

# Photometer System AL400



### Important steps before using the photometer

Please carry out the following steps as described in the Instruction manual. Become familiar with your new photometer before starting with the first tests:

- Unpacking and inspection of delivery contents, see page 338.
- Install the batteries, see page 286, 287.

Perform the following settings in the Mode-Menu; Instruction manual from page 297 and following:

• MODE 10: select language

• MODE 12: set date and time

• MODE 34: perform "Delete data"

• MODE 69: perform "User m. init" to initialise the user polynomial system

If required set other functions.



### (DE)

### Wichtige Information

### Um die Qualität unserer Umwelt zu erhalten, beschützen und zu verbessern Entsorgung von elektronischen Geräten in der Europäischen Union

Aufgrund der Europäischen Verordnung 2012/19/EU darf Ihr elektronisches Gerät nicht mit dem normalen Hausmüll entsorgt werden!

Tintometer GmbH entsorgt ihr elektrisches Gerät auf eine professionelle und für die Umwelt verantwortungsvolle Weise. Dieser Service ist, **die Transportkosten nicht inbegriffen**, kostenlos. Dieser Service gilt ausschließlich für elektrische Geräte die nach dem 13.08.2005 erworben wurden. Senden Sie Ihre zu entsorgenden Tintometer Geräte frei Haus an Ihren Lieferanten.



### Important Information

### To Preserve, Protect and Improve the Quality of the Environment Disposal of Electrical Equipment in the European Union

Because of the European Directive 2012/19/EU your electrical instrument must not be disposed of with normal household waste!

Tintometer GmbH will dispose of your electrical instrument in a professional and environmentally responsible manner. This service, **excluding the cost of transportation** is free of charge. This service only applies to electrical instruments purchased after 13th August 2005. Send your electrical Tintometer instruments for disposal freight prepaid to your supplier.



#### Notice importante

### Conserver, protéger et optimiser la qualité de l'environnement Élimination du matériel électrique dans l'Union Européenne

Conformément à la directive européenne n° 2012/19/UE, vous ne devez plus jeter vos instruments électriques dans les ordures ménagères ordinaires!

La société Tintometer GmbH se charge d'éliminer vos instruments électriques de façon professionnelle et dans le respect de l'environnement. Ce service, qui ne comprend pas les frais de transport, est gratuit. Ce service n'est valable que pour des instruments électriques achetés après le 13 août 2005. Nous vous prions d'envoyer vos instruments électriques Tintometer usés à vos frais à votre fournisseur.



#### Belangrijke informatie

Om de kwaliteit van ons leefmilieu te behouden, te verbeteren en te beschermen is voor landen binnen de Europese Unie de Europese richtlijn 2012/19/EU voor het verwijderen van elektronische apparatuur opgesteld.

Volgens deze richtlijn mag elektronische apparatuur niet met het huishoudelijk afval worden afgevoerd.

Tintometer GmbH verwijdert uw elektronisch apparaat op een professionele en milieubewuste wijze. Deze service is, **exclusief de verzendkosten**, gratis en alleen geldig voor elektrische apparatuur die na 13 augustus 2005 is gekocht. Stuur uw te verwijderen Tintometer apparatuur franco aan uw leverancier.



(ES)

# Información Importante Para preservar, proteger y mejorar la calidad del medio ambiente Eliminación de equipos eléctricos en la Unión Europea

Con motivo de la Directiva Europea 2012/19/UE, ¡ningún instrumento eléctrico deberá eliminarse junto con los residuos domésticos diarios!

Tintometer GmbH se encargará de dichos instrumentos eléctricos de una manera profesional y sin dañar el medio ambiente. Este servicio, el cual escluye los gastos de transporte, es gratis y se aplicará únicamente a aquellos instrumentos eléctricos adquiridos después del 13 de agosto de 2005. Se ruega enviar aquellos instrumentos

(II)

### Informazioni importanti

eléctricos inservibles de Tintometer a carga pagada a su distribuidor.

### Conservare, proteggere e migliorare la qualità dell'ambiente Smaltimento di apparecchiature elettriche nell'Unione Europea

In base alla Direttiva europea 2012/19/UE, gli apparecchi elettrici non devono essere smaltiti insieme ai normali rifiuti domestici!

Tintometer GmbH provvederà a smaltire i vostri apparecchi elettrici in maniera professionale e responsabile verso l'ambiente. Questo servizio, **escluso il trasporto**, è completamente gratuito. Il servizio si applica agli apparecchi elettrici acquistati successivamente al 13 agosto 2005. Siete pregati di inviare gli apparecchi elettrici Tintometer divenuti inutilizzabili a trasporto pagato al vostro rivenditore.

(PT)

### Informação Importante

### Para Preservar, Proteger e Melhorar a Qualidade do Ambiente Remoção de Equipamento Eléctrico na União Europeia

Devido à Directiva Europeia 2012/19/UE, o seu equipamento eléctrico nao deve ser removido com o lixo doméstico habitual!

A Tintometer GmbH tratará da remoção do seu equipamento eléctrico de forma profissional e responsável em termos ambientais. Este serviço, **não incluindo** 

**os custos de transporte**, é gratuito. Este serviço só é aplicável no caso de equipamentos eléctricos comprados depois de 13 de Agosto de 2005. Por favor, envie os seus equipamentos eléctricos Tintometer que devem ser removidos ao seu fornecedor (transporte pago).

(PL)

#### Istotna informacja

### Dla zachowania, ochrony oraz poprawy naszego środowiska Usuwanie urządzeń elektronicznych w Unii Europejskiej

Na podstawie Dyrektywy Parlamentu Europejskiego 2012/19/UE nie jest dozwolone usuwanie zakupionych przez Państwo urządzeń elektronicznych wraz z normalnymi odpadami z gospodarstwa domowego!

Tintometer GmbH usunie urządzenia elektrycznego Państwa w sposób profesjonalny i odpowiedzialny z punktu widzenia środowiska. Serwis ten jest, za wyjątkiem kosztów transportu, bezpłatny. Serwis ten odnosi się wyłącznie do urządzeń elektrycznych zakupionych po 13.08.2005r. Przeznaczone do usunięcia urządzenia firmy Tintometer mogą Państwo przesyłać na koszt własny do swojego dostawcy.



#### Wichtiger Entsorgungshinweis zu Batterien und Akkus

Jeder Verbraucher ist aufgrund der Batterieverordnung (Richtlinie 2006/66/ EG) gesetzlich zur Rückgabe aller ge- und verbrauchten Batterien bzw. Akkus verpflichtet. Die Entsorgung über den Hausmüll ist verboten. Da auch bei Produkten aus unserem Sortiment Batterien und Akkus im Lieferumgang enthalten sind, weisen wir Sie auf folgendes hin:

Verbrauchte Batterien und Akkus gehören nicht in den Hausmüll, sondern können unentgeltlich bei den öffentlichen Sammelstellen Ihrer Gemeinde und überall dort abgegeben werden, wo Batterien und Akkus der betreffenden Art verkauft werden. Weiterhin besteht für den Endverbraucher die Möglichkeit, Batterien und Akkus an den Händler, bei dem sie erworben wurden, zurückzugeben (gesetzliche Rücknahmepflicht).



### Important disposal instructions for batteries and accumulators

EC Guideline 2006/66/EC requires users to return all used and worn-out batteries and accumulators. They must not be disposed of in normal domestic waste. Because our products include batteries and accumulators in the delivery package our advice is as follows:

Used batteries and accumulators are not items of domestic waste. They must be disposed of in a proper manner. Your local authority may have a disposal facility; alternatively you can hand them in at any shop selling batteries and accumulators. You can also return them to the company which supplied them to you; the company is obliged to accept them.



### Information importante pour l'élimination des piles et des accumulateurs

En vertu de la Directive européenne 2006/66/CE relative aux piles et accumulateurs, chaque utilisateur est tenu de restituer toutes les piles et tous les accumulateurs utilisés et épuisés. L'élimination avec les déchets ménagers est interdite. Etant donné que l'étendue de livraison des produits de notre gamme contient également des piles et des accumulateurs, nous vous signalons ce qui suit :

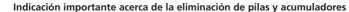
les piles et les accumulateurs utilisés ne sont pas des ordures ménagères, ils peuvent être remis sans frais aux points de collecte publics de votre municipalité et partout où sont vendus des piles et accumulateurs du type concerné. Par ailleurs, l'utilisateur final a la possibilité de remettre les piles et les accumulateurs au commerçant auprès duquel ils ont été achetés (obligation de reprise légale).



### Belangrijke mededeling omtrent afvoer van batterijen en accu's

Ledere verbruiker is op basis van de richtlijn 2006/66/EG verplicht om alle gebruikte batterijen en accu's in te leveren. Het is verboden deze af te voeren via het huisvuil. Aangezien ook onze producten geleverd worden met batterijen en accu's wijzen wij u op het volgende; Lege batterijen en accu's horen niet in het huisvuil thuis. Men kan deze inleveren bij inzamelpunten van uw gemeente of overal daar waar deze verkocht worden. Tevens bestaat de mogelijkheid batterijen en accu's daar in te leveren waar u ze gekocht heeft. (wettelijke terugnameplicht)





Basado en la norma relativa a pilas/ baterías (directiva 2006/66/CE), cada consumidor, está obligado por ley, a la devolución de todas las pilas/ baterías y acumuladores usados y consumidos. Está prohibida la eliminación en la basura doméstica. Ya que en productos de nuestra gama, también se incluyen en el suministro pilas y acumuladores, le sugerimos lo siquiente:

Las pilas y acumuladores usados no pertenecen a la basura doméstica, sino que pueden ser entregados en forma gratuita en cada uno de los puntos de recolección públicos de su comunidad en los cuales se vendan pilas y acumuladores del tipo respectivo. Además, para el consumidor final existe la posibilidad de devolver las pilas y baterías recargables a los distribuidores donde se hayan adquirido (obligación legal de devolución).

#### Indicazioni importanti sullo smaltimento di pile e accumulatori

IT

PL

In base alla normativa concernente le batterie (Direttiva 2006/66/CE) ogni consumatore è tenuto per legge alla restituzione di tutte le batterie o accumulatori usati ed esauriti. È vietato lo smaltimento con i rifiuti domestici. Dato che anche alcuni prodotti del nostro assortimento sono provvisti di pile e accumulatori, vi diamo di seguito delle indicazioni: Pile e accumulatori esauriti non vanno smaltiti insieme ai rifiuti domestici, ma depositati gratuitamente nei punti di raccolta del proprio comune o nei punti vendita di pile e accumulatori dello stesso tipo. Inoltre il consumatore finale può portare batterie e accumulatori al rivenditore presso il quale li ha acquistati (obbligo di raccolta previsto per legge).

### Instruções importantes para a eliminação residual de pilhas e acumuladores

Os utilizadores finais são legalmente responsáveis, nos termos do Regulamento relativo a pilhas e acumuladores (Directiva 2006/66/CE), pela entrega de todas as pilhas e acumuladores usados e gastos. É proibida a sua eliminação juntamente com o lixo doméstico. Uma vez que determinados produtos da nossa gama contêm pilhas e/ou acumuladores, alertamos para os seguintes aspectos:

As pilhas e acumuladores usados não podem ser eliminados com o lixo doméstico, devendo sim ser entregues, sem encargos, junto dos pontos de recolha públicos do seu município, ou em qualquer ponto de venda de pilhas e acumuladores. O utilizador final dispõe ainda da possibilidade de entregar as pilhas e/ou acumuladores no estabelecimento comerciante onde os adquiriu (dever legal de aceitar a devolução).

#### Istotna wskazówka dotycząca utylizacji baterii i akumulatorów

Każdy użytkownik na mocy rozporządzenia w sprawie baterii (wytyczna 2006/66/WE) jest ustawowo zobowiązany do oddawania wszystkich rozładowanych i zużytych baterii lub akumulatorów. Utylizacja wraz z odpadkami domowymi jest zabroniona. Ponieważ także w produktach z naszego asortymentu zawarte są w zakresie dostawy baterie i akumulatory, zwracamy uwagę na poniższe zasady:

zużyte baterie i akumulatory nie mogą być wyrzucane wraz z odpadkami domowymi, lecz powinny być bezpłatnie przekazywane w publicznych miejscach zbiórki wyznaczonych przez gminę lub oddawane w punktach, gdzie sprzedawane są baterie i akumulatory danego rodzaju. Poza tym użytkownik końcowy ma możliwość zwrócenia baterii i akumulatorów do przedstawiciela handlowego, u którego je nabył (ustawowy obowiązek przyjęcia).



### Safety precautions



### $\triangle$ CAUTION $\triangle$



Reagents are formulated exclusively for chemical analysis and must not be used for any other purpose. Reagents must not get into the hands of children. Some of the reagents contain substances which are not entirely harmless environmentally. Be aware of the ingredients and take proper care when disposing of the test solution.



### ♠ CAUTION ♠



Please read this instruction manual before unpacking, setting up or using the photometer. Please read the method description completely before performing the test. Be aware of the risks of using the required reagents by reading the MSDS (Material Safety Data Sheets). Failure could result in serious injury to the operator or damage to the instrument.

MSDS: www.aqualytic.de



### $\Lambda$ caution $\Lambda$



The accuracy of the instrument is only valid if the instrument is used in an environment with controlled electromagnetic disturbances according to DIN 61326. Wireless devices, e.g. wireless phones, must not be used near the instrument.

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### Part 1

**Methods** 

| No. | Analysis                  | Reagent            | Range         | Displayed as           | Method   | λ<br>[nm] | OTZ      | Page   |
|-----|---------------------------|--------------------|---------------|------------------------|--|-----------|----------|--------|
| 20  | Acid demand to pH 4.3 T   | tablet             | 0.1-4         | mmol/l                 | Acid/Indicator 1,2,5                                     | 610       | ✓        | 14     |
| 30  | Alkalinity, total T       | tablet             | 5-200         | mg/l CaCO <sub>3</sub> | Acid/Indicator 1,2,5                                     | 610       | ✓        | 16     |
| 31  | Alkalinity HR,<br>total T | tablet             | 5-500         | mg/l CaCO <sub>3</sub> | Acid/Indicator 1,2,5                                     | 610       | √        | 18     |
| 35  | Alkalinity-p T            | tablet             | 5-300         | mg/l CaCO₃             | Acid/Indicator 1,2,5                                     | 560       | ✓        | 20     |
| 40  | Aluminium T               | tablet             | 0.01-0.3      | mg/l Al                | Eriochrome<br>Cyanine R <sup>2</sup>                     | 530       | ✓        | 22     |
| 50  | Aluminium PP              | PP + liquid        | 0.01-<br>0.25 | mg/l Al                | Eriochrome<br>Cyanine R <sup>2</sup>                     | 530       | -        | 24     |
| 60  | Ammonia T                 | tablet             | 0.02-1        | mg/l N                 | Indophenol blue <sup>2,3</sup>                           | 610       | ✓        | 26     |
| 62  | Ammonia PP                | PP                 | 0.01-0.8      | mg/l N                 | Salicylate <sup>2</sup>                                  | 660       | _        | 28     |
| 65  | Ammonia LR TT             | tube test          | 0.02-2.5      | mg/l N                 | Salicylate <sup>2</sup>                                  | 660       | _        | 30     |
| 66  | Ammonia HR TT             | tube test          | 1-50          | mg/l N                 | Salicylate <sup>2</sup>                                  | 660       | _        | 32     |
| 85  | Boron T                   | tablet             | 0.1-2         | mg/l B                 | Azomethine <sup>3</sup>                                  | 430       | ✓        | 34     |
| 80  | Bromine T                 | tablet             | 0.05-13       | mg/l Br <sub>2</sub>   | DPD <sup>5</sup>   | 530       | ✓        | 36     |
| 81  | Bromine PP                | PP                 | 0.05-4.5      | mg/l Br <sub>2</sub>   | DPD 1,2  | 530       | ✓        | 38     |
| 90  | Chloride T                | tablet             | 0.5 -25       | mg/l Cl <sup>-</sup>   | Silver nitrate/<br>turbidity                             | 530       | ✓        | 40     |
| 92  | Chloride L                | liquid             | 0.5-20        | mg/l Cl <sup>-</sup>   | Mercurythiocyanate/<br>Iron nitrate                      | 430       | ✓        | 42     |
| 100 | Chlorine T *              | tablet             | 0.01-6        | mg/l Cl <sub>2</sub>   | DPD <sup>1,2,3</sup>                                     | 530       | ✓        | 44, 46 |
| 103 | Chlorine HR T *           | tablet             | 0.1-10        | mg/l Cl <sub>2</sub>   | DPD <sup>1,2,3</sup>                                     | 530       | ✓        | 44, 50 |
| 101 | Chlorine L *              | liquid             | 0.02-4        | mg/l Cl <sub>2</sub>   | DPD 1,2,3  | 530       | ✓        | 44, 54 |
| 110 | Chlorine PP *             | PP                 | 0.02-2        | mg/l Cl <sub>2</sub>   | DPD 1,2  | 530       | <b>√</b> | 44, 58 |
| 111 | Chlorine HR PP *          | PP                 | 0.1-8         | mg/l Cl <sub>2</sub>   | DPD 1,2  | 530       | -        | 44, 62 |
| 120 | Chlorine dioxide T        | tablet             | 0.02-11       | mg/l ClO <sub>2</sub>  | DPD, Glycine 1,2   | 530       | ✓        | 66     |
| 122 | Chlorine dioxide PP       | PP                 | 0.04-3.8      | mg/l ClO <sub>2</sub>  | DPD <sup>1,2</sup>                                       | 530       | ✓        | 72     |
| 105 | Chlorine HR (KI) T        | tablet             | 5-200         | mg/l Cl <sub>2</sub>   | KI/Acid <sup>5</sup>                                     | 530       | _        | 76     |
| 125 | Chromium PP               | PP                 | 0.02-2        | mg/l Cr                | 1,5-Diphenyl-<br>carbohydrazide <sup>1,2</sup>           | 530       | _        | 82     |
| 130 | COD LR TT                 | tube test          | 0 -150        | mg/l O <sub>2</sub>    | Dichromate/H <sub>2</sub> SO <sub>4</sub> 1,2            | 430       | _        | 88     |
| 131 | COD MR TT                 | tube test          | 0 -1500       | mg/l O <sub>2</sub>    | Dichromate/H <sub>2</sub> SO <sub>4</sub> <sup>1,2</sup> | 610       | _        | 90     |
| 132 | COD HR TT                 | tube test          | 0 -15         | g/l O <sub>2</sub>     | Dichromate/H <sub>2</sub> SO <sub>4</sub> <sup>1,2</sup> | 610       |          | 92     |
| 204 | Colour                    | direct<br>reading  | 0-500         | Pt-Co units            | Pt-Co-Scale <sup>1,2</sup><br>(APHA)                     | 430       | -        | 94     |
| 150 | Copper T *                | tablet             | 0.05-5        | mg/l Cu                | Biquinoline <sup>4</sup>                                 | 560       | ✓        | 96     |
| 151 | Copper L*                 | liquid +<br>powder | 0.05-4        | mg/l Cu                | Bicinchoninate   | 560       | ✓        | 100    |
| 153 | Copper PP                 | PP                 | 0.05-5        | mg/l Cu                | Bicinchoninate   | 560       | ✓        | 106    |

<sup>\* =</sup> free, combined, total; PP = powder pack; T = tablet; L = liquid; TT = tube test; LR = low range; MR = middle range; HR = high range; Vacu-vial® is a registered trade mark of CHEMetrics Inc.

| No. | Analysis                           | Reagent            | Range         | Displayed as                       | Method   | λ<br>[nm] | OTZ      | Page        |
|-----|------------------------------------|--------------------|---------------|------------------------------------|--|-----------|----------|-------------|
| 157 | Cyanide                            | Powder +<br>liquid | 0.01-0.5      | mg/l CN                            | Pyridine-<br>barbituric acid <sup>1</sup>            | 580       | ✓        | 108         |
| 160 | CyA-TEST T                         | tablet             | 0-160         | mg/l CyA                           | Melamine   | 530       | ✓        | 110         |
| 165 | DEHA T                             | tablet +<br>liquid | 20-500        | μg/l DEHA                          | PPST <sup>3</sup>                                    | 560       | ✓        | 1122        |
| 167 | DEHA PP                            | PP + liquid        | 20-500        | μg/l DEHA                          | PPST <sup>3</sup>                                    | 560       | -        | 114         |
| 170 | Fluoride L                         | liquid             | 0.05-2        | mg/l F                             | SPADNS <sup>2</sup>                                  | 580       | ✓        | 116         |
| 210 | H <sub>2</sub> O <sub>2</sub> T    | tablet             | 0.03-3        | mg/l H <sub>2</sub> O <sub>2</sub> | DPD/catalyst 5                                       | 530       | ✓        | 118         |
| 213 | H <sub>2</sub> O <sub>2</sub> LR L | liquid             | 1-50          | mg/l H <sub>2</sub> O <sub>2</sub> | Titanium<br>tetrachloride/acid                       | 430       |          | 120         |
| 214 | H <sub>2</sub> O <sub>2</sub> HR L | liquid             | 40-500        | mg/l H <sub>2</sub> O <sub>2</sub> | Titanium<br>tetrachloride/acid                       | 530       | -        | 122         |
| 190 | Hardness, Calcium T                | tablet             | 50-900        | mg/l CaCO <sub>3</sub>             | Murexide <sup>4</sup>                                | 560       | _        | 124         |
| 191 | Hardness, Calcium<br>2 T           | tablet             | 0-500         | mg/l CaCO <sub>3</sub>             | Murexide <sup>4</sup>                                | 560       | ✓        | 126         |
| 200 | Hardness, total T                  | tablet             | 2-50          | mg/l CaCO <sub>3</sub>             | Metallphthalein <sup>3</sup>                         | 560       | ✓        | 128         |
| 201 | Hardness, total<br>HR T            | tablet             | 20-500        | mg/l CaCO <sub>3</sub>             | Metallphthalein <sup>3</sup>                         | 560       | ✓        | 130         |
| 205 | Hydrazine P                        | powder             | 0.05-0.5      | mg/l N <sub>2</sub> H <sub>4</sub> | 4-(Dimethyl-<br>amino)-<br>benzaldehyde <sup>3</sup> | 430       | ✓        | 132         |
| 206 | Hydrazine L                        | liquid             | 0.005-<br>0.6 | mg/l N <sub>2</sub> H <sub>4</sub> | 4-(Dimethyl-<br>amino)-<br>benzaldehyde <sup>3</sup> | 430       | _        | 134         |
| 207 | Hydrazine C                        | Vacu-vial          | 0.01-0.7      | mg/l N <sub>2</sub> H <sub>4</sub> | PDMAB  | 430       | _        | 136         |
| 215 | Iodine T                           | tablet             | 0.05-3.6      | mg/l I                             | DPD <sup>5</sup>                                     | 530       | ✓        | 138         |
| 220 | Iron T                             | tablet             | 0.02-1        | mg/l Fe                            | PPST <sup>3</sup>                                    | 560       | ✓        | 140,<br>142 |
| 222 | Iron PP                            | PP                 | 0.02-3        | mg/l Fe                            | 1,10-Phenan-<br>troline <sup>3</sup>                 | 530       | ✓        | 140,<br>144 |
| 223 | Iron (TPTZ) PP                     | PP                 | 0.02-1.8      | mg/l Fe                            | TPTZ   | 580       | _        | 140,<br>146 |
| 224 | Iron (Fe in Mo) PP                 | PP                 | 0.01-1.8      | mg/l Fe                            | Fe in Mo   | 580       | _        | 140,<br>148 |
| 225 | Iron LR L                          | liquid             | 0.03-2        | mg/l Fe                            | Ferrozine /<br>Thioglycolate                         | 560       | ✓        | 140,<br>150 |
| 226 | Iron LR 2 L                        | liquid             | 0.03-2        | mg/l Fe                            | Ferrozine /<br>Thioglycolate                         | 560       | ✓        | 140,<br>154 |
| 227 | Iron HR L                          | liquid             | 0.1-10        | mg/l Fe                            | Thioglycolate  | 530       | <b>√</b> | 140,<br>158 |
| 240 | Manganese T                        | tablet             | 0.2-4         | mg/l Mn                            | Formaldoxime   | 530       | ✓        | 162         |
| 242 | Manganese LR PP                    | PP + liquid        | 0.01-0.7      | mg/l Mn                            | PAN  | 560       | _        | 164         |

<sup>\* =</sup> free, combined, total; PP = powder pack; T = tablet; L = liquid; TT = tube test; LR = low range; MR = middle range; HR = high range; Vacu-vial® is a registered trade mark of CHEMetrics Inc.

| No. | Analysis                 | Reagent            | Range    | Displayed as          | Method  | λ<br>[nm] | OTZ      | Page        |
|-----|--------------------------|--------------------|----------|-----------------------|---|-----------|----------|-------------|
| 243 | Manganese HR PP          | PP + liquid        | 0,1-18   | mg/l Mn               | Periodate oxidation <sup>2</sup>                      | 530       | ✓        | 166         |
| 245 | Manganese L              | liquid             | 0.05-5   | mg/l Mn               | Formaldoxime  | 430       | ✓        | 168         |
| 250 | Molybdate T              | tablet             | 1-50     | mg/l MoO <sub>4</sub> | Thioglycolate 4                                       | 430       | ✓        | 170         |
| 251 | Molybdate LR PP          | PP                 | 0,05-5   | mg/l MoO <sub>4</sub> | Ternary Complex                                       | 610       | ✓        | 172         |
| 252 | Molybdate HR PP          | PP                 | 0.5-66   | mg/l MoO <sub>4</sub> | Mercaptoacetic acid                                   | 430       | ✓        | 174         |
| 254 | Molybdate HR L           | liquid             | 1-100    | mg/l MoO <sub>4</sub> | Thioglycolate   | 430       | <b>√</b> | 176         |
| 257 | Nickel T                 | tablet             | 0.1-10   | mg/l Ni               | Nioxime   | 560       | ✓        | 178         |
| 260 | Nitrate                  | tablet +<br>powder | 0.08-1   | mg/l N                | Zinc reduction<br>/ NED                               | 530       | <b>√</b> | 180         |
| 265 | Nitrate TT               | tube test          | 1-30     | mg/l N                | Chromotropic acid                                     | 430       | _        | 182         |
| 270 | Nitrite T                | tablet             | 0.01-0.5 | mg/l N                | N-(1-Naphthyl)-<br>ethylendiamine <sup>2,3</sup>      | 560       | ✓        | 184         |
| 272 | Nitrite LR PP            | PP                 | 0.01-0.3 | mg/l N                | Diazotization   | 530       | ✓        | 186         |
| 280 | Nitrogen, total<br>LR TT | tube test          | 0.5-25   | mg/l N                | Persulfate<br>digestion method                        | 430       | _        | 188         |
| 281 | Nitrogen, total<br>HR TT | tube test          | 5-150    | mg/l N                | Persulfate<br>digestion method                        | 430       | _        | 190         |
| 290 | Oxygen, active T         | tablet             | 0.1-10   | mg/l O <sub>2</sub>   | DPD   | 530       | ✓        | 194         |
| 292 | Oxygen, dissolved        | Vacu-vial          | 10-800   | μg/l O <sub>2</sub>   | Rhodazine D™  | 530       | -        | 196         |
| 300 | Ozone (DPD) T            | tablet             | 0.02-2   | mg/l O <sub>3</sub>   | DPD/Glycine 5   | 530       | ✓        | 198         |
| 70  | PHMB T                   | tablet             | 2-60     | mg/l PHMB             | Buffer/Indicator                                      | 560       | ✓        | 204         |
| 320 | Phosphate, T<br>ortho LR | tablet             | 0.05-4   | mg/l PO <sub>4</sub>  | Ammonium-<br>molybdate <sup>2,3</sup>                 | 660       | ✓        | 206,<br>208 |
| 321 | Phosphate, ortho<br>HR T | tablet             | 1-80     | mg/l PO <sub>4</sub>  | Vanado-<br>molybdate <sup>2</sup>                     | 430       | ✓        | 206,<br>210 |
| 323 | Phosphate, PP ortho      | PP                 | 0.06-2.5 | mg/l PO <sub>4</sub>  | Molybdate/<br>Ascorbic acid <sup>2</sup>              | 660       | ✓        | 206,<br>212 |
| 324 | Phosphate, ortho<br>TT   | tube test          | 0.06-5   | mg/l PO <sub>4</sub>  | Molybdate/<br>Ascorbic acid <sup>2</sup>              | 660       | _        | 206,<br>214 |
| 327 | Phosphate 1 C, ortho     | Vacu-vial          | 5-40     | mg/l PO <sub>4</sub>  | Vanado-<br>molybdate <sup>2</sup>                     | 430       | -        | 206,<br>216 |
| 328 | Phosphate 2 C, ortho     | Vacu-vial          | 0.05-5   | mg/l PO <sub>4</sub>  | Stannous<br>chloride <sup>2</sup>                     | 660       | -        | 206,<br>218 |
| 325 | Phosphate, hydr.<br>TT   | tube test          | 0.02-1.6 | mg/l P                | Acid digestion,<br>Ascorbic acid <sup>2</sup>         | 660       | -        | 206,<br>220 |
| 326 | Phosphate, total<br>TT   | tube test          | 0.02-1.1 | mg/l P                | Acid persulf digestion,<br>Ascorbic acid <sup>2</sup> | 660       | -        | 206,<br>222 |
| 334 | Phosphate LR L           | liquid             | 0.1-10   | mg/l PO <sub>4</sub>  | Phosphomolybdic acid/Ascorbic acid                    | 660       | ✓        | 206,<br>224 |
| 335 | Phosphate HR L           | liquid             | 5-80     | mg/l PO <sub>4</sub>  | Vanado-<br>molybdate                                  | 430       | ✓        | 206,<br>228 |

<sup>\* =</sup> free, combined, total; PP = powder pack; T = tablet; L = liquid; TT = tube test; LR = low range; MR = middle range; HR = high range; Vacu-vial® is a registered trade mark of CHEMetrics Inc.

| No. | Analysis                 | Reagent            | Range    | Displayed as           | Method                                       | λ<br>[nm] | OTZ      | Page |
|-----|--------------------------|--------------------|----------|------------------------|--|-----------|----------|------|
| 316 | Phosphonate PP           | PP                 | 0-125    | mg/l                   | Persulfate<br>UV-Oxidation                   | 660       | -        | 232  |
| 329 | pH-Value LR T            | tablet             | 5.2-6.8  | _                      | Bromocresolpurple <sup>5</sup>               | 560       | ✓        | 236  |
| 330 | pH-Value T               | tablet             | 6.5-8.4  | _                      | Phenolred <sup>5</sup>                       | 560       | ✓        | 238  |
| 331 | pH-Value L               | liquid             | 6.5-8.4  | _                      | Phenolred <sup>5</sup>                       | 560       | ✓        | 240  |
| 332 | pH-Value HR T            | tablet             | 8.0-9.6  | _                      | Thymolblue 5                                 | 560       | ✓        | 242  |
| 338 | Polyacrylate L           | liquid             | 1-30     | mg/ l<br>Polyacryl     | Turbidity                                    | 660       | ✓        | 244  |
| 340 | Potassium T              | tablet             | 0.7-12   | mg/l K                 | Tetraphenylborate-<br>Turbidity <sup>4</sup> | 430       | 1        | 248  |
| 350 | Silica T                 | tablet             | 0.05-4   | mg/l SiO <sub>2</sub>  | Silicomolybdate <sup>2,3</sup>               | 660       | ✓        | 250  |
| 351 | Silica LR PP             | PP                 | 0.1-1.6  | mg/l SiO <sub>2</sub>  | Heteropolyblue <sup>2</sup>                  | 660       | -        | 252  |
| 352 | Silica HR PP             | PP                 | 1-90     | mg/l SiO <sub>2</sub>  | Silicomolybdate <sup>2</sup>                 | 430       | ✓        | 254  |
| 353 | Silica L                 | liquid +<br>powder | 0.1-8    | mg/l SiO <sub>2</sub>  | Heteropolyblue <sup>2</sup>                  | 660       | ✓        | 256  |
| 212 | Sodium<br>hypochlorite T | tablet             | 0.2-16   | % NaOCI                | Potassium iodide <sup>5</sup>                | 530       | <b>√</b> | 258  |
| 355 | Sulfate T                | tablet             | 5-100    | mg/l SO <sub>4</sub>   | Bariumsulfate-<br>Turbidity                  | 610       | ✓        | 260  |
| 360 | Sulfate PP               | PP                 | 5-100    | mg/l SO <sub>4</sub>   | Bariumsulfate-<br>Turbidity <sup>2</sup>     | 530       | ✓        | 262  |
| 365 | Sulfide                  | tablet             | 0.04-0.5 | mg/l S                 | DPD/Catalyst 3,4                             | 660       | ✓        | 264  |
| 370 | Sulfite T                | tablet             | 0.1-5    | mg/l SO <sub>3</sub>   | DTNB   | 430       | ✓        | 266  |
| 384 | Suspended Solids         | direct<br>reading  | 0-750    | mg/l TSS               | photometric                                  | 660       | -        | 268  |
| 386 | Turbidity                | direct<br>reading  | 0-1000   | FAU                    | Attenuated<br>Radiation Method               | 530       | -        | 270  |
| 388 | Tolyltriazole PP         | PP                 | 1-16     | mg/l Benzo<br>triazole | Catalysed UV photolysis                      | 430       | <b>√</b> | 272  |
| 390 | Urea T                   | tablet +<br>liquid | 0.1-2.5  | mg/l Urea              | Indophenol/<br>Urease                        | 610       | ✓        | 274  |
| 400 | Zinc T                   | tablet             | 0.02 -1  | mg/l Zn                | Zincon <sup>3</sup>                          | 610       | _        | 276  |
| 405 | Zinc L                   | liquid +<br>powder | 0.1 -2.5 | mg/l Zn                | Zincon / EDTA                                | 610       | ✓        | 278  |

<sup>\* =</sup> free, combined, total; PP = powder pack; T = tablet;

L = liquid; TT = tube test; LR = low range; MR = middle range; HR = high range; Vacu-vial® is a registered trade mark of CHEMetrics Inc.

The precision of Lovibond® Reagent Systems (tablets, powder packs and tube tests) is identical to the precision specified in standards literature such as American Standards (AWWA), ISO etc.

Most of the data referred to in these standard methods relates to Standard Solutions. Therefore they are not readily applicable to drinking-, boiler- or waste-water, since various interferences can have a major influence on the accuracy of the method.

For this reason we don't state such potentially misleading data.

Due to the fact that each sample is different, the only way to check the tolerances ('precision') is the Standard Additions Method.

According to this method, first the original sample is tested. Then further samples (2 to 4) are taken and small amounts of a Standard Solution are added, and further results are obtained. The amounts added range from approximately half, up to double the amount present in the sample itself.

These supplementary results make it possible to estimate the actual concentration of the original sample by comparison.

### Literature

Active Oxvaen

The reagent formulations are based on internationally recognised test methods. Some are described in national and/or international guidelines.

