinc** a pretreatment with Crack Set 10C, Cat.No. 252033, or Crack Set 10, Cat.No. 250496, and thermoreactor is necessary.

Result can be expressed as sum of zinc (Σ Zn).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) ready-for-use zinc standard solution, Cat.No. 250481, concentration 1000 mg/l Zn, can be used after diluting accordingly.

Applications

Available methods

Applications are special photometric procedures normally not based on test sets. The analysis specifications for these are given in the last part of the section, ANALYTICAL PROCEDURES. There you will find further information on auxiliaries and reagents. For applications, the method is selected manually, using the method number given in column 1. Instructions on how to select a method are given in the section, SELECTING A METHOD MANUALLY of the functional description of the photometer.

Method No.	Parameter	Total measuring range	Method
2518	ADMI	2.0 – 100.0	Inherent color
2517	ADMI	10 – 1000	Inherent color
2522	Ammonia, free	(0.010 – 0.500 mg/l NH ₄ -N)	as ammonium (with test 14752)
2521	Ammonia, free	(0.03 – 1.50 mg/l NH ₄ -N)	as ammonium (with test 14752)
2520	Ammonia, free	(0.05 – 3.00 mg/l NH ₄ -N)	as ammonium (with test 14752)
2523	Ammonia, free	(0.6 – 20.6 mg/l NH ₄ -N)	as ammonium (with test 14544)
130	Antimony in water and wastewater	0.10 - 8.00 mg/l Sb	Brilliant green
195	Bromate in water and drinking water	0.003 - 0.120 mg/l BrO ₃	3,3"-Dimethylnaphtidine
2525	Carbon dioxide	(0.4 - 8.00 mg/l OH)	Indicator reaction (with test 01758)
2509	Chlorophyll-a (DIN), 10 mm	result in µg/l Chl-a	Inherent color
2510	Chlorophyll-a (DIN), 20 mm	result in µg/l Chl-a	Inherent color
2511	Chlorophyll-a (DIN), 50 mm	result in µg/l Chl-a	Inherent color
2504	Chlorophyll-a (ASTM), 10 mm	result in mg/m3 Chl-a	Inherent color
2505	Chlorophyll-a (ASTM), 20 mm	result in mg/m3 Chl-a	Inherent color
2506	Chlorophyll-a (ASTM), 50 mm	result in mg/m3 Chl-a	Inherent color
2507	Chlorophyll-a,-b,-c (ASTM), 10 mm	result in mg/m3 Chl-a,-b,-c	Inherent color
2508	Chlorophyll-a,-b,-c (ASTM), 50 mm	result in mg/m3 Chl-a, -b, -c	Inherent color
020	Chromium Baths	4.0 – 400 g/l CrO ₃	Inherent color
015	Color α(436)(Color436) (Spectral Absorption Coefficient)	0.1 – 250 m ₋₁	Measurement at 436 nm
061	Color α(525)(Color525) (Spectral Absorption Coefficient)	0.1 – 250 m ₋₁	Measurement at 525 nm
078	Color α(620)(Color620) (Spectral Absorption Coefficient)	0.1 – 250 m ₋₁	Measurement at 620 nm
303	Color (410)(CU410) (EN 7887)	2 – 2500 mg/l Pt	Measurement at 410 nm
032	Color Hazen (CU340)*	0,2 – 500 CU	Platinum-cobalt-Standard Method, Measurement at 340 nm
179	Color Hazen (CU445)*	1 – 1000 CU	Platinum-cobalt-Standard Method, Measurement at 445 nm
180	Color Hazen (CU455)*	1 – 1000 CU	Platinum-cobalt-Standard Method, Measurement at 455 nm
181	Color (CU465)*	1 – 1000 CU	Platinum-cobalt-Standard Method, Measurement at 465 nm
083	Copper Baths	2.0 – 80.0 g/l Cu	Inherent color
033	Iodine color number (IodFa)	0.010 – 3.00 IFZ	Measurement at 340 nm
021	Iodine color number (IodFa)	0.2 – 50.0 IFZ	Measurement at 445 nm
135	Mercury in water and wastewater	0.025 – 1.000 mg/l Hg	Michler's ketone
057	Nickel Baths	2.0 – 120 g/l Ni	Inherent color
2503	Nitrate	0.0 − 7.0 mg/l NO ₃ -N	direct measurement in the UV range
133	Palladium in water and wastewater	0.05 – 1.25 mg/l Pd	Thio-Michler's ketone
134	Platinium in water and wastewater	0.10 – 1.25 mg/l Pt	o-Phenylendiamine
300	Spectral Absorption Coefficient $\alpha(254)$	0.5 – 250 m ₋₁	Measurement at 254 nm
301	Spectral Attenuation Coefficient μ(254)*	0.5 – 250 m ₋₁	Measurement at 254 nm

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Method No.	Parameter	Total measuring range	Method
302	Spectral Absorption Coefficient $\alpha(436)$	0.5 – 250 m-1	Measurement at 436 nm
182	Suspended Solids	25 – 750 mg/l Susp. solids	Measurement at 820 nm
077	Turbidity (T550)	1 – 100 FAU	Measurement at 550 nm

^{*} Turbidity correction possible

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Application · **ADMI color measurement**

a xylem brand

analog. to APHA 2120F (ADMI Weighted-Ordinate Spectrophotometric method)

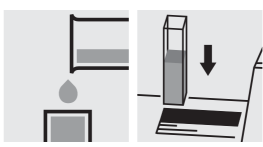
Measuring	10 – 1000	10-mm cell	Method No. 2517		
range:	2.0 - 100.0	50-mm cell	Method No. 2518		
Attention!	The measurement i	s carried out in a corr	esponding rectangular cell against a blank, prepared from distilled		
	water (Water for process analysis, Cat.No. 01051, is recommended).				

Preparation:



Filter turbid samples.

Determination at the original pH:



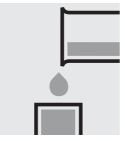
Transfer the solution into a corresponding cell.

Place the cell into the cell compartment.
Select method no. **2517** or **2518**.

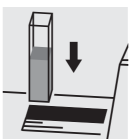
Determination at pH 7.0:



Check the pH of the sample, specified value: pH 7.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Transfer the solution into Place the cell into the a corresponding cell. Place the cell compartment.



Place the cell into the cell compartment. Select method no. **2517** or **2518**.

Note:

This method can be recalibrated by the user (one-point calibration). This method is activated by hitting the **Blank Zero** key and is subsequently menu-controlled (see the application for further details).

In the case of **serial measurements** the accuracy of the measurement can be enhanced by making a zero setting prior to **each** individual measurement.

Quality assurance:

To check the measurement system (measurement device, and handling) ready-for-use platinum-cobalt colour reference solution (Hazen 500) CertiPUR®, Cat.No. 00246 (Merck), concentration 500 mg/l Pt can be used after diluting ac-cordingly.



Application · Ammonia, free (as ammonium)

a xylem brand

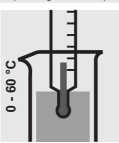
Measuring	Equivalent to 0.05 – 3.00 mg/l NH ₄ -N	Example*: 0.01 - 0.56 mg/l NH ₃	10 mm	Method No. 2520
range:	Equivalent to 0.03 – 1.50 mg/l NH ₄ -N	Example*: 0.01 - 0.28 mg/l NH ₃	20 mm	Method No. 2521
	Equivalent to 0.010 - 0.500 mg/l NH ₄ -N	Example*: 0.002 - 0.093 mg/l NH ₃	50 mm	Method No. 2522
	* Management for NIII on NIII NI dance d	line on all color and towns and the		

* Measuring ranges for NH₃ or NH₃-N depending on pH value and temperature.

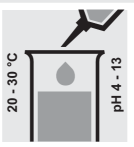
The example ranges refer to pH 8.5 and 25 °C.



Check the pH of the sample **and note**.



Check the temperature of the solution and note.



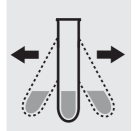
Pipette 5.0 ml of the sample into a test tube. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH and bring the sample to the appropriate temperature.



Add 0.60 ml of **NH₄-1** (from Ammonium Test, Cat. No. 250426 or 252081) with pipette and mix



Add 1 level blue microspoon of **NH₄-2** (from Ammonium Test, Cat. No. 250426 or 252081).



Shake vigorously to dissolve the solid substance.



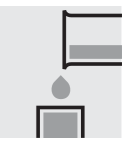
Reaction time: 5 minutes



Add 4 drops of **NH₄-3** (from Ammonium Test, Cat. No. 250426 or 252081) and mix.



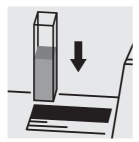
Reaction time: 5 minutes



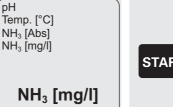
Transfer the solution into a corresponding cell.



Select method no. 2520, 2521, or 2518. Enter the pH and the temperature of the original sample.



Place the cell into the cell compartment.







NH₃-N [mg/l]

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

To measure in the 50-mm cell, the sample volume and the volume of the reagents have to be doubled for each. Alternatively, a semi-microcell can be used.



Application · Ammonia, free (as ammonium)

a xylem brand

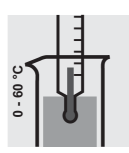
Measuring range Equivalent to 0.5 – 16.0 mg/l NH₄-N or 0.6 - 20.6 mg/l NH₄

Measuring ranges for NH₃ or NH₃-N depending on pH value and temperature.

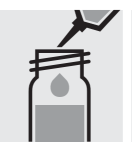
Example: $0.09 - 3.00 \text{ mg/l NH}_3$ at pH 8.5 and 25 °C.



Check the pH of the sample **and note**.



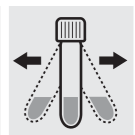
Check the temperature of the solution and note.



Pipette 0.50 ml of the sample into a reaction cell (from Ammonium Test, Cat. No. 250329) close with the screw cap, and mix.



Add 1 dose of **NH₄-1K** (from Ammonium Test, Cat. No. 250329) using the blue dose-metering cap, close the cell with the screw cap.



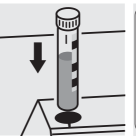
Shake the cell vigorously to dissolve the solid substance.



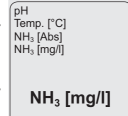
Reaction time: 15 minutes



Select method no. **2523**. Enter the pH and the temperature of the original sample.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.





pH Temp. [°C] NH₃ [Abs] NH₃ [mg/l] NH₃-N [mg/l]

 $NH_3-N [mg/l]$

Important:

Very high ammonium concentrations in the sample produce turquoise-coloured solutions (measurement solution should be yellow-green to green) and false-low readings are yielded. In such cases the sample must be diluted (plausibility check).

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) we recommended to use CombiCheck 20, Cat.No. 250483.

Ready-for-use ammonium standard solution, Cat.No. 250461, concentration 1000 mg/l NH₄, can also be used after diluting accordingly.

To check for sample-dependent effects the use of addition solutions (e.g. in CombiCheck 20) is highly recommended.



Application · Antimony in water and wastewater

Measuring range: 0.10-8.00 mg/l Sb 10-mm cell



Pipette 4.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add approx. 1.5 g of ammonium chloride hexahydrate extra pure, close the cell with the screw cap.



Shake the cell vigorously to dissolve the solid substance.



Add 1.0 ml phosphoric acid 85 % GR with pipette, close the cell with the screw cap, and mix.



Add 2 drops of reagent 1, close the cell with the screw cap, and mix.



Reaction time: 3 minutes



Add 2 drops of reagent 2, close the cell with the screw cap, and mix.



Reaction time: 2 minutes



Add 2 drops of reagent 3, close the cell with the screw cap, and mix.



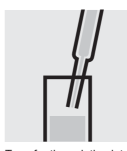
Add 5.0 ml toluene GR with pipette, close the cell with the screw cap.

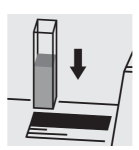


Shake the cell vigorously for 30 seconds. Leave to stand to allow phases to separate.



Aspirate the clear upper Transfer the solution into phase from the tube with a cell. pipette.





Place the cell into the cell compartment. Select method Antimony in the menu (method no. 130).

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Important:

The exact composition and preparation of the reagents 1, 2, and 3 used are given in the corresponding application, which also includes further information on the method employed. This application is available on request or else can be downloaded directly at http://photometry.merck.de

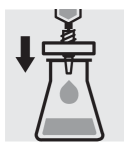


Application · Bromate in water and drinking water

a **xylem** brand

Measuring range: 0.003 – 0.120 mg/l BrO₃ 50-mm cell

Attention! The measurement is carried out at 550 nm in a 50-mm rectangular cell against a blank, prepared from distilled water and the reagents in an analogous manner.



Filter turbid samples.



Evaporate 200 ml of sample solution in a glass beaker almost to dryness on the hob.



Transfer the residue to a 20-ml volumetric glass using a little distilled water.



Make up the contents of the volumetric flask to the mark with distilled water and mix thoroughly: **pretreated sample.**



Pipette 10 ml of the pretreated sample into a test tube.



Add 0.10 ml of reagent 1 with pipette and mix.



Add 0.20 ml of reagent 2 with pipette and mix.



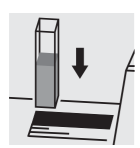
Add 0.20 ml perchloric acid 70–72 % GR with pipette and mix.



Reaction time: 30 minutes



Transfer the solution into a cell.



Place the cell into the cell compartment.
Select method **Bromate** in the menu (method no. **195**).

Important:

The exact composition and preparation of the reagents 1 and 2 used are given in the corresponding application, which also includes further information on the method employed. This application is available on request or else can be downloaded directly at http://photometry.merck.de



Application · Carbon dioxide

a xylem brand

Measuring range Equivalent to 0.40 – 8.00 mg/l OH

Measuring ranges for CO₂ depending on pH value and temperature.

Example: 14 – 275 mg/l CO₂ at pH 6.5 and 18.6 °C.



Check the pH of the sample **and note**.



Check the temperature of the solution **and note**.



Pipette 4.0 ml of **AC-1** (from Acid Capacity Test, Cat. No. 252087) into a round cell.



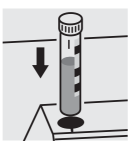
Add 1.0 ml of the sample with pipette, close the cell with the screw cap, and mix.



Add 0.50 ml of **AC-2** (from Acid Capacity Test, Cat. No. 252087) with pipette, close the cell with the screw cap, and mix.



Select method no. **2525**. Enter the pH and the temperature of the original sample.



Place the cell into the cell compartment. Align the mark on the cell with that on the photometer.

Quality assurance:

To check the measurement system (test reagents, measurement device, and handling) a sodium hydroxide solution 0.1 mol/l can be used after diluting accordingly (see section "Standard solutions").

a xylem brand

Determination of chlorophyll-a and phaeophytin-a analogous to DIN 38412

Page 1 of 2

Measuring	depending on the ratio of original sample to extract	10-mm cell	Method No. 2509
range:	in μg/l Chl-a or Phaeo	20-mm cell	Method No. 2510
		50-mm cell	Method No. 2511
Attention!	The measurement is carried out in a corresponding re	ctangular cell agair	nst a blank, prepared from
	ethanol (w = 90 %).		



Sufficiently homogenize 0.5 - 2 I of sample. Note the sample volume.



a suitable filter (e.g. glass-fibre filter).



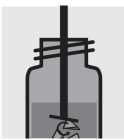
Filter the sample through Fold the loaded filter and tear into small pieces.



Place the pieces of the filter in an extraction vessel (e.g. 100-ml amber glass bottle).



Add approx. 30 ml of boiling ethanol (w = 90 %) and allow to cool to room temperature.



Disintegrate the filter in the homogenizer. Rinse together with a small por- to take place. tion of ethanol.



Allow to stand for 6 - 24 hours for the extraction



Filter the extract protected from light through a paper filter ("Blauband") into a volumetric flask (for DIN 38412: 100 ml). Rinse the filter with a small portion of ethanol.



Make the contents of the volumetric flask up to the mark with ethanol, keeping them protected from light in the process!



Transfer the solution into a corresponding cell.

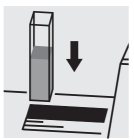
165



Select method no. 2509, 2510, or 2511. Enter the volumes of the original sample and extract (volumetric flask).

ba75728e11

03/2021



Place the cell into the cell compartment.

Vol (sample) [l] Vol (extr.) [ml] A (before acid.)[Abs] A (before acid.) [Abs]

a xylem brand

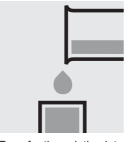
Determination of chlorophyll-a and phaeophytin-a analogous to DIN 38412

Page 2 of 2

Differentiation (chlorophyll-a - phaeophytin-a):



To differentiate the chlorophyll-a content and for the determination of the phaeophytin-a content, acidify a portion of the extract with hydrochloric acid for analysis (0.3 ml per 100 ml of extract).



START • ENTER

Transfer the solution into a corresponding cell.



Place the cell into the cell compartment and measure anew.