- 1. Deutsche Einheitsverfahren zur Wasser-, Abwasser- und Schlammuntersuchung
- 2. Standard Methods for the Examination of Water and Wastewater; 18th Edition, 1992

Oxygen, activ

- Photometrische Analysenverfahren, Schwedt, Wissenschaftliche Verlagsgesellschaft mbH, Stuttgart 1989
- 4. Photometrische Analyse, Lange / Vejdelek, Verlag Chemie 1980
- 5. Colorimetric Chemical Analytical Methods, 9th Edition, London

### Notes for searching:

Index (Water Balance)

### OTZ (OneTimeZero) switching on and off, see Mode 55, page 321

| Langelier Saturation | -> | Mode function 70  |
|----------------------|----|-------------------|
| total Hardness       | -> | Hardness, total   |
| total Alkalinity     | -> | Alkalinity, total |
| Silicon dioxide      | -> | Silica            |
| p-Value              | -> | Alkalinity-p      |
| Total Hardness       | -> | Hardness, total   |
| m-Value              | -> | Alkalinity, total |
| Monochloramine       | -> | Chloramine, mono  |
| $H_2O_2$             | -> | Hydrogen peroxide |
| Cyanuric acid        | -> | CyA-TEST          |
| Calcium Hardness     | -> | Hardness, Calcium |
| Biguanide            | -> | PHMB              |
| Alkalinity-m         | -> | Alkalinity, total |
| , ,                  |    | , ,               |





### Acid demand to pH 4.3 with Tablet

 $0.1 - 4 \, \text{mmol/l}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one ALKA-M-PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display as Acid demand to  $pH\ 4.3$  in mmol/l.

### Notes:

- 1. The terms total Alkalinity, Alkalinity-m, m-Value and Acid demand to pH 4.3 are identical.
- 2. For accurate results exactly 10 ml of water sample must be taken for the test.

| Reagent           | Form of reagent/Quantity | Order-No. |  |
|-------------------|--------------------------|-----------|--|
| ALKA-M-PHOTOMETER | Tablet / 100             | 4513210BT |  |





# Alkalinity, total = Alkalinity-m = m-Value with Tablet

5 - 200 mg/l CaCO<sub>3</sub>



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one ALKA-M-PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display as total Alkalinity.

#### Notes:

- 1. The terms total Alkalinity, Alkalinity-m, m-Value and Alkalinity to pH 4.3 are identical.
- 2. For accurate results exactly 10 ml of water sample must be taken for the test.
- 3. Conversion table:

|                          | Acid demand to pH 4.3 | German | English | French |
|--------------------------|-----------------------|--------|---------|--------|
|                          | DIN 38 409 (Ks4.3)    | °dH*   | °eH*    | °fH*   |
| 1 mg/l CaCO <sub>3</sub> | 0.02                  | 0.056  | 0.07    | 0.1    |

<sup>\*</sup>Carbonate hardness (reference = Hydrogencarbonate-anions)

Example:

10 mg/l CaCO<sub>3</sub> = 10 mg/l x 0.056 = 0.56 °dH 10 mg/l CaCO<sub>3</sub> = 10 mg/l x 0.02 = 0.2 mmol/l

4. ▲ CaCO<sub>3</sub> °dH °eH °fH

°aH

| Reagent           | Form of reagent/Quantity | Order-No. |
|-------------------|--------------------------|-----------|
| ALKA-M-PHOTOMETER | Tablet / 100             | 4513210BT |





# Alkalinity HR, total = Alkalinity-m HR = m-Value HR with Tablet

5 - 500 mg/l CaCO<sub>3</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one ALKA-M-HR PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved

### Countdown 1:00 start:

- 7. Press [4] key.
  Wait for a **reaction period of 1 minute.**
- 8. Remix the solution.
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

10. Press TEST key.

The result is shown in the display as total Alkalinity.

#### Notes:

1. For verification of the result look carefully at the bottom of the vial. If a thin yellow layer forms, then mix the vial again. This ensures that reaction is complete. Reread the result.

2. Conversion table:

|                          | Acid demand to pH 4.3 | German | English | French |
|--------------------------|-----------------------|--------|---------|--------|
|                          | DIN 38 409 (Ks4.3)    | °dH*   | °eH*    | °fH*   |
| 1 mg/l CaCO <sub>3</sub> | 0.02                  | 0.056  | 0.07    | 0.1    |

<sup>\*</sup>Carbonate hardness (reference = Hydrogencarbonate-anions)

Example:

10 mg/l 
$$CaCO_3 = 10$$
 mg/l  $\times 0.056 = 0.56$  °dH  
10 mg/l  $CaCO_3 = 10$  mg/l  $\times 0.02 = 0.2$  mmol/l

3. CaCO<sub>3</sub> °dH °eH °fH

°aH

| Reagent              | Form of reagent/Quantity | Order-No. |
|----------------------|--------------------------|-----------|
| ALKA-M-HR PHOTOMETER | Tablet / 100             | 4513240BT |





### Alkalinity-p = p-value with Tablet

5 - 300 mg/l CaCO<sub>3</sub>



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one ALKA-P-PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

### Countdown 5:00

8. Press **TEST** key. Wait for a **reaction period of 5 minutes.** 

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as Alkalinity-p.

#### Notes

- 1. The terms Alkalinity-p, p-Value and Alkalinity to pH 8.2 are identical.
- 2. For accurate test results exactly 10 ml of water sample must be taken for the test.
- 3. This method was developed from a volumetric procedure for the determination of Alkalinity-p. Due to undefined conditions, the deviations from the standardised method may be greater.
- 4. Conversion table:

|                          | mg/l CaCO₃ | °dH   | °fH  | °eH  |
|--------------------------|------------|-------|------|------|
| 1 mg/l CaCO <sub>3</sub> |            | 0.056 | 0.10 | 0.07 |
| 1 °dH                    | 17.8       |       | 1.78 | 1.25 |
| 1 °fH                    | 10.0       | 0.56  |      | 0.70 |
| 1 °eH                    | 14.3       | 0.80  | 1.43 |      |



6. By determining Alkalinity-p and Alkalinity-m it is possible to classify the alkalinity as Hydroxide, Carbonate and Hydrogencarbonate.

The following differentiation is only valid if:

a) no other alkalis are present and

b) Hydroxide und Hydrogen are not present in the same water sample. If condition b) is not fulfilled please get additional information from "Deutsche Einheitsverfahren zur Wasser-, Abwasser- und Schlammuntersuchung, D 8".

Case 1: Alkalinity-p = 0 Hydrogen carbonate = m

Carbonate = 0

Hydroxide = 0 Case 2: Alkalinity-p > 0 and Alkalinity-m > 2p

Hydrogen carbonate = m - 2p

Carbonate = 2p Hvdroxide = 0

Case 3: Alkalinity-p > 0 and Alkalinity-m < 2p

Hydrogen carbonate = 0Carbonate = 2m - 2pHydroxide = 2p - m

| Reagent           | Form of reagent/Quantity | Order-No. |
|-------------------|--------------------------|-----------|
| ALKA-P-PHOTOMETER | Tablet / 100             | 4513230BT |





### Aluminium with Tablet

0.01 - 0.3 mg/l Al



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one ALUMINIUM No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod (dissolve the tablet).
- Add one ALUMINIUM No. 2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl gently several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

### Countdown 5:00

Press TEST key.Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Aluminium.

#### Notes:

- 1. Before use, clean the vials and the accessories with Hydrochloric acid (approx. 20%). Rinse them thoroughly with deionised water.
- 2. To get accurate results the sample temperature must be between 20°C and 25°C.
- 3. A low test result may be given in the presence of Fluorides and Polyphosphates. The effect of this is generally insignificant unless the water has fluoride added artificially. In this case, the following table should be used:

| Fluoride |      | Displayed | l value: A | luminium | [mg/l Al] |      |
|----------|------|-----------|------------|----------|-----------|------|
| [mg/l F] | 0.05 | 0.10      | 0.15       | 0.20     | 0.25      | 0.30 |
| 0.2      | 0.05 | 0.11      | 0.16       | 0.21     | 0.27      | 0.32 |
| 0.4      | 0.06 | 0.11      | 0.17       | 0.23     | 0.28      | 0.34 |
| 0.6      | 0.06 | 0.12      | 0.18       | 0.24     | 0.30      | 0.37 |
| 0.8      | 0.06 | 0.13      | 0.20       | 0.26     | 0.32      | 0.40 |
| 1.0      | 0.07 | 0.13      | 0.21       | 0.28     | 0.36      | 0.45 |
| 1.5      | 0.09 | 0.20      | 0.29       | 0.37     | 0.48      |      |

Example: If the result of Aluminium determination is 0.15 mg/l Al and the Fluoride concentration is known to be 0.4 mg/l F, the true concentration of Aluminium is 0.17 mg/l Al.

4. A special tablet ingredient prevents effects on the measurement due to iron and manganese.



| Reagent                        | Form of reagent/Quantity                | Order-No. |
|--------------------------------|---|-----------|
| Set<br>ALUMINIUM No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517601BT |
| ALUMINIUM No. 1                | Tablet / 100                            | 4515460BT |
| ALUMINIUM No. 2                | Tablet / 100                            | 4515470BT |





### Aluminium with Vario Powder Pack

0.01 - 0.25 mg/l Al



Use two clean vials (24 mm  $\varnothing$ ) and mark one as blank for zeroing.

- 1. Fill 20 ml of the water sample in a 100 ml beaker.
- Add the contents of one Vario Aluminum ECR F20 Powder Pack straight from the foil to the water sample.
- 3. Dissolve the powder using a clean stirring rod.

### Countdown 1 0:30 start:

4. Press [ ] key.

Wait for a reaction period of 30 seconds.

After the reaction period is finished proceed as follows:



- Add the contents of one Vario Hexamine F20 Powder Pack straight from the foil to the same water sample.
- 6. Dissolve the powder using a clean stirring rod.
- Add 1 drop of Vario Aluminum ECR Masking Reagent in the vial marked as blank.
- 8. Add 10 ml of the prepared water sample to the vial (this is the blank).
- 9. Add the remaining 10 ml of the prepared water sample in the second clean vial (this is the sample).
- 10. Close the vials tightly with the caps and swirl several times to mix the contents.

Countdown 2 5:00 start:

11. Press [₄] key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished proceed as follows:

12. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

prepare Zero press ZERO

- 13. Press **ZERO** key.
- 14. Remove the vial from the sample chamber.
- 15. Place the vial (the sample) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

16. Press TEST key.

The result is shown in the display in mg/l Aluminium.

#### Notes:

- 1. Before use, clean the vials and the accessories with Hydrochloric acid (approx. 20%). Rinse them thoroughly with deionised water.
- 2. To get accurate results the sample temperature must be between 20°C and 25°C.
- 3. A low test result may be given in the presence of Fluorides and Polyphosphates. The effect of this is generally insignificant unless the water has fluoride added artificially. In this case, the following table should be used:

| Fluoride | Displayed value: Aluminium [mg/l Al] |      |      |      |      |      |
|----------|--------------------------------------|------|------|------|------|------|
| [mg/l F] | 0.05                                 | 0.10 | 0.15 | 0.20 | 0.25 | 0.30 |
| 0.2      | 0.05                                 | 0.11 | 0.16 | 0.21 | 0.27 | 0.32 |
| 0.4      | 0.06                                 | 0.11 | 0.17 | 0.23 | 0.28 | 0.34 |
| 0.6      | 0.06                                 | 0.12 | 0.18 | 0.24 | 0.30 | 0.37 |
| 0.8      | 0.06                                 | 0.13 | 0.20 | 0.26 | 0.32 | 0.40 |
| 1.0      | 0.07                                 | 0.13 | 0.21 | 0.28 | 0.36 | 0.45 |
| 1.5      | 0.09                                 | 0.20 | 0.29 | 0.37 | 0.48 |      |

Example: If the result of Aluminium determination is 0.15 mg/l Al and the Fluoride concentration is known to be 0.4 mg/l F, the true concentration of Aluminium is 0.17 mg/l Al.



| Reagent                             | Form of reagent/Quantity | Order-No. |
|-------------------------------------|--------------------------|-----------|
| Set                                 |                          | 4535000   |
| VARIO Aluminium ECR F20             | Powder Pack / 100        |           |
| VARIO Aluminium Hexamine F 20       | Powder Pack / 100        |           |
| VARIO Aluminium ECR Masking Reagent | Liquid reagent / 25 ml   |           |





### Ammonia with Tablet

0.02 - 1 mg/l N



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one AMMONIA No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Add **one AMMONIA No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 Press TEST key.
 Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Ammonia as N.

#### Notes:

- 1. The tablets must be added in the correct sequence.
- 2. The AMMONIA No. 1 tablet will only dissolve completely after the AMMONIA No. 2 tablet has been added.
- 3. The temperature of the sample is important for full colour development. At a temperature below 20°C the reaction period is 15 minutes.
- 4. Sea water samples:

Ammonia conditioning reagent is required when testing sea water or brackish water samples to prevent precipitation of salts.

Fill the test tube with the sample to the 10 ml mark and add one level spoonful of Conditioning Powder. Mix to dissolve, then continue as described in the test instructions.

5. Conversion:

$$mg/l NH_4 = mg/l N \times 1.29$$
  
 $mg/l NH_3 = mg/l N \times 1.22$ 

6. ▲ N NH.

▼ NH<sub>3</sub>

| Reagent  | Form of reagent/Quantity                | Order-No. |
|--|---|-----------|
| Set<br>AMMONIA No. 1 / No. 2                     | Tablet / per 100 inclusive stirring rod | 4517611BT |
| AMMONIA No. 1                                    | Tablet / 100                            | 4512580BT |
| AMMONIA No. 2                                    | Tablet / 100                            | 4512590BT |
| Ammonia conditioning reagent (Sea water samples) | (approx. 50 tests)<br>powder / 15 g     | 460170    |





### **Ammonia** with Vario Powder Pack

0.01 - 0.8 mg/l N



Ø 24 mm



Countdown 1 3:00

start: 🔟

### Countdown 2 15:00 start: 🚽

prepare Zero press ZERO

Zero accepted prepare Test press TEST

Use two clean vials (24 mm Ø) and mark one as blank for zeroina.

- 1. Fill a clean vial (24 mm Ø) with 10 ml of deionised water (this is the blank)
- 2. Fill the other clean vial (24 mm Ø) with 10 ml of the water sample (this is the sample).
- 3. Add the contents of one Vario Ammonia Salicylate F10 Powder Pack straight from the foil to each vial.
- 4. Close the vials with the caps and shake to mix the contents.
- 5. Press [ ] key. Wait for a reaction period of 3 minutes.

After the reaction period is finished proceed as follows:

- 6. Add the contents of one Vario Ammonia Cyanurate F10 Powder Pack straight from the foil to each sample.
- 7. Close the vials tightly with the caps and shake to mix the contents
- 8. Press [ ] key. Wait for a reaction period of 15 minutes.

After the reaction period is finished proceed as follows:

- 9. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 10. Press ZERO key.
- 11. Remove the vial from the sample chamber.
- 12. Place the vial (the sample) in the sample chamber making sure that the  $\chi$  marks are aligned.
- 13. Press **TEST** key.

The result is shown in the display in mg/l Ammonia as N.

#### Notes:

- 1. Extremely basic or acidic water samples should be adjusted with 0.5 mol/l (1 N) Sulfuric acid solution or 1 mol/l (1 N) Sodium hydroxide solution to pH 7.
- 2. Interferences:

| Interfering substance                    | Interference levels and treatments   |
|--|--|
| Calcium                                  | greater than 1000 mg/l CaCO <sub>3</sub>   |
| Iron                                     | Interferes at all levels. Correct as follows:  a) determine the concentration of iron present in the sample by performing a total Iron test b) add the same iron concentration as determined to the deionised water (step 1). The interference will be blanked out successfully. |
| Magnesium                                | greater than 6000 mg/l CaCO <sub>3</sub>   |
| Nitrate                                  | greater than 100 mg/l NO <sub>3</sub> -N   |
| Nitrite                                  | greater than 12 mg/l NO <sub>2</sub> -N  |
| Phosphate                                | greater than 100 mg/l PO <sub>4</sub> -P   |
| Sulfate                                  | greater than 300 mg/l SO <sub>4</sub>  |
| Sulfide                                  | intensifies the colour   |
| Glycine, Hydrazine,<br>Colour, Turbidity | Less common interferences such as Hydrazine and Glycine will cause intensified colours in the prepared sample. Turbidity and colour will give erroneous high values. Samples with severe interferences require distillation.   |

3. **A** N

 $NH_4$ 

▼ NH<sub>3</sub>

| Reagent  | Form of reagent/Quantity | Order-No. |
|--|--------------------------|-----------|
| Set VARIO Ammonia Salicylate F10 VARIO Ammonia Cyanurate F10 | Powder Pack / per 100 PP | 4535500   |





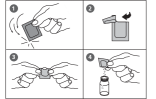
# Ammonia LR with Vario Tube Test

0.02 - 2.5 mg/l N



Insert the adapter for 16 mm Ø vials.

- Open one white capped reaction vial and add 2 ml deionised water (this is the blank).
- 2. Open another white capped reaction vial and add **2 ml of the water sample** (this is the sample).



- Add the contents of one Vario Ammonia Salicylate F5 Powder Pack straight from the foil into each vial.
- Add the contents of one Vario Ammonia Cyanurate
   F5 Powder Pack straight from the foil into each vial.
- 5. Close the vials tightly with the caps and swirl several times to dissolve the powder.

Countdown 1 20:00 start:

Press [ ] key.
 Wait for a reaction period of 20 minutes.

After the reaction period is finished proceed as follows:

7. Place the vial (the blank) in the sample chamber making sure that the marks are  $\lambda$  aligned.

prepare Zero press ZERO

- 8. Press **ZERO** key.
- 9. Remove the vial from the sample chamber.
- 10. Place the vial (the sample) in the sample chamber making sure that the marks are  $\frac{1}{\lambda}$  aligned.

Zero accepted prepare Test press TEST

11. Press TEST key.

The result is shown in the display in mg/l Ammonia as N.

#### Notes:

- 1. Strong alkaline or acidic water samples must be adjusted to approx. pH 7 before analysis (use 1 mol/l Hydrochloric acid resp. 1 mol/l Sodium hydroxide).
- 2. Iron interferes with the test. The interferences will be eliminated as follows:

  Determine the amount of total iron present in the water sample. To produce the blank add an iron standard solution with the same iron concentration to the vial (point 1) instead of deionised water
- 3. Conversion:

 $mg/l NH_4 = mg/l N x 1.29$  $mg/l NH_3 = mg/l N x 1.22$ 

4. ▲ N
NH<sub>4</sub>
NH<sub>3</sub>

| Reagent                     | Form of reagent/Quantity | Order-No. |
|-----------------------------|--------------------------|-----------|
| Set                         | Set                      | 4535600   |
| VARIO Ammonia Salicylate F5 | Powder Pack / 50         |           |
| VARIO Ammonia Cyanurate F5  | Powder Pack / 50         |           |
| VARIO Am Diluent Reagent LR | Reaction tube / 50       |           |
| VARIO deionised water       | 100 ml                   |           |





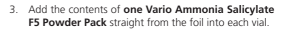
# Ammonia HR with Vario Tube Test

1 - 50 mg/l N



Insert the adapter for 16 mm Ø vials.

- Open one white capped reaction vial and add 0.1 ml deionised water (this is the blank).
- Open another white capped reaction vial and add0.1 ml of the water sample (this is the sample).





5. Close the vials tightly with the caps and swirl several times to dissolve the powder.



Countdown 1 20:00 start:

Press [4] key.
 Wait for a reaction period of 20 minutes.

After the reaction period is finished proceed as follows:

7. Place the vial (the blank) in the sample chamber making sure that the marks are  $\lambda$  aligned.

#### prepare Zero press ZERO

- 8. Press ZERO key.
- 9. Remove the vial from the sample chamber.
- 10. Place the vial (the sample) in the sample chamber making sure that the marks are  $\lambda$  aligned.

Zero accepted prepare Test press TEST

11. Press TEST key.

The result is shown in the display in mg/l Ammonia as N.

#### Notes:

- 1. Strong alkaline or acidic water samples must be adjusted to approx. pH 7 before analysis (use 1 mol/l Hydrochloric acid resp. 1 mol/l Sodium hydroxide).
- 2. If chlorine is known to be present, add one drop of 0.1 mol/l Sodium thiosulfate for each 0.3 mg/l Cl, in a one litre water sample.
- 3. Iron interferes with the test. The interferences will be eliminated as follows:

  Determine the amount of total iron present in the water sample. Add an iron standard solution with the same concentration to the vial (point 1) instead of deionised water to produce the blank.
- 4. Conversion:

 $mg/l NH_4 = mg/l N \times 1.29$  $mg/l NH_3 = mg/l N \times 1.22$ 

5. A N

▼ NH<sub>4</sub> NH<sub>3</sub>

| Reagent  | Form of reagent/Quantity                         | Order-No. |
|--|--|-----------|
| Set<br>VARIO Ammonia Salicylate F5   | Set<br>Powder Pack / 50                          | 4535650   |
| VARIO Ammonia Cyanurate F5<br>VARIO Am Diluent Reagent HR<br>VARIO deionised water | Powder Pack / 50<br>Reaction tube / 50<br>100 ml |           |





### Boron with Tablet

0.1 - 2 mg/l B



1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

### 3. Press **ZERO** key.

- 4. Remove the vial from the sample chamber.
- Add one BORON No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod and dissolve the tablet.
- 6. Add **one BORON No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### Zero accepted prepare Test press TEST

Countdown 20:00

### 9. Press **TEST** key.

Wait for a reaction period of 20 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Boron.

#### Notes:

- 1. The tablets must added in the correct sequence.
- 2. The sample solution should have a pH value between 6 and 7.
- 3. Interferences are prevented by the presence of EDTA in the tablets.
- 4. The rate of colour development depends on the temperature. The temperature of the sample must be 20°C  $\pm$  1°C.



| Reagent                  | Form of reagent/Quantity                | Order-No. |
|--------------------------|---|-----------|
| Set<br>Bor No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517681BT |
| BORON No. 1              | Tablet / 100                            | 4515790   |
| BORON No. 2              | Tablet / 100                            | 4515800BT |





# Bromine with Tablet

0.05 - 13 mg/l Br<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- 5. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod (note 5).
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

9. Press TEST key.

The result is shown in the display in mg/l Bromine.

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Bromine may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand. Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) forone hour, then rinse all glassware thoroughly with deionised water.

- 2. Preparing the sample:
  - When preparing the sample, the lost of Bromine, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment. Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the reagent is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range:
  - Concentrations above 22 mg/l Bromine can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Bromine. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. Depending on the preparation of the dosed bromine, bromine compounds may not react completely with the DPD No.1 tablet. In this case, the DPD No.3 tablet should be added under observation with a reaction time of 2 minutes. Please follow the directions of the bromine compound manufacturer where necessary.
- 6. Oxidising agents such as Chlorine, Ozone etc. interfere as they react in the same way as Bromine.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| DPD No. 1 | Tablet / 100             | 4511050BT |





### Bromine with Vario Powder Pack

 $0.05 - 4.5 \text{ mg/l Br}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.



- Add the contents of one VARIO Chlorine TOTAL-DPD / F10 Powder Pack straight from the foil to the water sample (note 5).
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 3:00

8. Press **TEST** key. Wait for a **reaction period of 3 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Bromine.

#### Notes:

- 1. Vial cleaning:
  - As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Bromine may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand. Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) forone hour, then rinse all glassware thoroughly with deionised water.
- Preparing the sample: When preparing the sample, the lost of Bromine, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment. Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the reagent is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range: Concentrations above 4.5 mg/l Bromine can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Bromine. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. Alternatively a VARIO Chlorine FREE-DPD / F10 powder pack may be used for the determination of some bromine compounds. Please follow the directions of the bromine compound manufacturer where necessary.
- 6. Oxidising agents such as Chlorine, Ozone etc. interfere as they react in the same way as Bromine

| Reagent                      | Form of reagent/Quantity | Order-No. |
|------------------------------|--------------------------|-----------|
| Vario Clorine Free-DPD/F10   | Powder Pack / 100        | 4530100   |
| VARIO Chlorine Total-DPD/F10 | Powder Pack / 100        | 4530120   |





### Chloride with Tablet

0.5 - 25 mg/l Cl



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one CHLORIDE T1 tablet straight from the foil to the water sample, crush the tablet using a clean stirring rod and dissolve the tablet.
- Add one CHLORIDE T2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl gently several times until the tablet is dissolved (Note 1).
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

9. Press **TEST** key.

Wait for a **reaction period of 2 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Chloride.

#### Notes:

1. Ensure that all particles of the tablet are dissolved – Chloride causes an extremely fine distributed turbidity with a milky appearance.

#### Heavy shaking leads to bigger sized particles which can cause false readings.

- 2. High concentrations of electrolytes and organic compounds have different effects on the precipitation reaction.
- 3. Ions which also form deposits with Silver nitrate in acidic media, such as Bromides, lodides and Thiocyanates, interfere with the analysis.
- 4. Highly alkaline water should if necessary be neutralised using Nitric acid before analysis.
- 5. Conversion: mg/l NaCl = mg/l Cl<sup>-</sup> x 1,65
- 6. CI NaCl

| Reagent                 | Form of reagent/Quantity                | Order-No. |
|-------------------------|---|-----------|
| Set<br>CHLORIDE T1 / T2 | Tablet / per 100 inclusive stirring rod | 4517741BT |
| CHLORIDE T1             | Tablet / 100                            | 4515910BT |
| CHLORIDE T2             | Tablet / 100                            | 4515920BT |





# Chloride with Liquid Reagent

 $0.5 - 20 \text{ mg/l Cl}^-$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

### 20 drops KS251 (Chloride Reagent A)

- 6. Close the vial tightly with the cap and invert several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 20 drops KS253 (Chloride Reagent B)

- 8. Close the vial tightly with the cap and invert several times to mix the contents.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

Press TEST key.Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Chloride.

#### Notes:

- 1. Chloride causes an extremely fine distributed turbidity with a milky appearance. **Heavy shaking leads to bigger sized particles which can cause false readings**.
- 2. Conversion:  $mg/l NaCl = mg/l Cl^{-} x 1,65$
- 3. CI NaCI

| Reagent                    | Form of reagent/Quantity | Order-No. |
|----------------------------|--------------------------|-----------|
| KS251 (Chloride Reagenz A) | Liquid reagent / 65 ml   | 56L025165 |
| KS253 (Chloride Reagenz B) | Liquid reagent / 65 ml   | 56L025365 |

Chlorine

>>

total











| Chlorine >> diff free total | The following selection is shown in the display:                           |
|-----------------------------|--|
| >> diff                     | for the differentiated determination of free, combined and total Chlorine. |
|                             |  |
| >> free                     | for the determination of free Chlorine.                                    |

Select the desired determination with the arrow keys [▲] and [▼]. Confirm with [₄] key.

for the determination of total Chlorine.

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Chlorine may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand.

- Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) for one hour, then rinse all glassware thoroughly with deionised water.
- 2. For individual testing of free and total Chlorine, the use of different sets of glassware is recommended (EN ISO 7393-2, 5.3)
- 3. Preparing the sample:
  - When preparing the sample, the lost of Chlorine, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 4. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagents therefore contain a buffer for the pH adjustment.
  - Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the reagent is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 5. Exceeding the measuring range:

Concentrations above:

- 10 mg/l Chlorine using tablets (method 100)
- 4 mg/l Chlorine using liquid reagents (method 101)
- 2 mg/l using powder packs (method 110)
- 8 mg/l using powder packs (method 111)
- can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Chlorine. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 6. Turbidity (can lead to errors):
  - The use of the DPD No. 1 tablet (method 100) in samples with high Calcium ion contents\* and/or high conductivity\* can lead to turbidity of the sample and therefore incorrect measurements. In this case, the reagent tablet DPD No. 1 High Calcium should be used as an alternative. If turbidity does occur after the DPD No. 3 tablet has been added, this can be prevented by using the DPD No. 1 High Calcium tablet and the DPD No. 3 High Calcium tablet.
  - The DPD No. 1 High Calcium should only be used in combination with the DPD No. 3 High Calcium.
  - \* it is not possible to give exact values, because the development of turbidity depends on the nature of the sample.
- 7. If ??? is displayed at a differentiated test result see page 342.
- 8. Oxidizing agents such as Bromine, Ozone etc. interfere as they react in the same way as Chlorine.







# Chlorine, free with Tablet

 $0.01 - 6 \text{ mg/l Cl}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty** it, leaving a few drops remaining in the vial.
- 5. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

9. Press TEST key.

The result is shown in the display in mg/l free Chlorine.

#### Notes:

See page 45







# Chlorine, total with Tablet

 $0.01 - 6 \text{ mg/l Cl}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- Add one DPD No. 1 tablet and one DPD No. 3 tablet straight from the foil and crush the tablets using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved
- 8. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00 Press TEST key.
 Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chlorine.

#### Notes:

See page 45







# Chlorine, differentiated determination with Tablet

0.01 - 6 mg/l Cl<sub>2</sub>



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- 5. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare T1 press TEST

- 9. Press **TEST** key.
- 10. Remove the vial from the sample chamber.
- Add one DPD No. 3 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 12. Close the vial tightly with the cap and swirl several times until the tablet is dissolved

13. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 2:00 14. Press **TEST** key.

Wait for a **reaction period of 2 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l free Cl \*,\*\* mg/l comb Cl \*,\*\* mg/l total Cl

mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

Notes: See page 45

| Reagent   | Form of reagent/Quantity                | Order-No. |
|---|---|-----------|
| Set<br>DPD No. 1 / No. 3                                  | Tablet / per 100 inclusive stirring rod | 4517711BT |
| DPD No. 1   | Tablet / 100                            | 4511050BT |
| DPD No. 3   | Tablet / 100                            | 4511080BT |
| Set<br>DPD No. 1 HIGH CALCIUM /<br>DPD No. 3 HIGH CALCIUM | Tablet / per 100 inclusive stirring rod | 4517781BT |
| DPD No. 1 HIGH CALCIUM                                    | Tablet / 100                            | 4515740BT |
| DPD No. 3 HIGH CALCIUM                                    | Tablet / 100                            | 4515730BT |







# Chlorine HR, free with Tablet

0.1 - 10 mg/l Cl<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- Remove the vial from the sample chamber and empty it, leaving a few drops remaining in the vial.
- 5. Add **one DPD No. 1 HR tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

9. Press **TEST** key.

The result is shown in the display in mg/l free Chlorine.

#### Notes:

See page 45







# Chlorine HR, total with Tablet

0.1 - 10 mg/l Cl<sub>3</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber and **empty** it, leaving a few drops remaining in the vial.
- Add one DPD No. 1 tablet and one DPD No. 3 tablet straight from the foil and crush the tablets using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00 9. Press **TEST** key. Wait for a **reaction period of 2 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chlorine.

Notes:

See page 45







# Chlorine HR, differentiated determination with Tablet

0.1 - 10 mg/l Cl<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- 5. Add **one DPD No. 1 HR tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare T1 press TEST

- 9. Press TEST key.
- 10. Remove the vial from the sample chamber.
- 11. Add **one DPD No. 3 HR tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.

12. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.

13. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 2:00

14. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l free Cl \*,\*\* mg/l comb Cl \*,\*\* mg/l total Cl mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

Notes: See page 45

| Reagent      | Form of reagent/Quantity | Order-No. |
|--------------|--------------------------|-----------|
| DPD No. 1 HR | Tablet / 100             | 4511500BT |
| DPD No. 3 HR | Tablet / 100             | 4511590BT |







# Chlorine, free with Liquid Reagent

0.02 - 4 mg/l Cl<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- Remove the vial from the sample chamber and empty the vial.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

6 drops of DPD 1 buffer solution 2 drops of DPD 1 reagent solution

- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times to mix the contents.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

9. Press TEST key.

The result is shown in the display in mg/l free Chlorine.

#### Notes (free and total Chlorine):

- 1. After use replace the bottle caps securely noting the colour coding.
- 2. Store the reagent bottles in a cool, dry place ideally between 6°C and 10°C.

3. Also see page 45







# Chlorine, total with Liquid Reagent

 $0.02 - 4 \text{ mg/l Cl}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty**the vial
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

6 drops of DPD 1 buffer solution 2 drops of DPD 1 reagent solution 3 drops of DPD 3 solution

- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times to mix the contents.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

Press TEST key.Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chlorine.







# Chlorine, differentiated determination with Liquid Reagent

 $0.02 - 4 \text{ mg/l Cl}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- Remove the vial from the sample chamber and empty the vial.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

6 drops of DPD 1 buffer solution2 drops of DPD 1 reagent solution

- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times to mix the contents.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare T1 press TEST

- 9. Press **TEST** key.
- 10. Remove the vial from the sample chamber.
- Add 3 drops of DPD 3 solution to the same water sample.
- 12. Close the vial tightly with the cap and swirl several times to mix the contents.

13. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 2:00 14. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l free Cl \*,\*\* mg/l comb. Cl \*,\*\* mg/l total Cl mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

#### Notes:

- 1. After use replace the bottle caps securely noting the colour coding.
- 2. Store the reagent bottles in a cool, dry place ideally between 6°C and 10°C.
- 3. Also see page 45

| Reagent   | Form of reagent/Quantity   | Order-No. |
|---|--|-----------|
| DPD No. 1 buffer solution<br>DPD No. 1 reagent solution<br>DPD No. 3 solution | (approx. 300 tests) 3 x Liquid reagent / 15 ml 1 x Liquid reagent / 15 ml 2 x Liquid reagent / 15 ml | 471056    |
| DPD No. 1 buffer solution   | Liquid reagent / 15 ml   | 471010    |
| DPD No. 1 reagent solution  | Liquid reagent / 15 ml   | 471020    |
| DPD No. 3 solution  | Liquid reagent / 15 ml   | 471030    |







# Chlorine, free with Vario Powder Pack

0.02 - 2 mg/l Cl<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.



- Add the contents of one VARIO Chlorine FREE-DPD / F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press **TEST** key.

The result is shown in the display in mg/l free Chlorine.

Notes:

See page 45







# Chlorine, total with Vario Powder Pack

 $0.02 - 2 \text{ mg/l Cl}_{3}$ 



- 1. Fill a clean vial (24 mm Ø) with **10 ml of the water** sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO



4. Remove the vial from the sample chamber.



- Add the contents of one VARIO Chlorine TOTAL-DPD / F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 3:00 8. Press **TEST** key. Wait for a **reaction period of 3 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chlorine.

#### Notes:

See page 45







# Chlorine, differentiated determination with Vario Powder Pack

 $0.02 - 2 \text{ mg/l Cl}_{2}$ 



- 1. Fill a clean vial (24 mm Ø) with **10 ml of the water** sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press ZERO key.



4. Remove the vial from the sample chamber.

- Add the contents of one VARIO Chlorine FREE-DPD/ F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare T1 press TEST

- 8. Press **TEST** key.
- 9. Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times and then fill the vial with **10 ml of the water sample**.
- Add the contents of one VARIO Chlorine TOTAL-DPD / F10 Powder Pack straight from the foil to the water sample.
- 11. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).

12. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 3:00

13. Press **TEST** key.

Wait for a **reaction period of 3 minutes.** 

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l free Cl \*,\*\* mg/l comb. Cl \*,\*\* mg/l total Cl mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

Notes: See page 45

| Reagent                      | Form of reagent/Quantity | Order-No. |
|------------------------------|--------------------------|-----------|
| Vario Clorine Free-DPD/F10   | Powder Pack / 100        | 4530100   |
| VARIO Chlorine Total-DPD/F10 | Powder Pack / 100        | 4530120   |



### Chlorine HR, free with Vario Powder Pack plastic vial (type 3) ⊔ 10 mm



 $0.1 - 8 \text{ mg/l Cl}_{2}$ 

- 1. Fill a clean vial (10 mm Ø) with 5 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.