START • ENTER

Vol (sample) [l]
Vol (extr.) [ml]
A (before acid.) [Abs]
A (after acid.) [Abs]
Chl-a [µg/l]
Phaeo [µg/l]

Phaeo [µg/l]

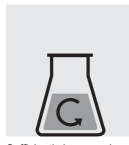


a xylem brand

Determination of chlorophyll-a and phaeophytin-a analogous to ASTM D3731-87

Page 1 of 2

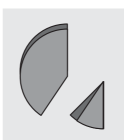
Measuring	depending on the ratio of original sample to extract	10-mm cell	Method No. 2504
range:	in mg/m ³ Chl-a or Phaeo-a	20-mm cell	Method No. 2505
		50-mm cell	Method No. 2506
Attention!	The measurement is carried out in a corresponding re	ectangular cell agair	nst a blank, prepared from
	extracting agent.		



Sufficiently homogenize the sample. Note the sample volume.



Filter the sample through Fold the loaded filter and a suitable filter (e.g. glass-fibre filter).



tear into small pieces.



Place the pieces of the filter in an extraction vessel (protected from light).



Add 2 - 3 ml of extracting agent.



Disintegrate the filter in the homogenizer.



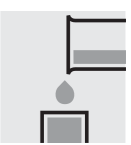
Make up to 10 ml with extracting agent.



Allow to stand at +4 °C for at least 2 hours for the extraction to take



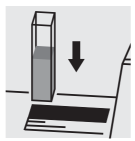
Filter the extract protected from light through a suitable filter.



Transfer the solution into a corresponding cell.



Select method no. 2504, 2505, or 2506. Enter the volumes of the original sample and extract (here: 10 ml).



Place the cell into the cell compartment.

Vol (sample) [l] Vol (extr.) [ml] A (before acid.) [Abs] A (before acid.) [Abs]

a xylem brand

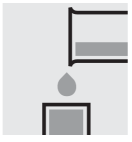
Determination of chlorophyll-a and phaeophytin-a analogous to ASTM D3731-87

Page 2 of 2

Differentiation (chlorophyll-a - phaeophytin-a):

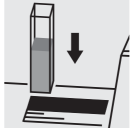


To differentiate the chlorophyll-a content and for the determination of the phaeophytin-a content, acidify a portion of the extract with hydrochloric acid 0.1 mol/l for analysis (0.15 ml per 5 ml of extract).

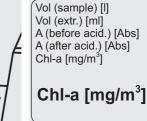


START • ENTER

Transfer the solution into a corresponding cell.



Place the cell into the cell compartment and measure anew.



START • ENTER

Vol (sample) [I] Vol (extr.) [ml] A (before acid.) [Abs] A (after acid.) [Abs] Chl-a [mg/m³] Phaeo-a [mg/m³]

Phaeo-a [mg/m³]

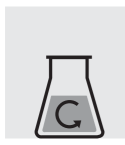


Application · Chlorophyll-a,-b,-c

a **xylem** brand

Trichromatic method analogous to ASTM D3731-87

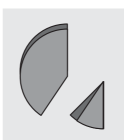
Method No. 2507 Measuring depending on the ratio of original sample to extract 10-mm cell in mg/m³ Chl-a, -b, -c Method No. 2508 range: 50-mm cell Attention! The measurement is carried out in a corresponding rectangular cell against a blank, prepared from extracting agent.



Sufficiently homogenize the sample. Note the sample volume.



Filter the sample through Fold the loaded filter and a suitable filter (e.g. glass-fibre filter).



tear into small pieces.



Place the pieces of the filter in an extraction vessel (protected from light).



Add 2 - 3 ml of extracting agent.



Disintegrate the filter in the homogenizer.



Make up to 10 ml with extracting agent.



Allow to stand at +4 °C for at least 2 hours for the extraction to take place.



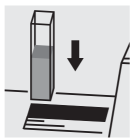
Filter the extract protected from light through a suitable filter.



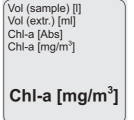
Transfer the solution into a corresponding cell.



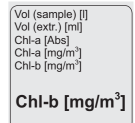
Select method no. 2507 or 2508. Enter the volumes of the original sample and extract (here: 10 ml).



Place the cell into the cell compartment.









Vol (sample) [l] Vol (extr.) [ml] Chl-a [Abs] Chl-a [mg/m³] Chl-b [mg/m³] Chl-c [mg/m³] Chl-c [mg/m³]



Application · Chromium in electroplating baths

a xylem brand

Inherent colour

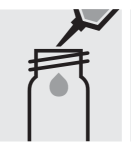
Measuring	20 -400 g/I CrO ₃	10-mm cell
range:	10 −200 g/l CrO ₃	20-mm cell
	4.0- 80.0 g/I CrO ₂	50-mm cell



Pipette 5.0 ml of the sample into a 100-ml volumetric flask, fill to the mark with distilled water and mix thoroughly.



Pipette 4.0 ml of the dilute sample into a 100-ml volumetric flask, fill to the mark with distilled water and mix thoroughly.



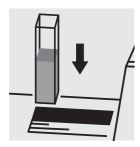
Pipette 5.0 ml of the 1:500 dilute sample into an empty round cell (Empty cells, Cat. No. 250621).



Add 5.0 ml of sulfuric acid 40%, close the cell with the screw cap, and mix.



Transfer the solution into a corresponding cell.



Place the cell into the cell compartment. Select method **Cr-bath** in the menu (method no. **20**).

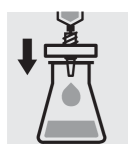


Application · Color (Spectral Absorption Coefficient)

a **xylem** brand

analogous to EN ISO 7887

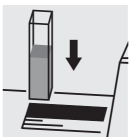
Measuring	1 -250 m ⁻¹	436 nm	10-mm cell	Method No. 015 α (436)
range:	0.3 – 125,0 m ⁻¹	436 nm	20-mm cell	Method No. 015 α (436)
	0.1 – 50.0 m ⁻¹	436 nm	50-mm cell	Method No. 015 α (436)
	1 -250 m ⁻¹	525 nm	10-mm cell	Method No. 061 α (525)
	0,3 – 125,0 m ⁻¹	525 nm	20-mm cell	Method No. 061 α (525)
	0.1 – 50.0 m ⁻¹	525 nm	50-mm cell	Method No. 061 α (525)
	1 -250 m ⁻¹	620 nm	10-mm cell	Method No. 078 α (620)
	0.3 – 125,0 m ⁻¹	620 nm	20-mm cell	Method No. 078 α (620)
	0.1 – 50.0 m ⁻¹	620 nm	50-mm cell	Method No. 078 α (620)



Filter sample solution through a membrane filter with 0.45 μ m pore size.



Transfer the solution into a corresponding cell.



Place the cell into the cell compartment, select method no. 15, 61, or 78.

Notes:

Filtered sample = true color. Unfiltered sample = apparent color.

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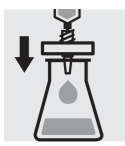


Applikation · Color (True Color - 410 nm)

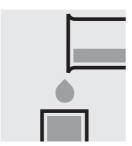
a **xylem** brand

analogous to EN ISO 7887

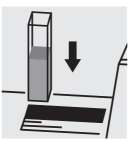
Measuring	10 – 2500 mg/l Pt	10 – 2500 mg/l Pt/Co	10 – 2500 CU	10-mm cell
range:	5 – 1250 mg/l Pt	5 – 1250 mg/l Pt/Co	5 – 1250 CU	20-mm cell
	2 - 500 mg/l Pt	2 - 500 mg/l Pt/Co	2 - 500 CU	50-mm cell



Filter sample solution through a membrane filter with 0.45 µm pore size.



Transfer the solution into a corresponding



Place the cell into the cell compartment, select method no. **303**.

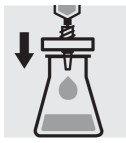


Application · Color Hazen (Platinum-Cobalt Standard Method)

a **xylem** brand

analogous to APHA 2120B, DIN EN ISO 6271-2, Water Research Vol. 30, No. 11, 2771-2775, 1996

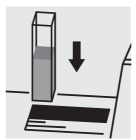
Measuring	1 - 500 mg/l Pt/Co	1 - 500 mg/l Pt	1 - 500 Hazen 1 - 500 CU	340 nm 10-mm cell
range:	1 - 250 mg/l Pt/Co	1 - 250 mg/l Pt	1 - 250 Hazen 1 - 250 CU	340 nm 20-mm cell
	0.2-100.0 mg/l Pt/Co	0.2-100.0 mg/l Pt	0.2-100.0 Hazen 0.2-100.0 CU	340 nm 50-mm cell



Filter sample solution through a membrane filter with 0.45 µm pore size.



Transfer the solution into a corresponding cell



Place the cell into the cell compartment, select method no. **32**.

Notes:

Filtered sample = true color.
Unfiltered sample = apparent color.

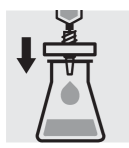


Application · Color Hazen (Platinum-Cobalt Standard Method)

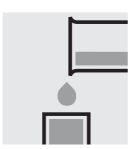
a **xylem** brand

analogous to APHA 2120B, DIN 53409, Water Research Vol. 30, No. 11, 2771-2775, 1996

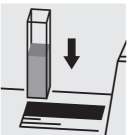
Measuring	1-1000 mg/l Pt/Co	1-1000 mg/l Pt	1-1000 Hazen	1-1000 CU	445 nm	50-mm cell	Method No. 179
range:	1-1000 mg/l Pt/Co	1-1000 mg/l Pt	1-1000 Hazen	1-1000 CU	455 nm	50-mm cell	Method No. 180
	1-1000 mg/l Pt/Co	1-1000 mg/l Pt	1-1000 Hazen	1-1000 CU	465 nm	50-mm cell	Method No. 181



Filter sample solution through a membrane filter with 0.45 µm pore size.



Transfer the solution into the cell.



Place the cell into the cell compartment, select method no. 179, 180, or 181.

Notes:

Filtered sample = true color. Unfiltered sample = apparent color.



Application · Copper in electroplating baths

a **xylem** brand

Inherent colour

Measuring	10.0-80.0 g/I Cu	10-mm cell	
range:	5.0-40.0 g/I Cu	20-mm cell	
	2.0-16.0 g/l Cu	50-mm cell	



Pipette 25 ml of the sample into a 100-ml volumetric flask, fill to the mark with distilled water and mix thoroughly.



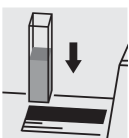
Pipette 5.0 ml of the 1:4 dilute sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 5.0 ml of sulfuric acid 40 %, close the cell with the screw cap, and mix.



Transfer the solution into Place the cell into the a corresponding cell. Place the cell compartment. Selection into the cell compartment into the cell compartment into the cell compartment into the cell into the



Place the cell into the cell compartment. Select method **Cu-bath** in the menu (method no. **83**).

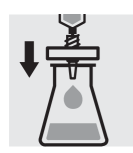


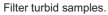
Application · lodine colour number

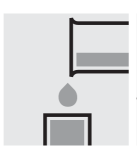
a **xylem** brand

analogous to DIN 6162A

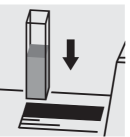
Measuring	0.05 -3.00 IFZ	340 nm	10-mm cell
range:	0.03 -1.50 IFZ	340 nm	20-mm cell
	0.010-0.600 IFZ	340 nm	50-mm cell







Transfer the solution into Place the cell into the a corresponding cell. Place the cell compartment, selections are considered to the cell compartment, selections are considered to the cell compartment.



Place the cell into the cell compartment, select method in the menue (method no. 33).



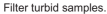
Application · lodine colour number

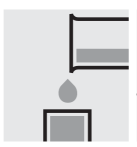
a **xylem** brand

analogous to DIN 6162A

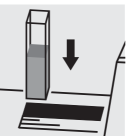
Measuring	1.0-50.0 IFZ	445 nm	10-mm cell	
range:	0.5-25.0 IFZ	445 nm	20-mm cell	
	0.2-10.0 IFZ	445 nm	50-mm cell	







Transfer the solution into Place the cell into the a corresponding cell. Place the cell compartment, selections are considered to the cell compartment, selections are considered to the cell compartment.



Place the cell into the cell compartment, select method in the menue (method no. 21).



Application · Mercury in water and wastewater

a xylem brand

Measuring range: 0.025-1.000 mg/l Hg 50-mm cell



Check the pH of the sample, specified range: pH 3-7. If required, add dilute sodium hydroxide solution or acetic acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into a test tube.



Add 1.0 ml of **reagent 1** with pipette and mix.



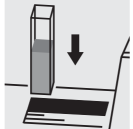
Add 1.5 ml of **reagent 2** with pipette and mix.



Reaction time: 5 minutes



Transfer the solution into a cell.



Place the cell into the cell compartment.
Select method **Mercury** in the menu (method no. **135**).

Important:

The exact composition and preparation of the reagents 1 and 2 used are given in the corresponding application, which also includes further information on the method employed. This application is available on request or else can be downloaded directly at http://photometry.merck.de



Application · Nickel in electroplating baths

a **xylem** brand

Inherent colour

Measuring	10 -120 g/l Ni	10-mm cell
range:	5.0- 60.0 g/l Ni	20-mm cell
	2.0 - 24.0 g/l Ni	50-mm cell



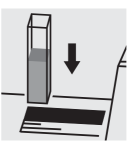
Pipette 5.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 5.0 ml of sulfuric acid 40%, close the cell with the screw cap, and mix.



Transfer the solution into Place the cell into the a corresponding cell. Place the cell into the cell compartment.



Place the cell into the cell compartment.
Select method **Ni-bath** in the menu (method no. **57**).

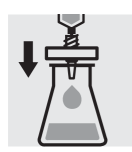


Application · Nitrate

a **xylem** brand

Direct measurement in the UV range analogous to APHA 4500-NO₃⁻ B

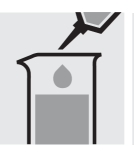
Measuring range: $0.0 - 7.0 \text{ mg/I NO}_3\text{-N}$ 10-mm quartz cell



Filter turbid samples.



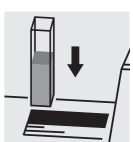
Place 50 ml of sample into a glass vessel.



Add 1 ml of hydrochloric acid 1mol/l for analy- the quartz cell. sis with pipette and mix.



Transfer the solution into Place the cell into the



cell compartment. Select method no. 2503.

Important:

If "Condition not met" appears on the display, this is due to a sample-dependent interference (matrix effect). In this case an evaluation is not possible.



Application · Palladium in wastewater

a xylem brand

Measuring range: 0.05 – 1.25 mg/l Pd 10-mm cell



Check the pH of the sample, specified range: pH 2-5. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1.0 ml of **reagent 1** with pipette, close the cell with the screw cap, and mix.



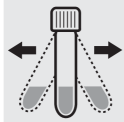
sample, specified value: pH 3.0. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust



Add 0.20 ml of reagent 2 with pipette, close the cell with the screw cap, and mix.



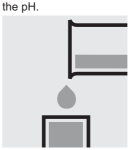
Add 5.0 ml **isoamyl alcohol GR** with pipette, close the cell with the screw cap.



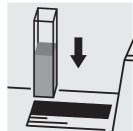
Shake the cell vigorously for 1 minute. Leave to stand to allow phases to separate.



Aspirate the organicclear upper phase from the tube with pipette and dry over sodium sulfate anhydrous.



Transfer the dried solution into a cell.



Place the cell into the cell compartment. Select method **Palladium** in the menu (method no. **133**).

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Important:

The exact composition and preparation of the reagents 1 and 2 used are given in the corresponding application, which also includes further information on the method employed. This application is available on request or else can be downloaded directly at http://photometry.merck.de

03/2021



Application · Platinum in water and wastewater

a xylem brand

Measuring range: 0.10–1.25 mg/l Pt 10-mm cell

Attention! The measurement is carried out at 690 nm in a 10-mm rectangular cell against a blank, prepared from distilled water and the reagents in an analogous manner.



Check the pH of the sample, specified range: pH 2-5. If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



Pipette 5.0 ml of the sample into an empty round cell (Empty cells, Cat.No. 250621).



Add 1.0 ml of **reagent 1** with pipette, close the cell with the screw cap, and mix.



Add 0.50 ml of reagent 2 with pipette, close the cell with the screw cap, and mix.



Check the pH of the sample, specified value: pH 6.5.
If required, add dilute sodium hydroxide solution or sulfuric acid drop by drop to adjust the pH.



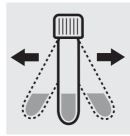
Heat the cell in the thermoreactor at 100 °C for 5 minutes.



Remove the cell from the thermoreactor and place in a test-tube rack to cool to room temperature.



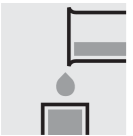
Add 5.0 ml Isobutylmethylketone GR with pipette, close the cell with the screw cap.



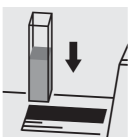
Shake the cell vigorously for 1 minute. Leave to stand to allow phases to separate.



Aspirate the organicclear upper phase from the tube with pipette and dry over **sodium sulfate anhydrous**.



Transfer the dried solution into a cell.



Place the cell into the cell compartment.
Select method **Platinum** in the menu (method no. **134**).

Note:

Empty cells with screw caps, Cat.No. 250621 are recommended for the preparation. These cells can be sealed with the screw caps, thus enabling a hazard-free mixing of the sample.

Important:

The exact composition and preparation of the reagents 1 and 2 used are given in the corresponding application, which also includes further information on the method employed. This application is available on request or else can be downloaded directly at http://photometry.merck.de

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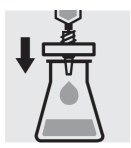


Application - Spectral Absorption Coefficient $\alpha(254)$

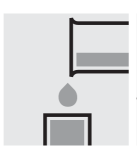
a xylem brand

analogous to DIN 38404

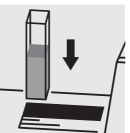
Measuring range:	3 –250 m ⁻¹	254 nm	10-mm cell
	1 –125 m ⁻¹	254 nm	20-mm cell
	0.5 - 50.0 m ⁻¹	254 nm	50-mm cell



Filter sample solution through a membrane filter with 0.45 μ m pore size.



Transfer the solution into the cell.



Place the cell into the cell compartment, select method in the menue (method no. **300**).

Cell type:

Use only quartz cells. Plastic cells cannot normally be used for the UV range because they do not cover this wavelength measuring range.



Application · Spectral Attenuation Coefficient µ(254)

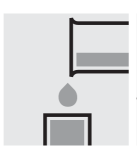
a xylem brand

analogous to DIN 38404

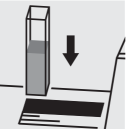
Measuring range:	3 –250 m ⁻¹	254 nm	10-mm cell
	1 –125 m ⁻¹	254 nm	20-mm cell
	0.5 - 50.0 m ⁻¹	254 nm	50-mm cell



Shake the unfiltered sample solution to evenly suspend the turbidity-causing substances. Do not disperse the contents, measure immediately.



Transfer the solution into the cell.



Place the cell into the cell compartment, select method in the menue (method no. 301).

Note:

When the turbidity correction function is activated (see Description of Function, section 4.5.9 "Automatic Turbidity correction"), the **corrected spectral attenuation coefficient \mu(254)korr** can be determined.

The turbidity correction is carried out as per DIN 38404 at $550\ \mathrm{nm}$.

Cell type:

Use only quartz cells. Plastic cells cannot normally be used for the UV range because they do not cover this wavelength measuring range.

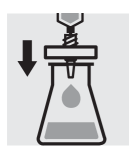


Application · Spectral Absorbtion Coefficient α (436)

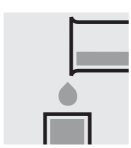
a xylem brand

analogous to EN ISO 7887

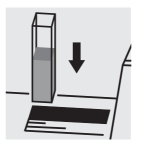
Measuring range:	3 -250 m ⁻¹	436 nm	10-mm cell
	1 –125 m ⁻¹	436 nm	20-mm cell
	0.5 – 50.0 m ⁻¹	436 nm	50-mm cell



Filter sample solution through a membrane filter with 0.45 µm pore size.



Transfer the solution into the cell.



Place the cell into the cell compartment, select method in the menue (method no. **302**).

Notes:

Filtered sample = true colour.
Unfiltered sample = apparent colour.

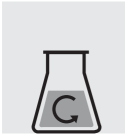
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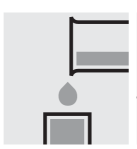
Application · Suspended Solids

a **xylem** brand

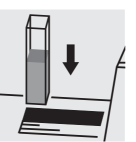
Measuring range: 25-750 mg/l Susp. solids 20-mm cell



Homogenize 500 ml of sample for 2 minutes in a mixer running at high speed.



Transfer the solution into a cell.



Place the cell into the cell compartment, select method in the menue (method no. 182).

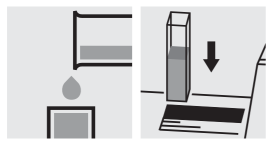


Application · Turbidity

a **xylem** brand

analogous to EN ISO 7027

Measuring range: 1–100 FAU 550 nm 50-mm cell



Transfer the sample into a cell.

Place the cell into the cell compartment, select method in the menue (method no. 77).

Test kits without barcode

Available methods

The analysis specifications for these test sets are given in Appendix 4. Here, the method is selected manually, using the method number given in column 5. Instructions on how to select a method are given in the section, SELECTING A METHOD MANUALLY of the functional description of the photometer.

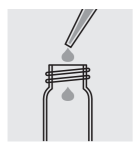
Parameter / Name	Modell	Artikel- Nr.	Gesamtmessbereich	Metho- den-Nr.	Тур**	Blindwertmessung
Alkalinity-M	KsM-1	#	5 - 200 mg/l CaCO₃	7339	TT	required
Alkalinity-P	KsP-1	#	5 - 300 mg/l CaCO₃	7340	TT	required
Ammonium vario	NH4-1 TP	251 408	0.01 - 0.50 mg/l NH ₄ -N	7324	PP	required
Ammonium vario LR	NH4-2 TC (LR)	251 997	0.02 - 2.50 mg/l NH ₄ -N	7312	KT	required
Ammonium vario HR	NH4-3 TC (HR)	251 998	0.4 - 50.0 mg/l NH ₄ -N	7313	KT	required
Chlor (free) vario	Cl2-1 TP	251 401	0.02 - 2.00 mg/l Cl ₂	7325	PP	required
Chlor (free) vario	CI2-2 TP	251 402	0.50 - 5.00 mg/l Cl ₂	7326	PP	required
Chlor (total) vario	CI2-3 TP	251 414	0.02 - 2.00 mg/l Cl ₂	7327	PP	required
Chlor (total) vario	CI2-4 TP	251 415	0.5 - 5.0 mg/l Cl ₂	7328	PP	required
COD LR	COD1 TC (LR)	251 990	3 - 150 mg/l COD	7309	KT	required
COD MR	COD2 TC (MR)	251 991	20 - 1500 mg/l COD	7310	KT	required
COD HR	COD3 TC (HR)	251 992	200 - 15000 mg/l COD	7311	KT	required
Copper vario	Cu-1 TP	251 403	0.04 - 5.00 mg/l Cu	7302	PP	required
DEHA vario	DEHA-1 TP	251 421	0.004 - 0.450 mg/l DEHA	7335	PP	required
Iron vario TPTZ	Fe-1 TP	251 404	0.012 - 1.800 mg/l Fe	7300	PP	required
Iron vario	Fe-2 TP	251 405	0.02 - 3.00 mg/l Fe	7301	PP	required
Hydrazine vario	N2H4-1 TP	251 416	0.004 - 0.600 mg/l N ₂ H ₄	7329	PP	required
Manganese vario	Mn-1 TP	251 406	0.2 - 20.0 mg/l Mn	7303	PP	required
Manganese vario	Mn-2 TP	251 417	0.007 - 0.700 mg/l Mn	7330	PP	required
Molybdate vario	Mo-1 TP	251 407	0.3 - 35.0 mg/l Mo	7304	PP	required
Molybdenum vario	Mo-2 TP	251 418	0.3 - 40.0 mg/l Mo	7331	PP	required
Nitrate	NO3-1 TC	251 993	0.2 - 30.0 mg/l NO ₃ -N	7314	KT	required
Nitrite vario	NO2-1 TP	251 409	0.002 - 0.300 mg/l NO ₂ -N	7305	PP	required
Nitrite LR	NO2-2 TC (LR)	251 994	0.03 - 0.60 mg/l NO ₂ -N	7318	KT	required
Nitrite HR	NO2-2 TC (HR)	251 994	0.3 - 3.0 mg/l NO ₂ -N	7317	KT	required
Nitrite vario	NO2-3 TP	251 420	0.002 - 0.300 mg/l NO ₂ -N	7334	PP	required
Nitrogen, total LR	Ntot1 TC (LR)	251 995	0.5 - 25.0 mg/l N _{tot}	7319	KT	required
Nitrogen, total HR	Ntot2 TC (HR)	251 996	10 - 150 mg/l N _{tot}	7320	KT	required
Phosphate vario (ortho)	PO4-1 TP	251 410	0.02 - 2.50 mg/l PO ₄	7306	KT	required
Phosphate, ortho	PO4-2 TC	251 989	0.06 - 5.00 mg/l PO ₄	7315	KT	required
Phosphat, total	PO4-3 TC	251 988	0.06 - 3.50 mg/l PO ₄	7316	KT	required
Phosphat, total	PO4-4 TP	251 987	0.06 - 3.50 mg/l PO ₄	7336	KT	required
Phosphat, acid hydrolyzable	PO4-4 TP	251 987	0.06 - 3.50 mg/l PO ₄	7336	KT	required
Silica HR vario	Si-3 TP (HR)	251 422	1 - 200 mg/l SiO ₂	7337	PP	required
Silica LR vario	Si-1 TP (LR)	251 411	0.01 - 1.60 mg/l SiO ₂	7321	PP	required
Silica HR vario	Si-2 TP (HR)	251 412	1 - 100 mg/l SiO ₂	7308	PP	required
Sulfate vario	SO4-2 TP	251 423	2 - 70 mg/l SO ₄	7338	PP	required

 $^{^{\}star}$ KT = reaction cell test (16 mm round cell); RT = reagent test; TP = powder pillow test

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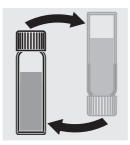
WTW model no.:	KsM-1
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	5 - 200 mg/l CaCO ₃



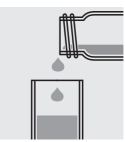
Pipette 10.0 ml of sample into the empty cell.



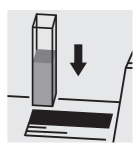
Add 1 tablet **ALKA-M-PHOTOMETER** directly from the foil; crush it with a clean stirring rod and close the cell with the screw cap.



Mix the contents by swirling the cell until the tablet has dissolved.



Fill the solution into the measuring cell.

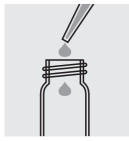


Insert the cell in the photometer cell shaft and start measurement.

- We recommend that you determine a new reagent blank value (H2O dist instead of the sample) when starting a new package.
- The coloring that has developed is not long-term stable. Therefore, measure the sample speedily after the tablet has dissolved.
- The coloring is very intensive and can discolor the stirring rod and cells. If possible, clean the utensils immediately after measuring.



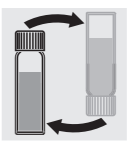
WTW model no.:	KsP-1
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	5 - 300 mg/l CaCO₃



Pipette 10.0 ml of sample into the empty cell.



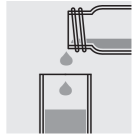
Add 1 tablet ALKA-P-PHOTOMETER directly from the foil; crush it with a clean stirring rod and close the cell with the screw cap.



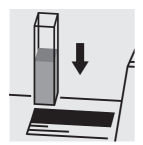
Mix the contents by swirling the cell until the tablet has dissolved.



Allow to react for 5 minutes.



Fill the solution into the measuring cell.

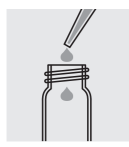


Insert the cell in the photometer cell shaft and start measurement.

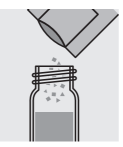
- We recommend that you determine a new reagent blank value (H2O dist instead of the sample) when starting a new package.
- The coloring that has developed is not long-term stable. Therefore, measure the sample speedily after the reaction time is
 over.
- The coloring is very intensive and can discolor the stirring rod and cells. If possible, clean the utensils immediately after measuring.



WTW model no.:	NH4-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.01 - 0.50 mg/l NH ₄ -N
	0.01 - 0.64 mg/l NH ₄
	Display in mmol/l possible



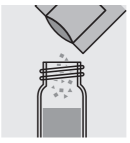
Pipette 10.0 ml of sample into the empty cell.



Add the contents of a VARIO AMMONIA Salicylate F10 powder pack and close the cell with the screw cap.



Allow to react for 3 minutes (reaction time).



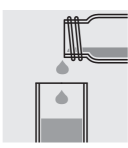
Add the contents of a VARIO AMMONIA
Cyanurate F10 powder pack and close the cell with the screw cap.



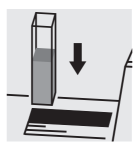
Shake the cell vigorously to dissolve solids.



Allow to react for 15 minutes (reaction time).



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If NH₄-N is present in the sample, the solution will turn green after the VARIO AMMONIA Cyanurate F10 was added.
- If chlorine is present, the sample has to be treated with sodium thiosulfate immediately after sampling. To 1 liter of the sample, add one drop of a 0.1 mol/l sodium thiosulfate solution per 0.3 mg/l chlorine.



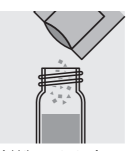
WTW model no.:	NH4-2 TC (LR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.02 - 2.50 mg/l NH ₄ -N
	0.03 - 3.20 mg/l NH ₄
	Display in mmol/l possible



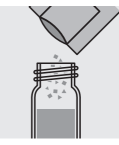
Check the pH value of the sample. Required value: approx. pH 7. Correct with diluted sodium hydroxide solution or hydrochloric acid as necessary.



Pipette 2.0 ml of sample into a reaction cell.



Add the contents of a VARIO AMMONIA Salicylate F5 powder pack.



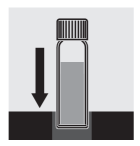
Add the contents of a VARIO AMMONIA Cyanurate F5 powder pack and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



Allow to react for 20 minutes.



Insert the cell in the photometer cell shaft and start measurement.

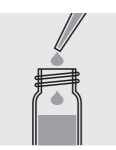
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If NH₄-N is present in the sample, the solution will turn green after the VARIO AMMONIA Cyanurate F5 was added.
- If chlorine is present, the sample has to be treated with sodium thiosulfate immediately after sampling. To 1 liter of the sample, add one drop of a 0.1 mol/l sodium thiosulfate solution per 0.3 mg/l chlorine.
- Iron interferes with the determination and can be eliminated as follows: Determine the total iron concentration and prepare
 an iron standard solution with the determined concentration. Use this solution to determine the reagent blank value for the
 determination of ammonium (instead of distilled water).



WTW model no.:	NH4-3 TC (HR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.4 - 50.0 mg/l NH ₄ -N
	0.5 - 64.4 mg/l NH ₄
	Display in mmol/l possible



Check the pH value of the sample. Required value: approx. pH 7. Correct with diluted sodium hydroxide solution or hydrochloric acid as necessary.



Pipette 0.1 ml of sample into a reaction cell.



Add the contents of a VARIO AMMONIA Salicylate F5 powder pack.



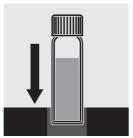
Add the contents of a VARIO AMMONIA Cyanurate F5 powder pack and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



Allow to react for 20 minutes.

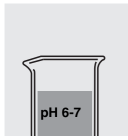


Insert the cell in the photometer cell shaft and start measurement.

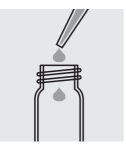
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If NH₄-N is present in the sample, the solution will turn green after the VARIO AMMONIA Cyanurate F5 was added.
- If chlorine is present, the sample has to be treated with sodium thiosulfate immediately after sampling. To 1 liter of the sample, add one drop of a 0.1 mol/l sodium thiosulfate solution per 0.3 mg/l chlorine.
- Iron interferes with the determination and can be eliminated as follows: Determine the total iron concentration and prepare
 an iron standard solution with the determined concentration. Use this solution to determine the reagent blank value for the
 determination of ammonium (instead of distilled water).



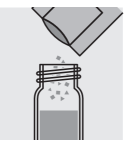
WTW model no.:	Cl2-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.02 - 2.00 mg/l Cl ₂
	Display in mmol/l possible



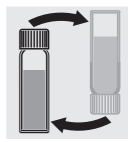
Check the pH value of the sample. Required range: pH 6-7. Correct with diluted sodium hydroxide solution or sulfuric acid as necessary.



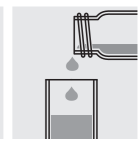
Pipette 10.0 ml of sample into the empty cell.



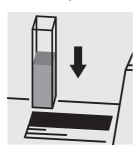
Add the contents of a **Chlorine Free-DPD F10** powder pack and close the cell with the screw cap.



Mix the contents by swaying the cell (20 seconds).



Fill the solution into the measuring cell.

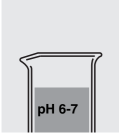


Within one minute, insert the cell in the cell shaft of the photometer and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The test sample should be pink. Very high chlorine concentrations in the sample cause yellow solutions and too low measured values. Dilute the sample in this case.



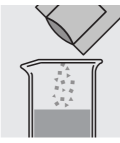
WTW model no.:	CI2-2 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.5 - 5.0 mg/l Cl ₂
	Display in mmol/l possible



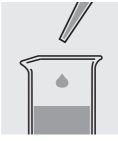
Check the pH value of the sample. Desired range: pH 6-7. Correct with diluted sodium hydroxide solution or sulfuric acid as necessary.



Pipette 10.0 ml of sample into an empty beaker.



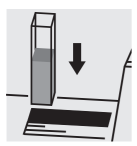
Add the contents of a VARIO Chlorine Free-DPD F25 powder pack and dissolve them by stirring.



Add 15.0 ml deionized water with a pipette and mix.