- Add the contents of two Vario Chlorine Free-DPD/ F10 Powder Pack straight from the foil into the water sample.
- 6. Close the vial tightly with the cap and invert several times to mix the contents (20 sec.).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

8. Press the **TEST** key.

The result is shown in the display in mg/l free Chlorine.

#### Notes:

See page 45





# Chlorine HR, total with Vario Powder Pack plastic vial (type 3) □ 10 mm

0.1 - 8 mg/l Cl<sub>2</sub>

- 1. Fill a clean vial (10 mm Ø) with 5 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO





- 4. Remove the vial from the sample chamber.
- Add the contents of two Vario Chlorine TOTAL-DPD/ F10 Powder Pack straight from the foil into the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

### Countdown 3:00

Press TEST key. Wait for a reaction period of 3-6 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chlorine.

#### Notes:

See page 45









# Chlorine HR, differentiated determination with Vario Powder Pack plastic vial (type 3) □ 10 mm

0.1 - 8 mg/l Cl<sub>2</sub>

- 1. Fill a clean vial (10 mm Ø) with 5 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press ZERO key.



- 4. Remove the vial from the sample chamber.
- Add the contents of two Vario Chlorine Free-DPD/ F10 Powder Pack straight from the foil into the water sample.
- 6. Close the vial tightly with the cap and invert several times to mix the contents (20 sec.).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare T1 press TEST

- 8. Press the **TEST** key.
- 9. Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times and then fill the vial with 5 ml of the water sample.
- Add the contents of two Vario Chlorine TOTAL-DPD/ F10 Powder Pack straight from the foil into the water sample.

- 11. Close the vial tightly with the cap and invert several times to mix the contents (20 sec.).
- 12. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 3:00

13. Press TEST key.

Wait for a reaction period of 3 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l free Cl \*,\*\* mg/l comb. Cl \*,\*\* mg/l total Cl

mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

Notes: See page 45

| Reagent                      | Form of reagent/Quantity | Order-No. |
|------------------------------|--------------------------|-----------|
| Vario Clorine Free-DPD/F10   | Powder Pack / 100        | 4530100   |
| VARIO Chlorine Total-DPD/F10 | Powder Pack / 100        | 4530120   |







# Chlorine dioxide with Tablet

0.02 - 11 mg/l ClO<sub>2</sub>

#### Chlorine dioxide

>> with Cl without Cl

The following selection is shown in the display:

>> with Cl

for the determination of Chlorine dioxide in the presence of Chlorine.

>> without Cl

for the determination of Chlorine dioxide in the absence of Chlorine.

Select the desired determination with the arrow keys  $[\blacktriangle]$  and  $[\blacktriangledown]$ . Confirm with  $[\clubsuit]$  key.

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Chlorine dioxide may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand. Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) for one hour, then rinse all glassware thoroughly with deionised water.

- 2. Preparing the sample:
  - When preparing the sample, the lost of Chlorine dioxide, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment.

  Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the tablet is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range:
  - Concentrations above 19 mg/l Chlorine dioxide can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Chlorine dioxide. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. If ??? is displayed at a differentiated test result see page 342.
- 6. Oxidising agents such as Chlorine, Ozone etc. interfere as they react in the same way as Chlorine dioxide.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| DPD No. 1 | Tablet / 100             | 4511050BT |
| GLYCINE   | Tablet / 100             | 4512170BT |







# Chlorine dioxide in the presence of Chlorine with Tablet

0.02 - 11 mg/l CIO<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample.
- 2. Add **one GLYCINE tablet** straight from the foil and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 4. **Fill a second clean vial with 10 ml of water sample** and close tightly with the cap.
- 5. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 6. Press ZERO key.
- Remove the vial from the sample chamber and empty the vial.
- 8. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- Transfer the contents of the first vial (Glycine solution) into the prepared vial (point 8).
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare T1 press TEST

12. Press TEST key.

- 13. Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times. Fill with a few drops of water sample.
- 14. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 15. Add water sample to the 10 ml mark.
- 16. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 17. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

#### 18. Press **TEST** key.

- 19. Remove the vial from the sample chamber.
- Add one DPD No. 3 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 21. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 22. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### Press TEST key. Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

Chlorine dioxide in mg/l CIO<sub>2</sub>.

mg/l free Chlorine mg/l combined Chlorine mg/l total Chlorine

#### \*,\*\* mg/l ClO<sub>2</sub>

T2 accepted prepare T3

press TEST

Countdown

2:00

\*,\*\* mg/l free Cl \*,\*\* mg/l comb. Cl \*,\*\* mg/l total Cl

#### Notes:

See next page.

prepare T2 press TEST

T1 accepted

#### Notes: (Chlorine dioxide in the presence of Chlorine)

- 1. The conversion factor to convert Chlorine dioxide (display) to Chlorine dioxide as Chlorine is 2.6315.
  - $mg/I CIO_{2} [CI] = mg/I CIO_{2} \cdot 2,6315$
  - Chlorine dioxide displayed as Chlorine units CIO<sub>2</sub> [CI] has its origin in swimming poolwater treatment according to DIN 19643.
- The total Chlorine result given includes the contribution of the chlorine dioxide as Chlorine reading. For true Chlorine value add the free and combined Chlorine readings.
- 3. See also page 67.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| DPD No. 1 | Tablet / 100             | 4511050BT |
| DPD No. 3 | Tablet / 100             | 4511080BT |
| GLYCINE   | Tablet / 100             | 4512170BT |







# Chlorine dioxide in absence of Chlorine with Tablet

0.02 - 11 mg/l ClO<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- 5. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

9. Press **TEST** key.

\*,\*\* mg/l ClO,

The result is shown in the display as Chlorine dioxide in mg/l ClO<sub>2</sub>.

#### Notes:

See page 67







#### Chlorine dioxide in absence of Chlorine with Vario Powder Pack

0.04 - 3.8 mg/l CIO<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press **ZERO** key.



- 4. Remove the vial from the sample chamber.
- Add the contents of one VARIO Chlorine FREE-DPD / F10 Powder Pack straight from the foil to the water sample (Note 5).
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display in mg/l Chlorine dioxide

Notes:

See page 74







# Chlorine dioxide in the presence of Chlorine with Vario Powder Pack

0.04 - 3.8 mg/l CIO<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one GLYCINE tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl gently several times until the tablet is dissolved



- Add the contents of one VARIO Chlorine FREE-DPD / F10 Powder Pack straight from the foil into the pre prepared vial.
- 8. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 20 seconds).
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

10. Press **TEST** key.

The result is shown in the display in mg/l Chlorine dioxide.

#### Notes:

See page 74

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Chlorine dioxide may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand. Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) for one hour, then rinse all glassware thoroughly with deionised water.

- 2. Preparing the sample:
  - When preparing the sample, the lost of Chlorine dioxide, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment.

  Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the tablet is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range:
  - Concentrations above 3.8 mg/l Chlorine dioxide can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Chlorine dioxide. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. Oxidising agents such as Chlorine, Ozone etc. interfere as they react in the same way as Chlorine dioxide.

| Reagent                    | Form of reagent/Quantity | Order-No. |
|----------------------------|--------------------------|-----------|
| Vario Clorine Free-DPD/F10 | Powder Pack / 100        | 4530100   |
| GLYCINE                    | Tablet / 100             | 4512170BT |







# Chlorine HR (KI) with Tablet

5 - 200 mg/l Cl<sub>2</sub>



Insert the adapter for 16 mm Ø vials.

- 1. Fill a clean vial (16 mm Ø) with **8 ml of the water sample**, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks are √ aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one CHLORINE HR (KI) tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Add one ACIDIFYING GP tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the marks are  $\lambda$  aligned.

Zero accepted prepare Test press TEST

9. Press **TEST** key.

The result is shown in the display in mg/l Chlorine.

#### Notes:

1. Oxidizing agents interfere as they react in the same way as Chlorine.

| Reagent                                | Form of reagent/Quantity                | Order-No. |
|--|---|-----------|
| Set ACIDIFYING GP/<br>CHLORINE HR (KI) | Tablet / per 100 inclusive stirring rod | 4517721BT |
| CHLORINE HR (KI)                       | Tablet / 100                            | 4513000BT |
| ACIDIFYING GP                          | Tablet / 100                            | 4515480BT |







## Chlorite in presence of Chlorine and Chlorine dioxide

0,01 - 6 mg/l Cl<sub>2</sub>

Firstly, the glycine method is used to measure the concentration of Chlorine Dioxide. This is then followed by the determination of the free and total chlorine, from which the Combined Chlorine can be calculated. A third test is performed which measures the Total Chlorine concentration plus any Chlorite present. Finally, the Chlorite concentration can be calculated from the three recorded results.

# Chlorine >> diff free

total

>> free

The following selection is shown in the display:

select for the determination of free Chlorine



- 1. Fill a clean vial with 10 ml of water sample.
- Add one GLYCINE tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 3. Close the vial tightly with the cap and swirl gently several times until the tablet is dissolved.
- 4. **Fill a second clean vial with 10 ml of water sample,** close tightly with the cap.
- 5. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 6. Press ZERO key.
- 7. Remove the vial from the sample chamber and empty the vial.
- 8. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.

- Transfer the contents of the first vial (Glycine solution) into the prepared vial (point 8).
- 10. Close the vial tightly with the cap and swirl several times until the tablets are dissolved
- 11. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

Zero accepted prepare Test press TEST

12. Press **TEST** key.

Record the displayed test result (G).

- 13. Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times. Fill with **a few drops of water sample**.
- 14. Add **one DPD No. 1 tablet** straight from the foil and crush the tablet using a clean stirring rod.
- 15. Add water sample to the 10 ml mark.
- 16. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 17. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

Zero accepted prepare Test press TEST

18. Press **TEST** key.

Record the displayed test result (A).

- 19. Remove the vial from the sample chamber.
- Add one DPD No. 3 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.

- 21. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 22. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 23. Wait for a reaction period of 2 minutes.

Zero accepted prepare Test press TEST

24. Press **TEST** key.

Record the displayed test result (C).

- 25. Remove the vial from the sample chamber.
- Add one DPD ACIDIFYING tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 27. Wait for a reaction period of 2 minutes.
- 28. Add **one DPD NEUTRALISING tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 29. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 30. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

31. Press TEST key.

Record the displayed test result (D).

#### Calculations:

mg/l Chlorine dioxide = result G x 1,9 mg/l free Chlorine = result A – result G mg/l combined Chlorine = result C – result A

mg/l Chlorite = result D – ( result C +  $4 \times 10^{-1}$  x result G )

#### **Tolerances:**

1. By calculation of non direct analysable parameters it is necessary to consider the error propagation besed on the possible tolerances of the single test tesults.

2. see Notes Chlorine, page 45.

| Reagent                  | Form of reagent/Quantity                | Order-No. |
|--------------------------|---|-----------|
| Set<br>DPD No. 1 / No. 3 | Tablet / per 100 inclusive stirring rod | 4517711BT |
| DPD No. 1                | Tablet / 100                            | 4511050BT |
| DPD No. 3                | Tablet / 100                            | 4511080BT |
| GLYCINE                  | Tablet / 100                            | 4512170BT |
| DPD ACIDIFYING           | Tablet / 100                            | 4512120   |
| DPD NEUTRALISING         | Tablet / 100                            | 4511020BT |





### **↑** Chromium with Powder Pack

0.02 - 2 mg/l Cr

Chrom
>> diff
Cr (VI)

Cr (III + VI)

The following selection is shown in the display:

>> diff

for the differentiated determination of Chromium (VI), Chromium (III) and total Chromium

>> Cr (VI)

for the determination of Chromium (VI)

>> Cr (III + VI)

for the determination of total Chromium (sum Cr (III) + Cr (VI))

Select the desired determination with the arrow keys [A] and [V]. Confirm with the [L] key.

#### Note:

1. If ??? is displayed at the diffentiated test result see page 342.







## Chromium, differentiated determination with Powder Pack

0.02 - 2 mg/l Cr



#### Digestion:

- 1. Fill a clean vial (16 mm Ø) with **10 ml of water sample**.
- Add the contents of one PERSULF.RGT FOR CR Powder Pack straight from the foil into the vial.
- 3. Close the vial tightly with the cap and swirl several times to mix the contents.
- Heat the vial for 120 minutes in a preheated thermoreactor at a temperature of 100°C.
- Remove the vial from the thermoreactor. (CAUTION: The vials are hot!).
   Invert the vial and allow to cool to room temperature.

#### Performing test procedure:

Insert the adapter for 16 mm Ø vials.

6. Place the pre prepared vial in the sample chamber making sure that the marks  $\frac{1}{\Lambda}$  are aligned.

#### prepare Zero press ZERO

- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.
- Add the contents of one CHROMIUM HEXAVALENT Powder Pack straight from the foil into the prepared vial.
- 10. Close the vial tightly with the cap and swirl several times to mix the contents.
- 11. Place the vial in the sample chamber making sure that the marks  ${\stackrel{L}{\Lambda}}$  are aligned.
- 12. Press TEST key.

press TEST Countdown 5:00

Zero accepted prepare T1

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.



13. Fill a second clean vial (16 mm Ø) with 10 ml of the water sample.

14. Add the contents of one CHROMIUM HEXAVALENT Powder Pack straight from the foil to the water sample.

15. Close the vial tightly with the cap and swirl several times to mix the contents.

16. Place the vial in the sample chamber making sure that the marks  $\frac{1}{\lambda}$  are aligned.

T1 accepted prepare T2 press TEST

Countdown 5:00

17. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l Cr (VI) \*,\*\* mg/l Cr (III)

\*,\*\* mg/l Cr (iii)

mg/l Cr (VI)

mg/l Cr (III)

mg/l Cr total Chromium

#### Notes:

- 1. Performing steps 1–12 determines concentration of total chromium and steps 13–17 determines concentration of Chromium (VI). The concentration of Chromium (III) results out of the difference.
- 2. pH value of the water sample should be between 3 and 9.
- 3. For information about interferences especially in waste water and chemical waste water through metals and reductive or oxidic agents see DIN 38 405 D 24 and Standard Methods of Water and Wastewater, 20th Edition; 1998.

| Reagent             | Form of reagent/Quantity | Order-No. |
|---------------------|--------------------------|-----------|
| PERSULF.RGT FOR CR  | Powder Pack / 100        | 4537300   |
| CHROMIUM HEXAVALENT | Powder Pack / 100        | 4537310   |







# Chromium (VI) with Powder Pack

0.02 - 2 mg/l Cr



Insert the adapter for 16 mm Ø vials.

- Fill a clean vial (16 mm Ø) with 10 ml of the water sample.
- 2. Place the vial in the sample chamber making sure that the marks  $\frac{1}{\lambda}$  are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add the contents of one CHROMIUM HEXAVALENT Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the marks  $\frac{1}{\lambda}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Chromium (VI).

Notes: see previous page

| Reagent             | Form of reagent/Quantity | Order-No. |
|---------------------|--------------------------|-----------|
| PERSULF.RGT FOR CR  | Powder Pack / 100        | 4537300   |
| CHROMIUM HEXAVALENT | Powder Pack / 100        | 4537310   |







# Chromium, total (Cr(III) + Cr(VI)) with Powder Pack

0.2 - 2 mg/l Cr



#### Digestion:

- 1. Fill a clean vial (16 mm Ø) with **10 ml of water sample**.
- Add the contents of one PERSULF.RGT FOR CR Powder Pack straight from the foil into the vial.
- 3. Close the vial tightly with the cap and swirl several times to mix the contents
- Heat the vial for 120 minutes in a preheated thermoreactor at a temperature of 100°C.
- Remove the vial from the thermoreactor. (CAUTION: The vials are hot!).
   Invert the vial and allow to cool to room temperature.

#### Performing test procedure:

Insert the adapter for 16 mm Ø vials.

- 6. Place the pre prepared vial in the sample chamber making sure that the marks  $\frac{1}{\Lambda}$  are aligned.
- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.
- Add the contents of one CHROMIUM HEXAVALENT Powder Pack straight from the foil to the water sample.
- 10. Close the vial tightly with the cap and swirl several times to mix the contents.
- 11. Place the vial in the sample chamber making sure that the marks  $\frac{1}{\Lambda}$  are aligned.
- Press TEST key.
   Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Chromium.

#### prepare Zero press ZERO

Zero accepted prepare Test press TEST

Countdown 5:00







## COD LR with Vario Tube Test

 $0 - 150 \text{ mg/l } O_{2}$ 



Insert the adapter for 16 mm Ø vials.

- Open one white capped reaction vial and add 2 ml deionised water (this is the blank (Note 1)).
- Open another white capped reaction vial and add 2 ml of the water sample (this is the sample).
- 3. Close the vials with the cap tightly. Invert the vial gently several times to mix the contents.

(CAUTION: The vial will become hot during mixing!)

- Heat the vials for 120 minutes in the preheated reactor at a temperature of 150°C.
- 5. (CAUTION: The vials are hot!)

Remove the tubes from the heating block and allow them to cool to  $60^{\circ}\text{C}$  or less. Mix the contents by carefully inverting each tube several times while still warm. Then allow the tubes to cool to ambient temperature before measuring. (Note 2).

6. Place the vial (the blank (Note 3, 4)) in the sample chamber making sure that the marks are  $\frac{1}{\lambda}$  aligned.

### prepare Zero press ZERO

- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.
- 9. Place the vial (the sample (Note 3, 4)) in the sample chamber making sure that the marks are  $\lambda$  aligned.

Zero accepted prepare Test press TEST

10. Press **TEST** key.

The result is shown in the display in mg/l COD.

#### Notes:

- 1. Run samples and blanks with the same batch of vials. The blank is stable when stored in the dark and can be used for further measurements with vials of the same batch.
- 2. Do not place the hot vials in the sample chamber. Cool the vials to room temperature for final measurements.
- 3. Suspended solids in the vial lead to incorrect measurements. For this reason it is important to place the vials carefully in the sample chamber. The precipitate at the bottom of the sample should be not suspended.
- 4. Clean the outside of the vials with a towel. Finger prints or other marks will be removed.
- 5. Samples can be measured when the Chloride content does not exceed 1000 mg/l.
- 6. In exceptional cases, compounds contained in the water cannot be oxidized adequately, so results may be lower than reference methods.

| Reagent      |              | Form of reagent/Quantity | Order-No. |
|--------------|--------------|--------------------------|-----------|
| CSB VARIO LR | 0 - 150 mg/l | 1 Set (25 tests)         | 420720    |







#### COD MR with Vario Tube Test

 $0 - 1500 \text{ mg/l } O_{3}$ 



Insert the adapter for 16 mm Ø vials.

- Open one white capped reaction vial and add 2 ml deionised water (this is the blank (Note 1)).
- Open another white capped reaction vial and add 2 ml of the water sample (this is the sample).
- Close the vials with the cap tightly. Invert the vial gently several times to mix the contents.
   (CAUTION: The vial will become hot during mixing!)

4. Heat the vials for **120 minutes** in the preheated reactor

#### 5. (CAUTION: The vials are hot!)

at a temperature of 150°C.

Remove the tubes from the heating block and allow them to cool to  $60^{\circ}\text{C}$  or less. Mix the contents by carefully inverting each tube several times while still warm. Then allow the tubes to cool to ambient temperature before measuring. (Note 2).

6. Place the vial (the blank (Note 3, 4)) in the sample chamber making sure that the marks are  $\frac{1}{\lambda}$  aligned.

### prepare Zero press ZERO

- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.

Zero accepted prepare Test press TEST

10. Press **TEST** key.

The result is shown in the display in mg/l COD.

#### Notes:

- 1. Run samples and blanks with the same batch of vials. The blank is stable when stored in the dark and can be used for further measurements with vials of the same batch.
- Do not place the hot vials in the sample chamber. Cool the vials to room temperature for final measurements.
- 3. Suspended solids in the vial lead to incorrect measurements. For this reason it is important to place the vials carefully in the sample chamber. The precipitate at the bottom of the sample should be not suspended.
- 4. Clean the outside of the vials with a towel. Finger prints or other marks will be removed.
- 5. Samples can be measured when the Chloride content does not exceed 1000 mg/l.
- 6. In exceptional cases, compounds contained in the water cannot be oxidized adequately, so results may be lower than reference methods.
- 7. For samples under 100 mg/l COD it is recommended to repeat the test with the tube test for COD LR.

| Reagent                    | Form of reagent/Quantity | Order-No. |
|----------------------------|--------------------------|-----------|
| CSB VARIO MR 0 - 1500 mg/l | 1 Set (25 tests)         | 420721    |







# COD HR with Vario Tube Test

 $0 - 15 \text{ g/l O}_{2} (\triangleq 0 - 15 000 \text{ mg/l O}_{2})$ 



Insert the adapter for 16 mm Ø vials.

- Open one white capped reaction vial and add 0.2 ml deionised water (this is the blank (Note 1)).
- Open another white capped reaction vial and add 0.2 ml of the water sample (this is the sample).
- 3. Close the vials with the cap tightly. Invert the vial gently several times to mix the contents.

(CAUTION: The vial will become hot during mixing!)

 Heat the vials for 120 minutes in the preheated reactor at a temperature of 150°C.

#### 5. (CAUTION: The vials are hot!)

Remove the tubes from the heating block and allow them to cool to  $60^{\circ}\text{C}$  or less. Mix the contents by carefully inverting each tube several times while still warm. Then allow the tubes to cool to ambient temperature before measuring. (Note 2).

6. Place the vial (the blank (Note 3, 4)) in the sample chamber making sure that the marks are  $\frac{1}{\lambda}$  aligned.

### prepare Zero press ZERO

- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.
- 9. Place the vial (the sample (Note 3, 4)) in the sample chamber making sure that the marks are  $\frac{1}{\Lambda}$  aligned.

Zero accepted prepare Test press TEST

10. Press **TEST** key.

The result is shown in the display in **q/l** COD.

#### Notes:

- 1. Run samples and blanks with the same batch of vials. The blank is stable when stored in the dark and can be used for further measurements with vials of the same batch.
- 2. Do not place the hot vials in the sample chamber. Cool the vials to room temperature for final measurements.
- 3. Suspended solids in the vial lead to incorrect measurements. For this reason it is important to place the vials carefully in the sample chamber. The precipitate at the bottom of the sample should be not suspended.
- 4. Clean the outside of the vials with a towel. Finger prints or other marks will be removed.
- 5. Samples can be measured when the Chloride content does not exceed 10 000 mg/l.
- 6. In exceptional cases, compounds contained in the water cannot be oxidized adequately, so results may be lower than reference methods.
- 7. For samples under 1 g/l COD it is recommended to repeat the test with the test kit for COD MR or for samples under 0,1 g/l COD with the tube test COD LR.

| Reagent                  |            | n of reagent/Quantity | Order-No. |
|--------------------------|------------|-----------------------|-----------|
| CSB VARIO HR 0 - 15000 r | ng/l 1 Set | t (25 tests)          | 420722    |







# Colour, true and apparent (APHA Platinum-Cobalt Standard Method)

0 - 500 Pt-Co units

#### Sample preparation (Note 4):

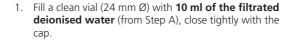
#### Step A

Filter approx. 50 ml deionised water through a membrane filter with a pore width of 0.45  $\mu m$ .

Discard the filtrate. Filter another **50 ml deionised water** and keep it for zeroing.

#### Step B

Filter approx.  $\bf 50~ml$  water sample using the same filter. Keep this filtrate for sample measurement.



2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and empty it completely.
- Rinse the vial with the filtrated water sample and fill with 10 ml filtrated water sample (from Step B).
- 6. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

# Zero accepted prepare Test press TEST

7. Press **TEST** key.

The result is shown in the display in Pt-Co units.

#### Notes:

- 1. This colour scale was originally developed by A. Hazen as a visual comparison scale. It is therefore necessary to ascertain whether the extinction maximum of the water sample is in the range from 420 to 470 nm, as this method is only suitable for water samples with yellowish to yellowish-brown coloration. Where applicable, a decision should be made based on visual inspection of the water sample.
- 2. This method 204 Colour (Hazen) is calibrated on the basis of the standards specified by "Standard Methods for the Examination of Water and Wastewater" (also see EN ISO 7887:1994)
  - 1 Pt-Co colour unit = 1 mg/L of platinum as chloroplatinate ion
- 3. The estimated detection limit is 15 mg/L Pt.
- 4. Colour may be expressed as "apparent" or "true" colour. The apparent colour is defined as the colour of a solution due to dissolved substances and suspended particles in the sample. This manual describes the determination of true colour by filtration of the water sample. To determine the apparent colour, non-filtrated deionised water and sample are measured.
- 5. Sample collection, preservation and storage:
  Pour the water sample into clean glass or plastic containers and analyse as soon as possible after the sample is taken. If this is not possible, fill the container right up to the top and seal tightly. Do not stir the sample; avoid lengthy contact with the air.
  The sample may be stored in a dark place at a temperature of 4°C for 24 hours. Before performing measurements, the water sample must be brought up to room temperature.







# Copper with Tablet

0.05 - 5 mg/l Cu

Copper >> dif

diff free total The following selection is shown in the display:

>> diff

for the differentiated determination of free, combined and total Copper.

>> free

for the determination of free Copper.

>> total

for the determination of total Copper.

Select the desired determination with the arrow keys [A] and [V]. Confirm with [A] key.

#### Note:

1. If ??? is displayed at the diffentiated test result see page 342.

| Reagent                     | Form of reagent/Quantity                | Order-No. |
|-----------------------------|---|-----------|
| Set<br>COPPER No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517691BT |
| COPPER No. 1                | Tablet / 100                            | 4513550BT |
| COPPER No. 2                | Tablet / 100                            | 4513560BT |







#### Copper, differentiated determination with Tablet

0.05 - 5 mg/l Cu



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water **sample**, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Add **one COPPER No. 1 tablet** straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

#### Zero accepted prepare T1 press TEST

- 8. Press **TEST** key.
- 9. Remove the vial from the sample chamber.
- 10. Add one COPPER No. 2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 11. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 12. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

The result is shown in the display in:

13. Press TEST key.

T1 accepted prepare T2 press TEST

\*,\*\* mg/l free Cu \*,\*\* mg/l comb Cu \*,\*\* mg/l total Cu

mg/l combined Copper mg/l total Copper

AL400\_8f 02/2016 97

mg/l free Copper







## Copper, free with Tablet

0.05 - 5 mg/l Cu



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press **ZERO** key.

- 4. Remove the vial from the sample chamber.
- Add one COPPER No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

8. Press **TEST** key.

The result is shown in the display in mg/l free Copper.







# Copper, total with Tablet

0.05 - 5 mg/l Cu



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one COPPER No. 1 tablet and one COPPER No. 2 tablet straight from the foil to the water sample and crush the tablets using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display in mg/l total Copper.







# Copper with Liquid reagent and powder

0.05 - 4 mg/l Cu

Copper
>> diff
free
total

The following selection is shown in the display:

>> diff

for the differentiated determination of free, combined and total Copper.

>> free

for the determination of free Copper.

>> total

for the determination of total Copper.

Select the desired determination with the arrow keys [A] and [V]. Confirm with [L] key.

#### Notes:

1. For correct dosage the spoon supplied with the reagents must be used.

2. If ??? is displayed at the diffentiated test result see page 342.

| Reagent                     | Form of reagent/Quantity | Order-No. |
|-----------------------------|--------------------------|-----------|
| KS240 – Coppercol Reagent 1 | Liquid reagent / 30 ml   | 56L024030 |
| KS241 – Coppercol Reagent 2 | Liquid reagent / 30 ml   | 56L024130 |
| KP242 – Coppercol Reagent 3 | Powder / 10 g            | 56L024210 |
| COPPER No. 2                | Tablet / 100             | 4513560BT |



prepare Zero

press ZERO





#### Copper, differentiated determination with Liquid reagent and powder

0.05 - 4 mg/l Cu



Ø 24 mm

1. Fill a clean vial (24 mm Ø) with **10 ml of the water sample**, close tightly with the cap.

- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS240 (Coppercol Reagent 1)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS241 (Coppercol Reagent 2)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents
- 9. Add 1 level spoon of reagent KP242 (Coppercol Reagent 3) (note 1, page 100).
- 10. Close the vial tightly with the cap and swirl several times to dissolve the powder.
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

12. Press **TEST** key.

Zero accepted prepare T1 press TEST

- 13. Remove the vial from the sample chamber.
- 14. Add **one COPPER No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 15. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 16. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

17. Press **TEST** key.

\*,\*\* mg/l free Cu \*,\*\* mg/l comb Cu

\*,\*\* mg/l comb Cu \*,\*\* mg/l total Cu The result is shown in the display in:

mg/l free Copper mg/l combined Copper mg/l total Copper







## Copper, free with Liquid reagent and powder

0.05 - 4 mg/l Cu



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS240 (Coppercol Reagent 1)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS241 (Coppercol Reagent 2)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- 9. Add 1 level spoon of reagent KP242 (Coppercol Reagent 3) (note 1, page 100).
- 10. Close the vial tightly with the cap and swirl several times to dissolve the powder.
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

12. Press TEST key.

The result is shown in the display in mg/l free Copper.







## Copper, total with Liquid reagent and powder

0.05 - 4 mg/l Cu



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS240 (Coppercol Reagent 1)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops of KS241 (Coppercol Reagent 2)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add 1 level spoon of reagent KP242 (Coppercol Reagent 3) (note 1, page 100).
- 10. Close the vial tightly with the cap and swirl several times to dissolve the powder.

- 11. Add **one COPPER No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 12. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 13. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

14. Press **TEST** key.

The result is shown in the display in mg/l total Copper.







## Copper, free (Note 1) with Vario Powder Pack

0.05 - 5 mg/l Cu



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press **ZERO** key.



- 4. Remove the vial from the sample chamber.
- Add the contents of one VARIO Cu 1 F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (Note 3).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

8. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Copper

#### Notes:

- 1. For determination of total Copper digestion is required.
- 2. Extremely acid water samples (pH 2 or less) must be adjusted between pH 4 and pH 6 before the reagent is added (with 8 mol/l Potassium hydroxide solution KOH). Caution: pH values above 6 can lead to Copper precipitation.
- 3. Accuracy is not affected by undissolved powder.
- 4. Interferences:

| Cyanide, CN | Cyanide prevents full colour development.  Add 0.2 ml Formaldehyde to 10 ml water sample and wait for a reaction time of 4 minutes (Cyanide is masked). After this perform test as described. Multiply the result by 1.02 to correct the sample dilution by Formaldehyde. |
|-------------|---|
| Silver, Ag+ | If a turbidity remains and turns black, silver interference is likely.  Add 10 drops of saturated Potassium chloride solution to 75 ml of water sample. Filtrate through a fine filter. Use 10 ml of the filtered water sample to perform test.                           |

| Reagent        | Form of reagent/Quantity | Order-No. |
|----------------|--------------------------|-----------|
| VARIO Cu 1 F10 | Powder Pack / 100        | 4530300   |







## Cyanide with Reagent Test

0.01 - 0.5 mg/l CN



- Fill a clean vial (24 mm Ø) with 2 ml of the water sample and 8 ml of deionised water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add two level spoons No. 4 (white) of Cyanide-11 into the prepared water sample, replace the cap tightly and invert the vial several times to mix the contents.
- Add two level spoons No. 4 (white) of Cyanide-12, replace the cap tightly and invert the vial several times to mix the contents
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 3 drops of Cyanide-13

- 8. Close the vial tightly with the cap and invert several times to mix the contents.
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## Zero accepted prepare Test press TEST

#### Countdown 10:00

10. Press TEST key.

Wait for a reaction **period of 10 minutes.** 

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Cyanide.

#### Notes:

- 1. Only free Cyanide and Cyanides that can be destroyed by Chlorine are determined by this test.
- 2. In the presence of Thiocyanate, heavy metal complexes, colorants or aromatic amines, the cyanide must be separated out by distillation before analysis is performed.
- 3. Store the reagents in closed containers at a temperature of + 15°C to + 25°C.

| Reagent | Form of reagent/Quantity                       | Order-No. |
|---------|--|-----------|
|         | Reagent test / 200<br>(Powder, Liquid reagent) | 2418875   |







### CyA-TEST (Cyanuric acid) with Tablet

0 - 160 mg/l CyA



 Fill a clean vial (24 mm Ø) with 5 ml of the water sample and 5 ml deionised water (Note 1), close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one CyA-TEST tablet straight from the foil to the prepared water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved (Note 2, 3).

## Zero accepted prepare Test press TEST

8. Press **TEST** key.

The result is shown in the display in mg/l Cyanuric acid.

#### Notes:

- 1. Use deionised water or tap water free of Cyanuric acid.
- 2. If Cyanuric acid is present a cloudy solution will occur.
  Small single particles are not necessarily caused by Cyanuric acid.
- 3. Dissolve the tablet completely (therefore swirl the vial approx. 1 minute). Un-dissolved particles of the tablet can cause results that are too high.

| Reagent  | Form of reagent/Quantity | Order-No. |
|----------|--------------------------|-----------|
| CyA-TEST | Tablet / 100             | 4511370BT |







## DEHA (N,N-Diethylhydroxylamine) with Tablet and Liquid Reagent

 $20 - 500 \mu g/l DEHA / 0.02 - 0.5 mg/l DEHA$ 



- 1. Fill a clean vial (24 mm Ø) with **10 ml of the water** sample, close tightly with the cap (Note 2).
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 6 drops (0.25ml) of DEHA solution

- Close the vial tightly with the cap and swirl several times to mix the contents.
- Add **one DEHA tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 8. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 10. Press TEST key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as DEHA.

#### Notes:

- 1. Application: Testing of residual corrosion inhibitors (Oxygen scavengers) in boiler feed water or condensate.
- 2. Before using clean the vials with Hydrochloric acid (approx. 20%). Rinse thoroughly with deionised water.
- 3. Keep the sample dark during colour development time. UV-light (sunlight) causes high measurement results.
- 4. Ideal temperature for full colour development is 20°C ± 2°C.
- 5. Interferences:
  - Iron (II) interferes at all concentrations:
     Repeat the test procedure but without adding the DEHA solution. If the displayed result is above 20 µg/l subtract this value from the DEHA test result.
  - Substances which reduce Iron (III) interfere. Substances which complex iron strongly may interfere also.
  - Substances which may interfere when present in concentrations at:

| Borate (as Na <sub>2</sub> B <sub>4</sub> O <sub>7</sub> ) | 500 mg/l   |
|--|------------|
| Cobalt   | 0.025 mg/l |
| Copper   | 8.0 mg/l   |
| Hardness (as CaCO <sub>3</sub> )                           | 1000 mg/l  |
| Lignosulfonates  | 0.05 mg/l  |
| Manganese  | 0.8 mg/l   |
| Molybdenum   | 80 mg/l    |
| Nickel   | 0.8 mg/l   |
| Phosphate  | 10 mg/l    |
| Phosphonates   | 10 mg/l    |
| Sulfate  | 1000 mg/l  |
| Zinc   | 50 mg/l    |

6. There is an option to change the unit from mg/l to  $\mu$ g/l.



| Reagent                     | Form of reagent/Quantity | Order-No. |
|-----------------------------|--------------------------|-----------|
| DEHA solution ca. 60 Tests  | Liquid reagent / 15 ml   | 461185    |
| DEHA solution ca. 400 Tests | Liquid reagent / 100 ml  | 461181    |
| DEHA                        | Tablet / 100             | 4513220BT |







# DEHA (N,N-Diethylhydroxylamin) with Vario Powder Pack and Liquid Reagent

 $20 - 500 \mu g/l DEHA / 0.02 - 0.5 mg/l DEHA$ 



Use two clean vials (24 mm  $\varnothing$ ) and mark one as blank for zeroing (Note 2).