Fill the solution into the measuring cell.

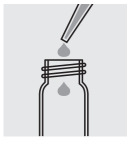


Insert the cell in the photometer cell shaft and start measurement.

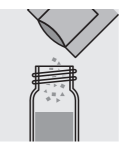
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The test sample should be pink. Very high chlorine concentrations in the sample cause yellow solutions and too low measured values. Dilute the sample in this case.



WTW model no.:	CI2-3 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.02 - 2.00 mg/l Cl ₂
	Display in mmol/l possible



Pipette 10.0 ml of sample into the empty cell.



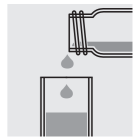
Add the contents of a Chlorine Total-DPD F10 powder pack and close the cell with the screw cap.



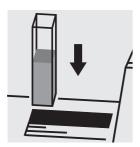
Shake the cell vigorously to dissolve solids. A small amount of solid matter may remain undissolved.



Allow to react for 3 minutes.



Fill the solution into the measuring cell.

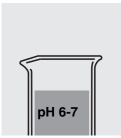


Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The test sample should be pink. Very high chlorine concentrations in the sample cause yellow solutions and too low measured values. Dilute the sample in this case.
- Each time after determining total chlorine, rinse the cell with sulfuric acid 25 % and then several times with distilled water.



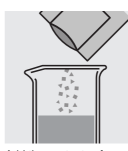
WTW model no.:	CI2-4 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.5 - 5.0 mg/l Cl ₂
	Display in mmol/l possible



Check the pH value of the sample. Required range: pH 6-7. Correct with diluted sodium hydroxide solution or sulfuric acid as necessary.



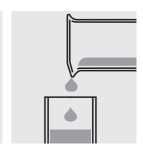
Pipette 10.0 ml of sample into an empty beaker.



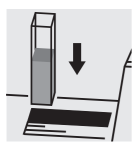
Add the contents of a VARIO Chlorine Total-DPD F25 ml powder pack and dissolve them by stirring.



Add 15.0 ml deionized water with a pipette and mix.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The test sample should be pink. Very high chlorine concentrations in the sample cause yellow solutions and too low measured values. Dilute the sample in this case.
- Each time after determining total chlorine, rinse the cell with sulfuric acid 25 % and then several times with distilled water.

Method no.

7309



WTW model no.:	COD1 TC (LR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	3 - 150 mg/l COD

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Shake the reaction cell so that the sediment is suspended.



Carefully pipette 2.0 ml of sample into the cell, close with screw cap and mix vigorously.

Caution, cell becomes very hot!



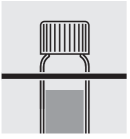
Heat the cell in the thermoreactor for two hours at 148 °C.



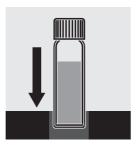
Remove the cell from the thermoreactor and let it cool down in a cell rack.



After approx. 10 min cooling time sway the cell.



Place the cell in the cell rack again and let it cool down to room temperature.



Carefully insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The chloride content of the sample must not exceed 1000 mg/l.
- Homogenize samples containing suspended matter with a disperser.
- Before being inserted in the thermoreactor and for photometric measurements the outside of the cell must be free of any contamination (e.g. fingerprints or drops of water). Wipe the cell with a dry cloth as necessary.
- Let the cell cool down long enough (at least 45 min) before inserting it in the photometer cell shaft. The cells remain stable for a long time after reaction and can also be left overnight and then measured.
- After cooling do not rock the cell until the measurement takes place in order not to suspend the solids that formed during the reaction. Suspended matter disturbs the photometric measurement.

Method no.

7310



WTW model no.:	COD2 TC (MR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	20 - 1500 mg/l COD

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Shake the reaction cell so that the sediment is suspended.



Carefully pipette 2.0 ml of sample into the cell, close with screw cap and mix vigorously.

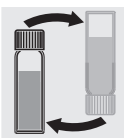
Caution, cell becomes very hot!



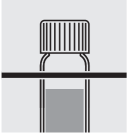
Heat the cell in the thermoreactor for two hours at 148 °C.



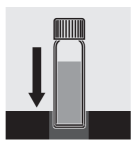
Remove the cell from the thermoreactor and let it cool down in a cell rack.



After approx. 10 min cooling time sway the cell.



Place the cell in the cell rack again and let it cool down to room temperature.



Carefully insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The chloride content of the sample must not exceed 1000 mg/l.
- Homogenize samples containing suspended matter with a disperser.
- Before being inserted in the thermoreactor and for photometric measurements the outside of the cell must be free of any contamination (e.g. fingerprints or drops of water). Wipe the cell with a dry cloth as necessary.
- Let the cell cool down long enough (at least 45 min) before inserting it in the photometer cell shaft. The cells remain stable for a long time after reaction and can also be left overnight and then measured.
- After cooling do not rock the cell until the measurement takes place in order not to suspend the solids that formed during the reaction. Suspended matter disturbs the photometric measurement.

Method no. **7311**



WTW model no.:	COD3 TC (HR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	200 - 15000 mg/l COD

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Shake the reaction cell so that the sediment is suspended.



Carefully pipette 0.2 ml of sample into the cell, close with screw cap and mix vigorously.

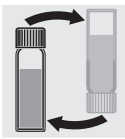
Caution, cell becomes very hot!



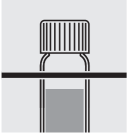
Heat the cell in the thermoreactor for two hours at 148 °C.



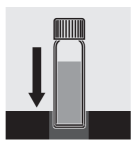
Remove the cell from the thermoreactor and let it cool down in a cell rack.



After approx. 10 min cooling time sway the cell.



Place the cell in the cell rack again and let it cool down to room temperature.



Carefully insert the cell in the photometer cell shaft and start measurement.

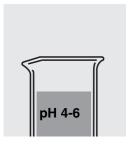
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The chloride content of the sample must not exceed 10,000 mg/l.
- Homogenize samples containing suspended matter with a disperser.
- Before being inserted in the thermoreactor and for photometric measurements the outside of the cell must be free of any contamination (e.g. fingerprints or drops of water). Wipe the cell with a dry cloth as necessary.
- Let the cell cool down long enough (at least 45 min) before inserting it in the photometer cell shaft. The cells remain stable for a long time after reaction and can also be left overnight and then measured.
- After cooling do not rock the cell until the measurement takes place in order not to suspend the solids that formed during the reaction. Suspended matter disturbs the photometric measurement.

7302

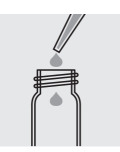


WTW model no.:	Cu-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.04 - 5.00 mg/l Cu
	Display in mmol/l possible

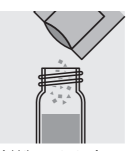
Note: Before using the test with your photometer for the first time, determine the reagent blank value.



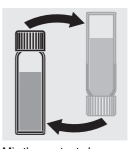
Check the pH value of the sample. Required range: pH 4-6. Correct with diluted sodium hydroxide solution or caustic potash solution as necessary.



Pipette 10.0 ml of sample into the empty cell.



Add the contents of a **VARIO Cu1 F10** powder pack and close the cell with the screw cap.

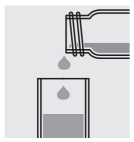


Mix the contents by carefully swaying the cell (10 times). Any undissolved powder does not adversely affect

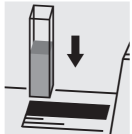
the measurement.



Allow to react for 2 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

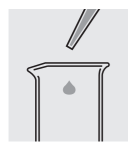
· We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

7335



WTW model no.:	DEHA-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.004 - 0.450 mg/l DEHA
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 25.0 ml of sample into an empty beaker.



Add the contents of a **VARIO Oxyscav 1 RGT** powder pack and dissolve it by stirring.



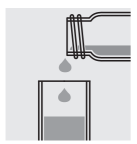
Add 0.5 ml **VARIO DEHA 2 RGT** with a pipette and mix.



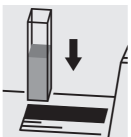
Fill an empty cell with the prepared sample, close it with the screw cap and put it in a dark place.



Allow the sample to react for ten minutes in a dark place. Then measure immediately.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

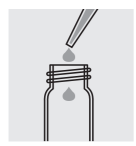
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- Clean all laboratory glassware with hydrochloric acid (approx. 20 %), then thoroughly rinse with deionized water.
- Avoid excessive movements and exposure to sun light during sampling. Store the samples hermetically sealed.
- The temperature of the samples must be 25±3 °C.

7329



WTW model no.:	N2H4-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	$0.004 - 0.600 \text{ mg/l N}_2\text{H}_4$
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 10.0 ml of sample into the empty cell.



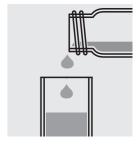
With a pipette add 0.5 ml VARIO Hydra2 Reagent carefully swaying the Solution and close the cell with the screw cap.



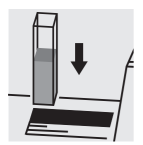
Mix the contents by cell.



Allow to react for 12 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

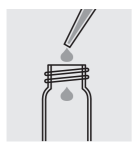
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If any hydrazine is present, the solution develops a yellow color after the reagent is added.
- The temperature of the samples must be 21±4 °C.
- Avoid moving the sample too much or too long exposure to air.

7300

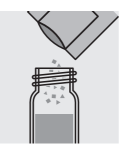


WTW model no.:	Fe-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.012 - 1.800 mg/l Fe
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 10.0 ml of sample into the empty cell.



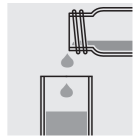
Add the contents of a VARIO Iron TPTZ F10 powder pack and close the cell with the screw cap.



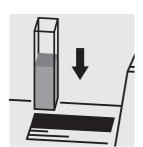
To dissolve the solids, shake the cell vigorously for approx. 30 seconds.



Allow to react for 3 minutes (reaction time).



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

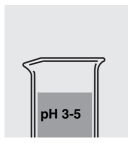
• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

7301

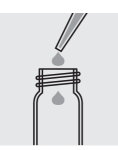


WTW model no.:	Fe-2 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.02 - 3.00 mg/l Fe
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



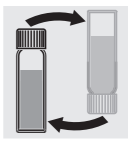
Check the pH value of the sample. Required range: pH 3-5. Correct with diluted sodium hydroxide solution or hydrochloric acid as necessary.



Pipette 10.0 ml of sample into the empty cell.



Add the contents of a **VARIO Ferro F10** powder pack and close the cell with the screw cap.

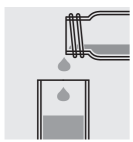


Mix the contents by carefully swaying the cell (10 times). Any undissolved powder does not adversely affect

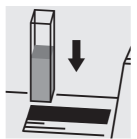
the measurement.



Allow to react for 3 minutes (reaction time).



Fill the solution into the measuring cell.

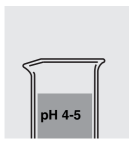


Insert the cell in the photometer cell shaft and start measurement.

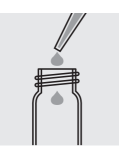
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- This method measures all types of dissolved iron and most types of undissolved iron.
- If there is visible rust in the sample the reaction time should be at least 5 minutes.



WTW model no.:	Mn-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.2 - 20.0 mg/l Mn
	Display in mmol/l possible



Check the pH value of the sample. Required range: pH 4-5. Correct with diluted nitric acid or sodium hydroxide solution as necessary.



Pipette 10.0 ml of sample into the empty cell.



Add the contents of a VARIO Manganese Citrate Buffer F10 powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell (10 times).



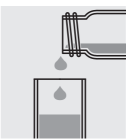
Add the contents of a VARIO Sodium
Periodate F10 powder pack and close the cell with the screw cap.



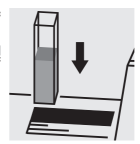
Mix the contents by carefully swaying the cell (10 times).



Allow to react for 2 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

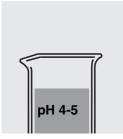
• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

7330

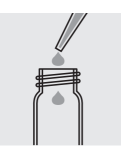


WTW model no.:	Mn-2 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.007 - 0.700 mg/l Mn
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



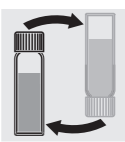
Check the pH value of the sample. Required range: pH 4-5. Correct with diluted nitric acid or sodium hydroxide solution as necessary.



Pipette 10.0 ml of sample into the empty cell.



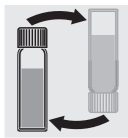
Add the contents of a **VARIO Ascorbic Acid** powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



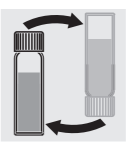
Add 15 drops of VARIO Alkaline-Cyanide Reagent Solution and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



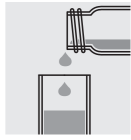
Add 21 drops of VARIO PAN Indicator Solution 0.1% and close the cell with the screw cap.



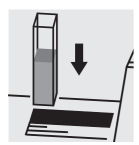
Mix the contents by carefully swaying the cell.



Allow to react for 2 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

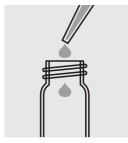
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- Clean all laboratory glassware with nitric acid, then thoroughly rinse with deionized water.

7304

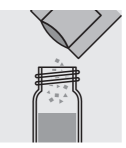


WTW model no.:	Mo-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.3 - 35.0 mg/l Mo
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



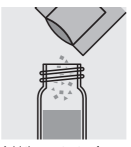
Pipette 10.0 ml of sample into the empty cell.



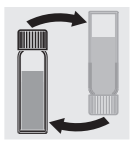
Add the contents of a **VARIO Molybdenum HR1 F10** powder pack and close the cell with the screw cap.



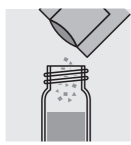
Mix the contents by carefully swaying the cell (10 times).



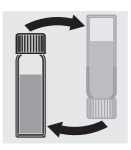
Add the contents of a **VARIO Molybdenum HR2 F10** powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell (10 times).



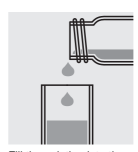
Add the contents of a **VARIO Molybdenum HR3 F10** powder pack and close the cell with the screw cap.



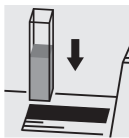
Mix the contents by carefully swaying the cell (10 times).
Any undissolved powder does not adversely affect the measurement.



Allow to react for 5 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

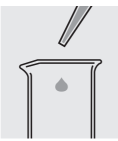
• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.



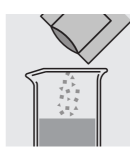
WTW model no.:	Mo-2 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.3 - 40.0 mg/l Mo
	Display in mmol/l possible



Check the pH value of the sample. Required value: approx. pH 7. Correct with diluted sodium hydroxide solution or nitric acid as necessary.



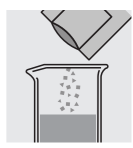
Pipette 25.0 ml of sample into an empty beaker.



Add the contents of a VARIO Molybdenum HR 1 F25 ml powder pack and dissolve them by stirring.



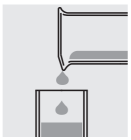
Add the contents of a VARIO Molybdenum HR 2 F25 ml powder pack and dissolve them by stirring.



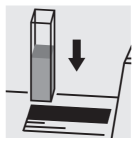
Add the contents of a VARIO Molybdenum HR 3 F25 ml powder pack and dissolve them by stirring.



Allow to react for 5 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If any molybdenum is present, the solution develops a yellow color after all reagents have been added.

7314

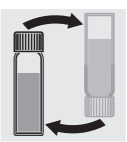


WTW model no.:	NO3-1 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.2 - 30.0 mg/l NO ₃ -N
	1.9 - 133.0 mg/l NO₃
	Display in mmol/l possible

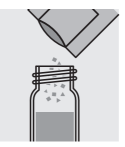
Note: Before using the test with your photometer for the first time, determine the reagent blank value.



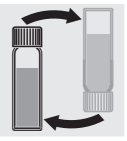
Pipette 1.0 ml of sample into a reaction cell and close the cell with the screw cap.



Mix the contents by carefully swaying the cell (10 times).



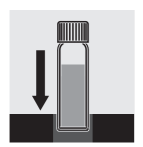
Add the contents of a **Nitrate Chromotropic** powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell (10 times). A small amount of solid matter may remain undissolved.



Allow to react for 5 minutes.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

7318

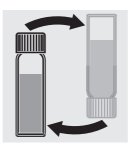


WTW model no.:	NO2-2 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.03 - 0.60 mg/l NO ₂ -N
	0.10 - 1.97 mg/l NO ₂
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 2.0 ml sample into a reaction cell.



Mix the contents by carefully swaying the cell.



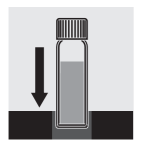
Add 1 level black measuring spoon of No. 8 **Nitrit 101** and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



Allow to react for 10 minutes.



Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- \bullet Store the reagents closed at a temperature of +4 ... +8 °C.

7317

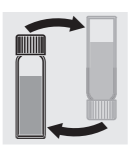


WTW model no.:	NO2-2 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.30 - 3.00 mg/l NO ₂ -N
	0.99 - 9.85 mg/l NO ₂
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 0.5 ml sample into a reaction cell.



Mix the contents by carefully swaying the cell.