- Fill a clean vial with 10 ml deionised water (this is the blank).
- Fill the second clean vial with 10 ml of the water sample (this is the sample).



- 3. Add the contents of **one VARIO OXYSCAV 1 Rgt Powder Pack** straight from the foil into each vial.
- 4. Close the vials tightly with the caps and swirl several times to mix the contents.
- Add 0.20 ml VARIO DEHA 2 Rgt Solution to each vial (Note 4).
- 6. Close the vials tightly with the caps and swirl several times to mix the contents

Countdown 1 10:00 start: 🔟

7. Press [ ] key.

Wait for a reaction **period of 10 minutes** (Note 5). After the reaction period is finished proceed as follows:

8. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

prepare Zero press ZERO

- 9. Press **ZERO** key.
- 10. Remove the vial from the sample chamber.
- 11. Place the vial (the sample) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

12. Press **TEST** key.

The result is shown in the display as DEHA.

#### Notes:

- Application: Testing of residual corrosion inhibitors (Oxygen scavengers) in boiler feed water or condensate.
- 2. Before using clean the vials with Hydrochloric acid (approx. 20%). Rinse thoroughly with deionised water.
- 3. Ideally temperature for full colour development is 25°C ± 3 °C.
- 4. Volume should always be metered by using suitable pipette (class A).
- 5. Keep blank and sample dark during colour development time. UV-light (sunlight) causes high measurement results.
- 6. Interferences:
  - Iron (II) interferes at all concentrations:
     Repeat the test procedure but without adding the VARIO DEHA Rgt 2 solution. If the displayed result is above 20 µg/l subtract this value from the DEHA test result.
  - Substances which reduce Iron (III) interfere. Substances which complex iron strongly may interfere also.
  - Substances who may interfere when present in concentrations at:

| 500 mg/l   |
|------------|
| 0.025 mg/l |
| 8.0 mg/l   |
| 1000 mg/l  |
| 0.05 mg/l  |
| 0.8 mg/l   |
| 80 mg/l    |
| 0.8 mg/l   |
| 10 mg/l    |
| 10 mg/l    |
| 1000 mg/l  |
| 50 mg/l    |
|            |

7. There is an option to change the unit from mg/l to  $\mu$ g/l.



| Reagent                   | Form of reagent/Quantity | Order-No. |
|---------------------------|--------------------------|-----------|
| Set (100 Tests)           |                          | 4536000   |
| VARIO OXYSCAV 1 Rgt       | Powder Pack / 200        |           |
| VARIO DEHA 2 Rgt solution | Liquid reagent / 100 ml  |           |







## Fluoride with Liquid Reagent

0.05 - 2 mg/l F



Caution: See notes!

- 1. Fill a clean vial (24 mm Ø) with **exactly 10 ml of water sample** (Note 4), close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Add **exactly 2 ml SPADNS reagent solution** (Note 4) to the water sample.

Caution: Vial is filled up to the top! (Note 8)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Press **TEST** key.

The result is shown in the display in mg/l Fluoride.

#### Notes:

- 1. The same batch of SPADNS reagent solution must be used for adjustment and test. The adjustment process needs to be performed for each new batch of SPADNS reagent solution (see Standard Methods 20th, 1998, APHA, AWWA, WEF 4500 F D., S. 4-82). The procedure is described in chapter 2.4.5 "Calibration Fluoride Method 170" on page 314.
- 2. During adjustment and test the same vial should be used for zeroing and test, as different vials may exhibit minor tolerances.
- 3. The calibration solution and the water samples to be tested should have the same temperature ( $\pm$  1°C).
- 4. As the test result is highly dependent on exact sample and reagent volumes, the sample and reagent volumes should always be metered by using a 10 ml resp. 2 ml volumetric pipette (class A).
- 5. The accuracy of the test methods decreases above a level of 1.2 mg/l Fluoride. Although the results are sufficiently accurate for most applications, even more exact results can be achieved by 1:1 dilution of the sample prior to use and subsequent multiplication of the result by 2.
- SPADNS reagent solution contains Arsenite.
   Chlorine concentrations up to 5 mg/l do not interfere.
- 7. Seawater and wastewater samples must be distilled.
- 8. It is convenient to use special vials with larger volume.

| Reagent                 | Form of reagent/Quantity | Order-No. |
|-------------------------|--------------------------|-----------|
| SPADNS reagent solution | Liquid reagent / 250 ml  | 4467481   |
| Fluoride standard       | Solution / 30 ml         | 205630    |







## H<sub>2</sub>O<sub>2</sub> (Hydrogen peroxide) with tablet reagent

 $0.03 - 3 \text{ mg/l H}_{2}O_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- Add one HYDROGENPEROXIDE LR tablet straight from the foil and crush the tablet using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00 9. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l H<sub>2</sub>O<sub>2</sub>.

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Hydrogen peroxide may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand. Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) for one hour, then rinse all glassware thoroughly with deionised water.

- 2. Preparing the sample:
  - When preparing the sample, the lost of Hydrogen peroxide, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment.

  Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the tablet is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range:
  - Concentrations above 5 mg/l Hydrogen peroxide can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Hydrogen peroxide. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. Oxidising agents such as Chlorine, Ozone etc. interfere as they react in the same way as Hydrogen peroxide.

| Reagent             | Form of reagent/Quantity | Order-No. |
|---------------------|--------------------------|-----------|
| Hydrogenperoxide LR | Tablet / 100             | 4512380BT |







## H<sub>2</sub>O<sub>2</sub> (Hydrogen peroxide) LR with Liquid Reagent

1 – 50 mg/l H<sub>2</sub>O<sub>2</sub>



Insert the adapter for 16 mm Ø vials.

- Fill a clean vial (16 mm Ø) with 10 ml of the water sample, close tightly with the cap. (Note 1, 2)
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press the **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the prepared vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 6 drops of H<sub>2</sub>O<sub>2</sub>-Reagent

- 6. Close the vial tightly with the cap and invert several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

#### Zero accepted prepare Test press TEST

8. Press the **TEST** key.

The result is shown in the display in mg/l  $H_2O_2$ .

#### Notes:

- 1. The hydrogen peroxide is determined in the form of yellow/orange coloured peroxotitanic acids in strongly acidic media. In connection with neutral to weakly alkaline (~pH 10) samples, the acid in the reagent is sufficient in order to produce a medium suitable for measurement. In the case of strongly alkaline samples (pH > 10), the samples must be acidified before measurement otherwise the results may be deficient. This is achieved by diluting the sample with a 5% sulphuric acid solution, for example, at a ratio of 1:1.
  - In contrast to many other colour reactions, in connection with the presence of hydrogen peroxide, discoloration with long-term stability is achieved that can still be measured after 24 h. Particles in the sample solution or turbidity distort the analysis and must be eliminated by centrifuging or simply filtering the sample solution prior to performing the measurement. Falsification of the measurement results should also be expected in connection with coloured solutions.
- 2. Oxidising agents such as chlorine, bromine, chlorine dioxide and ozone do not distort the analysis. On the other hand, however, water discoloration does distort the analysis. In this case, proceed as described in the following:
  - Fill a clean vial (16 mm Ø) with 10 ml of the water sample and perform zero calibration (see "Operation").
  - Measure the sample solution without the addition of drops of reagent (result B).
  - Then the same sample solution, measured with the addition of the reagent drops (result A).
  - Calculations: mg/l H<sub>2</sub>O<sub>2</sub> = result A result B
- 3. Attention: The reference reagent contains a 25% sulphuric acid solution. It is recommended to wear appropriate protective clothing (protective goggles/gloves).

| Reagent                                | Form of reagent/Quantity | Order-No. |
|--|--------------------------|-----------|
| H <sub>2</sub> O <sub>2</sub> -reagent | Liquid reagent / 15 ml   | 424991    |







## H<sub>2</sub>O<sub>2</sub> (Hydrogen peroxide) HR with Liquid Reagent

40 - 500 mg/l H<sub>2</sub>O<sub>2</sub>



Insert the adapter for 16 mm Ø vials.

- Fill a clean vial (16 mm Ø) with 10 ml of the water sample, close tightly with the cap. (Note 1, 2)
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press the **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the prepared vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 6 drops of H<sub>2</sub>O<sub>2</sub>-Reagent

- 6. Close the vial tightly with the cap and invert several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

#### Zero accepted prepare Test press TEST

8. Press the **TEST** key.

The result is shown in the display in mg/l  $H_2O_2$ .

#### Notes:

- 1. The hydrogen peroxide is determined in the form of yellow/orange coloured peroxotitanic acids in strongly acidic media. In connection with neutral to weakly alkaline (~pH 10) samples, the acid in the reagent is sufficient in order to produce a medium suitable for measurement. In the case of strongly alkaline samples (pH > 10), the samples must be acidified before measurement otherwise the results may be deficient. This is achieved by diluting the sample with a 5% sulphuric acid solution, for example, at a ratio of 1:1.
  - In contrast to many other colour reactions, in connection with the presence of hydrogen peroxide, discoloration with long-term stability is achieved that can still be measured after 24 h. Particles in the sample solution or turbidity distort the analysis and must be eliminated by centrifuging or simply filtering the sample solution prior to performing the measurement. Falsification of the measurement results should also be expected in connection with coloured solutions.
- 2. Oxidising agents such as chlorine, bromine, chlorine dioxide and ozone do not distort the analysis. On the other hand, however, water discoloration does distort the analysis. In this case, proceed as described in the following:
  - Fill a clean vial (16 mm Ø) with 10 ml of the water sample and perform zero calibration (see "Operation").
  - Measure the sample solution without the addition of drops of reagent (result B).
  - Then the same sample solution, measured with the addition of the reagent drops (result A).
  - Calculations: mg/l H<sub>2</sub>O<sub>2</sub> = result A result B
- 3. Attention: The reference reagent contains a 25% sulphuric acid solution. It is recommended to wear appropriate protective clothing (protective goggles/gloves).

| Reagent                                | Form of reagent/Quantity | Order-No. |
|--|--------------------------|-----------|
| H <sub>2</sub> O <sub>2</sub> -reagent | Liquid reagent / 15 ml   | 424991    |







#### Hardness, Calcium with Tablet

50 - 900 mg/l CaCO<sub>2</sub>



- 1. Fill a clean vial (24 mm Ø) with 10 ml deionised water.
- 2. Add one CALCHECK tablet straight from the foil to the deionised water and crush the tablet using a clean stirring rod.
- 3. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 4. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

#### prepare Zero press ZERO

#### Countdown 2:00

5. Press **ZERO** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

- 6. Remove the vial from the sample chamber.
- 7. Add 2 ml of the water sample to the prepared vial. Caution: Vial is filled up to the top! (Note 4)
- 8. Close the vial tightly with the cap and swirl several times (5x) to mix the contents.
- 9. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

Zero accepted prepare Test press TEST

10. Press TEST key.

The result is shown in the display as Calcium Hardness.

#### Notes:

- 1. Strong alkaline or acidic water samples must be adjusted between pH 4 and pH 10 before the tablet is added (use 1 mol/l Hydrochloric acid resp. 1mol/l Sodium hydroxide).
- 2. The tolerance of the method is increasing with higher concentrations. When diluting samples, this should be take into account, always measuring in the first third of the range.
- 3. This method was developed from a volumetric procedure for the determination of calcium. Due to undefined conditions, the deviations from the standardised method may be greater.
- 4. It is convenient to use special vials with larger volume.



| Reagent  | Form of reagent/Quantity | Order-No. |
|----------|--------------------------|-----------|
| CALCHECK | Tablet / 100             | 4515650   |







## Hardness, Calcium 2T with Tablet

0 - 500 mg/l CaCO<sub>3</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one CALCIO H No. 1 tablet straight from the foil to the 10 ml water sample, crush the tablet using a clean stirring rod and dissolve the tablet completely.
- 6. Add **one CALCIO H No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl gently several times until the tablet is completely dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

9. Press **TEST** key.

Wait for a **reaction period of 2 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as Calcium Hardness.

#### Notes:

- To optimise the readings an optional batch related calibration can be performed using Mode 40, see page 312.
- 2. Strong alkaline or acidic water samples must be adjusted to a pH-value between pH 4 and 10 before the tablets are added (use 1 mol/l Hydrochloride acid resp. 1 mol/l Sodium hydroxide).
- 3. For accurate test results exactly 10 ml of water sample must be taken for the test.
- 4. This method was developed from a volumetric procedure for the determination of Calcium Hardness. Due to undefined conditions, the deviations from the standardised method may be greater.
- 5. The tolerance of the method is increasing with higher concentrations. When diluting samples, this should be taken in account, always measuring in the first third of the range.

#### 6. Interferences:

- Magnesium hardness up to 200 mg/l CaCO<sub>2</sub> does not interfere.
- Iron concentration above 10 mg/l may cause low results.
- Zinc concentration above 5 mg/l may cause high results.
- 7. A CaCO<sub>3</sub>
  °dH
  °eH
  °fH
  •aH

| Reagent                    | Form of reagent/Quantity                | Order-No. |
|----------------------------|---|-----------|
| Set CALCIO H No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517761BT |



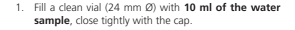




## Hardness, total with Tablet

2 - 50 mg/l CaCO<sub>3</sub>





2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one HARDCHECK P tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press TEST key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as total Hardness.

#### Notes:

1. Strong alkaline or acidic water samples must be adjusted between pH 4 and pH 10 before the tablet is added (use 1 mol/l Hydrochloric acid resp. 1mol/l Sodium hydroxide).

#### 2. Conversion table:

|                          | mg/l CaCO <sub>3</sub> | °dH   | °fH  | °eH  |
|--------------------------|------------------------|-------|------|------|
| 1 mg/l CaCO <sub>3</sub> |                        | 0.056 | 0.10 | 0.07 |
| 1 °dH                    | 17.8                   |       | 1.78 | 1.25 |
| 1 °fH                    | 10.0                   | 0.56  |      | 0.70 |
| 1 °eH                    | 14.3                   | 0.80  | 1.43 |      |

3. CaCO<sub>3</sub> °dH °eH °fH °aH

| Reagent     | Form of reagent/Quantity | Order-No. |
|-------------|--------------------------|-----------|
| HARDCHECK P | Tablet / 100             | 4515660BT |







## Hardness, total HR with Tablet

20 - 500 mg/l CaCO<sub>2</sub>



Ø 24 mm

- Fill a clean vial (24 mm Ø) with 1 ml of the water sample and 9 ml of deionised water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one HARDCHECK P tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## Zero accepted prepare Test press TEST

### Countdown 5:00

8. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as total Hardness.

#### Notes:

1. Strong alkaline or acidic water samples must be adjusted between pH 4 and pH 10 before the tablet is added (use 1 mol/l Hydrochloric acid resp. 1mol/l Sodium hydroxide).

#### 2. Conversion table:

|                          | mg/l CaCO₃ | °dH   | °fH  | °eH  |
|--------------------------|------------|-------|------|------|
| 1 mg/l CaCO <sub>3</sub> |            | 0.056 | 0.10 | 0.07 |
| 1 °dH                    | 17.8       |       | 1.78 | 1.25 |
| 1 °fH                    | 10.0       | 0.56  |      | 0.70 |
| 1 °eH                    | 14.3       | 0.80  | 1.43 |      |



| Reagent     | Form of reagent/Quantity | Order-No. |
|-------------|--------------------------|-----------|
| HARDCHECK P | Tablet / 100             | 4515660BT |







## Hydrazine with Powder Reagent

 $0.05 - 0.5 \text{ mg/l N}_2\text{H}_4 / 50 - 500 \text{ }\mu\text{g/l N}_2\text{H}_4$ 



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample (Note 1, 2), close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

3. Press **ZERO** key.

4. Remove the vial from the sample chamber.

Add 1 g HYDRAZINE test powder (Note 3) to the water sample.

6. Close the vial tightly with the cap and swirl several times to mix the contents.

#### Countdown 10:00 start:

7. Press [ key. Wait for a reaction period of 10 minutes.

After the reaction period is finished proceed as follows:

8. The slight turbidity that occurs when the reagent is added must be removed by filtration (Note 4).

9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

10. Press **TEST** key.

The result is shown in the display as Hydrazine.

#### Notes:

- 1. If the water sample is cloudy, you must filter it before performing the zero calibration.
- 2. The temperature of the water sample should not exceed 21  $^{\circ}\text{C}.$
- 3. Using the Hydrazine spoon: 1 g is equivalent to one level spoon.
- 4. Qualitative folded filter papers for medium precipitates are recommended.
- 5. In order to check whether the reagent has aged (if it has been stored for a lengthy period), perform the test as described above using tap water. If the result is above the detection limit of 0.05 mg/l, you should only use the reagent with reservations as there may be a major deviation in results.
- 6. There is an option to change the unit from mg/l to  $\mu$ g/l.



| Reagent / Accessories | Form of reagent/Quantity | Order-No. |
|-----------------------|--------------------------|-----------|
| Hydrazin Test Powder  | Powder / 30 g            | 462910    |
| Spoon                 |                          | 384930    |







## Hydrazine with Vario Liquid Reagent

0.005-0.6 mg/l  $\mathrm{N_2H_4}$  / 5-600  $\mu$ g/l  $\mathrm{N_2H_4}$ 



Ø 24 mm

Use two clean vials (24 mm  $\emptyset$ ) and mark one as blank for zeroing.

- Fill a clean vial with 10 ml deionised water (this is the blank).
- Add 1 ml VARIO Hydra 2 Rgt Solution into the vial (Note 3).
- 3. Close the vial tightly with the cap and swirl several times to mix the contents
- 4. Place the vial (the blank) in the sample chamber making sure that the  $\chi$  marks are aligned.

### prepare Zero press ZERO

- 5. Press ZERO key.
- 6. Remove the vial from the sample chamber.
- Fill the second clean vial with 10 ml of the water sample (this is the sample).
- 8. Add 1 ml VARIO Hydra 2 Rgt Solution into the vial.
- 9. Close the vial tightly with the cap and swirl several times to mix the contents.
- 10. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 12:00 11. Press **TEST** key.

Wait for a reaction period of 12 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as Hydrazine.

#### Notes:

- 1. Samples cannot be preserved and must be analysed immediately.
- 2. Sample temperature should be  $21^{\circ}C \pm 4^{\circ}C$ .
- 3. The blank may develop a faint yellow colour due to the reagent.
- 4. Interferences:
  - Ammonia causes no interferences up to 10 mg/l.
     At a concentration of 20 mg/l it is possible that the test result increases by 20%.
  - Morpholine does not interfere up to 10 mg/l.
  - Highly coloured or turbid samples:

Mix 1 part deionised water with 1 part household bleach. Add 1 drop of this mixture into 25 ml water sample and mix. Use 10 ml prepared sample in place of deionised water in point 1.

Note: at point 7 use the unprepared water sample.

Principle: Hydrazine is oxidised by household bleach. Colour interference will be eliminated by zeroing.

5. There is an option to change the unit from mg/L to  $\mu$ g/L.



| Reagent                    | Form of reagent/Quantity | Order-No. |
|----------------------------|--------------------------|-----------|
| VARIO Hydra 2 Rgt solution | Liquid reagent / 100 ml  | 4531200   |







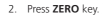
## Hydrazine with Vacu-vials® K-5003 (see Notes)

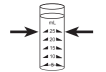
 $0.01 - 0.7 \text{ mg/l N}_2\text{H}_4 / 10 - 700 \text{ }\mu\text{g/l N}_2\text{H}_4$ 

Insert the adapter for 13 mm Ø vials.

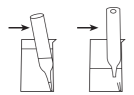
1. Place the blank in the sample chamber. The blank is part of the test kit.

### prepare Zero press ZERO





- 3. Remove the blank from the sample chamber.
- 4. Fill the sample container to the 25 ml mark with the water sample.



- 5. Place one Vacu-vial® in the sample container. Snap the tip by pressing the vial against the side of the sample container. The Vacu-vial® breaks at the neck and the vial fills automatically. A small volume of inert gas remains in the Vacu-vial®.
- 6. Mix the contents of the Vacu-vial® by inverting it several times, allowing the bubble to move from one end to the other. Dry the outside of the vial.
- 7. Place the Vacu-vial® in the sample chamber.

Zero accepted prepare Test press TEST

8. Press **TEST** key.

Countdown 10:00 Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display as Hydrazine.

#### Notes:

- 1. This method is adapted from CHEMetrics. The measuring range and wavelength used for this photometer may differ from the data specified by CHEMetrics.
- 2. Read the original test instruction and the MSDS (delivered with the test) before performing the test. MSDS also available at www.chemetrics.com.
- 3. Vacu-vials® is a registered trade mark of the company CHEMetrics, Inc. / Calverton, U.S.A.
- 4. There is an option to change the unit from mg/l to  $\mu$ g/l.



| Reagent                                     | Form of reagent/Quantity | Order-No. |
|---|--------------------------|-----------|
| Vacu-vials <sup>®</sup> / CHEMetrics K-5003 | Test-Kit / 30            | 380470    |







### lodine with Tablet

0.05 - 3.6 mg/l I



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber, **empty the vial leaving a view drops in.**
- Add one DPD No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

9. Press **TEST** key.

The result is shown in the display in mg/l lodine.

### Notes:

1. Oxidizing reagents, such as Chlorine, Bromine, etc. interfere as they react in the same way as lodine.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| DPD No. 1 | Tablet / 100             | 4511050BT |





0.02 - 3 mg/l Fe

Determination of all dissolved iron and most undissolved forms of iron. \*

# 2 2 3 Iron, total with Vario Powder Pack

0.02 - 1.8 mg/l Fe

Determination of all dissolved iron and most undissolved forms of iron; most undissolved iron oxides are recovered by the reagent. \*

# 2 4 Iron, total (Fe in Mo) with Vario Powder Pack

0.01 - 1.80 mg/l Fe

Determination of all dissolved iron and unsolved iron in the presence of high molybdate concentrations

# 2 2 5 Iron LR with Liquid Reagent

0.03 - 2 mg/l Fe

Determination of total soluble Iron  $Fe^{2+/3+}$  in presence of complexing agent (e.g. Molybdate) \*

# 2 2 6 Iron LR 2 with Liquid reagent

 $0.03 - 2 \text{ mg/l Fe}^{2+}$  and  $\text{Fe}^{3+}$ 

Determination of total soluble Iron Fe<sup>2+</sup> and Fe<sup>3+</sup> in presence of complexing agent (e.g. Molybdate) \*





# Iron HR with Liquid reagent

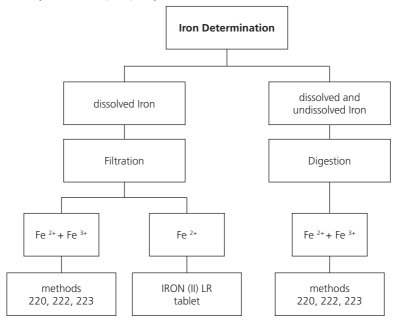
0.1 - 10 mg/l Fe

Determination of total soluble Iron Fe<sup>2+/3+</sup> in presence of complexing agent (e.g. Molybdate) \*

\*This information refers to analysis of the water sample without digestion.

Further information can be found in the method notes.

#### Notes (Methods 220, 222, 223):



**Digestion procedure** for the determination of total dissolved and undissolved iron.

- 1. Add 1 ml of concentrated sulfuric acid to 100 ml water sample. Heat and boil for 10 minutes or until all particles are dissolved. After cooling down, the sample is set to a pH-value of 3 to 6 by using ammonia solution. Refill with deionised water to the previous volume of 100 ml and mix well. 10 ml of this pre-treated solution is used for the following analysis. Perform as described by the selected test method.
- 2. Water which has been treated with organic compounds like corrosion inhibitors must be oxidised where necessary to break down the iron. Therefore add 1 ml concentrated sulfuric acid and 1 ml concentrated nitric acid to 100 ml water sample and boil to approx. half volume. After cooling down, proceed as described above.







## Iron (Note 1) with Tablet

0.02 - 1 mg/l Fe



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one IRON LR tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press **TEST** key. Wait for a **reaction period of 5 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Iron ( $Fe^{2+/3+}$ ).

#### Notes:

- 1. This method determines the total dissolved Iron as Fe<sup>2+</sup> and Fe<sup>3+</sup>.
- 2. The IRON (II) LR tablet is used for differentiation as described above instead of the IRON LR tablet.

$$Fe^{3+} = Fe^{2+/3+} - Fe^{2+}$$

3. For the determination of total dissolved and undissolved iron digestion is required. An example is described on page 141.

| Reagent      | Form of reagent/Quantity | Order-No. |
|--------------|--------------------------|-----------|
| IRON LR      | Tablet / 100             | 4515370BT |
| IRON (II) LR | Tablet / 100             | 4515420BT |







## Iron (Note 1) with Vario Powder Pack

0.02 - 3 mg/l Fe



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO



4. Remove the vial from the sample chamber.



- Add the contents of one Vario Ferro F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (Note 4).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 3:00 8. Press TEST key.

Wait for a reaction period of 3 minutes (Note 5).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Iron.

#### Notes:

- 1. The reagent reacts with all dissolved iron and most undissolved forms of iron in the water sample.
- 2. Iron oxide requires prior digestion: use mild, vigorous or Digesdahl digestion (e.g. for digestion with acid see page 141).
- 3. Very strong alkaline or acidic water samples must be adjusted to a pH value between 3 and 5 before analysis.
- 4. Accuracy is not affected by undissolved powder.
- 5. Water samples containing visible rust should be allowed to react for at least five minutes.

| Reagent         | Form of reagent/Quantity |         |
|-----------------|--------------------------|---------|
| VARIO Ferro F10 | Powder Pack / 100        | 4530560 |







## Iron, total (TPTZ, Note 1) with Vario Powder Pack

0.02 - 1.8 mg/l Fe



Ø 24 mm

Use two clean vials (24 mm  $\emptyset$ ) and mark one as blank for zeroing.

 Fill a clean vial with 10 ml deionised water (this is the blank).



- Fill the second clean vial with 10 ml of the water sample (this is the sample).
- Add the contents of one Vario IRON TPTZ F10 Powder Pack straight from the foil into each vial.
- 4. Close the vials tightly with the caps and swirl several times to mix the contents.

### Countdown 3:00 start:

5. Press [4] key.

Wait for a reaction **period of 3 minutes.** 

After the reaction period is finished proceed as follows:

6. Place the vial (the blank) in the sample chamber making sure that the  $\chi$  marks are aligned.

## prepare Zero press ZERO

- 7. Press **ZERO** key.
- 8. Remove the vial from the sample chamber.
- 9. Place the vial (the sample) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

10. Press TEST key.

The result is shown in the display in mg/l Iron.

#### Notes:

- 1. For determination of total Iron digestion is required.

  TPTZ reagent recovers most insoluble iron oxides without digestion.
- 2. Rinse all glassware with 1:1 Hydrochloric acid solution first and then rinse with deionised water to remove iron deposits that can cause slightly high results.
- 3. Strong alkaline or acidic water samples must be adjusted between pH 3 and pH 8 before the reagent is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Interferences:

When interferences occur, colour development is inhibited or a precipitate is formed. The values below refer to a standard with an iron concentration of 0.5 mg/l. The following substances do not interfere when present up to the levels given:

| Substance                | no interference to |
|--------------------------|--------------------|
| Cadmium                  | 4.0 mg/l           |
| Chromium <sup>(3+)</sup> | 0.25 mg/l          |
| Chromium (6+)            | 1.2 mg/l           |
| Cobalt                   | 0.05 mg/l          |
| Copper                   | 0.6 mg/l           |
| Cyanide                  | 2.8 mg/l           |
| Manganese                | 50 mg/l            |
| Mercury                  | 0.4 mg/l           |
| Molybdenum               | 4.0 mg/l           |
| Nickel                   | 1.0 mg/l           |
| Nitrite Ion              | 0.8 mg/l           |

| Reagent             | Form of reagent/Quantity | Order-No. |
|---------------------|--------------------------|-----------|
| VARIO IRON TPTZ F10 | Powder Pack / 100        | 4530550   |







### Iron, total (Fe in Mo) in the presence of Molybdate with Vario Powder Pack





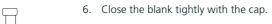


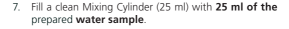


- 2. Add the contents of one Vario (Fe in Mo) Rgt 1 Powder Pack straight from the foil into the water sample (50 ml).
- Close the Mixing Cylinder tightly with a stopper and invert several times to dissolve the powder.



- 4. Use two clean vials (24 mm Ø) and mark one as blank for zeroing.
- 5. Add 10 ml of the prepared water sample to the vial (this is the blank).





- 8. Add the contents of one Vario (Fe in Mo) Rgt 2 **Powder Pack** straight from the foil into the prepared water sample (25 ml).
- 9. Close the Mixing Cylinder tightly with a stopper and invert several times to dissolve the powder (note 5).



Count-Down 1

3:00

Start:

10. Press [4] key.

- Wait for a reaction period of 3 minutes.
- 11. After the reaction period is finished proceed as follows: Fill the second prepared vial (point 4) with 10 ml of the sample .This is the sample.
- 12. Place **the blank** in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

| prepa | re | Zero |  |
|-------|----|------|--|
| press | ZE | RO   |  |

- 13. Press ZERO key.
- 14. Remove the vial from the sample chamber.
- 15. Place **the sample** in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

15. Press **TEST** key.

The result is shown in the display in mg/l Fe.

#### Notes:

- 1. Rinse all glassware with detergent, followed by tap water. Rinse again with 1:1 Hydrochloric acid solution and deionized water. These steps will remove deposits that can cause slightly high results.
- 2. Take the sample reading immediately after the instrument zero, If the sample contains 100 mg/l or more Molybdate (MoO<sub>4</sub> <sup>2</sup>-).
- 3. For more accurate results, a reagent blank value for each new lot of reagent is advisable. Follow the described procedure using deionized water instead of the sample. Subtract the obtained reading value from the final results.
- 4. Interference pH: A sample pH of less than 3 or more than 4 after addition of reagent, may inhibit colour formation, as the developed colour fades too quickly or results in turbidity. Adjust the sample pH to between 3 and 5 in the graduated cylinder before the addition of reagent:
  - Add by drops an applicable amount of Iron-free acid or base eg. 1 N Sulfuric acid solution or 1 N Sodium hydroxide solution.
  - If necessary make a volume correction if significant volumes of acid or base are used.
- 5. If Iron is present a blue colour developes. A small amount of undissolved reagent does not have an affect on the results of the test.

#### Sample collection and storage:

- Collect samples in clean glass or plastic bottles. These should have been cleaned with 6 N (1:1) Hydrochloric acid and rinsed with deionised water.
- To preserve samples for later analysis, adjust the sample pH to less than 2 with concentrated Hydrochloric acid by adding about 2 ml per liter. If the sample is tested immediately this acid addition is not necessary.
- If the dissolved Iron is required, filter the sample through a 0.45-micron filter or equivalent medium immediately after collection and before acidification.
- The preserved samples should be kept at room temperature for a maximum of 6 months
- Adjust the pH to 3 5 by adding 5 N Sodium hydroxide solution before analysis. Do not exceed pH 5 as Iron might precipitates.
- The test result needs to be corrected for the dilution caused by the volume additions.

| Reagent                | Form of reagent/Quantity | Order-No. |
|------------------------|--------------------------|-----------|
| Set                    |                          | 4536010   |
| Vario (Fe in Mo) Rgt 1 | Powder Pack / 100        |           |
| Vario (Fe in Mo) Rgt 2 | Powder Pack / 100        |           |







# Iron LR with Liquid reagent

 $0.03 - 2 \text{ mg/l Fe}^{2+/3+}$ 



This test is suitable for determining total soluble iron. The sample should be pre-filtered using a 0.45 µm membrane if total dissolved iron is required. Particulate or suspended iron will otherwise add to the result

- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS61 (Ferrozine/Thioglycolate)

- Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the marks  $\sqrt{}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press TEST key.

Wait for a reaction **period of 5 minutes** (note 1).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Iron.

#### Notes:

- 1. Complexed iron may be measured by increasing the development period until no further colour development is seen. Very strongly complexed iron may not be included in the measured iron. In this case the complexing agent must be destroyed by oxidation with acid/persulphate followed by neutralisation to pH 6–9. Follow the procedure on page 152.
- 2. For total iron (suspended and dissolved), boil sample with acid/persulphate. Neutralise back to pH 6–9 making back up to original volume with distilled or deionised water. Follow the procedure on page 152.
- 3. When using KS61 (Ferrozine/Thioglycolate), high levels of molybdate will produce an intense vellow colour.

In this case a reagent blank is required:

- Use two clean vials (24 mm Ø).
- Mark one as blank for zeroing.
- Fill a clean vial (24 mm Ø) with **10 ml of the water sample** (blank).
- Add 10 drops KS63 (Thioglycolate).
- Close the vial tightly with the cap and swirl gently several times.
- Place the blank in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.
- Press ZERO key.
- Remove the vial from the sample chamber.
- Fill a second clean 24 mm vial with **10 ml water sample** (this is the sample).
- Follow the procedure as described on page 150, point 5.

| Reagent / Accessories                          | Form of reagent/Quantity | Order-No. |
|--|--------------------------|-----------|
| KS61 (Ferrozine/ Thioglycolate)                | Liquid reagent / 65 ml   | 56L006165 |
| KS63 (Thioglycolate Reagent)                   | Liquid reagent / 65 ml   | 56L006365 |
| KP962 (Ammonium Persulphate Powder)            | Powder                   | 56P096240 |
| KS135 (Phenolphthalein Substitute<br>Indikator | Liquid reagent / 65 ml   | 56L013565 |
| KS144 (Calcium Hardness Puffer)                | Liquid reagent / 65 ml   | 56L014465 |
| Spoon  | 0,5 g Spoon              | 385340    |







# Iron, total LR with Liquid reagent

 $0.03 - 2 \text{ mg/l Fe}^{2+/3+}$ 





Total iron consists of soluble, complexed and suspended iron. Do not filter the sample but ensure the sample is homogeneous by vigorously shaking immediately prior to sampling. For Total Soluble (including all complexed) filtration will be necessary.

This procedure requires equipment and reagents not included in the standard test pack supplied.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- Add 5 ml 1:1 Hydrochloric acid and one KT274 (Ammonium Persulphate) tablet.
- Boil for 20 minutes, maintaining the sample volume above 25 ml with deionised water.
- 4. Cool the sample to room temperature.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:
  - 1 drop KS135 (Phenolphthalein Substitute Indicator)
- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pink/red colour just appears.
- 7. Fill the sample up to 50ml with deionised water.
- 8. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

9. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO

- 10. Press **ZERO** key.
- 11. Remove the vial from the sample chamber and empty the vial.
- 12. Add 10 ml prepared water sample to the same vial.
- 13. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

### 10 drops KS61 (Ferrozine/Thioglycolate)

- 14. Close the vial tightly with the cap and swirl several times to mix the contents.
- 15. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

# Zero accepted prepare Test press TEST

## Countdown 5:00

16. Press **TEST** key.

Wait for a reaction **period of 5 minutes** (note 1, page 151).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total iron or, if a filtered sample was used, in mg/l total soluble iron.