Add 1 level black measuring spoon of No. 8 **Nitrit 101** and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



Allow to react for 10 minutes.

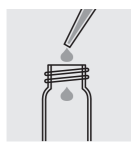


Insert the cell in the photometer cell shaft and start measurement.

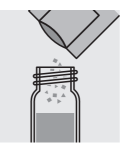
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- Store the reagents closed at a temperature of +4 ... +8 °C.



WTW model no.:	NO2-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.002 - 0.300 mg/l NO ₂ -N
	0.001 - 0.091 mg/l NO ₂
	Display in mmol/l possible



Pipette 10.0 ml of sample into the empty cell.



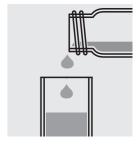
Add the contents of a **VARIO Nitri 3 F10** powder pack and close the cell with the screw cap.



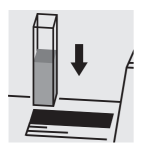
Shake the cell. Any undissolved powder does not adversely affect the measurement.



Allow to react for 15 minutes.



Fill the solution into the measuring cell.



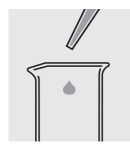
Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.



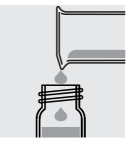
WTW model no.:	NO2-3 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.002 - 0.300 mg/l NO ₂ -N
	0.007 - 0.982 mg/l NO ₂
	Display in mmol/l possible



Pipette 25.0 ml of sample into an empty beaker.



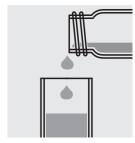
Add the contents of a Fill a VARIO Nitri 3 F25 ml prowder pack and close dissolve them by stirring. cap.



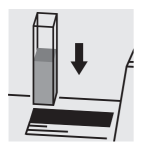
Fill an empty cell with the prepared sample and close it with the screw



Allow to react for 20 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.



WTW model no.:	Ntot2 TC (HR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	10 - 150 mg/l N



Put the contents of a
Total Nitrogen
Persulfate Rgt. powder
pack into a Total
Nitrogen Hydroxide HR
Tube digestion cell.



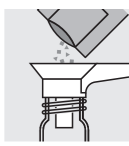
Add 0.5 ml sample with a pipette, close the cell with the screw cap and mix vigorously for at least 30 seconds. A small amount of solid matter may remain undissolved.



Heat the cell in the thermoreactor at 120 °C for 30 minutes.



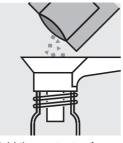
Remove the cell from the thermoreactor and let it cool down in a cell rack.



Add the contents of a VARIO Total Nitrogen Reagent A Powder pack. Close the cell with the screw cap and mix for at least 15 s.



Allow to react for 3 minutes.



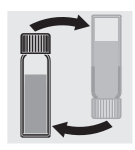
Add the contents of a **VARIO Total Nitrogen Reagent B** powder pack. Close the cell with the screw cap and mix for at least 15 s.



Allow to react for 2 minutes.



With a pipette add 2.0 ml of the prepared sample to a **Total Nitrogen Acid HR (Reagent C)** and close the cell with the screw cap.

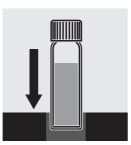


Mix the contents by carefully swaying the cell (10 x, i.e. for approx. 30 s).

Caution, cell grows hot!



Allow to react for 5 minutes.



Insert the cell in the photometer cell shaft and start measurement.

Note:

Clean the powder funnel thoroughly each time before adding the reagent!

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WTW model no.:	Ntot1 TC (LR)
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.5 - 25.0 mg/l N

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Put the contents of a

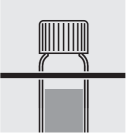
Total N Persulfate RGT
powder pack into a Total
Nitrogen Hydroxide LR
Tube digestion cell.



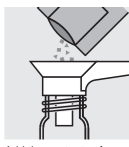
Add 2.0 ml sample with a pipette, close the cell with the screw cap and mix vigorously for at least 30 seconds. A small amount of solid matter may remain undissolved.



Heat the cell in the thermoreactor at 120 °C for 30 minutes.



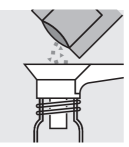
Remove the cell from the thermoreactor and let it cool down in a cell rack.



Add the contents of a VARIO Total Nitrogen Reagent A Powder pack. Close the cell with the screw cap and mix for at least 15 s.



Allow to react for 3 minutes.



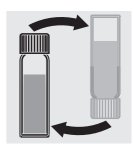
Add the contents of a VARIO Total Nitrogen Reagent B powder pack. Close the cell with the screw cap and mix for at least 15 s.



Allow to react for 2 minutes.



With a pipette add 2.0 ml of the prepared sample to a **Total Nitrogen Acid LR (Reagent C)** and close the cell with the screw cap.

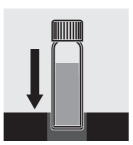


Mix the contents by carefully swaying the cell (10 x, i.e. for approx. 30 s).

Caution, cell grows hot!



Allow to react for 5 minutes.



Insert the cell in the photometer cell shaft and start measurement.

Note:

Clean the powder funnel thoroughly each time before adding the reagent!

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WTW model no.:	PO4-1 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.02 - 2.50 mg/l PO ₄
	0.007 - 0.800 mg/l PO ₄ -P
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 10.0 ml of sample into the empty cell.



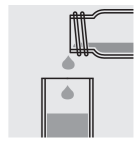
Add the contents of a VARIO Phosphate RGT F10 powder pack and close the cell with the screw cap.



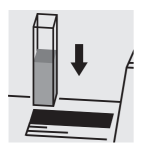
Shake the cell for 10 to 15 seconds. Any undissolved powder does not adversely affect the measurement.



Allow to react for 2 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

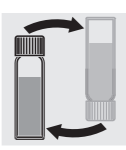


WTW model no.:	PO4-2 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.06 - 5.00 mg/l PO ₄
	0.02 - 1.63 mg/l PO ₄ -P
	Display in mmol/l possible

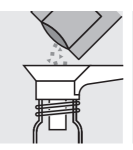
Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 5.0 ml of sample into a reaction cell and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



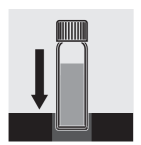
Add the contents of a VARIO Phosphate RGT F10 powder pack and close the cell with the screw cap.



To dissolve the solids, shake the cell for 10 to 15 seconds. A small amount of solid matter may remain undissolved.



Allow to react for 2 minutes.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

7316

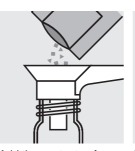


WTW model no.:	PO4-3 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.06 - 3.50 mg/l PO ₄
	0.020 - 1.141 mg/I PO ₄ -P
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 5.0 ml of sample into a reaction cell.



Add the contents of a VARIO Potassium Persulfate F10 ml powder pack and close the cell with the screw cap.



Shake the cell vigorously to dissolve solids.



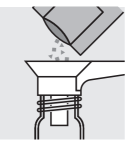
Heat the cell in the thermoreactor for 30 minutes at 120 °C.



Remove the cell from the thermoreactor and let it cool down in a cell rack.



With a pipette add 2.0 ml 1,54 N sodium hydroxide solution. Close the cell with the screw cap and mix the contents by carefully swaying the cell.



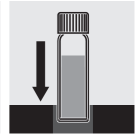
Add the contents of a VARIO Phosphate RGT F10 powder pack and close the cell with the screw cap.



To dissolve the solids, shake the cell for 10 to 15 seconds. A small amount of solid matter may remain undissolved.



Allow to react for 2 minutes.



Insert the cell in the photometer cell shaft and start measurement.

Notes:

• We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.

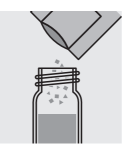


WTW model no.:	PO4-4 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.06 - 3.50 mg/l PO ₄
	0.020 - 1.141 mg/l PO ₄ -P
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



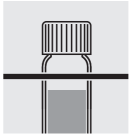
Pipette 5 ml of sample into a reaction cell.



Add the contents of a VARIO Potassium Persulfate F10 ml powder pack and close the cell with the screw cap.



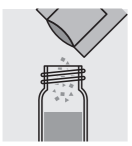
Heat the cell in the thermoreactor for 30 minutes at 120 °C.



Remove the cell from the thermoreactor and let it cool down in a cell rack.



With a pipette add 2.0 ml VARIO Sodium hydroxide 1.54N, close the cell with the screw cap and mix.



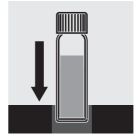
Add the contents of a VARIO Phosphate RGT F10 ml powder pack and close the cell with the screw cap.



Shake the cell for 10-15 s. A small amount of solid matter remains undissolved.



Allow to react for 2 minutes.



Within 8 minutes after the last reagent was added: Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- Clean all laboratory glassware with hydrochloric acid (approx. 20 %), then thoroughly rinse with deionized water. Do not used any detergents that contain phosphate!



WTW model no.:	PO4-4 TC
Category:	KT (reaction cell test)
Cell:	16 mm
Measuring range:	0.06 - 3.50 mg/l PO ₄
	0.020 - 1.141 mg/l PO ₄ -P
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 5 ml of sample into a reaction cell and close the cell with the screw cap.



Heat the cell in the thermoreactor for 30 minutes at 120 °C.



Remove the cell from the thermoreactor and let it cool down in a cell rack.



With a pipette add 2.0 ml VARIO Sodium hydroxide 1.00 N, close the cell with the screw cap and mix.



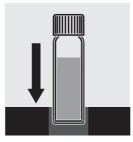
Add the contents of a VARIO Phosphate RGT F10 ml powder pack and close the cell with the screw cap.



Shake the cell for 10-15 s. A small amount of solid matter remains undissolved.



Allow to react for 2 minutes.



Within 8 minutes after the last reagent was added: Insert the cell in the photometer cell shaft and start measurement.

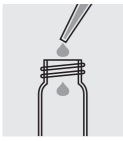
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- Clean all laboratory glassware with hydrochloric acid (approx. 20 %), then thoroughly rinse with deionized water. Do not used any detergents that contain phosphate!

7337

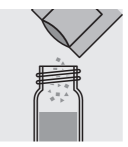


WTW model no.:	Si-3 TP (HR)
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	1 - 200 mg/l SiO ₂
	1 - 93 mg/l Si
	Display in mmol/l possible

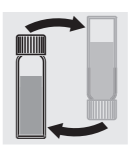
Note: Before using the test with your photometer for the first time, determine the reagent blank value.



Pipette 25.0 ml of sample into the empty cell.



Add the contents of a VARIO Silica HR Molybdate F25 powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



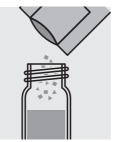
Add the contents of a VARIO Silica HR Acid Rgt F25 powder pack and close the cell with the screw cap.



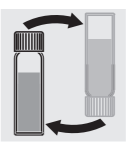
Mix the contents by carefully swaying the cell.



Allow to react for 10 minutes.



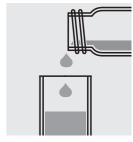
Add the contents of a VARIO Silica HR Citric Acid F25 powder pack and close the cell with the screw cap.



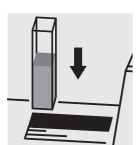
Mix the contents by carefully swaying the cell.



Allow to react for 2 minutes.



Fill the solution into the measuring cell.



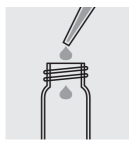
Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The temperature of the samples must be in the range 15 ... 25 °C.

7321



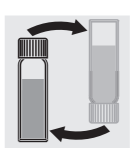
WTW model no.:	Si-1 TP (LR)
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	0.01 - 1.60 mg/l SiO ₂
	0.005- 0.748 mg/l Si
	Display in mmol/l possible



Pipette 10.0 ml of sample into the empty cell.



Add 15 drops of VARIO Molybdate 3 Reagent Solution and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



Allow to react for 4 minutes (temperature dependency, see note).



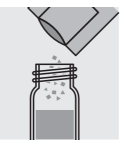
Add the contents of a VARIO Silica Citric Acid F10 powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



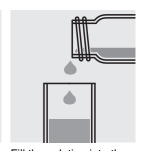
Allow to react for 1 minute (temperature dependency, see note).



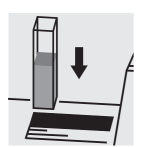
Add the contents of a VARIO Silica LR Amino Acid F F10 powder pack. Close the cell with the screw cap and mix.



Allow to react for 2 minutes. If SiO₂ is present in the sample the solution will turn blue.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

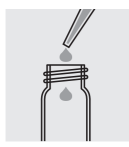
- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The reaction times mentioned above apply to a room temperature of 20 °C. At 10 °C the reaction time has to be doubled, at 30 °C reduced by half.

7308

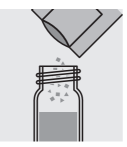


WTW model no.:	Si-2 TP (HR)
Category:	RS (reagent test)
Cell:	16 mm
Measuring range:	1 - 100 mg/l SiO ₂
	0.5 - 46.7 mg/l Si
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



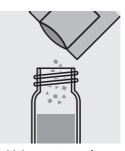
Pipette 10.0 ml of sample into the empty cell.



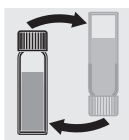
Add the contents of a VARIO Silica HR Molybdate F10 powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



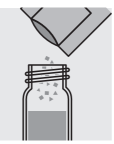
Add the contents of a VARIO Silica HR Acid Rgt F10 powder pack and close the cell with the screw cap.



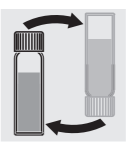
Mix the contents by carefully swaying the cell.



Allow to react for 10 minutes.



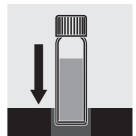
Add the contents of a VARIO Silica Citric Acid F10 powder pack and close the cell with the screw cap.



Mix the contents by carefully swaying the cell.



Allow to react for 2 minutes.



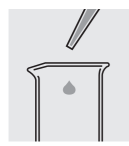
Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- The sample temperature has to be between 15 and 25 °C.

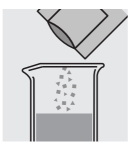


WTW model no.:	SO4-2 TP
Category:	RS (reagent test)
Cell:	20 mm
Measuring range:	2 - 70 mg/l SO ₄
	Display in mmol/l possible

Note: Before using the test with your photometer for the first time, determine the reagent blank value.



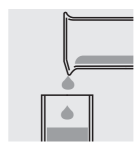
Pipette 25.0 ml of sample into an empty beaker.



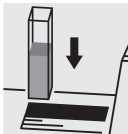
Add the contents of a **VARIO Sulfa 4** powder pack and dissolve them by stirring.



Allow to react for 5 minutes.



Fill the solution into the measuring cell.



Insert the cell in the photometer cell shaft and start measurement.

- We recommend to determine a new reagent blank value (deionized water instead of sample) for each test package started.
- If any sulfate is present, a white turbidity develops.
- Powder sedimented at the bottom does not affect the measurement result.