# Iron LR 2 with Liquid reagent

 $0.03 - 2 \text{ mg/l Fe}^{2+} \text{ and Fe}^{3+}$ 



This test is suitable for determining total soluble iron and differentiating between the ferrous and ferric state. The sample should be pre-filtered using a 0.45  $\mu$ m membrane if total dissolved iron is required. Particulate or suspended iron will otherwise add to the result.

- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO

3. Press **ZERO** key.

- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS60 (Acetate Buffer)

- Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

**10 drops KS63 (Thioglycolate)** (note 1)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS65 (Ferrozine)

- 10. Close the vial tightly with the cap and swirl several times to mix the contents.
- 11. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

12. Press TEST key.

Wait for a reaction **period of 5 minutes** (note 2).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l  $Fe^{2+/3+}$  or, if step 7 is omitted,  $Fe^{2+}$ .

$$Fe^{3+} = Fe^{2+/3+} - Fe^{2+}$$

#### Notes:

- 1. For soluble iron Fe<sup>2+</sup> omit step 7.
- 2. Complexed iron may be measured by increasing the development period until no further colour development is seen. Very strongly complexed iron may not be included in the measured iron. In this case the complexing agent must be destroyed by oxidation with acid/persulphate followed by neutralisation to pH 6–9. Follow the procedure on page 156.
- 3. For total iron (suspended and dissolved), boil sample with acid/persulphate. Neutralise back to pH 6–9 making back up to original volume with distilled or deionised water. Follow the procedure on page 156.
- 4. When using KS63 (Thioglycolate), high levels of molybdate will produce an intense yellow colour.

In this case a reagent blank is required:

- Use two clean vials (24 mm Ø).
- Mark one as blank for zeroing.
- Fill a clean vial (24 mm Ø) with **10 ml of the water sample** (blank).
- Add 10 drops KS63 (Thioglycolate).
- Close the vial tightly with the cap and swirl gently several times.
- Place the blank in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.
- Press ZERO key.
- Remove the vial from the sample chamber.
- Fill a second clean 24 mm vial with **10 ml water sample** (this is the sample).
- Follow the procedure as described on page 154, point 5.

| Reagent / Accessories                       | Form of reagent/Quantity | Order-No. |
|---|--------------------------|-----------|
| KS60 – Acetate Buffer                       | Liquid reagent / 65 ml   | 56L006065 |
| KS63 – Thioglycolate Reagent                | Liquid reagent / 65 ml   | 56L006365 |
| KS65 – Ferrozine Reagent                    | Liquid reagent / 65 ml   | 56L006565 |
| KP962 (Ammonium Persulphate Powder)         | Powder                   | 56P096240 |
| KS135 (Phenolphthalein Substitute Indikator | Liquid reagent / 65 ml   | 56L013565 |
| KS144 (Calcium Hardness Puffer)             | Liquid reagent / 65 ml   | 56L014465 |
| Spoon                                       | 0,5 g Spoon              | 385340    |







# Iron, total LR 2 with Liquid reagent

 $0.03 - 2 \text{ mg/l Fe}^{2+/3+}$ 

**Digestion procedure** for the determination of total iron.



Total iron consists of soluble, complexed and suspended iron. Do not filter the sample but ensure the sample is homogeneous by vigorously shaking immediately prior to sampling. For Total Soluble (including all complexed) filtration will be necessary. This procedure requires equipment and reagents not included in the standard test pack supplied.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- Add 5 ml 1:1 Hydrochloric acid and one KT274 (Ammonium Persulphate) tablet.
- 3. Boil for **20 minutes**, maintaining the sample volume above 25 ml with deionised water
- 4. Cool the sample to room temperature.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:
  - 1 drop KS135 (Phenolphthalein Substitute Indicator)
- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pink/red colour just appears.
- 7. Fill the sample up to 50ml with deionised water.
- 8. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 9. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

### prepare Zero press ZERO

- 10. Press ZERO key.
- 11. Remove the vial from the sample chamber and empty the vial
- 12. Add 10 ml prepared water sample to the same vial
- 13. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS60 (Acetate Buffer)

- 14. Close the vial tightly with the cap and swirl several times to mix the contents.
- 15. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

**10 drops KS63 (Thioglycolate)** (note 1, page 155)

- 16. Close the vial tightly with the cap and swirl several times to mix the contents.
- 17. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS65 (Ferrozine)

- 18. Close the vial tightly with the cap and swirl several times to mix the contents.
- 19. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

20. Press **TEST** key.

Wait for a reaction **period of 5 minutes** (note 2, page 155).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total iron or, if a filtered sample was used, in mg/l total soluble iron.







# Iron HR with Liquid reagent

 $0.1 - 10 \text{ mg/l Fe}^{2+/3+}$ 



This test is suitable for determining total soluble iron. The sample should be pre-filtered using a 0.45  $\mu$ m membrane if total dissolved iron is required. Particulate or suspended iron will otherwise add to the result

- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 10 drops KS63 (Thioglycolate)

- Close the vial tightly with the cap and swirl several times to mix the contents. Wait until purple coloration goes before continuing.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

### 10 drops KS160 (Total Hardness Buffer)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- 9. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

| Zero accepted |
|---------------|
| prepare Test  |
| press TEST    |

#### Countdown 15:00

10. Press TEST key.

Wait for a reaction **period of 15 minutes** (note 1).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Iron.

#### Notes:

- 1. Complexed iron may be measured by increasing the development period until no further colour development is seen. Very strongly complexed iron may not be included in the measured iron. In this case the complexing agent must be destroyed by oxidation with acid/persulphate followed by neutralisation to pH 6–9. Follow the procedure on page 160.
- 2. For total iron (suspended and dissolved), boil sample with acid/persulphate. Neutralise back to pH 6–9 making back up to original volume with distilled or deionised water. Follow the procedure on page 160.

| Reagent / Accessories               | Form of reagent/Quantity | Order-No. |
|-------------------------------------|--------------------------|-----------|
| KS160 – Total Hardness Buffer       | Liquid reagent / 65 ml   | 56L016065 |
| KS63 – Thioglycolate Reagent        | Liquid reagent / 65 ml   | 56L006365 |
| KP962 (Ammonium Persulphate Powder) | Powder                   | 56P096240 |
| KS144 (Calcium Hardness Puffer)     | Liquid reagent / 65 ml   | 56L014465 |
| Spoon                               | 0,5 g Spoon              | 385340    |







### Iron, total HR with Liquid reagent

 $0.1 - 10 \text{ mg/l Fe}^{2+/3+}$ 

**Digestion procedure** for the determination of total iron.



Total iron consists of soluble, complexed and suspended iron. Do not filter the sample but ensure the sample is homogeneous by vigorously shaking immediately prior to sampling. For Total Soluble (including all complexed) filtration will be necessary.

This procedure requires equipment and reagents not included in the standard test pack supplied.

- 1. Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- 2. Add 5 ml 1:1 Hydrochloric acid and one KT274 (Ammonium Persulphate) tablet.
- 3. Boil for 20 minutes, maintaining the sample volume above 25 ml with deionised water
- 4. Cool the sample to room temperature.
- 5. Add drops of KS144 (Calcium Hardness Buffer), two drop at a time with mixing, until a neutral or sligthly alkaline solution is obtained. Test periodically with a pH meter or dip-papers (take care not to add exessive buffer).
- 6. Fill the sample up to 50ml with deionised water.
- 7. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 8. Place the vial in the sample chamber making sure that the marks  $\chi$  are aligned.







### prepare Zero press ZERO

- 9. Press ZERO key.
- 10. Remove the vial from the sample chamber and empty the vial
- 11. Add 10 ml prepared water sample to the same vial.
- 12. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

### 10 drops KS63 (Thioglycolate)

- 13. Close the vial tightly with the cap and swirl several times to mix the contents.
- 14. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 10 drops KS160 (Total Hardness Buffer)

- 15. Close the vial tightly with the cap and swirl several times to mix the contents.
- 16. Place the vial in the sample chamber making sure that the marks  $\chi$  are aligned.

# Zero accepted prepare Test press TEST

#### Countdown 15:00

#### 17. Press **TEST** key.

Wait for a reaction **period of 15 minutes** (note 1, page 159).

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total iron or, if a filtered sample was used, in mg/l total soluble iron.







# Manganese with Tablet

0.2 - 4 mg/l Mn



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one MANGANESE LR 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod and dissolve the tablet.
- Add one MANGANESE LR 2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

9. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Manganese.

### Note:

1. **A** Mn

 $MnO_4$ 

▼ KMnO₄

| Reagent                        | Form of reagent/Quantity                | Order-No. |
|--------------------------------|---|-----------|
| Set MANGANESE LR No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517621BT |
| MANGANESE LR No. 1             | Tablet / 100                            | 4516080BT |
| MANGANESE LR No. 2             | Tablet / 100                            | 4516090BT |







### Manganese LR with Vario Powder Pack

0.01 - 0.7 mg/l Mn



2

Use two clean vials (24 mm Ø) and mark one as blank for zeroina (Note 1).

- 1. Fill a clean vial with 10 ml of deionised water (this is the hlank)
- 2. Fill the second clean vial with 10 ml of the water sample (this is the sample).
- 3 Add the contents of one Vario Ascorbic Acid Powder **Pack** straight from the foil into each vial (Note 2).
- 4. Close the vials tightly with the caps and swirl several times to mix the contents
- 5. Fill each vial with drops of the same size by holding the bottle vertically and squeeze slowly (Note 3): 15 drops of Alkaline Cyanide reagent solution
- 6. Close the vials tightly with the caps and swirl several times to mix the contents.
- 7. Fill each vial with drops of the same size by holding the bottle vertically and squeeze slowly: 21 drops of PAN Indicator solution
- 8. Close the vials tightly with the caps and swirl several times to mix the contents.

Countdown 1 2:00 start: 🔟

9. Press [4] key. Wait for a reaction period of 2 minutes (Note 4).

After the reaction period is finished proceed as follows:

10. Place the vial (the blank) in the sample chamber making sure that the marks are  $\chi$  aligned.

prepare Zero

- 11. Press ZERO key.
- 12. Remove the vial from the sample chamber.
- 13. Place the vial (the sample) in the sample chamber making sure that the marks are  $\chi$  aligned.
- 14. Press **TEST** key.

The result is shown in the display in mg/l Manganese.

press ZERO

Zero accepted prepare Test press TEST

#### Notes:

- 1. Rinse all glassware with 1:1 Nitric acid solution first and then rinse with deionised water.
- 2. Water samples that contain more than 300 mg/l  $CaCO_3$  hardness: after adding the Vario Ascorbic Acid powder pack add additionally 10 drops of Rochelle Salt Solution.
- 3. After addition of the reagent solution "Alkaline-Cyanide" a cloudy or turbid solution may form in some water samples. The turbidity should disappear after point 7.
- 4. Water samples containing more than 5 mg/l iron should be allowed to react for at least 10 minutes.
- 5. Conversion:  $mg/l MnO_4 = mg/l Mn \times 2.17$
- 6. ▲ Mn

  MnO₄

  KMnO₄

| Reagent   | Form of reagent/Quantity  | Order-No. |
|---|---|-----------|
| Set<br>VARIO Ascorbic Acid<br>VARIO Alkaline-Cyanide<br>VARIO PAN Indicator | Powder Pack / 100<br>Liquid reagent / 60 ml<br>Liquid reagent / 60 ml | 4535090   |
| VARIO Rochelle Salt Solution  | 30 ml   | 4530640   |







### Manganese HR with Vario Powder Pack

0.1 - 18 mg/l Mn



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.



- Add the contents of one Vario Manganese Citrate Buffer F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add the contents of one VARIO Sodium Periodate F10 Powder Pack straight from the foil to the same water sample.
- 8. Close the vial tightly with the cap and swirl several times to mix the contents
- 9. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned

Zero accepted prepare Test press TEST

Countdown 2:00 10. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Manganese.

#### Notes:

- 1. This test is applicable for the determination of soluble Manganese in water and wastewater.
- 2. Highly buffered water samples or extreme pH values may exceed the buffering capacity of the reagents and requires sample pre-treatment.

  If samples were acidified for storing, adjust the pH between 4 and 5 with 5 mol/l (5 N) Sodium hydroxide before test. Do not exceed pH 5, as manganese may precipitate.
- 3. Interferences:

| Interfering substance | Interference level        |  |
|-----------------------|---------------------------|--|
| Calcium               | greater than 700 mg/l     |  |
| Chloride              | greater than 70 000 mg/l  |  |
| Iron                  | greater than 5 mg/l       |  |
| Magnesium             | greater than 100 000 mg/l |  |

4. **M**n

▼ MnO<sub>4</sub> KMnO<sub>4</sub>

| Reagent                            | Form of reagent/Quantity | Order-No. |
|------------------------------------|--------------------------|-----------|
| Set                                |                          | 4535100   |
| VARIO Manganese Citrate Puffer F10 | Powder Pack / 100        |           |
| VARIO Sodiumperiodate F10          | Powder Pack / 100        |           |







# Manganese with Liquid reagent

0.05 - 5 mg/l Mn



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS265 (Manganese Reagent A)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS266 (Manganese Reagent B)

- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- 9. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

10 drops KS304 (Manganese Reagent C)

 Close the vial tightly with the cap and swirl several times to mix the contents.

11. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 3:00

12. Press **TEST** key.

Wait for a reaction period of 3 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Manganese.

#### Notes:

1. The following substances interfer with this test:

 Calcium
 > 500mg/l

 Sodium
 > 500mg/l

 Nickel
 > 0.5 mg/l

 Iron
 > 5 mg/l

 Chromium
 > 5 mg/l

| Reagent                     | Form of reagent/Quantity | Order-No. |
|-----------------------------|--------------------------|-----------|
| KS265 – Manganese Reagent A | Liquid reagent / 30 ml   | 56L026530 |
| KS266 – Manganese Reagent B | Liquid reagent / 30 ml   | 56L026630 |
| KS304 – Manganese Reagent C | Liquid reagent / 30 ml   | 56L030430 |







## Molybdate with Tablet

 $1 - 50 \text{ mg/l MoO}_4 / 0.6 - 30 \text{ mg/l Mo}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO





- Remove the vial from the sample chamber and empty the vial.
- 5. Fill **20 ml of the water sample** in a 100 ml beaker.
- Add one MOLYBDATE HR No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 7. Add **one MOLYBDATE HR No. 2 tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 8. Dissolve the tablets using a clean stirring rod.
- 9. Rinse out the vial with the prepared water sample and then fill to the 10 ml mark
- 10. Close the vial tightly with the cap.
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

12. Press **TEST** key.

The result is shown in the display in mg/l Molybdate / Molybdenum.

#### Notes:

- 1. The tablets must be added in the correct sequence.
- 2. Under test conditions (pH 3.8 3.9) iron does not interfere nor do other metals at levels likely to be found in industrial water systems.
- 3. Conversions: mg/l Mo = mg/l MoO<sub>4</sub> x 0.6 mg/l Na<sub>2</sub>MoO<sub>6</sub> = mg/l MoO<sub>4</sub> x 1.3
- 4. ▲ MoO<sub>4</sub>
  Mo
  Na<sub>2</sub>MoO<sub>4</sub>

| Reagent                        | Form of reagent/Quantity                | Order-No. |
|--------------------------------|---|-----------|
| Set MOLYBDATE HR No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517631BT |
| MOLYBDATE HR No. 1             | Tablet / 100                            | 4513060BT |
| MOLYBDATE HR No. 2             | Tablet / 100                            | 4513070BT |







## Molybdate / Molybdenum LR mit Vario Powder Pack

 $0.05 - 5.0 \text{ mg/l MoO}_4 / 0.03 - 3 \text{ mg/l Mo}$ 







- Add the contents of one Vario Molybdenum 1 LR F20 Powder Pack straight from the foil into the water sample (20 ml).
- 3. Close the Mixing Cylinder tightly with a stopper and swirl several times to dissolve the powder.



- 4. Use two clean vials (24 mm Ø) and mark one as blank for zeroing.
- 5. Fill each vial with 10 ml of pre prepared water sample.
- 6. Close the blank tightly with the cap.
- Add 0,5 ml of Vario Molybdenum 2 LR solution to the sample.
- 8. Close the vial tightly with the cap and invert several times to mix the contents.

Count-Down 1 2:00 Start:

- Press [4] key.
   Wait for a reaction period of 2 minutes.
- 10. After the reaction period is finished proceed as follows:
- 11. Place the blank in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 12. Press **ZERO** key.
- 13. Remove the vial from the sample chamber.
- 14. Place the sample in the sample chamber making sure that the  $\chi$  marks are aligned.

# Zero accepted prepare Test press TEST

15. Press **TEST** key.

The result is shown in the display in mg/l Molybdate / Molybdenum.

#### Notes:

- 1. Strong alkaline or acidic water samples must be adjusted between pH 3 and pH 5 before the reagent is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 2. Before using clean the vials with Hydrochloric acid (approx. 20%). Rinse thoroughly with deionised water.
- 3. MoO<sub>4</sub>
  - ▼ Na<sub>2</sub>MoO<sub>4</sub>

| Reagent                   | Form of reagent/Quantity | Order-No. |
|---------------------------|--------------------------|-----------|
| Set                       |                          | 4535450   |
| VARIO Molybdenum 1 LR F20 | Powder Pack / 100        |           |
| VARIO Molybdenum 2 LR     | Liquid reagent / 50 ml   |           |
| Mixing Cylinder           | 25 ml                    | 19802650  |







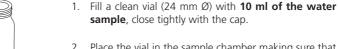
# Molybdate / Molybdenum HR with Vario Powder Pack

 $0.5 - 66 \text{ mg/l MoO}_4 / 0.3 - 40 \text{ mg/l Mo}$ 



Ø 24 mm

## prepare Zero press ZERO



- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.



- Add the contents of one Vario Molybdenum HR 1 F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add the contents of one Vario Molybdenum HR 2
   F10 Powder Pack straight from the foil to the same
   water sample.
- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add the contents of one Vario Molybdenum HR 3
   F10 Powder Pack straight from the foil to the same water sample.
- 10. Close the vial tightly with the cap and swirl several times to mix the contents
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

12. Press TEST key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Molybdate / Molybdenum.

#### Notes:

- 1. Filter turbid water samples using filter paper and funnel before analysis.
- 2. Highly buffered water samples or extreme pH values should be adjusted to a pH of nearly 7 with 1 mol/l Nitric acid or 1 mol/l Sodium hydroxide.
- 3. Concentrations above 10 mg/l Cu causes too high test values if the reaction time of 5 minutes is increased. So it is very important to perform the test procedure as described.
- 4. Substances which may interfere when present in concentrations at:

| Aluminium | 50 mg/l    |
|-----------|------------|
| Chromium  | 1000 mg/l  |
| Iron      | 50 mg/l    |
| Nickel    | 50 mg/l    |
| Nitrite   | all levels |

5. ▲ MoO₄ Mo Na₂MoO₄

| Reagent                   | Form of reagent/Quantity | Order-No. |
|---------------------------|--------------------------|-----------|
| Set                       |                          | 4535300   |
| VARIO Molybdenum HR 1 F10 | Powder Pack / 100        |           |
| VARIO Molybdenum HR 2 F10 | Powder Pack / 100        |           |
| VARIO Molybdenum HR 3 F10 | Powder Pack / 100        |           |







# Molybdate / Molybdenum HR with Liquid reagent

 $1 - 100 \text{ mg/l MoO}_{4} / 0.6 - 60 \text{ mg/l Mo}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 10 drops KS63 (Thioglycolate)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Molybdate / Molybdenum.

#### Notes:

1. Perform tests on sample water taken directly from the system. Molybdate will be absorbed onto the walls of sample containers and give low results.

2. MoO<sub>4</sub> Mo

▼ Na<sub>2</sub>MoO<sub>4</sub>

| Reagent                      | Form of reagent/Quantity | Order-No. |
|------------------------------|--------------------------|-----------|
| KS63 (Thoiglycolate Reagent) | Liquid reagent / 65 ml   | 56L006365 |







### Nickel with Tablet

0.1 - 10 mg/l Ni



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one NICKEL No. 1 tablet straight from the foil to the 10 ml water sample, crush the tablet using a clean stirring rod and dissolve the tablet completely (Note 1).
- Add one NICKEL No. 2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

# Zero accepted prepare Test press TEST

## 9. Press **TEST** key.

Wait for a reaction period of 2 minutes.

## Countdown 2:00

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Ni.

#### Notes:

- 1. If Iron is present in the sample, add one level spoonful of Nickel PT powder to the sample (after adding Nickel No. 1) and mix.
- 2. The presence of cobalt at 0.5 mg/l gives a positive response in the test.
- 3. The presence of higher levels of EDTA (at least 25 mg/l) complexes nickel and reduces response in the test. Complexing agents used in water treatment, such as polyphosphates, do not affect the results.

| Reagent      | Form of reagent/Quantity | Order-No. |
|--------------|--------------------------|-----------|
| NICKEL No. 1 | Tablet / 100             | 4515630BT |
| NICKEL No. 2 | Tablet / 100             | 4515640BT |







# Nitrate with Tablet and Powder

0.08 - 1 mg/l N



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and empty the vial.
- Fill the Nitrate test tube with 20 ml of the water sample.
- 6. Add 1 level spoon of Nitrate Test powder.
- 7. Close the tube tightly with the cap and swirl vigorously for one minute.
- 8. Add **one NITRATE TEST tablet** straight from the foil to the water sample.
- 9. Close the tube tightly with the cap and swirl vigorously for one minute
- 10. Stand the tube upright and after the reducing agent has settled to the bottom, gently invert it three to four times so as to complete the flocculation of the reducing agent. Then let the tube stand for a further 2 minutes. Open the tube and wipe around the top of the tube with a clean tissue to remove any residuals of the reducing agent.
- 11. Carefully decant 10 ml of the treated solution into the vial (24 mm Ø) used for zeroing, ensuring that no reducing agent is carried over.

- Add one NITRITE LR tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 13. Close the vial tightly with the cap and swirl several times until the tablet is dissolved
- 14. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 15. Press TEST key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Nitrate.

#### Notes:

- 1. If Nitrite is present in the sample as well as nitrate, it will react with the NITRITE LR-tablet, leading to a high result. For correction, carry out a nitrite determination using method 270 in NO<sub>2</sub>-N and subtract the result from the nitrate reading in NO<sub>3</sub>-N to give the corrected result.
- 2. Concentration of nitrate nitrogen above 1 mg/l (e.g. 50 mg/l) lead to an apricot colour instead of the reddish pink solution after the reaction time of 10 minutes. This colour cannot be correctly measured by the photometer. The result displayed does not show the concentration of nitrate nitrogen. The range of the test can be extended by first diluting the water sample with deionised water. One standard method is to dilute 1.0 ml of sample up to 100 ml (dilution factor of 100). The subsequent result of the test must then be multiplied by the dilution factor.
- 3. The following ions can produce interference as under the reaction conditions they can cause precipitation: antimony(III), iron(III), lead, mercury(I), silver, chloroplatinate, metavanadate and bismuth. Copper(II) ions may give a low result as they accelerate the decomposition of the diazonium salt. It is improbable in practice that these interfering ions will occur in such high concentrations that they cause significant errors.

| Reagent / Accessories | Form of reagent/Quantity | Order-No. |
|-----------------------|--------------------------|-----------|
| NITRATE TEST          | Powder 15 g              | 465230    |
| NITRATE TEST          | Tablet / 100             | 502810    |
| NITRITE LR            | Tablet / 100             | 4512310BT |
| Nitrate test tube     |                          | 366220    |







# Nitrate with Tube Test

1 - 30 mg/l N



Insert the adapter for 16 mm Ø vials.

- 1. Open one white capped vial (Reagent A), add 1 ml of the water sample and close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks are  $\lambda$  aligned.

## prepare Zero press ZERO





- 4. Remove the vial from the sample chamber.
- Add the contents of one Vario Nitrate Chromotropic Powder Pack straight from the foil into the same water sample.
- 6. Close the vial tightly with the cap and invert gently several times (10 x) to mix the contents (Note 1).
- 7. Place the vial in the sample chamber making sure that the marks are  $\frac{1}{\lambda}$  aligned.

# Zero accepted prepare Test press TEST

## Countdown 5:00

8. Press TEST key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Nitrate.

#### Notes:

- 1. Some solids may not dissolve.
- 2. To optimise the readings an optional batch related calibration can be performed. Follow the procedure using 1 ml deionised water in place of the sample and subtract the reagent blank value from the final result.
- 3. Conversion:  $mg/l NO_3 = mg/l N \times 4.43$
- 4. N NO3

| Reagent   | Form of reagent/Quantity               | Order-No. |
|---|--|-----------|
| Set   | Set                                    | 4535580   |
| VARIO Nitrate Chromotropic VARIO Nitra X Reagent tube | Powder Pack / 50<br>Reaction tube / 50 |           |
| VARIO deionised water                                 | 100 ml                                 |           |







### Nitrite with Tablet

0.01 - 0.5 mg/l N



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one NITRITE LR tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 Press TEST key.
 Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Nitrite.

#### Notes:

1. The following ions can produce interferences since under the reaction conditions they cause precipitation:

Antimony (III), Iron (III), Lead, Mercury (I), Silver, Chloroplatinate, Metavanadate and Bismuth.

Copper (II)-ions may cause lower test results as they accelerate the decomposition of the Diazonium salt.

It is unlikely in practice that these interfering ions will occur in such high concentrations that they cause significant reading errors.

2. Conversion:

 $mg/l NO_3 = mg/l N \times 3.29$ 



| Reagent    | Form of reagent/Quantity | Order-No. |
|------------|--------------------------|-----------|
| NITRITE LR | Tablet / 100             | 4512310BT |







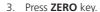
# Nitrite LR with Vario Powder Pack

0.01 - 0.3 mg/l N



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO





- 4. Remove the vial from the sample chamber.
- Add the contents of one VARIO Nitri 3 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 20:00

8. Press **TEST** key.

Wait for a reaction period of 20 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Nitrite.

#### Notes:

- 1. Interferences:
  - Strong oxidizing and reducing substances interfere.
  - Cupric and ferrous ions cause low results.
  - Antimonous, Auric, Bismuth, Chloroplatinate, Ferric, Lead, Mercurous, Metavanadate, Silver ions interfere by causing precipitation.
  - In samples with very high concentrations of Nitrate (> 100 mg/L N) a small amount of Nitrite will be found. Such high levels of Nitrate appear to undergo a slight amount of reduction to Nitrite, either spontaneously or during the reaction time of the test.



| Reagent           | Form of reagent/Quantity | Order-No. |
|-------------------|--------------------------|-----------|
| Vario Nitri 3 F10 | Powder Pack / 100        | 4530980   |







# Nitrogen, total LR with Vario Tube Test

0.5 - 25 mg/l N



Insert the adapter for 16 mm Ø vials.

- Open two TN Hydroxide LR digestion vials and add the contents of one Vario TN Persulfate Rgt. Powder Pack (Note 2, 3).
- 2. Add **2 ml deionised water** to the prepared vial (this is the blank, Note 4, 5).
- 3. Add **2 ml of the water sample** to the other prepared vial (this is the sample).
- 4. Close the vials with the caps and shake to mix the contents (at least 30 seconds, Note 6).
- Heat the vials for 30 minutes in the preheated reactor at a temperature of 100°C (Note 7).
- After 30 minutes remove the vials from the reactor. (CAUTION: The vials are hot!)
   Allow the vials to cool to room temperature.
- Open the cooled digestion vials and add the contents of one Vario TN Reagent A Powder Pack to each vial (Note 2).
- 8. Close the vials with the caps and shake to mix the contents (at least 15 seconds).

Countdown 3:00 start: 🚽

- Press [4] key.
   Wait for a reaction period of 3 minutes.
   After the reaction period is finished proceed as follows:
- Open the digestion vials and add the contents of one Vario TN Reagent B Powder Pack to each vial (Note 2).

11. Close the vials with the caps and shake to mix the contents (at least 15 seconds, Note 8).

#### Countdown 2:00 start:

12. Press [ ] key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished proceed as follows:

- Open two TN Acid LR/HR (Reagent C) vials and add 2 ml of the digested, treated blank to one vial (this is the blank).
- 14. Add **2 ml of the digested, treated water sample** to the other TN Acid LR/HR vial (this is the sample).
- 15. Close the vials with the caps and swirl the vials gently several times to mix the contents (10 x, Note 9). **(CAUTION: Vials warm up).**
- 16. Place the vial (the blank) in the sample chamber making sure that the marks  $\lambda$  are aligned.

# prepare Zero press ZERO

## Countdown 5:00

17. Press ZERO key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

- 18. Remove the vial from the sample chamber.

Zero accepted prepare Test press TEST

20. Press TEST key.

The result is shown in the display in mg/l Nitrogen.

Notes and Reagent: see page 192







# Nitrogen, total HR with Vario Tube Test

5 - 150 mg/l N



Insert the adapter for 16 mm Ø vials.

- Open two TN Hydroxide HR digestion vials and add the contents of one Vario TN Persulfate Rgt. Powder Pack (Note 2, 3).
- 2. Add **0.5 ml deionised water** to the prepared vial (this is the blank, Note 4, 5).



- 3. Add **0.5 ml of the water sample** to the other prepared vial (this is the sample).
- 4. Close the vials with the caps and shake to mix the contents (at least 30 seconds, Note 6).
- Heat the vials for 30 minutes in the preheated reactor at a temperature of 100°C (Note 7).
- After 30 Minutes remove the vials from the reactor. (CAUTION: The vials are hot!)
   Allow the vials to cool to room temperature.
- 7. Open the cooled digestion vials and add the contents of **one Vario TN Reagent A Powder Pack** to each vial (Note 2).
- 8. Close the vials with the caps and shake to mix the contents (at least 15 seconds)

Countdown 3:00 start: 🚽

- Press [ ] key.
   Wait for a reaction period of 3 minutes.
   After the reaction period is finished proceed as follows:
- Open the digestion vials and add the contents of one Vario TN Reagent B Powder Pack to each vial (Note 2).

11. Close the vials with the caps and shake to mix the contents (at least 15 seconds, Note 8).

#### Countdown 2:00 start:

12. Press [ ] key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished proceed as follows:

- 13. Open **two TN Acid LR/HR (Reagent C) vials** and add **2 ml of the digested, treated blank** to one vial (this is the blank).
- 14. Add **2 ml of the digested, treated water sample** to the other TN Acid LR/HR vial (this is the sample).
- 15. Close the vials with the caps and swirl the vials gently several times to mix the contents (10 x, Note 9). **(CAUTION: Vials warm up)**.
- 16. Place the vial (the blank) in the sample chamber making sure that the  $\lambda$  marks are aligned.

## prepare Zero press ZERO

## Countdown 5:00

17. Press **ZERO** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

- 18. Remove the vial from the sample chamber.
- 19. Place the vial (the sample, Note 10) in the sample chamber making sure that the  $\frac{1}{4}$  marks are aligned.

Zero accepted prepare Test press TEST

20. Press TEST key.

The result is shown in the display in mg/l Nitrogen.

Notes and Reagent: see page 192

#### Notes:

- Appropriate safety precautions and a good lab technique should be used during the whole procedure.
- 2. Use a funnel to add the reagent.
- 3. Wipe off any Persulfate reagent that may get on the lid or the tube threads.
- 4. Nitrogen, total HR:

Volumes for samples and blank should always be metered by using 2 ml volumetric pipettes (class A).

Nitrogen, total HR:

Volumes for samples and blank should always be metered by using suitable pipettes (class A).

- 5. One blank is sufficient for each set of samples.
- 6. The reagent may not dissolve completely.
- 7. It is very important to remove the vials from the reactor after exactly 30 minutes.
- 8. The reagent will not completely dissolve.
- 9. Hold the vial in a vertical position with the cap pointing up. Turn the vial upside-down. Wait for all of the solution to flow down to the cap. Return the vial to the upright position. Wait for all the solution to flow to the bottom of the vial. This process is one inversion; 10 inversions = approx. 30 seconds.
- 10. The zero (stored in the dark) can be used for 7 days, if the measured samples were prepared with the same batch of reagent.
- 11. Large quantities of nitrogen free, organic compounds which are included in some water samples may reduce the effectiveness of the digestion by reacting with the Persulfate reagent. Samples which are well known to contents large quantities of organic compounds must be diluted and digestion and measurement must be repeated for checking the effectiveness of the digestion.
- 12. Application: for water, wastewater and seawater
- 13. Interferences:

Interfering substances that resulted in a concentration change of 10%: Bromide more than 60 mg/l and Chloride more than 1000 mg/l produce positive interferences.

TN = Total Nitrogen

14.

N NH,

 $\blacksquare$ 

NH,

## Nitrogen, total LR with Vario Tube Test

| Reagent                  | Form of reagent/Quantity | Order-No. |
|--------------------------|--------------------------|-----------|
| Tube test contains:      | Set                      | 4535550   |
| VARIO TN HYDROX LR Tube  | Digestion tube / 50      |           |
| VARIO PERSULFATE Reagent | Powder Pack / 50         |           |
| VARIO TN Reagent A       | Powder Pack / 50         |           |
| VARIO TN Reagent B       | Powder Pack / 50         |           |
| VARIO TN ACID LR/HR Tube | Reaction tube / 50       |           |
| VARIO deionised water    | 100 ml                   |           |

### Nitrogen, total HR with Vario Tube Test

| Reagent   | Form of reagent/Quantity  | Order-No. |
|---|---|-----------|
| Tube test contains: VARIO TN HYDROX HR Tube VARIO PERSULFATE Reagent VARIO TN Reagent A VARIO TN Reagent B VARIO TN ACID LR/HR Tube VARIO deionised water | Set Digestion tube / 50 Powder Pack / 50 Powder Pack / 50 Powder Pack / 50 Powder Pack / 50 Reaction tube / 50 100 ml | 4535560   |







# Oxygen, active \* with Tablet

 $0.1 - 10 \text{ mg/l O}_{2}$ 



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one DPD No. 4 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

8. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l active Oxygen.

#### Notes:

- \* Active Oxygen is a synonym for a common disinfectant (based on "Oxygen") in Swimming Pool Treatment.
- 1. When preparing the sample, the lost of Oxygen, e.g. by pipetting or shaking, must be avoided.
- 2. The analysis must take place immediately after taking the sample.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| DPD No. 4 | Tablet / 100             | 4511220BT |







# Oxygen, dissolved with Vacu-vials® K-7553 (see Notes)

 $10 - 800 \mu g/l O_{2}$ 

Insert the adapter for 13 mm Ø round vials.

1. Place the blank in the sample chamber. The blank is part of the test kit.

## prepare Zero press ZERO

2. Press ZERO key.

- 3. Remove the blank from the sample chamber.
- 4. Water should flow through the special sample container for several minutes to remove any air bubbles sticking at the surface.

The water must flow from the bottom to the top.



 When the sample container is bubble-free press one Vacu-vial® into the lower edge of the sample container. The Vacu-vial® breaks at the neck and the vial fills automatically.

A small volume of inert gas remains in the Vacu-vial®.

 Remove the Vacu-vial® point downwards from the sample container immediately.

As the contents of the vial has a higher density than water, it is important to remove the vial from the sample container within 5 seconds to prevent any loss of reagent.

- The Vacu-vial® is closed with one finger (covered with a glove) to prevent entry of air. Invert the vial several times. Dry the outside of the vial.
- 8. Place the Vacu-vial® in the sample chamber.

Zero accepted prepare Test press TEST

9. Press **TEST** key.

The result is shown in the display in µg/l Oxygen.

#### Notes:

- 1. This method is adapted from CHEMetrics. The measuring range and wavelength used for this photometer may differ from the data specified by CHEMetrics.
- 2. Read the original test instruction and the MSDS (delivered with the test) before performing the test. MSDS also available at www.chemetrics.com.
- 3. Vacu-vials® should be stored in the dark and at room temperature.
- 4. Vacu-vials $^{\circ}$  is a registered trade mark of the company CHEMetrics, Inc. / Calverton, U.S.A.

| Reagent                         | Form of reagent/Quantity | Order-No. |
|---------------------------------|--------------------------|-----------|
| Vacu-vials® / CHEMetrics K-7553 | Test-Kit / 30 Tests      | 380450    |
| 13-mm-adapter Ø                 |                          | 19802192  |







# Ozone with Tablet

 $0.02 - 2 \text{ mg/l O}_{3}$ 

Ozon

>> with Cl without Cl

The following selection is shown in the display:

>> with Cl

for the determination of Ozone in the presence of Chlorine.

>> without Cl

for the determination of Ozone in the absence of Chlorine.

Select the desired method with the arrow keys [A] and [V]. Confirm with [A] key.

#### Notes:

1. Vial cleaning:

As many household cleaners (e.g. dishwasher detergent) contain reducing substances, the subsequent determination of Ozone may show lower results. To avoid any measurement errors, only use glassware free of Chlorine demand.

Preparation: Put all applicable glassware into Sodium hypochlorite solution (0.1 g/l) for one hour, then rinse all glassware thoroughly with deionised water.

- 2. Preparing the sample:
  - When preparing the sample, the lost of Ozone, e.g. by pipetting or shaking, must be avoided. The analysis must take place immediately after taking the sample.
- 3. The DPD colour development is carried out at a pH value of 6.2 to 6.5. The reagent tablet therefore contains a buffer for the pH adjustment.

  Strong alkaline or acidic water samples must be adjusted between pH 6 and pH 7 before the tablet is added (use 0.5 mol/l Sulfuric acid resp. 1 mol/l Sodium hydroxide).
- 4. Exceeding the measuring range:
  Concentrations above 6 mg/l Ozone can lead to results showing 0 mg/l. In this case, the water sample must be diluted with water free of Ozone. 10 ml of the diluted sample should be mixed with the reagent and the measurement repeated.
- 5. If ??? is displayed at the diffentiated test result see page 342.
- 6. Oxidising agents such as Bromine, Chlorine etc. interfere as they react in the same way as Ozone.







# Ozone, in the presence of Chlorine with Tablet

 $0.02 - 2 \text{ mg/l O}_{3}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty** it, leaving a few drops remaining in the vial.
- Add one DPD No. 1 tablet and one DPD No. 3 tablet straight from the foil and crush the tablets using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare T1 press TEST

Countdown 2:00 9. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

- 10. Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times.
- 11. Fill a second clean vial with 10 ml of water sample.
- 12. Add **one GLYCINE tablet** straight from the foil and crush the tablet using a clean stirring rod.

- 13. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 14. Add **one DPD No. 1 tablet** and **one DPD No. 3 tablet** straight from the foil into the first cleaned vial and crush the tablets using a clean stirring rod.
- 15. Transfer the contents of the second vial (Glycine solution) into the prepared vial (point 14).
- 16. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 17. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

T1 accepted prepare T2 press TEST

Countdown 2:00 18. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in:

\*,\*\* mg/l O<sub>3</sub>
\*,\*\* mg/l total Cl

mg/l Ozone mg/l total Chlorine

Notes: See page 199

| Reagent                | Form of reagent/Quantity                | Order-No. |
|------------------------|---|-----------|
| Set<br>DPD No. 1/No. 3 | Tablet / per 100 inclusive stirring rod | 4517711BT |
| DPD No. 1              | Tablet / 100                            | 4511050BT |
| DPD No. 3              | Tablet / 100                            | 4511080BT |
| GLYCINE                | Tablet / 100                            | 4512170BT |







### Ozone, in absence of Chlorine with Tablet

 $0.02 - 2 \text{ mg/l O}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and **empty it, leaving a few drops remaining in the vial**.
- Add one DPD No. 1 tablet and one DPD No. 3 tablet straight from the foil and crush the tablets using a clean stirring rod.
- 6. Add water sample to the 10 ml mark.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

Press TEST key. Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Ozone.

Notes: See page 199

| Reagent             | Form of reagent/Quantity                   | Order-No. |
|---------------------|--|-----------|
| Set DPD No. 1/No. 3 | Tablet / per 100<br>inclusive stirring rod | 4517711BT |
| DPD No. 1           | Tablet / 100                               | 4511050BT |
| DPD No. 3           | Tablet / 100                               | 4511080BT |
| GLYCINE             | Tablet / 100                               | 4512170BT |





# PHMB (Biguanide) with Tablet

2 - 60 mg/l PHMB



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one PHMB PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the 

  √ marks are aligned.

Zero accepted prepare Test press TEST

8. Press **TEST** key.

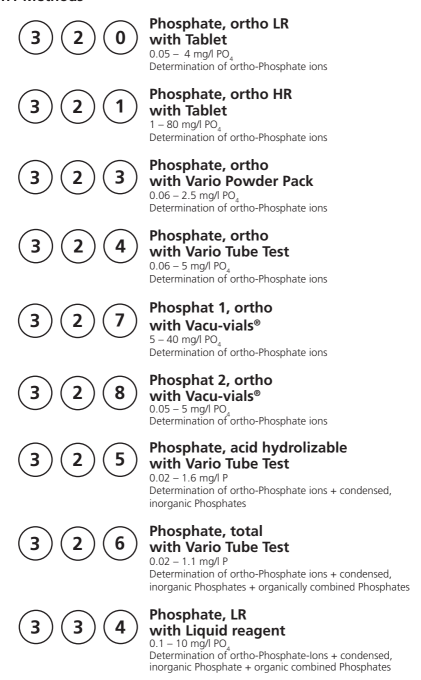
The result is shown in the display in mg/l PHMB.

#### Notes:

- 1. Clean vials with the brush immediately after analysis.
- 2. Vials and stirring rods may turn blue after prolonged use. In this case clean vials and stirring rods with a laboratory detergent (see chapter 1.2.2 Cleaning of vials and accessories for analysis). Rinse vials and caps thoroughly with tap water and then with deionised water.
- 3. The test result is influenced by Hardness and Total Alkalinity.
  The calibration of this method was done using water with the following concentration:
  Ca-Hardness: 200 mg/l CaCO.

Ca-Hardness: 200 mg/l CaCO<sub>3</sub> Total Alkalinity: 120 mg/l CaCO<sub>3</sub>

| Reagent         | Form of reagent/Quantity | Order-No. |
|-----------------|--------------------------|-----------|
| PHMB PHOTOMETER | Tablet / 100             | 4516100BT |









# Phosphate, HR with Liquid reagent

5 - 80 mg/l PO

Determination of ortho-Phosphate-Ions + condensed, inorganic Phosphate + organic combined Phosphates

Additional information can be found in the notes for each method

#### General:

Ortho-Phosphate ions react with the reagent to form an intense blue colour (methods **320**, **323**, **324**, **325** and **326**).

Phosphate in organic and condensed inorganic forms (meta-, pyro- and polyphosphates) must be converted to ortho-Phosphate ions before analysis.

Pretreatment of the sample with acid and heat provides the conditions for hydrolysis of the condensed inorganic forms. Organically combined phosphates are converted to ortho-Phosphate ions by heating with acid and persulfate.

The amount of organically combined phosphates can be calculated:

mg/l Phosphate, organic = mg/l Phosphate, total – mg/l Phosphate, acid hydrolysable

In methods **321** and **327** the ortho-Phosphate ions react with the Vanadate-molybdate-reagent under acid conditions to form a yellow coloured product.

## Notes – only for tube tests and tests with powder packs: 323, 324, 325, 326

1. Application: for water, wastewater and seawater.

Intentantantan automas

2. Highly buffered samples or samples with extreme pH values should be adjusted between pH 6 and pH 7 before analysis (with 1 mol/l Hydrochloric acid or 1 mol/l Sodium hydroxide).

Intenference level

3. Interferences:

Large amounts of turbidity may cause inconsistent results.

| interfering substance     | interference level:   |  |
|---------------------------|-----------------------|--|
| Aluminium                 | greater than 200 mg/l |  |
| Arsenate                  | at any level          |  |
| Chromium                  | greater than 100 mg/l |  |
| Copper                    | greater than 10 mg/l  |  |
| Iron                      | greater than 100 mg/l |  |
| Nickel                    | greater than 300 mg/l |  |
| Silica (Silicium dioxide) | greater than 50 mg/l  |  |
| Silicate                  | greater than 10 mg/l  |  |
| Sulfide                   | at any level          |  |
| Zinc                      | greater than 80 mg/l  |  |

### Phosphate, ortho ≜ Phosphorus, reactive







# Phosphate, ortho LR with Tablet

0.05 - 4 mg/l PO<sub>4</sub>



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close the cap tightly.

2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

## prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one PHOSPHATE No. 1 LR tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Add **one PHOSPHATE No. 2 LR tablet** straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 8. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.

Zero accepted prepare Test press TEST

9. Press **TEST** key.

Wait for a reaction period of 10 minutes.

#### Countdown 10:00

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. Only ortho-Phosphate ions react.
- 2. The tablets must be added in the correct sequence.
- 3. The test sample should have a pH-Value between 6 and 7.
- 4. Interferences:

Higher concentrations of Cu, Ni, Cr (III), V (V) and W (VI) interfere due to their colour. Silicates do not interfere (masked by Citric acid in the tablets).

- 5. see also page 207
- 6. Conversion:  $mg/l P = mg/l PO_4 \times 0.33$  $mg/l P_2O_5 = mg/l PO_4 \times 0.75$
- 7. ▲ PO<sub>4</sub>
  P
  P<sub>2</sub>O<sub>5</sub>

| Reagent                           | Form of reagent/Quantity                | Order-No. |
|-----------------------------------|---|-----------|
| Set<br>PHOSPHATE LR No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517651BT |
| PHOSPHATE LR No. 1                | Tablet / 100                            | 4513040BT |
| PHOSPHATE LR No. 2                | Tablet / 100                            | 4513050BT |







## Phosphate HR, ortho with Tablet

1 - 80 mg/l PO<sub>4</sub>



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one PHOSPHATE HR P1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Add one PHOSPHATE HR P2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 9. Press **TEST** key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. For samples under 5 mg/l  $PO_4$  it is reccommended to analyse the water sample with method 320 "Posphate LR, ortho with Tablet".
- 2. Only ortho-Phosphate ions react.
- 3. see also page 207
- 4. Conversions:  $mg/l P = mg/l PO_4 \times 0.33$  $mg/l P_2O_5 = mg/l PO_4 \times 0.75$
- 5. A PO<sub>4</sub> P P<sub>2</sub>O<sub>5</sub>

| Reagent                       | Form of reagent/Quantity                | Order-No. |
|-------------------------------|---|-----------|
| SET<br>PHOSPHATE HR P 1 / P 2 | Tablet / per 100 inclusive stirring rod | 4517661BT |
| PHOSPHATE HR P1               | Tablet / 100                            | 4515810BT |
| PHOSPHATE HR P2               | Tablet / 100                            | 4515820BT |







## Phosphate, ortho with Vario Powder Pack

 $0.06 - 2.5 \text{ mg/l PO}_{4}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO





4. Remove the vial from the sample chamber.

- Add the contents of one VARIO Phosphate Rgt. F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 10-15 sec., Note 1).
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press **TEST** key.

Countdown 2:00

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. The reagent does not dissolve completely.
- 2. see also page 207
- 3. Conversions:  $mg/l \ P = mg/l \ PO_4 \ x \ 0.33$   $mg/l \ P_2O_5 = mg/l \ PO_4 \ x \ 0.75$
- 4. A PO<sub>4</sub>
  P
  P<sub>2</sub>O<sub>5</sub>

| Reagent                | Form of reagent/Quantity                         | Order-No. |
|------------------------|--|-----------|
| Set<br>VARIO PHOS3 F10 | Powder Pack /<br>2 x 50 VARIO PHOSPHATE RGT. F10 | 4531550   |







## Phosphate, ortho with Vario Tube Test

 $0.06 - 5 \text{ mg/l PO}_{4}$ 



Insert the adapter for 16 mm Ø vials.

- Open the white cap of one tube PO<sub>4</sub>-P Dilution and add 5 ml of the water sample.
- 2. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.

### prepare Zero press ZERO





- 4. Remove the vial from the sample chamber.
- Add the contents of one VARIO Phosphate Rgt. F10 Powder Pack straight from the foil to the water sample (Note 1).
- 6. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 10-15 sec., Note 2).
- 7. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press TEST key.

Countdown 2:00

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. Use a funnel to add the reagent.
- 2. The reagent does not dissolve completely.
- 3. see also page 207
- 4. Conversions: mg/l P = mg/l PO $_4$  x 0.33 mg/l P $_2$ O $_5$  = mg/l PO $_4$  x 0.75
- 5. ▲ PO<sub>4</sub>
  P
  P,O<sub>5</sub>

| Reagent                    | Form of reagent/Quantity | Order-No. |
|----------------------------|--------------------------|-----------|
| Set                        | Set                      | 4535200   |
| VARIO Dilution Vial        | Reaction tube / 50       |           |
| VARIO PHOSPHATE RGT F10 PP | Powder Pack / 50         |           |
| VARIO deionised water      | 100 ml                   |           |







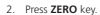
## Phosphate 1, ortho with Vacu-vials® K-8503 (see Notes)

5 - 40 mg/l PO<sub>4</sub>

Insert the adapter for 13 mm Ø vials.

1. Place the blank in the sample chamber. The blank is part of the test kit.

### prepare Zero press ZERO



sample.



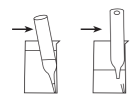


- ·
- Place one Vacu-vial® in the sample container. Snap the tip by pressing the vial against the side of the sample container.

4. Fill the sample container to the 25 ml mark with the water

The Vacu-vial® breaks at the neck and the vial fills automatically.

A small volume of inert gas remains in the Vacu-vial®.



- 6. Mix the contents of the Vacu-vial® by inverting it several times, allowing the bubble to move from one end to the other. Dry the outside of the vial.
- 7. Place the Vacu-vial® in the sample chamber.

Zero accepted prepare Test press TEST

8. Press TEST key.

Wait for a reaction period of 5 minutes.

Countdown 5:00

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. This method is adapted from CHEMetrics. The measuring range and wavelength used for this photometer may differ from the data specified by CHEMetrics.
- 2. Read the original test instruction and the MSDS (delivered with the test) before performing the test. MSDS also available at www.chemetrics.com.
- 3. Vacu-vials® is a registered trade mark of the company CHEMetrics, Inc. / Calverton, U.S.A.
- 4. Only ortho-Phosphate ions react.
- 5. Sulfide, Thiosulfate and Thiocyanate cause low test results.



| Reagent                         | Form of reagent/Quantity | Order-No. |
|---------------------------------|--------------------------|-----------|
| Vacu-vials® / CHEMetrics K-8503 | Test-Kit / 30            | 380460    |







## Phosphate 2, ortho with Vacu-vials® K-8513 (see Notes)

 $0.05 - 5 \text{ mg/l PO}_{4}$ 

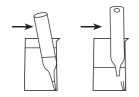
Insert the adapter for 13 mm Ø vials.

 Place the blank in the sample chamber. The blank is part of the test kit.

### prepare Zero press ZERO







## Zero accepted prepare Test press TEST

#### Countdown 3:00

#### 2. Press **ZERO** key.

- 3. Remove the blank from the sample chamber.
- 4. Fill the sample container to the 25 ml mark with the water sample.
- 5. Fill the sample container with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 2 drops A-8500 Activator Solution

- Close the sample container with the cap tightly and swirl several times to mix the contents.
- 7. Place one Vacu-vial® in the sample container. Snap the tip by pressing the vial against the side of the sample container. The Vacu-vial® breaks at the neck and the vial fills automatically. A small volume of inert gas remains in the Vacu-vial®.
- 8. Mix the contents of the Vacu-vial® by inverting it several times, allowing the bubble to move from one end to the other. Dry the outside of the vial.
- 9. Place the Vacu-vial® in the sample chamber.

#### 10. Press **TEST** key.

Wait for a reaction period of 3 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l ortho-Phosphate.

#### Notes:

- 1. This method is adapted from CHEMetrics. The measuring range and wavelength used for this photometer may differ from the data specified by CHEMetrics.
- 2. Read the original test instruction and the MSDS (delivered with the test) before performing the test. MSDS also available at www.chemetrics.com.
- 3. Vacu-vials® is a registered trade mark of the company CHEMetrics, Inc. / Calverton, U.S.A.
- 4. Only ortho-Phosphate ions react.
- 5. Sulfide, Thiosulfate and Thiocyanate cause low test results.



▼ P<sub>2</sub>O<sub>5</sub>

| Reagent                                     | Form of reagent/Quantity | Order-No. |
|---|--------------------------|-----------|
| Vacu-vials <sup>®</sup> / CHEMetrics K-8513 | Test-Kit / 30            | 380480    |







### Phosphate, acid hydrolyzable with Vario Tube Test

 $0.02 - 1.6 \text{ mg/l P} (\triangleq 0.06 - 5 \text{ mg/l PO}_a)$ 



Insert the adapter for 16 mm Ø vials.

- Open the white cap of one digestion tube PO4-P Acid reagent and add 5 ml of the water sample.
- 2. Close the vial tightly with the cap and invert gently several times to mix the contents.
- Heat the vials for 30 minutes in the preheated reactor at a temperature of 100°C.
- After 30 minutes remove the vial from the reactor. (CAUTION: The vials are hot!)
   Allow the vials to cool to room temperature.
- Open the cooled digestion vial and add 2 ml 1.00 N Sodium Hydroxide solution to the vial.
- 6. Close the vial with the cap and invert gently several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.
- 8. Press **ZERO** key.
- 9. Remove the vial from the sample chamber.
- 10. Add the contents of **one VARIO Phosphate Rgt. F10 Powder Pack** straight from the foil to the vial (Note 2).
- 11. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 10-15 sec., Note 3).
- 12. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.
- 13. Press TEST key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l acid hydrolyzable Phosphate.

### prepare Zero press ZERO



Zero accepted prepare Test press TEST

Countdown 2:00

#### Notes:

- 1. Appropriate safety precautions and a good lab technique should be used during the whole procedure.
- 2. Use a funnel to add the reagent.
- 3. The reagent does not dissolve completely.
- 4. see also page 207
- 5. Conversions:  $mg/l PO_4 = mg/l P \times 3.07$  $mg/l P_2O_5 = mg/l P \times 2.29$
- 6. ▲ PO<sub>4</sub> P

| $\mathrm{P_2O_5}$ |
|-------------------|

| Reagent                        | Form of reagent/Quantity | Order-No. |
|--------------------------------|--------------------------|-----------|
| Set:                           | Set                      | 4535250   |
| VARIO Acid Reagent Vial        | Reaction tube / 50       |           |
| VARIO PHOSPHATE RGT F10 PP     | Powder Pack / 50         |           |
| VARIO Potassium F10 Persulfate | Powder Pack / 50         |           |
| VARIO Natriumhydroxid 1,54 N   | Solution / 100 ml        |           |
| VARIO deionised water          | 100 ml                   |           |
| VARIO Natriumhydroxid 1,00 N   | Solution / 100 ml        |           |







## Phosphate, total with Vario Tube Test

 $0.02 - 1.1 \text{ mg/l P} (\triangleq 0.06 - 3.5 \text{ mg/l PO}_a)$ 





Insert the adapter for 16 mm Ø vials.

- Open the white cap of one digestion tube PO4-P Acid reagent and add 5 ml of the water sample.
- Add the contents of one Vario Potassium Persulfate F10 Powder Pack straight from the foil to the vial (Note 2).
- 3. Close the vial tightly with the cap and invert several times to mix the contents.
- Heat the vials for 30 minutes in the preheated reactor at a temperature of 100°C.
- After 30 minutes remove the vial from the reactor. (CAUTION: The vials are hot!)
   Allow the vials to cool to room temperature.
- Open the cooled digestion vial and add 2 ml 1.54 N Sodium Hydroxide Solution to the vial.
- 7. Close the vial with the cap and invert gently several times to mix the contents.
- 8. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.

### prepare Zero press ZERO

- 9. Press **ZERO** key.
- 10. Remove the vial from the sample chamber.
- 11. Add the contents of **one VARIO Phosphate Rgt. F10 Powder Pack** straight from the foil to the vial (Note 2).
- 12. Close the vial tightly with the cap and swirl several times to mix the contents (approx. 10-15 sec., Note 3).
- 13. Place the vial in the sample chamber making sure that the  $\frac{1}{\lambda}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00 14. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l total Phosphate.

#### Notes:

- 1. Appropriate safety precautions and a good lab technique should be used during the whole procedure.
- 2. Use a funnel to add the reagent.
- 3. The reagent does not dissolve completely.
- 4. see also page 207
- 5. Conversions:  $mg/l PO_4 = mg/l P \times 3.07$  $mg/l P_2O_5 = mg/l P \times 2.29$
- 6. ▲ P
  PO<sub>4</sub>
  P<sub>2</sub>O<sub>5</sub>

| Reagent  | Form of reagent/Quantity  | Order-No. |
|--|---|-----------|
| VARIO Acid Reagent Vial VARIO PHOSPHATE RGT F10 PP VARIO Potassium F10 Persulfate VARIO Natriumhydroxid 1,54 N VARIO deionised water | Set Reaction tube / 50 Powder Pack / 50 Powder Pack / 50 Solution / 100 ml 100 ml | 4535210   |







## Phosphate LR with Liquid reagent

0.1 - 10 mg/l PO<sub>4</sub>

This test is suitable for determining ortho-Phosphate in boiler waters and potable water supplies. Samples should be filtered prior to analysis to remove any suspended insoluble phosphate. A GF/C filter is suitable.

Unscrew the two halves of the filter holder and place one GF/C filter circle onto the base section. Screw the two parts together again, **ensuring the O ring is correctly located**.

- Fill a clean 20 ml syringe with approx. 14 ml water sample.
- Connect the syringe to the filtration assembly and discharge the syringe to waste, down to the 10 ml mark.
- Fill a clean vial (24 mm Ø) with 10 ml of water sample from the prepared syringe, close tightly with the cap.
- 4. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 5. Press **ZERO** key.
- 6. Remove the vial from the sample chamber.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

50 drops KS80 (CRP)

8. Close the vial tightly with the cap and invert several times to mix the contents

- Add one level spoon of reagent KP119 (Ascorbic Acid) to the same water sample (note 1).
- 10. Close the vial tightly with the cap and swirl several times to dissolve the powder.
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 12. Press **TEST** key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Phosphate.

#### Notes:

- 1. For correct dosage the spoon supplied with the reagents must be used.
- 2. For the analysis of Polyphosphate and total Phosphate a prior digestion is required (see page 226).
- 3. Sample temperature should be between 15 and 30°C.
- 4. Conversions:  $mg/l P = mg/l PO_4 \times 0.33$  $mg/l P_2O_5 = mg/l PO_4 \times 0.75$
- 5. ▲ P
  PO<sub>4</sub>
  P<sub>2</sub>O<sub>5</sub>

| Reagent   | Form of reagent/Quantity  | Order-No.  |
|---|---|--|
| KS80 – CRP Reagent<br>KP119 – Ascorbic Acid   | Liquid reagent / 2 x 65 ml<br>Powder / 20 g   | 56L008065<br>56P011920                           |
| For digestion method:<br>KS278 (50% Sulphuric Acid)<br>KS135 (Phenolphthalein Substitute<br>Indikator)<br>KS144 (Calcium Hardness Puffer)<br>KP962 (Ammonium Persulfate Powder) | Liquid reagent / 65 ml<br>Liquid reagent / 65 ml<br>Liquid reagent / 65 ml<br>Powder / 20 g | 56L027865<br>56L013565<br>56L014465<br>56P096240 |







## Polyphosphate LR with Liquid reagent

0.1 - 10 mg/l PO<sub>4</sub>

This test will give total inorganic phosphate. Polyphosphate being determined by the difference of total inorganic phosphate and ortho-Phosphate.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- Add 15 drops of KS278 (50% Sulphuric Acid) to the same water sample.
- Boil for 20 minutes, maintaining the sample volume above 25 ml with deionised water.
- 4. Swirl gently several times to mix the contents and allow the Erlenmeyer flask to cool to room temperature.
- 5. Fill the Erlenmeyer flask with drops of the same size by holding the bottle vertically and squeeze slowly:
  - 2 drops KS135 (Phenolphthalein Substitute Indicator)
- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pale pink colour just appears.
- 7. Fill the sample up to 50ml with deionised water.
- 8. Proceed as in **point 3** of the method before (page 224).

The result is shown in the display in mg/l inorganic total Phosphate (ortho-Phosphate or Polyphosphate).







## Total Phosphate LR with Liquid reagent

0.1 - 10 mg/l PO<sub>4</sub>

This test will measure all phosphorous containing compounds present in the sample, including ortho-Phosphate, Polyphosphate and organic phosphorous compounds.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- 2. Add one **KT274 (Ammonium Persulfate) tablet** to the prepared water sample
- Add 15 drops of KS278 (50% Sulphuric Acid) to the same water sample.
- 4. Boil for **20 minutes**, maintaining the sample volume above 25 ml with deionised water.
- 5. Swirl gently several times to mix the contents and allow the Erlenmeyer flask to cool to room temperature.
- 6. Fill the Erlenmeyer flask with drops of the same size by holding the bottle vertically and squeeze slowly:

2 drops KS135 (Phenolphthalein Substitute Indicator)

- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pale pink colour just appears.
- 8. Fill the sample up to 50ml with deionised water.
- 9. Proceed as in **point 3** of the method before (page 224).

The result is shown in the display in mg/l total-Phosphate.







## Phosphate HR with Liquid reagent

5 - 80 mg/l PO<sub>4</sub>

This test is suitable for determining ortho-Phosphate in boiler waters and potable water supplies. Samples should be filtered prior to analysis to remove any suspended insoluble phosphate. A GF/C filter is suitable.

Unscrew the two halves of the filter holder and place one GF/C filter circle onto the base section. Screw the two parts together again, **ensuring the O ring is correctly located**.

- Fill a clean 20 ml syringe with approx. 14 ml water sample.
- Connect the syringe to the filtration assembly and discharge the syringe to waste, down to the 10 ml mark.
- Fill a clean vial (24 mm Ø) with 10 ml of water sample from the prepared syringe, close tightly with the cap.
- 4. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 5. Press **ZERO** key.
- 6. Remove the vial from the sample chamber.
- 7. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

25 drops KS228 (Ammonium Molybdate)

8. Close the vial tightly with the cap and invert several times to mix the contents

- 9. Add **25 drops of KS229 (Ammonium Metavanadate)** solution to the same water sample.
- 10. Close the vial tightly with the cap and invert several times to mix the contents
- 11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 12. Press **TEST** key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Phosphate.

#### Notes:

- 1. For the analysis of Polyphosphate and total Phosphate a prior digestion is required (see page 230).
- 2. Reagents and accessories available on request.
- 3. Conversions:  $mg/l P = mg/l PO_4 \times 0.33$  $mg/l P_2O_5 = mg/l PO_4 \times 0.75$
- 4. ▲ P
  PO<sub>4</sub>
  P<sub>2</sub>O<sub>5</sub>

| Reagent   | Form of reagent/Quantity  | Order-No.  |
|---|---|--|
| KS228 (Ammonium Molybdate)<br>KS229 (Ammonium Metavanadate)   | Liquid reagent / 65 ml<br>Liquid reagent / 65 ml  | 56L022865<br>56L022965                           |
| For digestion method:<br>KS278 (50% Sulphuric Acid)<br>KS135 (Phenolphthalein Substitute<br>Indikator)<br>KS144 (Calcium Hardness Puffer)<br>KP962 (Ammonium Persulfate Powder) | Liquid reagent / 65 ml<br>Liquid reagent / 65 ml<br>Liquid reagent / 65 ml<br>Powder / 20 g | 56L027865<br>56L013565<br>56L014465<br>56P096240 |







## Polyphosphate HR with Liquid reagent

5 - 80 mg/l PO<sub>4</sub>

This test will give total inorganic phosphate. Polyphosphate being determined by the difference of total inorganic phosphate and ortho-Phosphate.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- Add 15 drops of KS278 (50% Sulphuric Acid) to the same water sample.
- Boil for 20 minutes, maintaining the sample volume above 25 ml with deionised water.
- 4. Swirl gently several times to mix the contents and allow the Erlenmeyer flask to cool to room temperature.
- 5. Fill the Erlenmeyer flask with drops of the same size by holding the bottle vertically and squeeze slowly:
  - 2 drops KS135 (Phenolphthalein Substitute Indicator)
- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pale pink colour just appears.
- 7. Fill the sample up to 50ml with deionised water.
- 8. Proceed as in **point 3** of the method before (page 228).

The result is shown in the display in mg/l inorganic total Phosphate (ortho-Phosphate or Polyphosphate).







## Total Phosphate HR with Liquid reagent

5 – 80 mg/l PO<sub>4</sub>

This test will measure all phosphorous containing compounds present in the sample, including ortho-Phosphate, Polyphosphate and organic phosphorous compounds.

- Fill a clean 100-ml-Erlenmeyer flask with 50 ml homogenized sample.
- 2. Add one **KT274 (Ammonium Persulfate) tablet** to the prepared water sample
- Add 15 drops of KS278 (50% Sulphuric Acid) to the same water sample.
- Boil for 20 minutes, maintaining the sample volume above 25 ml with deionised water.
- 5. Swirl gently several times to mix the contents and allow the Erlenmeyer flask to cool to room temperature.
- 6. Fill the Erlenmeyer flask with drops of the same size by holding the bottle vertically and squeeze slowly:

2 drops KS135 (Phenolphthalein Substitute Indicator)

- Add drops of KS144 (Calcium Hardness Buffer), one drop at a time with mixing, until a pale pink colour just appears.
- 8. Fill the sample up to 50ml with deionised water.
- 9. Proceed as in **point 3** of the method before (page 228).

The result is shown in the display in mg/l total-Phosphate.

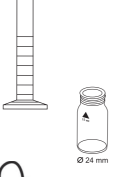




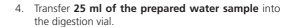


# Phosphonates Persulfate UV oxidation method with Vario Powder Pack

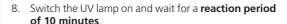
0 - 125 mg/l (see Table 1)

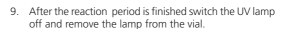


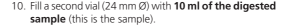
- 1. Choose the appropriate sample volume from table 1 (see following pages).
- Pipette the chosen sample volume into a clean 50 ml graduated cylinder. If necessary fill up with deionised water to the 50 ml mark and mix well.
- 3. Fill a clean vial (24 mm Ø) with 10 ml of the prepared water sample (this is the blank).



- Add the contents of one Vario Potassium Persulfate F10 Powder Pack straight from the foil to the digestion vial.
- Close the digestion vial tightly with the cap and swirl until the reagent is dissolved completely.
- 7. Insert the UV lamp into the digestion vial (Note 3, 4, 5). **CAUTION: Wear UV safety goggles!**









Countdown 1 10:00 start:



- Add the contents of one Vario Phosphate Rgt. F10 Powder Pack straight from the foil into each vial (blank and sample).
- 12. Close the vials tightly with the cap and swirl gently several times (30 sec.). (Note 6)

prepare Zero press ZERO

Countdown 2:00

13. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

14. Press ZERO key.

Wait for a reaction period of 2 minutes (Note 7).

After the reaction period is finished the measurement starts automatically.

15. Remove the vial from the sample chamber.

16. Place the vial (the sample) in the sample chamber making sure that the  $\chi$  marks are aligned.

Zero accepted prepare Test press TEST

17. Press **TEST** kev.

The result is shown in the display in mg/L PO<sub>4</sub>3-.

To calculate the actual phosphonate concentration multiply the reading with the corresponding dilution factor from table 1.

To calculate the active phosphonate concentration multiply the actual phosphonate concentration using the appropriate factor from table 2.

#### Notes:

- 1. Rinse all glassware with 1:1 Hydrochloric acid first and then rinse with deionised water. Do not use detergents with phosphates.
- 2. During UV digestion Phosphonates are converted to ortho-Phosphates.

  This step is normally completed in 10 minutes. High organic loaded samples or a weak lamp can cause incomplete phosphate conversion.
- 3. UV lamp available on request.
- 4. While the UV lamp is on UV safety goggles must be worn.
- 5. For handling of the UV lamp see manufacturer's manual.

  Do not touch the surface of the UV lamp. Fingerprints will etch the glass.

  Wipe the UV lamp with a soft and clean tissue between measurements.
- 6. The reagent does not dissolve completely.
- 7. The given reaction time of 2 minutes refers to a water sample temperature of more than 15°C. At a sample temperature lower than 15°C a reaction time of 4 minutes is required.

Tables and Reagent: see next page

#### Table 1:

| Expected range (mg/L Phosphonate) | Sample volume<br>in ml | Factor |
|-----------------------------------|------------------------|--------|
| 0 – 2.5                           | 50                     | 0.1    |
| 0 – 5.0                           | 25                     | 0.2    |
| 0 – 12.5                          | 10                     | 0.5    |
| 0 – 25                            | 5                      | 1.0    |
| 0 – 125                           | 1                      | 5.0    |

#### Table 2:

| Phosphonate type | Conversion factor for active phosphonate |  |
|------------------|--|--|
| PBTC             | 2.840                                    |  |
| NTP              | 1.050                                    |  |
| HEDPA            | 1.085                                    |  |
| EDTMPA           | 1.148                                    |  |
| HMDTMPA          | 1.295                                    |  |
| DETPMPA          | 1.207                                    |  |
| HPA              | 1.490                                    |  |

| Reagent                        | Form of reagent/Quantity | Order-No. |
|--------------------------------|--------------------------|-----------|
| Set:                           |                          | 4535220   |
| VARIO Potassium F10 Persulfate | Powder Pack / 50         |           |
| VARIO PHOSPHATE RGT F10 PP     | Powder Pack / 100        |           |

Interference levels decrease with increasing sample volume. Example: Iron interferes above 200 mg/L if a sample volume of 5 ml is used. At a sample volume of 10 ml the interference level decreases to 100 mg/L.

Table 3:

| Interfering substances                       | Interference level using 5 ml of sample   |
|--|---|
| Aluminium                                    | 100 mg/l  |
| Arsenate                                     | interferes at all concentrations  |
| Benzotriazole                                | 10 mg/l   |
| Bicarbonate                                  | 1000 mg/l   |
| Bromide                                      | 100 mg/l  |
| Calcium                                      | 5000 mg/l   |
| CDTA   | 100 mg/l  |
| Chloride                                     | 5000 mg/l   |
| Chromate                                     | 100 mg/l  |
| Copper                                       | 100 mg/l  |
| Cyanide                                      | 100 mg/l; increase the UV digestion to 30 minutes                                 |
| Diethanoldithiocarbamate                     | 50 mg/l   |
| EDTA   | 100 mg/l  |
| Iron   | 200 mg/l  |
| Nitrate                                      | 200 mg/l  |
| NTA  | 250 mg/l  |
| ortho-Phosphate                              | 15 mg/l   |
| Phosphite and organophosphorus compounds     | reacts quantitatively;<br>Meta- and Polyphosphates do not interfere               |
| Silica                                       | 500 mg/l  |
| Silicate                                     | 100 mg/l  |
| Sulfate                                      | 2000 mg/l   |
| Sulfide                                      | interferes at all concentrations  |
| Sulfite                                      | 100 mg/l  |
| Thiourea                                     | 10 mg/l   |
| highly buffered samples or extreme sample pH | may exceed the buffering capacity of the reagents and require sample pretreatment |







### pH value LR 5.2 – 6.8 with Tablet



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one BROMOCRESOLPURPLE PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press **TEST** key.