Appendix

Suitability of test kits for testing seawater

п	imit	of	t۸	lerance.	ealte	in %

Test kit	Model	Seawater	NaCl	NaNO ₃	Na ₂ SO ₄
Acid Capacity KT	1758	no	_	_	_
Aluminium KT	594	yes	20	20	20
Aluminium RT	14825	yes	10	20	20
Ammonium KT	14544	yes	20	15	20
Ammonium KT	14558	yes	20	10	15
Ammonium KT	14559	yes	20	20	20
Ammonium KT	14739	no	5	5	5
Ammonium KT	A6/25	yes	20	10	15
Ammonium RT	683	yes	20	20	20
Ammonium RT	14752	no *	10	10	20
AOX KT	675	no	0.4	20	20
Arsenic RT	1747	no	10	10	10
BOD KT	687	yes	20	20	20
Boron KT	826	yes	10	20	20
Boron RT	14839	no	20	5	20
Bromine RT	605	no	10	10	10
Cadmium KT	14834	no	1	10	1
Cadmium RT	1745	no	1	10	1
Calcium KT	858	no	2	2	1
Calcium RT	49	no	_	-	-
Calcium RT	14815	yes	20	20	10
Chloride KT	14730	yes	_	20	1
Chloride RT	14897	yes	_	10	0.1
Chlorine dioxide RT	608	no	10	10	10
Chlorine KT	595	no	10	10	10
Chlorine KT	597	no	10	10	10
Chlorine KT (liquid reagent) (free)	00086/00087	no	10	10	10
Chlorine KT (liquid reagent) (total)	00086/ 00087/00088	no	10	10	10
Chlorine RT	598	no	10	10	10
Chlorine RT	599	no	10	10	10
Chlorine RT	602	no	10	10	10
Chlorine RT (liquid reagent) (total)	00086/ 00087/00088	no	10	10	10
ChlorineTest (liquid reagent) (free)	00086/00087	no	10	10	10
Chromate KT	14552	yes	10	10	10
Chromate RT	14758	yes	10	10	10
Chromium total	14552	no	1	10	10
COD KT	1796	no	0.4	10	10
COD KT	1797	no	10	20	20
COD KT	14540	no	0.4	10	10
COD KT	14541	no	0.4	10	10
COD KT	14555	no	1.0	10	10
COD KT	14560	no	0.4	10	10
COD KT	14690	no	0.4	20	20
COD KT	14691	no	0.4	20	20
COD KT	14895	no	0.4	10	10

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Test lift Media Seawater NeCo Nation 104 10 10 COD KT C425 no 0.4 10				Lillit Of	iolei allice, sai	13 111 /0
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COD KT (Hg free) 9772 no 0 10 10 COD KT (Hg free) 9773 no 0 10 10 COpper RT 14553 yes 15 15 15 Copper RT 14767 yes 15 15 15 Cyanide KT 14561 no 10 10 10 Cyanide RT 19253 yes - - - - Fluoride RT 19253 yes - - - - - Fluoride RT 14598 yes 20 20 20 20 Fluoride RT 14598 yes 20 20 20 20 Formaldehyde RT 14678 no 5 0 10	COD KT	C3/25	no	0.4	10	10
COD KT (Hg free) 9773 no 0 10 10 Copper KT 14553 yes 15 15 15 Copper RT 14767 yes 15 15 15 Copper RT 14767 yes 15 15 15 Coparlor RT 14561 no 10 10 10 Cyanide RT 19253 yes - - - - Fluoride RT 19253 yes - - - - Fluoride RT 19588 yes 20 20 20 Formaldehyde KT 14500 no 5 0 10 Formaldehyde RT 14678 no 5 0 10 Gold BT 14821 yes 10 20 5 2 Hydragenperoxide RT 14731 yes 20 20 20 20 Hydrogenperoxide RT 18789 no 0.1 1 5	COD KT	C4/25	no	0.4	10	10
Copper KT 14553 yes 15 15 15 Copper RT 14767 yes 15 15 15 Cypanide KT 14561 no 10 10 10 Cyanide RT 19253 yes - - - Fluoride KT 14598 yes 20 20 20 Fluoride KT 14598 yes 20 20 20 Formaldehyde KT 14590 no 5 0 10 Formaldehyde RT 14678 no 5 0 10 Gold RT 14821 yes 10 20 5 Hydrogenperoxide KT 14731 yes 20 20 20 Hydrogenperoxide RT 18789 no 0.1 1 5 5 2 Hydrogenperoxide RT 18789 no 0.1 10 10 10 10 10 10 10 10 10 10 10	COD KT (Hg free)	9772	no	0	10	10
Copper RT 14767 yes 15 15 15 Cyanide KT 14561 no 10 10 10 Cyanide RT 9701 no 10 10 10 Cyanide RT 19253 yes — — — — Fluoride RT 14598 yes 20 20 20 Fluoride RT 14590 no 5 0 10 Formaldehyde KT 14600 no 5 0 10 Gold RT 14678 no 5 0 10 Gold RT 14731 yes 10 20 5 Hydrogenperoxide KT 14731 yes 20 20 20 Hydrogenperoxide RT 16789 no 0.1 1 5 Ion KT 14589 yes 20 20 20 Iron KT 14886 no 5 5 5 Iron KT 14761 yes<	COD KT (Hg free)	9773	no	0	10	10
Cyanide KT 14561 no 10 10 10 Cyanide RT 9701 no 10 10 10 Cyanuric acid RT 19253 yes - - - - Fluoride RT 1898 yes 20 20 20 Fluoride RT 14598 yes 20 20 20 Formaldehyde KT 14590 no 5 0 10 Gold RT 14678 no 5 0 10 Gold RT 14821 yes 10 20 5 Hydrogenperoxide KT 14731 yes 20 20 20 Hydrogenperoxide RT 18789 no 0.1 1 1 10 Iron KT 14549 yes 20 20 20 Iron KT 14589 no 5 5 5 Iron RT 14761 yes 20 20 20 Iron RT	Copper KT	14553	yes	15	15	15
Cyanide RT 9701 no 10 10 10 Cyanuide acid RT 19283 yes - - - Fluoride KT 809 no 10 10 10 Fluoride RT 14598 yes 20 20 20 Formaldehyde KT 14500 no 5 0 10 Formaldehyde RT 14678 no 5 0 10 Gold RT 14821 yes 10 20 5 2 Hydrogenperoxide KT 14731 yes 20 20 20 20 Hydrogenperoxide KT 14731 yes 20 20 20 20 Hydrogenperoxide KT 14731 yes 20 <td>Copper RT</td> <td>14767</td> <td>yes</td> <td>15</td> <td>15</td> <td>15</td>	Copper RT	14767	yes	15	15	15
Cyanuric acid RT 19253 yes - - - Fluoride RT 14598 yes 20 20 20 Fluoride RT 14598 yes 20 20 20 Formaldehyde KT 14678 no 5 0 10 Gold RT 14678 no 5 0 10 Hydrazine RT 19711 no 20 20 20 Hydrogenperoxide RT 14731 yes 20 20 20 Hydrogenperoxide RT 18789 no 0.1 1 5 Iodine RT 606 no 10 10 10 Iron KT 14549 yes 20 20 20 Iron RT 14761 yes 20 20 20 Iron RT 14761 yes 20 20 20 Lead KT 14333 no 20 20 1 Manganese RT 1846 no	Cyanide KT	14561	no	10	10	10
Fluoride KT 809 no 10 10 10 10 Fluoride RT 14598 yes 20 20 20 20 Formaldehyde KT 14500 no 5 0 10 Fluoride RT 14678 no 5 0 10 10 Gold RT 14821 yes 10 20 5 E Hydrazine RT 14678 no 20 5 2 E Hydragenperoxide KT 14731 yes 20 20 20 20 E Hydrogenperoxide KT 14731 yes 20 20 20 20 E Hydrogenperoxide RT 18789 no 0.1 1 5 5 E E E E E E E E E E E E E E E E	Cyanide RT	9701	no	10	10	10
Fluoride RT	Cyanuric acid RT	19253	yes	_	_	_
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Formaldehyde RT	Fluoride RT	14598	yes	20	20	20
Formaldehyde RT	Formaldehyde KT	14500	no	5	0	10
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Nitrogen (total) KT 14763 no 2 - 20 Oxygen KT 14694 no 10 5 1					-	
Oxygen KT 14694 no 10 5 1			no		-	
			no			
Ozone RT 607 no 10 10 10			no		5	
	Ozone RT	607	no	10	10	10

Limit of tolerance, salts in %

			2	ororanoo, cano	, , o
Test kit	Model	Seawater	NaCl	NaNO₃	Na ₂ SO ₄
pH KT	1744	yes	_	_	-
Phenol KT	14551	yes	20	20	15
Phenol RT	856	yes	20	20	20
Phosphate KT	616	yes	20	20	20
Phosphate KT	14543	yes	5	10	10
Phosphate KT	14546	yes	20	20	20
Phosphate KT	14729	yes	20	20	20
Phosphate KT	P6/25	yes	5	10	10
Phosphate KT	P7/25	yes	20	20	20
Phosphate RT	798	yes	15	20	10
Phosphate RT	14842	yes	20	20	20
Phosphate RT	14848	yes	5	10	10
Phosphorus total	14543	no	1	10	10
Phosphorus total	14729	yes	5	20	20
Phosphorus total	P6/25	no	1	10	10
Phosphorus total	P7/25	yes	5	20	20
Potassium KT	615	yes	20	20	20
Potassium KT	14562	yes	20	20	20
Residual Hardness KT	14683	no	0.01	0.01	0.01
Silicate (Silicic Acid) RT	857	no	5	10	02. Mai
Silicate (Silicic Acid) RT	14794	yes	5	10	5
Silver RT	14831	no	0	1	5
Sodium KT	885	no	_	10	1
Sulfate KT	617	yes	10	20	-
Sulfate KT	14548	yes	10	20	-
Sulfate KT	14564	yes	10	20	_
Sulfate RT	14791	no	0.2	0.2	_
Sulfide RT	14779	no	0.5	1	1
Sulfite KT	14394	no	20	20	20
Sulfite RT	1746	no	20	20	20
Surfactants (anionic) KT	14697	no	0.1	0.01	10
Surfactants (cationic) KT	1764	no	0.1	0.1	20
Surfactants (nonionic) KT	1787	no	2	5	2
Tin KT	14622	yes	20	20	20
TOC KT	14878	no	0.5	10	10
TOC KT	14879	no	5	20	20
Total Hardness KT	961	no	2	2	1
Volatile Organic Acids KT	1749	no	20	20	10
Volatile Organic Acids KT	1809	no	20	20	10
Zinc KT	861	no	20	20	1
Zinc KT	14566	no	10	10	10
			- •	· -	

¹⁾ This test kit is also suitable for testing seawater after the addition of sodium hydroxide solution (see package insert).

CombiCheck and standard solutions

Acid Capacity KT 1758 - OH 5.00 mmol/l* ± 0.50 mmol/l see preparation instr Aluminium KT 594 - Al 0.25 mg/l* ± 0.03 mg/l SL Al 19770 Aluminium RT 14825 CombiCheck 40, 14692 Al 0.75 mg/l ± 0.08 mg/l SL Al 19770 Ammonium KT 14544 CombiCheck 20, 14675 NH4-N 12.0 mg/l ± 1.0 mg/l SL NH4 19812 Ammonium KT 14558 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium KT 14739 CombiCheck 70, 14689 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 AMMONIUM RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 AMMONIUM RT 14752 CombiCheck 50, 14695 NH4-N<
Aluminium RT 14825 CombiCheck 40, 14692 Al 0.75 mg/l ± 0.08 mg/l SL Al 19770 Ammonium KT 14544 CombiCheck 20, 14675 NH4-N 12.0 mg/l ± 1.0 mg/l SL NH4 19812 Ammonium KT 14558 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium KT 14559 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium KT 14739 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium KT 14544 CombiCheck 20, 14675 NH4-N 12.0 mg/l ± 1.0 mg/l SL NH4 19812 Ammonium KT 14558 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium KT 14559 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium KT 14739 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium KT 14558 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium KT 14559 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium KT 14739 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium KT 14559 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium KT 14739 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium KT 14739 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812 Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium KT A6/25 CombiCheck 10, 14676 NH4-N 4.00 mg/l ± 0.30 mg/l SL NH4 19812 Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium RT 683 CombiCheck 70, 14689 NH4-N 50.0 mg/l ± 5.0 mg/l SL NH4 19812 Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
Ammonium RT 14752 CombiCheck 50, 14695 NH4-N 1.00 mg/l ± 0.10 mg/l SL NH4 19812
AOX KT 675 – AOX 1.00 mg/l* ± 0.10 mg/l AOX 00680
Arsenic RT 1747 – As 0.050 mg/l* ± 0.005 mg/l 19773 (Merck-No.)**
BOD KT 687 – O2 210 mg/l ± 20 mg/l BSB 00718
Boron KT 826 – B 1.00 mg/l* ± 0.15 mg/l SL B 19500
Boron RT 14839 – B 0.400 mg/l* ± 0.040 mg/l SL B 19500
Bromine RT 605 - Br2 5,00 mg/l* \pm 0.50 mg/l see preparation instr
Cadmium KT 14834 CombiCheck 30, 14677 Cd 0.500 mg/l ± 0.060 mg/l SL Cd 19777
Cadmium RT 1745 – Cd 0.250 mg/l ± 0.010 mg/l SL Cd 19777
Calcium KT 858 – Ca 75 mg/l* ± 7 mg/l SL Ca 19778
Calcium RT 49 – Ca 2.00 mg/l* ± 0.20 mg/l SL Ca 19778
Calcium RT 14815 – Ca 80 mg/l* ± 8 mg/l SL Ca 19778
Chloride KT 14730 CombiCheck 20, 14675 Cl 60 mg/l ± 10 mg/l SL Cl 19897
Chloride KT 14730 CombiCheck 10, 14676 Cl 25 mg/l ± 6 mg/l SL Cl 19897
Chloride RT 14897 CombiCheck 60, 14696 Cl 125 mg/l ± 13 mg/l SL Cl 19897
Chloride RT 14897 – Cl 12.5 mg/l* ± 0.13 mg/l SL Cl 19897
Chlorine Dioxide RT 608 – CIO2 5.00 mg/l* ± 0.50 mg/l see preparation instr
Chlorine KT 595 – Cl2 3.00 mg/l* ± 0.30 mg/l see preparation instr
Chlorine KT 597 – Cl2 3.00 mg/l^* $\pm 0.30 \text{ mg/l}$ see preparation instr
Chlorine KT (liquid 00086/ – Cl2 3.00 mg/l* \pm 0.30 mg/l see preparation instragent) = 00087 \pm 0.30 mg/l see preparation instragence \pm 0.30 mg/l see p
Chlorine KT (liquid 00086/ – Cl2 3.00 mg/l* \pm 0.30 mg/l see preparation instreagent) 00087/ 00088
Chlorine RT 598 – Cl2 3.00 mg/l^* $\pm 0.30 \text{ mg/l}$ see preparation instr
Chlorine RT 599 – Cl2 3.00 mg/l^* $\pm 0.30 \text{ mg/l}$ see preparation instr
Chlorine RT 602 – Cl2 3.00 mg/l^* $\pm 0.30 \text{ mg/l}$ see preparation instr
Chlorine RT (liquid $00086/$ – Cl2 0.500 mg/l^* $\pm 0.050 \text{ mg/l}$ see preparation instraction reagent) 00087
Chlorine RT (liquid 00086/ - Cl2 0.500 mg/l* \pm 0.050 mg/l see preparation instreagent) 00087/ 00088
Chromate KT 14552 – Cr 1.00 mg/l* ± 0.10 mg/l SL CrO3 19780
Chromate RT 14758 – Cr 1.00 mg/l* ± 0.10 mg/l SL CrO3 19780
COD KT 1796 CombiCheck 50, 14695 COD 20.0 mg/l ± 4.0 mg/l see preparation instr
COD KT 9772 CombiCheck 10, 14676 CSB 80 mg/l ± 12 mg/l see preparation instr
COD KT 9773 CombiCheck 20, 14675 CSB 750 mg/l ± 75 mg/l see preparation instr
COD KT 14540 CombiCheck 10, 14676 COD 80 mg/l ± 12 mg/l see preparation instr
COD KT 14541 CombiCheck 20, 14675 COD 750 mg/l ± 75 mg/l see preparation instr
COD KT 14555 CombiCheck 70, 14689 COD 5000 mg/l ± 400 mg/l see preparation instr

RTkit	Cat.No.	CombiCheck, Model	Evaluation as	Confidence interval Spec.value for the standard	Tolerance	Other standards** Model
COD KT	14560	CombiCheck 50, 14695	COD	20.0 mg/l	± 4.0 mg/l	see preparation instr.
COD KT	14690	CombiCheck 60, 14696	COD	250 mg/l	± 25 mg/l	see preparation instr.
COD KT	14691	CombiCheck 80, 14738	COD	1500 mg/l	± 150 mg/l	see preparation instr.
COD KT	14895	CombiCheck 60, 14696	COD	250 mg/l	± 20 mg/l	see preparation instr.
COD KT	C3/25	CombiCheck 10, 14676	COD	80 mg/l	± 12 mg/l	see preparation instr.
COD KT	C4/25	CombiCheck 20, 14675	COD	750 mg/l	± 75 mg/l	see preparation instr.
Copper KT	14553	CombiCheck 30, 14677	Cu	2.00 mg/l	± 0.20 mg/l	SL Cu 19786
Copper RT	14767	CombiCheck 30, 14677	Cu	2.00 mg/l	± 0.20 mg/l	SL Cu 19786
Cyanide KT	14561	_	CN	0.250 mg/l*	± 0.030 mg/l	19533 (Merck-No.)***
Cyanide RT	9701		CN	0.250 mg/l*	± 0.030 mg/l	19533 (Merck-No.)***
Cyanuric Acid RT	19253	_	Cyan Acid	80 mg/l*	± 10 mg/l	see preparation instr.
Fluoride KT	809	_	F	0.75 mg/l*	± 0.08 mg/l	SL F 19814
Fluoride RT	14598	_	F	1.00 mg/l*	± 0.15 mg/l	SL F 19814
Fluoride RT	14598	_	F	10.0 mg/l*	± 1.2 mg/l	SL F 19814
Formaldehyde KT	14500	_	НСНО	5.00 mg/l*	± 0.50 mg/l	see preparation instr.
Formaldehyde RT	14678	_	НСНО	4.50 mg/l*	± 0.50 mg/l	see preparation instr.
Gold RT	14821	_	Au	6.0 mg/l*	± 0.6 mg/l	70216 (Merck-No.)***
Hardness see Total F	lardness or	Residual Hardness				
Hydrazine RT	9711	_	N2H4	1.00 mg/l*	± 0.10 mg/l	see preparation instr.
Hydrogenperoxide KT	14731	-	H2O2	10.0 mg/l*	± 1.0 mg/l	see preparation instr.
Hydrogenperoxide RT	18789	-	H2O2	2.00 mg/l*	± 0.20 mg/l	see preparation instr.
Iodine RT	606	_	12	5.00 mg/l*	± 0.50 mg/l	see preparation instr.
Iron KT	14549	CombiCheck 30, 14677	Fe	1.00 mg/l	± 0.15 mg/l	SL Fe 19781
Iron KT	14896	_	Fe	25.0 mg/l*	± 2.5 mg/l	SL Fe 19781
Iron RT	796	CombiCheck 30, 14677	Fe	1.00 mg/l	± 0.15 mg/l	SL Fe 19781
Iron RT	14761	CombiCheck 30, 14677	Fe	1.00 mg/l	± 0.15 mg/l	SL Fe 19781
Lead KT	14833	CombiCheck 40, 14692	Pb	2.00 mg/l	± 0.20 mg/l	SL Pb 19776
Lead RT	9717	CombiCheck 40, 14692	Pb	2.00 mg/l	± 0.20 mg/l	SL Pb 19776
Magnesium KT	815	-	Mg	40.0 mg/l*	± 4.0 mg/l	see preparation instr.
Manganese KT	816	CombiCheck 30, 14677	Mn	1.00 mg/l	± 0.15 mg/l	SL Mn 19789
Manganese RT	1846	_	Mn	1.00 mg/l*	± 0.10 mg/l	SL Mn 19789
Manganese RT	14770	CombiCheck 30, 14677	Mn	1.00 mg/l	± 0.15 mg/l	SL Mn 19789
Molybdenum KT	860	_	Мо	0,50 mg/l*	± 0.05 mg/l	70227 (Merck-No.)***
Molybdenum RT	19252	_	Мо	25.0 mg/l*	± 2.5 mg/l	70227 (Merck-No.)***
Monochloramine RT	1632	_	Cl2	5.00 mg/l*	± 0.50 mg/l	see preparation instr.
Nickel KT	14554	CombiCheck 40, 14692	Ni	2.00 mg/l	± 0.20 mg/l	SL Ni 19792
Nickel RT	14785	CombiCheck 40, 14692		2.00 mg/l	± 0.20 mg/l	SL Ni 19792
Nitrat KT	614		NO3-N	100 mg/l*	± 10 mg/l	SL NO3 19811
Nitrate KT	14542	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
Nitrate KT	14556	CombiCheck 10, 14676	NO3-N	2.50 mg/l	± 0.25 mg/l	SL NO3 19811
Nitrate KT	14563	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
Nitrate KT	14764	CombiCheck 80, 14738	NO3-N	25.0 mg/l	± 2.5 mg/l	SL NO3 19811
Nitrate KT	N2/25	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
Nitrate RT	9713	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
Nitrate RT	14773	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
Nitrate RT	14942	CombiCheck 20, 14675	NO3-N	9.0 mg/l	± 0.9 mg/l	SL NO3 19811
	609	_	NO2-N	45.0 mg/l*	± 5 mg/l	SL NO2 19899
Nitrite KT					9,.	

- - CombiCheck 50, 14695		the standard		Model
- CombiCheck 50, 14695	NO2-N	0.300 mg/l*	± 0.030 mg/l	SL NO2 19899
CombiCheck 50, 14695	NO2-N	0.50 mg/l*	± 0.05 mg/l	SL NO2 19899
	N	5.0 mg/l	± 0.7 mg/l	see preparation instr.
CombiCheck 50, 14695	N	5.0 mg/l	± 0.7 mg/l	see preparation instr.
CombiCheck 70, 14689	N	50 mg/l	± 7 mg/l	see preparation instr.
-	02	-	± 0.6 mg/l	compare with O2-Sensor
-	DEHA	0,250 mg/l*	± 0,030 mg/l	s. Arbeitsvorschrift
-	O3	2.00 mg/l*	± 0.20 mg/l	see preparation instr.
-	рН	7.0	± 0.2	STP 7
-	C6H5OH	1.25 mg/l*	± 0.13 mg/l	see preparation instr.
-	C6H5OH	2.50 mg/l*	± 0.25 mg/l	see preparation instr.
-	PO4-P	50.0 mg/l*	± 5.0 mg/l	SL PO4 19898
CombiCheck 10, 14676	PO4-P	0.80 mg/l	± 0.08 mg/l	SL PO4 19898
-	PO4-P	15.0 mg/l*	± 1.0 mg/l	SL PO4 19898
CombiCheck 80, 14738	PO4-P	15.0 mg/l	± 1.0 mg/l	SL PO4 19898
CombiCheck 20, 14675	PO4-P	8.0 mg/l	± 0.7 mg/l	SL PO4 19898
CombiCheck 10, 14676	PO4-P	0.80 mg/l	± 0.08 mg/l	SL PO4 19898
CombiCheck 80, 14738	PO4-P	15.0 mg/l	± 1.0 mg/l	SL PO4 19898
CombiCheck 20, 14675	PO4-P	8.0 mg/l	± 0.7 mg/l	SL PO4 19898
_	PO4-P	50.0 mg/l*	± 5.0 mg/l	SL PO4 19898
_	PO4-P	15.0 mg/l*	± 1.0 mg/l	SL PO4 19898
CombiCheck 10, 14676	PO4-P	0.80 mg/l	± 0.08 mg/l	SL PO4 19898
_	K	150 mg/l*	± 15 mg/l	SL K 70230
_	K	25.0 mg/l*	± 4.0 mg/l	SL K 70230
-	Ca	2.50 mg/l*	± 0.30 mg/l	SL Ca 19778
-	Si	25.0 mg/l*	± 2.5 mg/l	SL Si 70236
_	Si	2.50 mg/l*	± 0.25 mg/l	SL Si 70236
_		0.375 mg/l*	± 0.040 mg/l	SL Si 70236
_	Ag	1.50 mg/l*	± 0.20 mg/l	SL Ag 19797
_	Na	100 mg/l*	± 10 mg/l	see preparation instr.
CombiCheck 10, 14676	SO4	100 mg/l	± 15 mg/l	SL SO4 19813
CombiCheck 10, 14676	SO4	100 mg/l	± 15 mg/l	SL SO4 19813
CombiCheck 20, 14675	SO4	500 mg/l	± 75 mg/l	SL SO4 19813
CombiCheck 10, 14676	SO4	100 mg/l	± 15 mg/l	SL SO4 19813
_	S	0.75 mg/l*	± 0.08 mg/l	see preparation instr.
_	SO3	12.5 mg/l*	± 1.5 mg/l	see preparation instr.
_	SO3	30.0 mg/l*	± 1.0 mg/l	see preparation instr.
_	MBAS	1.00 mg/l*	± 0.20 mg/l	see preparation instr.
	k-Ten	1.00 mg/l*	± 0.10 mg/l	see preparation instr.
_				see preparation instr.
_				70242 (Merck-No.)***
_				SL TOC 09017 SL TOC 09017
	- -	- n-Ten - Sn - TOC	- n-Ten 4.00 mg/l* - Sn 1.25 mg/l* - TOC 40.0 mg/l*	- n-Ten 4.00 mg/l* ± 0.40 mg/l - Sn 1.25 mg/l* ± 0.13 mg/l - TOC 40.0 mg/l* ± 3.0 mg/l

RTkit	Cat.No.	CombiCheck, Model	Evaluation as	Confidence interval Spec.value for the standard	Tolerance	Other standards** Model
Total Hardnes KT	961	_	Ca	75 mg/l*	± 7 mg/l	SL Ca 19778
Volatile Organic Acids KT	1749	_	СНЗСООН	1500 mg/l*	± 80 mg/l	see preparation instr.
Volatile Organic Acids KT	1809	_	СНЗСООН	1500 mg/l*	± 80 mg/l	see preparation instr.
Zinc KT	861	_	Zn	0.500 mg/l*	± 0.050 mg/l	SL Zn 19806
Zinc KT	14566	CombiCheck 40, 14692	Zn	2.00 mg/l	± 0.40 mg/l	SL Zn 19806
Zinc RT	14832	_	Zn	1.25 mg/l*	± 0.20 mg/l	SL Zn 19806

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 $^{^{\}star}$ Self prepared, recommended concentration ** c = 1000 $\,$ mg/l analyte *** The reagents are available from Merck under the stated number.

Instructions for the preparation of standard solutions

Standard solution of acid capacity

Preparation of a standard solution:

A sodium hydroxide solution of 0.1 mol/l (corresponds to 100 mmol/l) is used. 1.16754.9010 Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

* The reagents

Stability:

When stored in a cool place (refrigerator), the diluted investigational solutions remain stable for one week.

Reagents required:*

1.09141.1000 Sodium hydroxide solution 0.1 mol/l

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solution of bromine analogous to DIN EN ISO 7393 Reagents required:*

Preparation of a KIO₃ stock solution:

Dissolve 1.006 g of KIO₃ in 250 ml of distilled water in a calibrated or conformity-checked 1000-ml volumetric flask. Subsequently make up to the mark with distilled water.

1.05043.0250
1.09072.1000

Preparation of a KIO₃/KI standard solution:

Transfer 11.13 ml of the KIO₃ stock solution to a calibrated or conformity-che- 1.16754.9010 cked 1000-ml volumetric flask, add approx. 1 g of Kl and make up to the mark with distilled water.

1 ml of this solution is equivalent to 0.025 mg of bromine.

Preparation of the bromine standard solution:

Pipette 20.0 ml (full pipette) KlO $_3$ /Kl standard solution into a calibrated or conformity-checked 100-ml volumetric flask, add 2.0 ml of H $_2$ SO $_4$ 0.5 mol/l, leave to stand for 1 min, and then add NaOH 2 mol/l dropwise (approx. 1 ml) until the solution just loses its colour. Subsequently make up the solution to the mark with distilled water.

The concentration of the solution is 5.00 mg/l bromine.

Stability:

The KIO₃ stock solution remains stable for 4 weeks when stored in a cool place (refrigerator). The KIO₃/KI standard solution can be used for 5 hours when stored in a cool place (refrigerator). The dilute bromine standard solution is not stable and must be used immediately.

volum. standard

1.05043.0250 Potassium iodide GR for analysis

1.09072.1000 Sulfuric acid 0.5 mol/l

1.09136.1000 Sodium hydroxide solution 2 mol/l

1.16754.9010 Water GR for analy-

1.02404.0100 Potassium iodate.

* The reagents are available from Merck under the stated number.

Standard solution of calcium

Preparation of a standard solution:

Dissolve 2.946 g of calcium nitrate tetrahydrate GR with distilled water in a calibrated or conformity-checked 500-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l calcium.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

The standard solution of 1000 mg/l remains stable for one week. The diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

1.02121.0500 Calcium nitrate

tetrahydrate GR for

analysis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solutions of free chlorine

All standard solutions described here for free chlorine yield equivalent results and are identically suited for the determination of chlorine.

Standard solution of free chlorine

Preparation of a standard solution:

Dissolve 1.85 g of dichloroisocyanuric acid sodium salt dihydrate GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l free chlorine.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/l and the diluted standard solutions (investigational concentrations) remain stable for one day.

Note:

This is a standard solution that can be prepared particularly rapidly and easily.

Reagents required:*

1.10888.0250 Dichloroisocyanuric

acid sodium salt dihydrate GR for analysis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solution of free chlorine analogous to DIN EN ISO Reagents required:*

Preparation of a KIO₃ stock solution:

Dissolve 1.006 g of KIO₃ in 250 ml of distilled water in a calibrated or conformity-checked 1000-ml volumetric flask. Subsequently make up to the mark with distilled water.

Preparation of a KIO₃/KI standard solution:

Transfer 15.00 ml (5.00 ml) of the KIO₃ stock solution to a calibrated or conformity-checked 1000-ml volumetric flask, add approx. 1 g of Kl and make up * The reagents are available from to the mark with distilled water.

1 ml of this solution is equivalent to 0.015 mg (0.005 mg) of free chlorine.

Preparation of the chlorine standard solution:

Pipette 20.0 ml (10.0 ml) (full pipette) KIO₃/KI standard solution into a calibrated or conformity-checked 100-ml volumetric flask, add 2.0 ml of H₂SO₄ 0.5 mol/l, leave to stand for 1 min, and then add NaOH 2 mol/l dropwise (approx. 1 ml) until the solution just loses its colour. Subsequently make up the solution to the mark with distilled water.

The concentration of the solution is 3.00 mg/l (0.500 mg/l) free chlorine.

Stability:

The KIO3 stock solution remains stable for 4 weeks when stored in a cool place (refrigerator). The KIO₃/KI standard solution can be used for 5 hours when stored in a cool place (refrigerator). The dilute chlorine standard solution is not stable and must be used immediately.

Note:

This procedure involves the preparation according to a standardized method.

	1.02404.0100	Potassium iodate, volum. standard
_	1.05043.0250	Potassium iodide GR for analysis
	1.09072.1000	Sulfuric acid 0.5 mol/l
	1.09136.1000	Sodium hydroxide solution 2 mol/l
	1.16754.9010	Water GR for analy-

Merck under the stated number.

sis

Standard solution of free chlorine

Preparation of a stock solution:

First prepare a 1:10 dilution using a sodium hypochlorite solution containing approx. 13% of active chlorine. For this pipette 10 ml of sodium hypochlorite solution into a calibrated or conformity-checked 100-ml volumetric flask and then make up to the mark with distilled water.

Precise assay of the stock solution:

Pipette 10.0 ml of the stock solution into a 250-ml ground-glassstoppered conical flask containing 60 ml of distilled water. Subsequently add to this solution 5 ml of hydrochloric acid 25% GR and 3 g of potassium iodide. Close the conical flask with the ground-glass stopper, mix thoroughly, and leave to stand for 1 min.

Titrate the eliminated iodine with sodium thiosulfate solution 0.1 mol/l until a weakly yellow colour emerges. Add 2 ml of zinc iodide-starch solution and titrate from blue to colourless.

Calculation and preparation of a standard solution:

1 ml sodium thiosulfate solution = 3.55 mg free chlorine

Further investigational concentrations may be prepared from the stock solution prepared according to the procedure described above by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), a standard solution remains stable for approx. one week. The diluted standard solutions (investigational concentrations) are stable for approx. 2 hours.

Note:

This is a standard solution that is <u>absolutely</u> necessary for the preparation of the monochloramine standard.

Standard solution of total chlorine

Preparation of a standard solution:

Dissolve 4.00 g of chloramine T trihydrate GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l total chlorine.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/ I and the diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

	_	-
	1.00316.1000	Hydrochloric acid 25 % GR for analysis
)	1.05614.9025	Sodium hypochlorite solution techn. approx. 13% active chlorine
	1.09147.1000	Sodium thiosulfate solution 0.1 mol/l
	1.05043.0250	Potassium iodide GR for analysis
)	1.05445.0500	Zinc iodidestarch solution GR for ana- lysis
	1.16754.9010	Water GR for analy-

The reagents are available from Merck under the stated number.

sis

Reagents required:*

1.02426.0250	Chloramine T trihydrate GR for analysis
1.16754.9010	Water GR for analysis

* The reagents are available from Merck under the stated number.

Standard solution of chlorine dioxide analogous to DIN EN Reagents required:* **ISO 7393**

Preparation of a KIO₃ stock solution:

Dissolve 1.006 g of KIO₃ in 250 ml of distilled water in a calibrated or conformity-checked 1000-ml volumetric flask. Subsequently make up to the mark with distilled water.

Preparation of a KIO₃/KI standard solution:

Transfer 13.12 ml of the KIO₃ stock solution to a calibrated or conformity-checked 1000-ml volumetric flask, add approx. 1 g of Kl and make up to the mark with distilled water.

1 ml of this solution is equivalent to 0.025 mg of chlorine dioxide.

Preparation of the chlorine dioxide standard solution:

Pipette 20.0 ml (full pipette) KIO₃/KI standard solution into a calibrated or conformity-checked 100-ml volumetric flask, add 2.0 ml of $H_2SO_4\,0.5\,mol/l$, leave to stand for 1 min, and then add NaOH 2 mol/l dropwise (approx. 1 ml) until the solution just loses its colour. Subsequently make up the solution to the mark with distilled water.

The concentration of the solution is 5.00 mg/l chlorine dioxide.

Stability:

The KIO₃ stock solution remains stable for 4 weeks when stored in a cool place (refrigerator). The KIO₃/KI standard solution can be used for 5 hours when stored in a cool place (refrigerator). The dilute chlorine dioxide standard solution is not stable and must be used immediately.

	1.02404.0100	Potassium iodate, volum. standard
_	1.05043.0250	Potassium iodide GR for analysis
	1.09072.1000	Sulfuric acid 0.5 mol/l
	1.09136.1000	Sodium hydroxide solution 2 mol/l
	1.16754.9010	Water GR for analy-

The reagents are available from Merck under the stated number.

sis

Standard solution of COD

Preparation of a standard solution:

Dissolve 0.850 g of potassium hydrogen phthalate GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/I COD.

Further investigational concentrations may be prepared from this stock solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution remains stable for one month. When stored under appropriate cool conditions (refrigerator), the diluted standard solutions (investigational concentrations) remain stable - depending on the respective concentration - for approx. one week to one month.

1.02400.0080 Potassium hydrogen

phthalate GR for analysis, volum. standard

1.16754.9010 Water GR for analy-

* The reagents are available from Merck under the stated number.