The result is shown in the display as pH-value.

#### Notes:

- 1. For photometric determination of pH values only use BROMOCRESOLPURPLE tablets in black printed foil pack and marked with PHOTOMETER.
- 2. pH values below 5.2 and above 6.8 can produce results inside the measuring range. A plausibility test (pH-meter) is recommended.
- 3. The accuracy of the colorimetric determination of pH-values depends on various boundary conditions (buffer capacity of the sample, salt contents etc.).
- 4. Salt error

Correction of test results (average values) for samples with salt contents of:

| Indicator        |         | Salt content |         |
|------------------|---------|--------------|---------|
| Bromcresolpurple | 1 molar | 2 molar      | 3 molar |
|                  | – 0.26  | – 0.33       | – 0.31  |

The values of Parson and Douglas (1926) are based on the use of Clark and Lubs buffers. 1 Mol NaCl = 58.4 g/l = 5.8 %

| Reagent                      | Form of reagent/Quantity | Order-No. |
|------------------------------|--------------------------|-----------|
| BROMOCRESOLPURPLE PHOTOMETER | Tablet / 100             | 4515700BT |







### pH value 6.5 – 8.4 with Tablet



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one PHENOL RED PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display as pH-value.

#### Notes:

- 1. For photometric determination of pH values only use PHENOL RED tablets in black printed foil pack and marked with PHOTOMETER.
- 2. Water samples with low values of Alkalinity-m (below 35 mg/l  $CaCO_3$ ) may give wrong pH readings.
- 3. pH values below 6.5 and above 8.4 can produce results inside the measuring range. A plausibility test (pH-meter) is recommended.
- 4. The accuracy of the colorimetric determination of pH values depends on various boundary conditions (buffer capacity of the sample, salt contents etc.).
- 5. Salt error

Correction of test results (average values) for samples with salt contents of:

| Indicator  |         | Salt content |         |
|------------|---------|--------------|---------|
| Phenol red | 1 molar | 2 molar      | 3 molar |
|            | – 0.21  | – 0.26       | – 0.29  |

The values of Parson and Douglas (1926) are based on the use of Clark and Lubs buffers. 1 Mol NaCl = 58.4 g/l = 5.8 %

| Reagent               | Form of reagent/Quantity | Order-No. |
|-----------------------|--------------------------|-----------|
| PHENOL RED PHOTOMETER | Tablet / 100             | 4511770BT |







## pH value 6.5 – 8.4 with Liquid Reagent



1. Fill a clean vial (24 mm  $\varnothing$ ) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

#### 6 drops of PHENOL RED solution

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

## Zero accepted prepare TEST press Test

8. Press **TEST** key.

The result is shown in the display as pH-value.

#### Notes:

- 1. When testing chlorinated water the residual chlorine contents can influence the colour reaction of the liquid reagent. This can be avoided (without interfering with the pH measurement) by adding a small crystal of Sodiumthiosulfate (Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> · 5 H<sub>2</sub>O) to the sample before adding the PHENOL RED solution. PHENOL RED tablets already contain Thiosulfate.
- 2. Due to differing drop sizes results can show a discrepancy in accuracy by comparison with tablets. This can be minimised by using a pipette (0.18 ml PHENOLRED solution is equivalent to 6 drops).
- 3. After use replace the bottle cap securely.
- 4. Store the reagent in a cool, dry place ideally at between 6°C and 10°C.

| Reagent             | Form of reagent/Quantity | Order-No. |
|---------------------|--------------------------|-----------|
| PHENOL RED solution | Liquid reagent / 15 ml   | 471040    |







### pH value HR 8.0 – 9.6 with Tablet



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one THYMOLBLUE PHOTOMETER tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare TEST press Test

8. Press **TEST** key.

The result is shown in the display as pH-value.

#### Notes:

- 1. For photometric determination of pH values only use THYMOLBLUE tablets in black printed foil pack and marked with PHOTOMETER.
- 2. pH values below 8.0 and above 9.6 can produce results inside the measuring range. A plausibility test (pH-meter) is recommended.
- 3. The accuracy of the colorimetric determination of pH values depends on various boundary conditions (buffer capacity of the sample, salt contents etc.).
- 4. Salt error

Correction of test results (average values) for samples with salt contents of:

| Indicator  |         | Salt content |         |
|------------|---------|--------------|---------|
| Thymolblue | 1 molar | 2 molar      | 3 molar |
|            | – 0.22  | – 0.29       | - 0.34  |

The values of Parson and Douglas (1926) are based on the use of Clark and Lubs buffers. 1 Mol NaCl = 58.4 g/l = 5.8 %

| Reagent               | Form of reagent/Quantity | Order-No. |
|-----------------------|--------------------------|-----------|
| THYMOLBLUE PHOTOMETER | Tablet / 100             | 4515710   |







## Polyacrylate with Liquid reagent

1 - 30 mg/l



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

### prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Add 1 ml (25 drops) KS255 (Polyacrylate reagent 1) to the water sample (note 1).
- 6. Close the vial tightly with the cap and swirl gently several times
- 7. Add 1 ml (25 drops) KS256 (Polyacrylate reagent 2) to the water sample (note 1).
- 8. Close the vial tightly with the cap and swirl gently several times
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 10. Press TEST key.

Wait for a reaction period of 10 minutes

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Polyacrylic Acid 2'100 sodium salt.

#### Notes:

- 1. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly.
- 2. If little or no turbidity is present at correct dose concentrations, the sample will need a pre-concentration step in order to detect this level of polyacrylate/polymer. Carry out this procedure as directed then test the pre-concentrated sample as above (see next page).
- 3. Anomalous results occur when interferences are present as part of the product blend or from sample contaminants. In these instances follow the interference removal steps detailed below and test this treated sample as above (see next page).
- 4. This test has been calibrated using polyacrylic acid 2'100 sodium salt in the range 1-30 mg/l. Other polyacrylates/polymers will give differing responses and therefore the test range will vary.

| Reagent  | Form of reagent/Quantity                         | Order-No.                           |
|--|--|-------------------------------------|
| Set:<br>KS255 (Polyacrylate Reagenz 1)<br>KS256 (Polyacrylate Reagenz 2) | Liquid reagent / 65 ml<br>Liquid reagent / 65 ml | 56R019165<br>56L025565<br>56L025665 |

#### Pre-Concentration

Pre-concentration uses exactly the same procedure as interference removal, except a greater volume of sample is used in step 1, instead of deionised/tap water. For calculation of the original sample concentration a concentration factor should be considered:

If a 50 ml sample is used the concentration factor is 20/50 = 0.4If a 100 ml sample is used the concentration factor is 20/100 = 0.2

This can be extended as required in order to concentrate the polyacrylate/polymer sufficiently for analysis.

#### **Example:**

If the reading is 20 mg/l and 50 ml are taken for pre-concentration the original concentration should be calculated as 20 \* 0.4 = 8 mg/l.

#### Note:

Samples exceeding 10,000 TDS should be diluted prior to loading onto the cartridge. Take this dilution into consideration when working out the overall concentration factor.

### **Cartridge Preparation**

- 1. Remove the plunger of the 20 ml syringe from the barrel and attach the C18 cartridge.
- 2. Add 5 ml of KS336 (Propan-2-ol) to the syringe barrel, attach the plunger and pass dropwise through the cartridge. Discard the eluent to waste.
- 3. Remove plunger and fill the syringe barrel with 20 ml of deionised/tap water. Attach the plunger and pass dropwise through the cartridge. Discard the eluent to waste. The cartridge is now ready to be used/reused.

| Reagent                              | Form of reagent/Quantity | Order-No.                 |
|--------------------------------------|--------------------------|---------------------------|
| KS336 (Propan-2-ol)<br>C18-Cartridge | Liquid reagent / 65 ml   | 56L033665<br>AS-K22811-KW |
| KS173 (2,4 Dinitrophenol)            | Liquid reagent / 65 ml   | 56L017365                 |
| KS183 (Nitric Acid)                  | Liquid reagent / 65 ml   | 56L018365                 |

#### Interference removal

- 1. Transfer exactly 20 ml of sample water to a 100 ml sample bottle and dilute to approximately 50-60 ml with deionised water or tap water.
- 2. Add drops of KS173 (2,4 Dinitrophenol) until a pale yellow colour is observed in the sample.
- 3. Add drops of KS183 (Nitric Acid) until the yellow colour **JUST** disappears.
- 4. Remove the plunger from the barrel of the 60ml plastic syringe and firmly attach the prepared C18 cartridge (see page 246) to the end of the barrel.
- 5. Transfer the 50-60 ml of sample from the bottle to the syringe barrel and attach the plunger. Depress the plunger and allow the sample to flow dropwise from the cartridge. Do not use excessive force to elute the sample quickly. **LEAVE THE C18 CARTRIDGE ATTACHED** and remove the plunger. Discard all of eluted sample to waste.
- 6. Using the 20 ml syringe, add exactly 20 ml of deionised/tap water to the 60 ml syringe barrel attached to the cartridge followed by 1 ml (25 drops) of KS255 (Polyacrylate Reagent 1). Gently swirl the syringe to mix.
- 7. Attach the plunger and depress. Collect the eluted sample in a clean vessel. Allow the sample to flow dropwise from the cartridge. Do not use excessive force to elute the sample quickly.
- 8. Add 10 ml of the eluted water sample into clean vial (24 mm Ø).
- 9. Using this vial perform the measurement of the method polyacrylate (see page 244).







# Potassium with Tablet

0.7 - 12 mg/l K



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one Potassium T tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display in mg/l Potassium.

### Notes:

1. If Potassium is present a cloudy solution will appear. Single particles are not necessarily caused by Potassium.

| Reagent     | Form of reagent/Quantity | Order-No. |
|-------------|--------------------------|-----------|
| Potassium T | Tablet / 100             | 4515670   |







# Silica/Silicon dioxide with Tablet

0.05 - 4 mg/l SiO<sub>2</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one SILICA No. 1 tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.

# Countdown 5:00 start: 🔟

7. Press [ ] key.
Wait for a **reaction period of 5 minutes**.

After the reaction period is finished proceed as follows:

- Add one SILICA PR tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- Add one SILICA No. 2 tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- Close the cap tightly and swirl several times until the tablets are dissolved.

11. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 2:00

12. Press TEST key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Silica.

### Notes:

- 1. The tablets must be added in the correct sequence.
- 2. Phosphate ions do not interfere under the given reaction conditions.
- 3. Conversion:  $mg/l Si = mg/l SiO_2 \times 0.47$



| Reagent                     | Form of reagent/Quantity                | Order-No. |
|-----------------------------|---|-----------|
| Set<br>SILICA No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517671BT |
| SILICA No. 1                | Tablet / 100                            | 4513130BT |
| SILICA No. 2                | Tablet / 100                            | 4513140BT |
| SILICA PR                   | Tablet / 100                            | 4513150BT |







# Silica LR / Silicon dioxide LR with Vario Powder Pack and Liquid Reagent

 $0.1 - 1.6 \text{ mg/l SiO}_{2}$ 



Use two clean vials (24 mm Ø) and mark one as blank for zeroing.

- 1. Fill each vial with 10 ml of the water sample.
- 2. Add **0.5 ml Vario Molybdate 3 reagent solution** into each vial.
- 3. Close the vials tightly with the caps and swirl several times to mix the contents (Note 1).

Countdown 4:00 start: 🔟







4. Press [ ] key.

Wait for a reaction period of 4 minutes (Note 2). After the reaction period is finished proceed as follows:

- 5. Add the contents of **one Vario Silica Citric Acid F10 Powder Pack** straight from the foil into each vial.
- 6. Close the vials tightly with the caps and swirl several times to mix the contents

Countdown 1:00 start: 🔟

7. Press [ ] key.

Wait for a reaction period of 1 minute (Note 3). After the reaction period is finished proceed as follows:

- 8. Place the vial (the blank) in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 9. Add the contents of one Vario LR Silica Amino Acid F F10 Powder Pack straight from the foil into the vial (the sample).
- 10. Close the vial tightly with the cap and swirl several times to mix the contents.

| prepare Zero |  |
|--------------|--|
| press ZERO   |  |

# Countdown 2:00

11. Press **ZERO** key (blank is already placed in the sample chamber – see point 8).

Wait for a reaction period of 2 minutes.

After the reaction period is finished the zero-reading starts automatically.

- 12. Remove the vial from the sample chamber.
- 13. Place the vial (the sample) in the sample chamber making sure that the  $\sqrt{\phantom{a}}$  marks are aligned.

Zero accepted prepare Test press TEST

14. Press **TEST** key.

The result is shown in the display in mg/l Silica.

#### Notes:

- 1. Close the vials with the cap immediately after adding the Vario Molybdate 3 reagent solution, otherwise low readings may result.
- 2. The given reaction time of 4 minutes refers to a water sample temperature of 20°C. At 30°C a reaction time of 2 minutes, at 10°C a reaction time of 8 minutes are required.
- 3. The given reaction time of 1 minute refers to a water sample temperature of 20°C. At 30°C a reaction time of 30 seconds, at 10°C a reaction time of 2 minutes are required.
- 4 Interferences:

| Substance | Interference   |
|-----------|--|
| Iron      | large amounts interfere  |
| Phosphate | does not interfere at concentrations less than 50 mg/l $PO_4$ at 60 mg/l $PO_4$ the interference is approx. – 2% at 75 mg/l $PO_4$ the interference is approx. – 11% |
| Sulfide   | interferes at all levels   |

Occasionally water samples contain forms of silica which reacts very slowly with Molybdate. The nature of these forms is not known.

A pre-treatment with Sodium hydrogencarbonate and then with Sulfuric Acid will make these forms reactive to Molybdate (pre-treatment is given in "Standard Methods for the Examination of Water and Wastewater" under "Silica Digestion with Sodium Bicarbonate").



| Reagent                        | Form of reagent/Quantity  | Order-No. |
|--------------------------------|---------------------------|-----------|
| Set:                           |                           | 4535690   |
| VARIO LR Silica Amino Acid F10 | Powder Pack / 100         |           |
| VARIO Silica Citric Acid F10   | Powder Pack / 200         |           |
| VARIO Molybdate 3              | Liquid reagent / 2x 50 ml |           |







# Silica HR / Silicon dioxide HR with Vario Powder Pack

1 - 90 mg/l SiO<sub>3</sub>



Ø 24 mm

# prepare Zero press ZERO



# Countdown 10:00 start: 🚽

1. Fill a clean vial (24 mm Ø) with **10 ml of the water** sample (Note 1), close tightly with the cap.

- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add the contents of one Vario Silica HR Molybdate F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add the contents of one Vario Silica HR Acid Rgt. F10
   Powder Pack straight from the foil to the same water sample (Note 2).
- 8. Close the vial tightly with the cap and swirl several times to mix the contents.
- Press [4] key.
   Wait for a reaction period of 10 minutes.

After the reaction period is finished proceed as follows:

- Add the contents of one Vario Silica Citric Acid F10
   Powder Pack straight from the foil to the water sample (Note 3).
- 11. Close the vial tightly with the cap and swirl several times to mix the contents.
- 12. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 13. Press **TEST** key.

Wait for a reaction period of 2 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Silica.

# Zero accepted prepare Test press TEST

# Countdown 2:00

#### Notes:

- 1. Temperature of the sample should be  $15^{\circ}\text{C} 25^{\circ}\text{C}$ .
- 2. If Silica or Phosphate is present a yellow colour is developed
- 3. In this step any yellow colour due to Phosphate is removed.
- 4. Interferences:

| Substance | Interference  |
|-----------|---|
| Iron      | large amounts interfere   |
| Phosphate | does not interfere at concentrations less than 50 mg/l $PO_4$ at 60 mg/l $PO_4$ the interference is approx. – 2% at 75 mg/l $PO_4$ the interference is approx. – 11 % |
| Sulfide   | interferes at all levels  |

Occasionally water samples contain forms of silica which reacts very slowly with Molybdate. The nature of these forms is not known.

A pre-treatment with Sodium hydrogencarbonate and then with Sulfuric Acid will make these forms reactive to Molybdate (pre-treatment is given in "Standard Methods for the Examination of Water and Wastewater" under "Silica Digestion with Sodium Bicarbonate").



| Reagent                         | Form of reagent/Quantity | Order-No. |
|---------------------------------|--------------------------|-----------|
| Set:                            |                          | 4535700   |
| VARIO Silica HR Molybdate F10   | Powder Pack / 100        |           |
| VARIO Silica HR Acid Rgt F10    | Powder Pack / 100        |           |
| VARIO Silica HR Citric Acid F10 | Powder Pack / 100        |           |







# Silica / Silicon dioxide with Liquid reagent and powder

 $0.1 - 8 \text{ mg/l SiO}_{2}$ 



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

20 drops KS104 (Silica Reagent 1)

Close the vial tightly with the cap and swirl several times to mix the contents.

# Countdown 5:00

- 7. Wait for a reaction period of 5 minutes.
- 8. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

20 drops KS105 (Silica Reagent 2)

- Close the vial tightly with the cap and swirl several times to mix the contents
- Add 1 level spoon of reagent KP106 (Silica Reagent 3) (note 1).
- 11. Close the vial tightly with the cap and swirl several times to dissolve the powder.

12. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted press ZERO press TEST

Countdown 10:00 13. Press **TEST** key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Silica.

#### Notes:

- 1. For correct dosage the spoon supplied with the reagents must be used.
- 2. For accurate results, ensure that the water being tested is between 20 °C and 30 °C.
- 3. At temperatures under 20°C the reaction does not proceed to completion and low results are obtained.



| Reagent                  | Form of reagent/Quantity | Order-No. |
|--------------------------|--------------------------|-----------|
| KS104 – Silica Reagent 1 | Liquid reagent / 65 ml   | 56L010465 |
| KS105 – Silica Reagent 2 | Liquid reagent / 65 ml   | 56L010565 |
| KP106 – Silica Reagent 3 | Powder / 10 g            | 56P010610 |

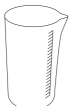






# Sodium hypochlorite (Soda bleaching lye) with Tablet

0.2 - 16 % w/w NaOCI



# Preparation:

- Fill a 5 ml plastic syringe with the test solution, ensuring that all air bubbles are expelled. Transfer the 5 ml test solution slowly into a 100 ml beaker and dilute to the 100 ml mark with chlorine-free water. Mix thoroughly.
- Fill a 5 ml plastic syringe with the diluted test solution (step 1) to the 1 ml mark, ensuring that all air bubbles are expelled. Transfer the 1 ml test solution slowly into a 100 ml beaker and dilute to the 100 ml mark with chlorine-free water. Mix thoroughly.

# Performing test procedure:



- Fill a clean vial (24 mm Ø) with 10 ml of the prepared water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one CHLORINE HR (KI) tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Add one ACIDIFYING GP tablet straight from the foil to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.

8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

9. Press TEST key.

The result is shown in the display in % w/w as available chlorine present in the original sample of Sodium hypochlorite.

#### Notes:

- 1. Please pay attention when handling sodium hypochlorite. The material has a very strong alkalinity and can cause corrosion. Contact with eyes, skin and clothes etc.has to be avoided. Refer to the detailed information the producer supplied with the product.
- 2. The tablets must be added in the correct sequence.
- 3. This method provides a fast and simple test. The test can be performed on site but the result will not be as precise as a laboratory method.
- 4. By strictly following the test procedure, an accuracy of  $\pm$ 1 weight % can be achieved.

| Reagent                                 | Form of reagent/Quantity                | Order-No. |
|---|---|-----------|
| Set: ACIDIFYING GP/<br>CHLORINE HR (KI) | Tablet / per 100 inclusive stirring rod | 4517721BT |
| CHLORINE HR (KI)                        | Tablet / 100                            | 4513000BT |
| ACIDIFYING GP                           | Tablet / 100                            | 4515480BT |







# Sulfate with Tablet

5 - 100 mg/l SO<sub>4</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- Add one SULFATE T tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

8. Press **TEST** key.

The result is shown in the display in mg/l Sulfate.

### Notes:

1. If Sulfate is present a cloudy solution will appear.

| Reagent   | Form of reagent/Quantity | Order-No. |
|-----------|--------------------------|-----------|
| SULFATE T | Tablet / 100             | 4515450BT |







# Sulfate with Vario Powder Pack

5 - 100 mg/l SO<sub>4</sub>



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO





- 4. Remove the vial from the sample chamber.
- Add the contents of one VARIO Sulpha 4/ F10 Powder Pack straight from the foil to the water sample.
- 6. Close the vial tightly with the cap and swirl several times to mix the contents
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

Wait for a reaction period of 5 minutes.

8. Press **TEST** key.

# Countdown 5:00

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Sulfate.

### Note:

1. If Sulfate ions are present a cloudy solution will appear.

| Reagent              | Form of reagent/Quantity | Order-No. |
|----------------------|--------------------------|-----------|
| VARIO Sulpha 4 / F10 | Powder Pack / 100        | 4532160   |







# Sulfide with Tablet

0.04 - 0.5 mg/l S



 Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.

2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one SULFIDE No. 1 tablet to the water sample and crush the tablet using a clean stirring rod and dissolve the tablet.
- 6. Add **one SULFIDE No. 2 tablet** to the same water sample and crush the tablet using a clean stirring rod.
- 7. Close the vial tightly with the cap and swirl several times until the tablets are dissolved
- 8. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 9. Press **TEST** key.

Wait for a reaction period of 10 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Sulfide.

#### Notes:

- 1. The tablets must be added in the correct sequence.
- 2. Chlorine and other oxidizing agents which react with DPD do not interfere with the test.
- 3. To avoid loss of Sulfide collect the sample carefully with a minimum of aeration. It is essential to test the sample immediately after collection.
- 4. The sample temperature should be 20°C. A different temperature can lead to higher or lower results.
- 5. Conversion:

 $H_2S = mg/l S \times 1.06$ 



| Reagent       | Form of reagent/Quantity | Order-No. |
|---------------|--------------------------|-----------|
| SULFIDE No. 1 | Tablet / Bottle / 100    | 502930    |
| SULFIDE No. 2 | Tablet / Bottle / 100    | 502940    |







# Sulfite with Tablet

 $0.1 - 5 \text{ mg/l SO}_{\circ}$ 



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add one SULFITE LR tablet straight from the foil to the water sample and crush the tablet using a clean stirring rod.
- 6. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 5:00

8. Press **TEST** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Sulfite.

# Notes:



| Reagent    | Form of reagent/Quantity | Order-No. |
|------------|--------------------------|-----------|
| SULFITE LR | Tablet / 100             | 4518020BT |







# **Suspended Solids**

0 - 750 mg/l TSS

# A to m

Ø 24 mm

### Sample preparation:

Blend approx. 500 ml of the water sample in a blender at high speed for 2 minutes.

- Fill a clean vial (24 mm Ø) with 10 ml of deionised water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

3. Press **ZERO** key.

- 4. Remove the vial from the sample chamber and empty the vial completely.
- Stir the blended water sample. Immediately rinse the vial with the water sample and fill with 10 ml water sample.
- 6. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

7. Press **TEST** key.

The result is shown in the display in mg/l TSS (Total Suspended Solids).

#### Note:

- 1. The photometric determination of Suspended Solids is based on a gravimetric method. In a lab this is usually done by evaporation of the filter residue of a filtrated water sample in an oven at 103°C 105°C and weighing of the dried residue.
- 2. When higher accuracy is required perform a gravimetric determination of a water sample. The result can be used to calibrate the photometer with the same water sample.
- 3. The estimated detection limit is 20 mg/L TSS.
- 4. Collect water samples in clean plastic or glass bottles and analyse the water sample as soon as possible. It is possible to store the sample at 4°C for 7 days. Before measurement warm up the sample to the temperature at collection time.
- 5. Interferences:
  - Air bubbles interfere and can be removed by swirling the vial gently.
  - Colour interferes if light is absorbed at 660 nm.







# **Turbidity**

0 - 1000 FAU



- 1. Fill a clean vial (24 mm Ø) with **10 ml of deionised** water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber and empty the vial completely.
- Stir the water sample. Immediately rinse the vial with the water sample and fill with 10 ml water sample.
- 6. Close the vial tightly with the cap and swirl gently several times.
- 7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

8. Press TEST key.

The result is shown in the display in FAU.

#### Note:

- This test uses an attenuated radiation method for the reading of FAU (Formazin Attenuation Units). The results can not be used for USEPA reporting purposes, but may be used for routine measurements. The attenuated radiation method is different from the Nephelometric method.
- 2. The estimated detection limit is 20 FAU.
- 3. Collect water samples in clean plastic or glass bottles and analyse the water sample as soon as possible. It is possible to store the sample at 4°C for 48 hours. Before measurement warm up the sample to the temperature at collection time. Temperature differences between measurement and sample collection can effect the turbidity of the sample.
- 4. Colour interferes if light is absorbed at 530 nm. For strong coloured water samples a filtrated portion of the sample can be used for zeroing instead of the deionised water.
- 5. Air bubbles interfere and can be removed using an ultrasonic bath.



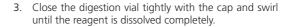




# Triazole Benzotriazole / Tolyltriazole with Vario Powder Pack

 $1 - 16 \, \text{mg/l} / 1.1 - 17.8$ 

- 1. Transfer **25 ml of the water sample** into the digestion vial
- 2. Add the contents of one Vario Triazole Rgt Powder Pack F25 straight from the foil into the water sample (note 1).



- 4. Insert the UV lamp into the digestion vial (notes 1, 2, 3). **CAUTION: Wear UV safety goggles!**
- 5. Switch the UV lamp on



Countdown 1 5:00 start: 🔟

6. Press [4] key.

Wait for a reaction period of 5 minutes (notes 10, 11). After the reaction period is finished proceed as follows:

7. Switch the UV lamp off and remove the lamp from the vial.



- 9. Fill a clean vial (24 mm Ø) with 10 ml of the deionised water, close tightly with the cap.
- 10. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.



# prepare Zero press ZERO

- 11. Press ZERO key.
- 12. Remove the vial from the sample chamber and empty the vial.
- 13. Add the digested water sample to the 10 ml mark.
- 14. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# Zero accepted prepare Test press TEST

15. Press TEST key.

The result is shown in the display in mg/L Benzotriazole or Tolyltriazole (note 4).

#### Notes:

- 1. UV lamp and Triazole Powder Pack available on request.
- 2. While the UV lamp is on UV safety goggles must be worn.
- 3. For handling of the UV lamp see manufacturer's manual.

  Do not touch the surface of the UV lamp. Fingerprints will etch the glass.

  Wipe the UV lamp with a soft and clean tissue between measurements.
- 4. The test will not distinguish between benzotriazole and tolyltriazole.
- 5. The analysis should take place immediately after taking the sample.
- 6. Strong oxidising or reducting agents in the vial lead to incorrect measurements.
- 7. To get accurate results the sample temperature must be between 20°C and 25°C.
- 8. If sample contains nitrite or borax (sodium borate), adjust the pH between 4 and 6 with 1 N sulfuric acid.
- 9. If the sample contains more than 500 mg/l  $CaCO_3$  hardness  $(CaCO_3)$ , add 10 drops of Rochelle Salt Solution.
- 10. A yellow colour will form if Triazol is present.
- 11. Low results will occur if photolysis (lamp on) takes place for more than or less than five minutes.
- 12. A Benzotriazole Tolyltriazole

| Reagent                | Form of reagent/Quantity | Order-No. |
|------------------------|--------------------------|-----------|
| VARIO TRIAZOLE Rgt F25 | Powder Pack / 100        | 4532200   |



prepare Zero

press ZERO





# Urea with Tablet and Liquid Reagent

**sample**, close tightly with the cap.

0.1 - 2.5 mg/l (NH<sub>2</sub>)<sub>2</sub>CO / mg/l Urea



- - 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

1. Fill a clean vial (24 mm Ø) with 10 ml of the water

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. In the presence of free Chlorine (HOCI), add one UREA PRETREAT tablet straight from the foil and crush the tablet using a clean stirring rod (Note 10).
- 6. Close the vial tightly with the cap and swirl several times to mix the contents
- 7. Add **2 drops of Urea reagent 1** to the water sample (Note 9).
- 8. Close the vial tightly with the cap and swirl several times to mix the contents
- 9. Add 1 drop of Urea Reagent 2 (Urease) to the same water sample (Note 9).
- 10. Close the vial tightly with the cap and swirl several times to mix the contents.

Countdown 5:00

start: 🔟

11. Press [₄] key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished proceed as follows:

- 12. Add one AMMONIA No. 1 tablet straight from the foil to the prepared water sample and mix to dissolve with a clean stirring rod.
- 13. Add one AMMONIA No. 2 tablet straight from the foil to the same water sample and mix to dissolve with a clean stirring rod.

- 14. Close the vial tightly with the cap and swirl several times until the tablets are dissolved.
- 15. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted prepare Test press TEST

Countdown 10:00 16. Press **TEST** key.

Wait for a **reaction period of 10 minutes**.

After the reaction period is finished the measurement starts automatically.

The result is shown in the display in mg/l Urea.

#### Notes:

- 1. The sample temperature should be between 20°C and 30°C.
- 2. Carry out the test at the latest one hour after sample taking.
- 3. Concentrations above 2 mg/l Urea can produce results inside the measuring range. In this case, the water sample should be diluted with Urea free water and remeasured.
- 4. The tablets must be added in the correct sequence.
- 5. The AMMONIA No. 1 tablet will only dissolve completely after the AMMONIA No. 2 tablet has been added.
- Do not store reagent 1 (Urease) below 10°C; granulation is possible.
   Store reagent 2 (Urease) in the refrigerator at a temperature of 4°C to 8°C.
- 7. Ammonia and chloramines are also measured during urea measurement.
- 8. Before analysing seawater samples, a measuring spoon of Ammonia Conditioning Powder must be added to the sample and swirled to dissolve before AMMONIA No. 1 tablet is added.
- 9. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly.
- 10. One UREA PRETREAT tablet compensates for the interference of free Chlorine up to 2 mg/l (two tablets up to 4 mg/l, three tablets up to 6 mg/l).

| Reagent                       | Form of reagent/Quantity                | Order-No. |
|-------------------------------|---|-----------|
| UREA PRETREAT                 | Tablet / 100                            | 4516110BT |
| UREA Reagent 1                | Liquid reagent / 15 ml                  | 459300    |
| UREA Reagent 2                | Liquid reagent / 10 ml                  | 459400    |
| Set:<br>AMMONIA No. 1 / No. 2 | Tablet / per 100 inclusive stirring rod | 4517611BT |
| AMMONIA No. 1                 | Tablet / 100                            | 4512580BT |
| AMMONIA No. 2                 | Tablet / 100                            | 4512590BT |







# Zinc with Tablet

0.02 - 1 mg/l Zn



- 1. Fill a clean vial (24 mm Ø) with 10 ml of the water sample.
- 2. Add one COPPER / ZINC LR tablet straight from the foil to the water sample, crush the tablet using a clean stirring rod.
- 3. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 4. Place the vial in the sample chamber making sure that the  $\chi$  marks are aligned.

### prepare Zero press ZERO

Countdown 5:00

5. Press **ZERO** key.

Wait for a reaction period of 5 minutes.

After the reaction period is finished the measurement starts automatically.

- 6. Remove the vial from the sample chamber.
- 7. Add one EDTA tablet straight from the foil to the prepared vial and crush the tablet using a clean stirring rod
- 8. Close the vial tightly with the cap and swirl several times until the tablet is dissolved.
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted press ZERO press TEST

10. Press **TEST** key.

The result is shown in the display in mg/l Zinc.

#### Notes:

- 1. The tablets must be added in the correct sequence.
- 2. In the case of high levels of residual chlorine, perform the analysis with a dechlorinated water sample. To dechlorinate add one DECHLOR tablet to the water sample (point 1). Crush and mix to dissolve the tablet. Then add the COPPER / ZINC LR tablet (point 2) and continue with the test procedure as described above.

| Reagent          | Form of reagent/Quantity | Order-No. |
|------------------|--------------------------|-----------|
| COPPER / ZINC LR | Tablette / 100           | 4512620BT |
| EDTA             | Tablette / 100           | 4512390BT |
| DECHLOR          | Tablette / 100           | 4512350BT |







# Zinc with Liquid reagent and powder

0.1 - 2.5 mg/l Zn



- Fill a clean vial (24 mm Ø) with 10 ml of the water sample, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

# prepare Zero press ZERO

- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- 5. Fill the vial with drops of the same size by holding the bottle vertically and squeeze slowly:

20 drops KS243 (Zinc Reagent 1)

- 6. Close the vial tightly with the cap and swirl several times to mix the contents.
- Add 1 level spoon of reagent KP244 (Zinc Reagent 2) (note 1).
- 8. Close the vial tightly with the cap and swirl several times to dissolve the powder.
- 9. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted press ZERO press TEST

10. Press TEST key.

The result is shown in the display in mg/l Zinc.

#### Notes:

- 1. For correct dosage the spoon supplied with the reagents must be used.
- 2. This test is suitable for determining free soluble Zinc. Zinc bound with strong complexing agents will not be measured.
- 3. Cationics such as quaternary ammonium compounds will cause the colour to change from rose red to purple, depending upon the level of copper present. In this event add drops of KS89 (cationic suppressor) one at a time, mixing between additions until the orange/blue colour is obtained.

| Reagent  | Form of reagent/Quantity                | Order-No.                           |
|--|---|-------------------------------------|
| Set:<br>KS243 (Zinc Reagent 1)<br>KP244 (Zinc Reagent 2) | Liquid reagent / 65 ml<br>Powder / 20 g | 56R023965<br>56L024365<br>56L024420 |
| KS89 (cationic suppressor)                               | Liquid reagent / 65 ml                  | 56L008965                           |

# 1.2 Important notes

# 1.2.1 Correct use of reagents

The reagents must be added in the correct sequence.

### **Tablet reagents:**

The tablet reagents should be added to the water sample straight from the foil without touching them with the fingers.

# Liquid reagents:

Add drops of the same size to the water sample by holding the bottle vertically and squeezing slowly.

After use replace the bottle caps securely noting the colour coding.

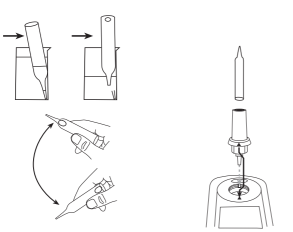
Note recommendation for storage (e.g. cool and dry).

#### **Powder Packs:**



# Vacu-vials® from CHEMetrics:

Vacu-vials® should be stored in the dark and at room temperature. For further information see MSDS.



## 1.2.2 Cleaning of vials and accessories for analysis

Vials, caps and stirring rods should be cleaned thoroughly **after each analysis** to prevent interferences.

### Procedure:

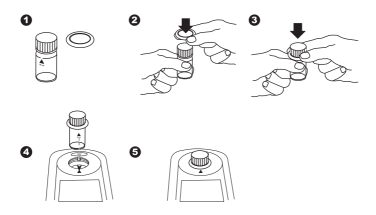
Clean vials and accessories after each analysis as soon as possible.