Standard solution of cyanuric acid

Preparation of a standard solution:

Dissolve 1.00 g of cyanuric acid with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water. The substance is slightly soluble and the dissolution process may take several hours.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l cyanuric acid.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/ I and the diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

8.20358.0005 Cyanuric acid for synthesis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solution of formaldehyde

Preparation of a stock solution:

In a calibrated or conformity-checked 1000-ml volumetric flask make up 2.50 ml of formaldehyde solution min. 37% GR to the mark with distilled water. The stock solution prepared according to this procedure has a concentration of approx. 1000 mg/l formaldehyde.

Precise assay of the stock solution:

Pipette 40.0 ml (full pipette) of the formaldehyde stock solution into a 300-ml ground-glass conical flask and add 50.0 ml (buret) of iodine solution 0.05 mol l and 20 ml of sodium hydroxide solution 1 mol/l.

Leave to stand for 15 minutes and subsequently add 8 ml of sulfuric acid 25 % GR. Subsequently titrate with sodium thiosulfate solution 0.1 mol/l until the yellow iodine colour has disappeared, add 1 ml of zinc iodide-starch solution, and continue to titrate until a milky, pure white colour emerge.

Calculation and preparation of a standard solution:

C1 = consumption of sodium thiosulfate solution 0.1 mol/l

C2 = quantity of iodine solution 0.05 mol/l (50,0 ml)

mg/l formaldehyde = $(C2 - C1) \times 37.525$

Further investigational concentrations may be prepared from the stock solution exactly determined according to the procedure described above by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the stock solution of approx. 1000 mg/l remains stable for one week. After this time, the stock solution must be determined anew. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Reagents required:*

)	1.04003.1000	Formaldehyde solution min. 37% GR for analysis
•	1.09099.1000	lodine solution 0.05 mol/l
•	1.09147.1000	Sodium thiosulfate solution 0.1 mol/l
	1.09137.1000	Sodium hydroxide solution 1 mol/l
I /	1.00716.1000	Sulfuric acid 25% GR for analysis
5 e	1.05445.0500	Zinc iodidestarch solution GR for ana- lysis
,	1.16754.9010	Water GR for analysis

* The reagents are available from Merck under the stated number.

Standard solution of hydrazine

Preparation of a standard solution:

Dissolve 4.07 g of hydrazinium sulfate GR with oxygen-low (boil previously) distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with oxygen-low distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l hydrazine.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with oxygen-low distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/ I and the diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

1.04603.0100 Hydrazinium sulfate GR for analysis

1.16754.9010 Water GR for analy-

* The reagents are available from Merck under the stated number.

Standard solution of hydrogen peroxide

Preparation of a stock solution:

Place 10.0 ml of Perhydrol® 30% H₂O₂ GR in a calibrated or conformity-checked 100-ml volumetric flask and make up to the mark with distilled water. Transfer 30.0 ml (full pipette) of this solution to a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water. The stock solution prepared according to this procedure has a concentration of approx. 1000 mg/l hydrogen peroxide.

Precise assay of the stock solution:

Pipette 50.0 ml (full pipette) of the hydrogen peroxide stock solution into a 500-ml conical flask, dilute with 200 ml of distilled water, and add 30 ml of sulfuric acid 25% GR.

Titrate with a 0.02 mol/l potassium permanganate solution until the colour changes to pink.

Calculation and preparation of a standard solution:

Consumption of potassium permanganate (ml) x 34.02 = content of hydrogen peroxide, in mg/l

Further investigational concentrations may be prepared from the stock solution exactly determined according to the procedure described above by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the stock solution of approx. 1000 mg/l and the diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

1.09122.1000 Potassium permanganate solution 0.02

mol/l

1.07209.0250 Perhydrol® 30 % GR

for analysis 1.00716.1000Sulfuric acid 25% GR for analysis

1.16754.9010 Water GR for analy-

SİS

* The reagents are available from Merck under the stated number.

Standard solution of iodine analogous to DIN EN ISO 7393

Preparation of a KIO₃ stock solution:

Dissolve 1.006 g of KIO₃ in 250 ml of distilled water in a calibrated or conformity-checked 1000-ml volumetric flask. Subsequently make up to the mark with distilled water.

1.05043.0250
1.09072.1000

Preparation of a KIO₃/KI standard solution:

Transfer 7.00 ml of the KIO_3 stock solution to a calibrated or conformity-che- 1.16754.9010 cked 1000-ml volumetric flask, add approx. 1 g of Kl and make up to the mark with distilled water.

1 ml of this solution is equivalent to 0.025 mg of iodine.

Preparation of the iodine standard solution:

Pipette 20.0 ml (full pipette) KlO $_3$ /Kl standard solution into a calibrated or conformity-checked 100-ml volumetric flask, add 2.0 ml of H $_2$ SO $_4$ 0.5 mol/l, leave to stand for 1 min, and then add NaOH 2 mol/l dropwise (approx. 1 ml) until the solution just loses its colour. Subsequently make up the solution to the mark with distilled water.

The concentration of the solution is 5.00 mg/l iodine.

Stability:

The KIO₃ stock solution remains stable for 4 weeks when stored in a cool place (refrigerator). The KIO₃/KI standard solution can be used for 5 hours when stored in a cool place (refrigerator). The dilute iodine standard solution is not stable and must be used <u>immediately</u>.

Standard solution of magnesium

Preparation of a standard solution:

Dissolve 1.055 g of magnesium nitrate hexahydrate GR with distilled water in a calibrated or conformity-checked 100-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l magnesium.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

The standard solution of 1000 mg/l remains stable for one week. The diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

1.02404.0100 Potassium iodate, volum. standard

1.05043.0250 Potassium iodide GR for analysis

1.09072.1000 Sulfuric acid 0.5 mol/l

1.09136.1000 Sodium hydroxide solution 2 mol/l

.16754.9010 Water GR for analysis

* The reagents are available from Merck under the stated number.

Reagents required:*

1.05853.0500 Magnesium nitrate

hexahydrate GR for

analysis

1.16754.9010 Water GR for analy-

SIS

* The reagents are available from Merck under the stated number.

Standard solution of monochloramine

Preparation of a standard solution:

Place 5.0 ml of chlorine standard solution 100 mg/l Cl_2 and 10.0 ml ammonium standard solution 10 mg/l NH_4 -N in a calibrated or conformity-checked 100-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 5.00 mg/l Cl₂ or 3.63 mg/l NH₂Cl.

Stability:

The standard solution is not stable and must be used immediately.

Reagents required:*

Chlorine standard solution 100 mg/l Cl₂

Preparation see "Standard solution of free chlorine" with hypochlorite solution (standard solution that is <u>absolutely</u> necessary for the preparation of the monochloramine standard)

Ammonium standard solution 10 mg/l NH₄-N

Preparation with Ammonium standard solution Certipur $^{\oplus}$, Cat.No. 1.19812.0500, 1000 mg/l NH₄ = = 777 mg/l NH₄-N

- 1.16754.9010 Water GR for analysis
- * The reagents are available from Merck under the stated number.

Standard solution of nitrogen (total)

Preparation of a standard solution:

Dissolve 5.36 g of glycine GR with distilled water in a calibrated or con formity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a con centration of 1000 mg/l total nitrogen.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/l remains stable for one week. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Reagents required:*

1.04201.0100 Glycine GR for analysis

1.16754.9010 Water GR for analysis

The reagents are available from Merck under the stated number.

Standard solution of oxygenscavengers

Preparation of a standard solution:

Dissolve 1.00 g of N,N-diethylhydroxylamine with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l N,N-diethylhydroxylamine (DEHA).

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/l and the diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

8.18473.0050 N,N-Diethylhydroxylamine for syn-

thesis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solution of ozone analogous to DIN EN ISO 7393

Preparation of a KIO₃ stock solution:

Dissolve 1.006 g of KIO₃ in 250 ml of distilled water in a calibrated or confor- 1.05043.0250 mity-checked 1000-ml volumetric flask. Subsequently make up to the mark with distilled water.

Preparation of a KIO₃/KI standard solution:

Transfer 14.80 ml of the KIO₃ stock solution to a calibrated or conformity-che- 1.16754.9010 cked 1000-ml volumetric flask, add approx. 1 g of Kl and make up to the mark with distilled water.

1 ml of this solution is equivalent to 0.010 mg of ozone.

Preparation of the ozone standard solution:

Pipette 20.0 ml (full pipette) KIO₃/KI standard solution into a calibrated or conformity-checked 100-ml volumetric flask, add 2.0 ml of H₂SO₄0.5 mol/l, leave to stand for 1 min, and then add NaOH 2 mol/l dropwise (approx. 1 ml) until the solution just loses its colour. Subsequently make up the solution to the mark with distilled water.

The concentration of the solution is 2.00 mg/l ozone.

Stability:

The KIO₃ stock solution remains stable for 4 weeks when stored in a cool place (refrigerator). The KIO₃/KI standard solution can be used for 5 hours when stored in a cool place (refrigerator). The dilute ozone standard solution is not stable and must be used immediately.

Standard solution of phenol

Preparation of a standard solution:

Dissolve 1.00 g of phenol GR with distilled water in a calibrated or conformitychecked 1000-ml volumetric flask and make up to the mark with distilled water

The standard solution prepared according to this procedure has a concentration of 1000 mg/l phenol.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with disilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/ I remains stable for one week. The diluted standard solutions (investigational concentrations) must be used immediately.

Reagents required:*

1.02404.0100 Potassium iodate, volum. standard

Potassium iodide GR

for analysis

Water GR for analy-

1.09072.1000 Sulfuric acid 0.5 mol/l 1.09136.1000 Sodium hydroxide

solution 2 mol/l

* The reagents are available from Merck under the stated number.

Reagents required:*

1.00206.0250 Phenol GR for analy-

1.16754.9010 Water GR for analy-

* The reagents are available from Merck under the stated number.

Standard solution of sodium

Preparation of a standard solution:

A chloride standard solution of 1000 mg/l is used.

1000 mg/l chloride corresponds to 649 mg/l sodium.

Further investigational concentrations may be prepared by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the diluted standard solutions (investigational concentrations) remain stable for one month.

Reagents required:*

1.19897.0500 Chloride standard

solution Certipur®

1.16754.9010 Water GR for analy-

sis

The reagents are available from Merck under the stated number.

Standard solution of sulfide	Reagents required:*		
Preparation of a stock solution:	1.06657.0500	Sodium sulfide hyd- rate GR for <u>analysis</u>	
Dissolve 7.2 g of glass-clear, if necessary washed crystals of sodium sulfide hydrate GR with distilled water in a calibrated or conformitychecked 1000-ml	1.09099.1000	lodine solution 0.05mol/l	
volumetric flask and make up to the mark with distilled water.	1.09147.1000	Sodium thiosulfate	
The stock solution prepared according to this procedure has a concentration		solution 0.1 mol/l	
of approx. 1000 mg/l sulfide.	1.00716.1000	Sulfuric acid 25% GR for analysis	
Precise assay of the stock solution:	1.05445.0500	Zinc iodidestarch	
Place 100 ml of distilled water and 5.0 ml (full pipette) of sulfuric acid		solution GR for ana- lysis	
25% GR in a 500-ml ground-glass-stoppered conical flask. To this solution add 25.0 ml (full pipette) of the sulfide stock solution and 25.0 ml (full pipette) of iodine solution 0.05 mol/l. Shake the contents of the flask	1.16754.9010	Water GR for analysis	
thoroughly for about 1 minute, subsequently titrate with sodium thiosulfate solution 0.1 mol/l until the yellow iodine colour has disappeared, add 1 ml of		are available from ne stated number.	

Calculation and preparation of the standard solution:

C1 = consumption of sodium thiosulfate 0.1 mol/l

C2 = quantity of iodine solution 0.05 mol/l (25.0 ml)

 $mg/l \ sulfide = (C2 - C1) \times 64.13$

Further investigational concentrations may be prepared from the stock solution exactly determined according to the procedure described above by diluting accordingly with distilled water.

zinc iodide-starch solution, and continue to titrate until a milky, pure white

Stability:

colour emerges.

When stored in a cool place (refrigerator), the stock solution of approx. 1000 mg/l remains stable for at most one day. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Preparation of a stock solution: Dissolve 1.57 g of sodium sulfite GR and 0.4 g of Titriplex® III GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of approx. 1000 mg/l sulfite.

Precise assay of the stock solution:

Place 50.0 ml (full pipette) of the sulfite stock solution and 5.0 ml (full pipette) of hydrochloric acid 25 % GR in a 300-ml conical flask.

To this solution add 25.0 ml (full pipette) of iodine solution 0.05 mol/l and process <u>immediately</u>. After mixing the contents of the flask, subsequently titrate with sodium thio-sulfate solution 0.1 mol/l until the yellow iodine colour has disappeared, add 1 ml of zinc iodide-starch solution, and continue to titrate from blue to colourless.

Calculation and preparation of the standard solution:

C1 = consumption of sodium thiosulfate 0.1 mol/l C2 = quantity of iodine solution 0.05 mol/l (25.0 ml)

 $mg/l \ sulfite = (C2 - C1) \times 80.06$

Further investigational concentrations may be prepared from the stock solution exactly determined according to the pro-cedure described above by diluting accordingly with distilled water and buffer solution pH 9.00. This is done in the following manner:

Withdraw the desired aliquot from the stock solution, place in a calibrated or conformity-approved 1000-ml volumetric flask, add 20 ml of buffer solution pH 9.00, make up to the mark with distilled water, and mix.

Stability:

When stored in a cool place (refrigerator), the stock solution of approx. 1000 mg/l remains stable for at most one day. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Standard solution of surfactants (anionic)

Preparation of a standard solution:

Dissolve 1.00 g of sodium 1-dodecanesulfonate with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l anionic surfactants.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/ I remains stable for one month. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Reagents required:*

	1.06657.0500	Sodium sulfite anhydrous GR for analysis
	1.08418.0100	Titriplex [®] III GR for analysis
	1.09099.1000	lodine solution 0.05mol/l
	1.09147.1000	Sodium thiosulfate solution 0.1 mol/l
	1.00316.1000	Hydrochloric acid 25 % GR for analysis
-	1.05445.0500	Zinc iodidestarch solution GR for ana- lysis
	1.09461.1000	Buffer solution pH 9.00 Certipur®
	1.16754.9010	Water GR for analy-

^{*} The reagents are available from Merck under the stated number.

sis

Reagents required:*

1.12146.0005	Sodium 1-dodecane- sulfonate
1.16754.9010	Water GR for analysis

The reagents are available from Merck under the stated number.

Standard solution of surfactants (cationic)

Preparation of a standard solution:

Dissolve 1.00 g of N-cetyl-N,N,N-trimethyl-ammonium bromide GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l cat-ionic surfactants.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/I remains stable for one month. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Reagents required:*

1.02342.0100 N-cetyl-N,N,Ntri-

methylammonium bromide GR for ana-

lysis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Standard solution of surfactants (nonionic)

Preparation of a standard solution:

Dissolve 1.00 g of Triton® X-100 with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1000 mg/l non-ionic surfactants.

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

When stored in a cool place (refrigerator), the standard solution of 1000 mg/l remains stable for one week. The diluted standard solutions (investigational concentrations) must be used <u>immediately</u>.

Reagents required:*

1.12298.0101 Triton® X-100 1.16754.9010 Water GR for analy-

.

* The reagents are available from Merck under the stated number.

Standard solution of total hardness

Preparation of a standard solution:

Dissolve 2.946 g of calcium nitrate tetrahydrate GR with distilled water in a calibrated or conformity-checked 500-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of $1000 \, \text{mg/l}$ calcium (corresponds to $175 \, ^{\circ}\text{e}$).

Further investigational concentrations may be prepared from this standard solution by diluting accordingly with distilled water.

Stability:

The standard solution of 1000 mg/l remains stable for one week. The diluted standard solutions (investigational concentrations) remain stable for one day.

Reagents required:*

1.02121.0500 Calcium nitrate

tetrahydrate GR for analysis

1.16754.9010 Water GR for analy-

SIS

* The reagents are available from Merck under the stated number.

Standard solution of volatile organic acids

Preparation of a standard solution:

Dissolve 2,05 g of sodium acetate anhydrous GR with distilled water in a calibrated or conformity-checked 1000-ml volumetric flask and make up to the mark with distilled water.

The standard solution prepared according to this procedure has a concentration of 1500 mg/l acetic acid.

Reagents required:*

1.06268.0250 Sodium acetate

anhydrous GR for

analysis

1.16754.9010 Water GR for analy-

sis

* The reagents are available from Merck under the stated number.

Stability:

When stored in a cool place (refrigerator), the standard solution remains stable for one week.

What can Xylem do for you?

We're a global team unified in a common purpose: creating innovative solutions to meet our world's water needs. Developing new technologies that will improve the way water is used, conserved, and re-used in the future is central to our work. We move, treat, analyze, and return water to the environment, and we help people use water efficiently, in their homes, buildings, factories and farms. In more than 150 countries, we have strong, long-standing relationships with customers who know us for our powerful combination of leading product brands and applications expertise, backed by a legacy of innovation.

For more information on how Xylem can help you, go to xyleminc.com.



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