- a. Clean vials and accessories with laboratory detergent (e.g. Extran® MA 02 (neutral, phosphatic), Extran® MA 03 (alkaline, phosphate-free) from Merck KGaA).
- b. Rinse thoroughly with tap water.
- c. On demand (see Notes) perform special cleaning as required, e.g.: rinse with diluted Hydrochloric acid solution.
- d. Rinse thoroughly with deionised water.

## 1.2.3 Guidelines for photometric measurements

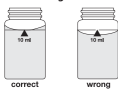
- 1. Vials, caps and stirring rods should be cleaned thoroughly after each analysis to prevent interferences. Even minor reagent residues can cause errors in the test result.
- 2. The outside of the vial must be clean and dry before starting the analysis. Clean the outside of the vials with a towel. Fingerprints or other marks will be removed.
- 3. If there is no defined vial for the blank, the zeroing and the test must be carried out with the same vial as there may be slight differences in optical performance between vials.
- 4. The vials must be positioned in the sample chamber for zeroing and test with the  $\Delta$  mark on the vial aligned with the  $\nabla$  mark on the instrument.

#### Correct position of the vial (Ø 24 mm):

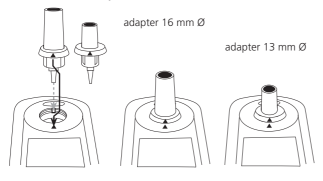


- 5. Always perform zeroing and test with closed vial cap. Only use cap with sealing ring.
- 6. Bubbles on the inside wall of the vial lead to incorrect measurements. To prevent this, remove the bubbles by swirling the vial before performing the test.
- 7. Avoid spillage of water in the sample chamber. If water should leak into the instrument housing, it can destroy electronic components and cause corrosion.
- 8. Contamination of the lens in the sample chamber can result in errors. Check at regular intervals and if necessary clean the light entry surfaces of the sample chamber using a moist cloth or cotton buds.
- 9. Large temperature differences between the instrument and the environment can lead to errors e.g. due to the formation of condensation in the area of the lens or on the vial.
- 10. To avoid errors caused by stray light do not use the instrument in bright sunlight.

### Correct filling of the vial:



### Insertion of the adapter:



## 1.2.4 Sample dilution techniques

Proceed as follows for accurate dilutions:

Pipette the water sample (see table) into a 100 ml volumetric flask and fill up to 100 ml mark with deionised water. Swirl to mix the contents.

| Water sample<br>[ml] | Multiplication<br>factor |
|----------------------|--------------------------|
| 1                    | 100                      |
| 2                    | 50                       |
| 5                    | 20                       |
| 10                   | 10                       |
| 25                   | 4                        |
| 50                   | 2                        |

Pipette the required volume of the diluted sample into the vial and proceed as described in the test methods.

### Caution:

- 1. Dilution decreases accuracy.
- 2. Do not dilute water samples for measurement of pH-values. This will lead to incorrect test results. If "Overrange" is displayed use another instrument (e.g. pH-meter).

## 1.2.5 Correcting for volume additions

If a larger volume of acid or base is used to pre-adjust the pH-value, a volume correction of the displayed result is necessary.

## **Example:**

For adjusting the pH-value of a 100 ml water sample 5 ml of acid had to be added. The corresponding displayed result is 10 mg/l.

Total volume = 100 ml + 5 ml = 105 ml

Correction factor = 105 ml / 100 ml = 1.05

Corrected result =  $10 \text{ mg/l} \times 1.05 = 10.5 \text{ mg/l}$ 

# Part 2

# **Instrument Manual**

## 2.1 Operation

### 2.1.1 Set up

Before working with the photometer insert the batteries (delivery contents). See chapter 2.1.2 Saving data – Important Notes, 2.1.3 Replacement of batteries.

### Before using the photometer perform the following settings in the Mode-Menu:

- MODE 10: select language
- MODE 12: set date and time
- MODE 34: perform "Delete data"
- MODE 69: perform "User m. init" to initialise the userpolynomial system

See chapter 2.4 Photometer settings.

## 2.1.2 Saving data - Important Notes

The batteries save data (stored results and photometer setting).

During battery change the data in the MD 600 is saved for 2 minutes. If the change time exceeds 2 minutes all stored data and settings are lost.

Recommendation: for replacement a screwdriver and new batteries must be available.

## 2.1.3 Replacement of batteries

See chapter 2.1.2 "Saving data - important notes" before replacing batteries.

- 1 Switch the instrument off
- 2. If necessary remove vial from the sample chamber.
- 3. Place the instrument upside down on a clean and even surface.
- 4. Unscrew the four screws (A) of the battery compartment cover (B).
- 5. Lift off battery compartment cover at the notch (C).
- 6. Remove old batteries (D).
- 7 Place 4 new batteries

### Ensuring the correct polarity!

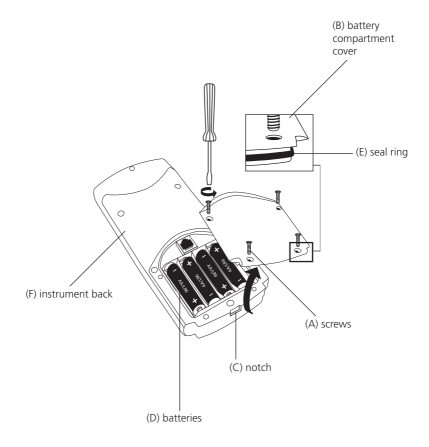
- 8. Replace the battery compartment cover. Check the seal ring (E) of the notch to make sure if is tight-fitting
- 9. Tighten the screws carefully.

### CAUTION

Dispose of used batteries in accordance with all federal, state and local regulations.

## 2.1.4 Instrument (explosion drawing):

- (A) screws
- (B) battery compartment cover
- (C) notch
- (D) batteries: 4 batteries (AA/LR6)
- (E) seal ring
- (F) instrument back



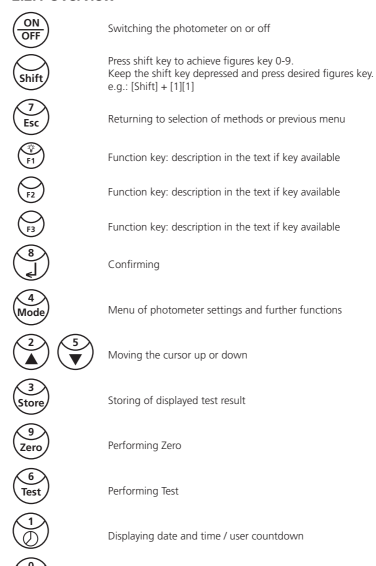
### **CAUTION:**

To ensure that the instrument is water proof:

- seal ring (E) must be in position
- battery compartment cover (B) must be fixed with the four screws

## 2.2 Overview of function keys

### 2.2.1 Overview



AL400\_8f 02/2016 289

Decimal point

## 2.2.2 Displaying time and date:



Press ["clock"] key.

19:30:22 2012-06-15

The display shows:





After 15 seconds the photometer reverts to the previous display automatically

or press [ع] key or [ESC].

### 2.2.3 User countdown

With this function the operator is able to define his own countdown.



Press ["clock"] key.

19.30.20 2012-06-15

The display shows time and date:



Press ["clock"] key.

Countdown

The display shows:

mm : ss 99 : 99

Either press  $[\[\[\]]$  key to accept the last used user countdown.

or



press any number key to start entering a new value



The entry comprises two digits each.

Enter minutes and seconds, e.g.: 2 minutes, 0 seconds = [Shift] + [0][2][0][0]. Confirm with [] key.

Countdown 02:00 start: \_ The display shows:

Start countdown with [4] key.

After countdown has finished the photometer reverts to the previous display automatically.

## 2.2.4 Display backlight





Press the [Shift] + [F1] key to turn the display backlight on or off. The backlight is switched off automatically during the measurement.

## 2.3 Operation mode



Switch the photometer on by pressing the [ON/OFF] key.

selftest ...

The photometer performs an electronic self-test.

### 2.3.1 Automatic switch off

The instrument switches off automatically after 20 minutes. This is indicated 30 seconds before by a beeper. Press any key to avoid the instrument switching off.

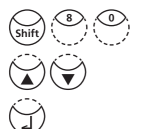
As long as the instrument is working (for example countdown or printing) the automatic switch off is inactive.

## 2.3.2 Selecting a method



The display shows a selection:

There are two possibilities to select the required method:



a) enter method-number directly e.g.: [Shift] + [8] [0] to select Bromine

b) press arrow key [▼] or [▲] to select the required method from the displayed list.

Confirm with [4] key.

## 2.3.2.1 Method Information (F1)

Use [F1] key to switch between the compact and the detailed list for method selection.

### Example:

100 Chlorine 0.02-6 mg/l Cl, **Tablet** 

24 mm DPD No 1

DPD No 3

Line 1. Method number, Method name

Line 2: Range

Line 3: Kind of reagent

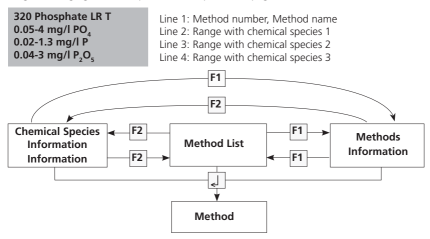
Line 4: Vial

Line 5-7: Used reagent

tube = reagent vial contained in tube test

### 2.3.2.2 Chemical Species Information

Pressing the [F2] key the display shows a list with available chemical species and corresponding ranges. Changing chemical species see chapter 2.3.7 page 294.



### 2.3.3 Differentiation



Differentiation is possible in some methods (e.g. Chlorine). The photometer then requires the type of determination.



Press arrow key [lacktrianglet] or [lacktrianglet] to select the required determination.



Confirm with [4] key.

## 2.3.4 Performing Zero

The display shows:

Prepare a clean vial as described in "Method" and place the vial in the sample chamber making sure that the ∑ marks are aligned.

Press [ZERO] key.

Zero accepted prepare Test press TEST

The display shows:

## 2.3.5 Performing Tests

When zero calibration is complete, remove the vial from the sample chamber and perform the tests as described under "Method".

When the results have been displayed:

- with some methods you can change between different chemical species
- you can store and/or print out the results
- perform further analysis with the same zero
- select a new method

## 2.3.6 Ensuring reaction periods (countdown)

To ensure compliance with reaction periods a time delay is incorporated: the countdown. There are two kinds of countdowns:



• Press [ ] key.

Prepare water sample, start countdown with [ ] key and

proceed as described in the mode description.

The vial must not be placed in the sample chamber.





Countdown 1:59 • Press [TEST] key.

Prepare the water sample as described in the method description and place the vial in the sample chamber. The display shows the countdown by pressing the [TEST] key and the countdown is started automatically. After the reaction period is finished the measurement starts automatically.

#### Notes:

1. It is possible to finish the working countdown by pressing the [4] key. Reading starts immediately. In this case the operator is responsible for ensuring the necessary reaction period.

Non-compliance with reaction periods leads to incorrect test results.

2. The time remaining is displayed continuously. The beeper indicates the last 10 seconds.

## 2.3.7 Changing chemical species

For some methods there is a possibility to change the chemical species of the test result. If the test result is displayed press arrow key  $[\blacktriangle]$  or  $[\blacktriangledown]$ .

### **Example:**

If the species of a test result is changed the displayed range is adjusted automatically. For an already stored result it is not possible to change the chemical species. The last displayed chemical species is kept by the instrument and will be displayed if this method is used the next time. If there is the possibility to change the chemical species for a method it is described in the manual. The arrows indicate the possible chemical species and are printed below the notes of the method:

- ▲ PO<sub>4</sub>
- ▼ P<sub>2</sub>O<sub>5</sub>

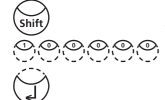
## 2.3.8 Storing results



Press [STORE] key while the test result is displayed.

Code-No.:

The display shows:



 We advise you to enter a numeric code (up to 6 places).
 (A Code No. can contain references to the operator or the sampling location.)

After entering confirm with [4] key.

If a code number is not necessary confirm by pressing
 [ ] directly. (The assignment for the Code No. is then 0
 automatically.)

The entire data set is stored with date, time, Code No., method and test result.

Stored!

The display shows:

The test result is then shown again.

### Note:

Storage: 900 free records left The display shows the number of free data sets.

Storage: only 29 free records left

If there are less than 30 data sets free the display shows:

Clear the memory as soon as possible (see "Deleting stored results"). If memory capacity is used up it is impossible to save additional test results.

## 2.3.9 Printing results (Infra-Red Interface Module) (optional)

If the IRIM (see chapter 2.5) is switched on and the printer is connected, it is possible to print out the test results (without saving it beforehand).



Press [F3] key.

The entire data set is printed with date, time, Code No., method and test result. Printing example:

100 Chlorine T 0.02-6 mg/l Cl<sub>2</sub> Profi-Mode: no 2009-07-01 14:53:09 Test No.: 1 Code-Nr.: 007 4.80 ma/l Cl2

The test No. is an internal number that is set automatically if a test result is stored. It appears only on the print out.

### 2.3.10 Perform additional measurements



To perform additional tests using the same method:

Zero accepted prepare Test

press TEST



or

• Press [TEST] key

The display shows:

Confirm with [TEST] key

• Press [ZERO] key to perform a new zero calibration.

prepare Zero press ZERO

The display shows:

## 2.3.11 Selecting a new method



Press [ESC] key to return to method selection.



Or enter the required method number directly, e.g. [Shift] + [1][6][0] for CyA-TEST (Cyanuric acid).

Confirm with [4] key.

### 2.3.12 Measure absorbance

Range: -2600 mAbs to +2600 mAbs

| Method-No. | Title       |
|------------|-------------|
| 900        | mAbs 430 nm |
| 910        | mAbs 530 nm |
| 920        | mAbs 560 nm |
| 930        | mAbs 580 nm |
| 940        | mAbs 610 nm |
| 950        | mAbs 660 nm |

Select the desired wavelength from the method list or by entering the corresponding method number directly.

900 mAbs 430 nm -2600 mAbs - + 2600 mAbs prepare Zero press ZERO

The display shows e.g.:

Always carry out zeroing using a filled (e.g. deionised water) vial.

Zero accepted prepare Test press TEST

The display shows:

Carry out measurement of the sample.

500 mAbs

The display shows e.g.:

TIP: To ensure complete reaction times the user countdown may be helpful (chapter 2.2.3, page 290).

## **2.4 Photometer settings: Table of Mode Functions**

| MODE-Function       | No. | Description   | Page |
|---------------------|-----|---|------|
| Calibration         | 40  | Special method calibration  | 312  |
| Clear calibration   | 46  | Deleting user calibration   | 319  |
| Clock               | 12  | Setting date and time   | 299  |
| Countdown           | 13  | Switching the countdown on/off to ensure reaction times           | 300  |
| Delete data         | 34  | Deleting all stored results                                       | 311  |
| Key beep            | 11  | Switching the acoustic signal on/off to indicate key-<br>pressing | 299  |
| Langelier           | 70  | Calculation of Langelier saturation Index<br>(Water Balance)      | 332  |
| Language            | 10  | Selecting language  | 298  |
| LCD contrast        | 80  | Setting the display contrast                                      | 334  |
| LCD brightness      | 81  | Setting the display brightness                                    | 334  |
| Method list         | 60  | User method list, adaption  | 322  |
| M list all on       | 61  | User method list, switching on all methods                        | 323  |
| M list all off      | 62  | User method list, switching off all methods                       | 323  |
| OTZ                 | 55  | One Time Zero (OTZ)   | 321  |
| Print               | 20  | Printing all stored results                                       | 302  |
| Print, code no.     | 22  | Print only results of a selected Code No. range                   | 304  |
| Print, date         | 21  | Print only results of a selected time period                      | 303  |
| Print, method       | 23  | Print only results of one selected method                         | 305  |
| Printing parameters | 29  | Setting of printing options                                       | 306  |
| Profi-Mode          | 50  | Switching the detailed operator instructions on/off               | 320  |
| Signal beep         | 14  | Switching the acoustic signal on/off to indicate end of reading   | 301  |
| Storage             | 30  | Displaying all stored results                                     | 307  |
| Stor., code         | 32  | Displaying only results of a selected Code No. range              | 309  |
| Stor., date         | 31  | Displaying only results of a selected time period                 | 308  |
| Stor., method       | 33  | Displaying only results of one selected method                    | 310  |
| System info         | 91  | Information about the instrument e.g. current software version    | 335  |

| MODE-Function      | No. | Description  |     |
|--------------------|-----|--|-----|
| Temperature        | 71  | Selection of °C or °F for Langelier Mode 70                                    | 333 |
| User calibration   | 45  | Storage of user calibration  | 318 |
| User concentration | 64  | Entering the data necessary to run a user concentration method                 | 324 |
| User polynoms      | 65  | Entering the data necessary to run a user polynomial                           | 326 |
| User methods clear | 66  | Delete all data of a user polynomial or of a concentration method              | 329 |
| User methods print | 67  | Print out all data stored with mode 64 (concentration) or mode 65 (polynomial) | 330 |
| User methods init  | 69  | Initialise the user method system (polynomial and concentration)               | 331 |

The selected settings are kept by the photometer even when switched off. To change photometer settings a new setting is required.

## 2.4.1 blank because of technical requirements

## 2.4.2 Instrument basic settings 1

## Selecting a language



Press [MODE], [Shift] + [1][0] keys.



Confirm with [4] key.

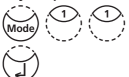
The display shows:

Press arrow key  $[ \mathbf{V} ]$  or  $[ \mathbf{A} ]$  to select the required language from the displayed list.



Confirm with [4] key.

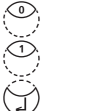




Press [MODE], [Shift] + [1][1] keys.

Confirm with [4] key.

<Key-Beep> ON: 1 OFF: 0 The display shows:



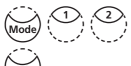
- Press [Shift] + [0] keys to switch the key beep off.
- Press [Shift] + [1] keys to switch the key beep on.

Confirm with [4] key.

#### Note:

In the case of methods with reaction periods, an acoustic signal still sounds during the last 10 seconds of the countdown even if the key beep is switched off.

### Setting date and time



Press [MODE], [Shift] + [1][2] keys.



Confirm with  $[\begin{cases} \begin{cases} \begin{cases}$ 



The display shows:

The entry comprises two digits each.

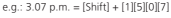
| yy-mm-dd | hh:mm |
|----------|-------|
| 09-05-14 | :     |

Enter year, month and day,

e.g.: 14. May 2009 = [Shift] + [0][9][0][5][1][4]

| yy-mm-dd | hh:mm |
|----------|-------|
| 09-05-14 | 15:07 |

Enter hours and minutes





Confirm with [4] key.

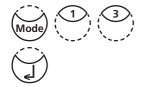
#### Note:

While confirming date and time with [4] key the seconds are adjusted to zero automatically.

### **Countdown (Ensuring reaction periods)**

Some methods require a reaction period. This reaction period is incorporated in the method as standard with the countdown function.

It is possible to switch the countdown off for all methods:



Press [MODE], [Shift] + [1][3] keys.

Confirm with  $[\ \ ]$  key.

<Countdown>
ON: 1 OFF: 0

The display shows:



- Press [Shift] + [0] keys to switch the countdown off.
- Press [Shift] + [1] keys to switch the countdown on.

Confirm with [4] key.

#### Notes:

1. It is possible to interrupt the working countdown by pressing the [4] key (application e.g. serial analysis).

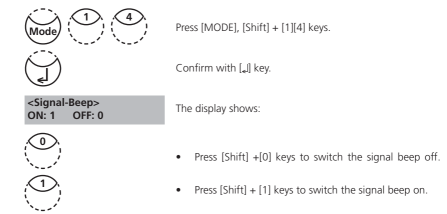
The "user countdown" is also available if the countdown is switched off.

2. If the countdown function is switched off, the operator is responsible for ensuring the necessary reaction period.

Non-compliance with reaction periods leads to incorrect test results.

## Signal beep

Performing a zero or a measurement takes 8 seconds. The photometer indicates the end of zeroing or measuring by a short beep.



#### Note:

In the case of methods with reaction periods, an acoustic signal still sounds during the last 10 seconds of the countdown even if the key beep is switched off.

Confirm with [4] key.

#### Printing of stored results 2.4.3

## **Printing all results**



Press [MODE], [Shift] + [2][0] keys.

Confirm with [4] key.



The display shows:

Press [ ] key for printing out all stored test results.



Test No.: The display shows e.g.:

> After printing the photometer goes back to <Mode-Menu> automatically.

### Note:

1. It is possible to cancel the entry by [ESC].

ESC

- 2. All stored data are printed out.
- 3. See chapter 2.5.1 Data Printing.

## Printing results of a selected time period



Press [MODE], [Shift] + [2][1] keys.



Confirm with [4] key.

<Print> sorted: date from yy-mm-dd The display shows:

Enter year, month and day for the first day of the required period, e.g.: 14 May 2009 = [Shift] + [0][9][0][5][1][4]



Confirm with [4] key.

to yy-mm-dd

The display shows:

Enter year, month and day for the last day of the required period, e.g.: 19 May 2009 = [Shift] + [0][9][0][5][1][9]



Confirm with [4] key.

The display shows:

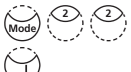
Press  $[\cline{L}]$  key and all stored results in the selected date range are printed.

After printing the photometer goes back to mode menu automatically.

#### Note

- 1. It is possible to cancel the entry by [ESC].
- 2. If you want to print only results of one day enter the same date twice to determine the period.

## Printing results of a selected Code No. range



Press [MODE], [Shift] + [2][2] keys.



Confirm with [4] key.

<Print> sorted: Code-No. from \_\_\_\_\_

The display shows:

Enter numeric code number (up to 6 places) for the first required Code No., e.g.: [Shift] + [1].



Confirm with [4] key.

The display shows:

Enter numeric code number (up to 6 places) for the last required Code No., e.g.: [Shift] + [1][0].



Confirm with [ ] key.

from 000001 000010 to Start: cancel: ESC

The display shows:

Press [4] key and all stored results in the selected code number range are printed.

After printing the photometer goes back to mode menu automatically.

### 1. Note:

2. It is possible to cancel the entry by [ESC].

If you want to print only results of one code number enter the same code number twice.

3. If you want to print all results without code no. (code no. is 0) enter Zero [0] twice.

## Printing results of one selected method



Press [MODE], [Shift] + [2][3] keys.



Confirm with [4] key.

<Print>
>>20 Acid demand
30 Alkalinity-tot
40 Aluminium T

The display shows:

Select the required method from the displayed list or enter the method-number directly.



Confirm with [4] key.

In case of differentiated methods select the required kind of determination and confirm with [』] key.

<Print>
method
30 Alkalinity-tot
Start: 
cancel: ESC

The display shows:

Press [4] key and all stored results of the selected method are printed.

After printing the photometer goes back to mode menu automatically.

### Note:

1. It is possible to cancel the entry by [ESC].

### **Printing Parameter**



Press [MODE], [Shift] + [2][9] keys.

Confirm with [ ] key.

<printing parameter>
2: Baud rate

The display shows:

\_\_\_\_\_

**ESC** 

cancel:

Press [Shift] + [2] keys to select "Baud rate".

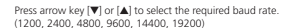
<Baud rate>

select: [▲] [▼]

save: cancel: ESC The display shows:









Confirm with [4] key.



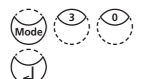
End with [ESC] key.

Back to Mode Menu with [ESC] key.

Back to method selection with [ESC] key.

### 2.4.4 Recall / delete stored results

### Recall all stored results



Confirm with [ ] key.

Press [MODE], [Shift] + [3][0] keys.

<Storage> display all data Start: ↓ cancel: ESC print: F3 print all: F2 The display shows:

The stored data sets are displayed in chronological order, starting with the latest stored test result. Press [4] key and all stored results are displayed.

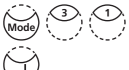
- Press [F3] key to print the displayed result.
- Press [F2] key to print all results.
- End with [ESC].
- Press arrow key [▼] to display the following test result.
- Press arrow key [▲] to display the previous test result.

If there are no test results in memory the display shows:



no data

## Recall results of a selected time period



Press [MODE], [Shift] + [3][1] keys.

Confirm with [4] key.

<Storage> sorted: date from yy-mm-dd The display shows:

Enter year, month and day for the first day of the required period, e.g.: 14 May 2009 = [Shift] + [0][9][0][5][1][4]



Confirm with [4] key.

to yy-mm-dd

The display shows:

Enter year, month and day for the last day of the required period, e.g.: 19 May 2009 = [Shift] +[0][9[0][5][1][9]



print all: F2

Confirm with [4] key.

from 2009-05-14 to 2009-05-19 Start: print: F3

The display shows:

- Press [ ] key and all stored results in the selected date range are displayed.
- Press [F3] key to print the displayed result.
- Press [F2] key to print all selected results.
- End with [ESC].

#### Note:

- 1. It is possible to cancel the entry by [ESC].
- 2. If you want to recall only results of one day enter the same date twice to determine the time period.

## Recall results of a selected Code No. range



Press [MODE], [Shift] + [3][2] keys.

Confirm with [هـ] key.

<Storage>
sorted: Code-No.
from \_ \_ \_ \_

The display shows:

Enter numeric code number (up to 6 places) for the first required Code No., e.g.: [Shift] + [1].



Confirm with [4] key.

to \_ \_ \_ \_

The display shows:

Enter numeric code number (up to 6 places) for the last required Code No., e.g.: [Shift] + [1][0].



Confirm with [ع] key.

The display shows:

- Press [
   ] key and all stored results in the selected Code No. range are displayed.
- Press [F3] key to print the displayed result.
- Press [F2] key to print all selected results.
- End with [ESC].

#### Note:

- 1. It is possible to cancel the entry by [ESC].
- 2. If you want to recall only results of one code number enter the same code number twice.
- 3. If you want to recall all results without code no. (code no. is 0) enter Zero [0] twice.

### Recall results of one selected method



Press [MODE], [Shift] + [3][3] keys.

Confirm with [4] key.

<Storage>
>>20 Acid demand
30 Alkalinity-tot
40 Aluminium T

The display shows:

Select the required method from the displayed list or enter the method number directly.



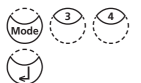
Confirm with [4] key.

In case of differentiated methods select the required kind of determination and confirm with [』] key.

<Storage> method 30 Alkalinity-tot Start: ↓ cancel: ESC print: F3 print all: F2 The display shows:

- Press [
   -|] key and all stored results of the selected method are displayed.
- Press [F3] key to print the displayed result.
- Press [F2] key to print all selected results.
- End with [ESC].

### **Delete stored results**



Press [MODE], [Shift] + [3][4] keys.

Confirm with  $[\c 4]$  key.

<Delete data>
Delete all data?
YES:1 NO:0

The display shows:



- Press [Shift] + [0] keys to retain the data sets in memory.
- After pressing keys [Shift] + [1] the following acknowledgment is displayed:

<Delete data> Delete data ↓ Do not delete: ESC

Press [ ] key to delete.

# ATTENTION: All stored test results are deleted

or cancel without deleting data by pressing [ESC] key.

### Note:

All stored test results are deleted.

### 2.4.5 Calibration

# Calcium Hardness Method 191 – Calibration of a method blank



Press [MODE], [Shift] + [4] [0] keys.

Confirm with [4] key.

<Calibration>

1: M191 Ca-hardness 2 2: M191 reset 0 cali.

3: M170 Fluoride L

The display shows:



Press [Shift] + [1] keys.

<Calibration>
M191 Ca-hardness 2T
prepare Zero
press ZERO

The display shows:



- Fill a clean vial (24 mm Ø) with exactly 10 ml of deionised water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.
- 3. Press ZERO key.
- 4. Remove the vial from the sample chamber.
- 5. Pipette 100 ml of water free of calcium to an appropriate beaker (note 2, 3).
- Add 10 CALCIO H No. 1 tablets straight from the foil to the 100 ml of water, crush the tablets using a clean stirring rod and dissolve the tablets completely.
- Add 10 CALCIO H No. 2 tablets straight from the foil to the same water, crush the tablets using a clean stirring rod and dissolve the tablets completely.
- 8. Press [ ] key.

Wait for a reaction period of 2 minutes.



Zero accepted countdown 2:00 Start:

After the reaction period is finished proceed as follows:

9. Rinse the vial (24 mm Ø) with the coloured sample from the beaker and fill with 10 ml of the sample.

### prepare Test press TEST

10. Press **TEST** kev.

#### stored

The batch related method blank is saved.

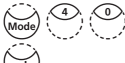


Press [ع] key, to go back to mode menu.

#### Notes:

- 1. If a new batch of CALCIO tablets is used a calibration of the method blank has to be performed to optimise the results.
- 2. Deionised or tap water
- 3. If no water free of Calcium is available these ions can be masked by using EDTA. Preparation: Add 50 mg (a spatula-tipful) EDTA to 100 ml water and dissolve.
- 4. To achieve the most accurate method blank it is important to adhere exactly to the sample volume of 100 ml.

## Calcium Hardness Method 191 -Reset method blank to factory calibration



Press [MODE], [Shift] + [4] [0] keys.



Confirm with [4] key.

<Calibration>

1: M191 Ca-hardness 2

2: M191 reset 0 cali.

3: M170 Fluoride L

The display shows:



Press [Shift] + [2] keys.

<Calibration> M191 Ca-hardness 2T

Reset?

YES: Shift + 1

NO: Shift + 0 The display shows:

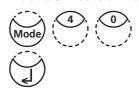


Press [Shift] + [0] keys to keep the method blank.

Press [Shift] + [1] keys to erase the method blank and set the value back to factory calibration.

The instrument goes back to mode menu automatically.

### Fluoride Method 170



Press [MODE], [Shift] + [4] [0] keys.

Confirm with [4] key.

<Calibration>

1: M191 Ca-hardness 2

2: M191 reset 0 cali.

3: M170 Fluoride L

3

Press [Shift] + [3] keys.

The display shows:

The display shows:

<Calibration>
M170 Fluoride L
ZERO: deionised water
press ZERO

- Fill a clean vial (24 mm Ø) with exactly 10 ml of deionised water, close tightly with the cap.
- 2. Place the vial in the sample chamber making sure that the marks  $\overline{\chi}$  are aligned.
- 3. Press **ZERO** key.
- 4. Remove the vial from the sample chamber.
- Add exactly 2 ml SPADNS reagent solution to the water sample. Caution: Vial is filled up to the top!
- 6. Close the vial tightly with the cap and swirl gently several times to mix the contents.

7. Place the vial in the sample chamber making sure that the  $\overline{\chi}$  marks are aligned.

Zero accepted T1: 0 mg/l F press TEST

- 8. Press TEST key.
- Remove the vial from the sample chamber, empty the vial, rinse vial and cap several times and then fill the vial with exactly 10 ml Fluoride standard (Concentration 1 mg/l F).
- 10. Add **exactly 2 ml SPADNS reagent solution** to the Fluoride standard.

Caution: Vial is filled up to the top!

 Place the vial in the sample chamber making sure that the \( \frac{1}{2} \) marks are aligned.

T1 accepted T2: 1 mg/l F press TEST

12. Press **TEST** key.



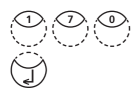
The display shows:



Confirm with [4] key.



Back to method selection with ESC key.



Select Fluoride method with keys [Shift] + [1][7][0] and [ $\downarrow$ ].



If an error message appears please repeat adjustment.

#### Notes:

- 1. The same batch of SPADNS reagent solution must be used for adjustment and test. The adjustment process needs to be performed for each new batch of SPADNS reagent solution (see Standard methods 20th, 1998, APHA, AWWA, WEF 4500 F D., S. 4-82).
- 2. As the test result is highly dependent on exact sample and reagent volumes, the sample and reagent volumes should always be metered by using a 10 ml resp. 2 ml volumetric pipette (class A).

### **User Calibration**

If a test method is user calibrated the method name is displayed inverse.

#### Procedure:

- Prepare a standard of known concentration and use this standard instead of the sample according to the test procedure.
- It is recommend to use well known standards which are formulated according to DIN EN, ASTM or other international norms or to use certified standards which are commercially available.
- After measuring this standard solution it is possible to change the displayed results to the required value.
- If a method uses a mathematic equation for the calculation of the result, it is only
  possible to calibrate the basic tests since all the other tests use the same polynomial.
- The same applies for some test procedures which use a polynomial from another test procedure.

### Return to factory calibration:

If the user calibration is deleted the factory calibration is automatically activated.

#### Remarks:

The method "Fluoride" cannot be calibrated with mode 45 since the test requires a calibration related to the batch of the liquid reagent (SPADNS) (mode 40, chapter "Fluoride Method 170").

| Table |                       |   |
|-------|-----------------------|---|
| No.   | Method                | Recommended range for user calibration        |
| 20    | Acid demand           | 1–3 mmol/l                                    |
| 35    | Alkalinity-p          | 100–300 mg/l CaCO <sub>3</sub>                |
| 30    | Alkalinity-total      | 50–150 mg/l CaCO <sub>3</sub>                 |
| 31    | Alkalinity-total HR T | 50–300 mg/l CaCO <sub>3</sub>                 |
| 40    | Aluminium T           | 0.1–0.2 mg/l Al                               |
| 50    | Aluminium PP          | 0.1-0.2 mg/l Al                               |
| 60    | Ammonia T             | 0.3-0.5 mg/l N                                |
| 62    | Ammonia PP            | 0.3-0.5 mg/l N                                |
| 65    | Ammonia LR TT         | 1 mg/l N                                      |
| 66    | Ammonia HR TT         | 20 mg/l N                                     |
| 85    | Boron                 | 1 mg/l B                                      |
| 80    | Bromine T             | Calibration with basic test 100 Chlorine free |
| 81    | Bromine PP            | Calibration with basic test 110 Chlorine free |
| 90    | Chloride              | 10–20 mg/l Cl <sup>-</sup>                    |
| 92    | Chloride L            | 10–15 mg/l Cl <sup>-</sup>                    |
| 100   | Chlorine T            | 0.5-1.5 mg/l Cl                               |
| 103   | Chlorine HR T         | 0.5–6 mg/l Cl                                 |
| 101   | Chlorine L            | Calibration with basic test 100 Chlorine free |
| 110   | Chlorine PP           | 0.5–1 mg/l Cl <sub>2</sub>                    |
| 111   | Chlorine HR PP        | 4–5 mg/l Cl <sub>2</sub>                      |
| 105   | Chlorine (KI) HR      | 70-150 mg/l Cl                                |
| 120   | Chlorine dioxide T    | Calibration with basic test 100 Chlorine free |
| 122   | Chlorine dioxide PP   | Calibration with basic test 110 Chlorine free |

| 125         Chromium         1 mg/l Cr           130         COD LR         100 mg/l O₂           131         COD MR         500 mg/l O₂           132         COD HR         5 g/l O₂ = 5000 mg/l O₂           150         Copper T         0.5 – 1.5 mg/l Cu           151         Copper L         2 – 3 mg/l Cu           153         Copper PP         0.5 – 1.5 mg/l Cu           157         Cyanide         0.1 – 0.3 mg/l CN           160         CyA-TEST         30 – 60 mg/l CyA           165         DEHA T         200 – 400 µg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           170         Fluoride         Calibration with basic test 100 Chlorine free           181         H,O₂ T         Calibration with basic test 100 Chlorine free           191         Hardness, Calcium         100 – 200 mg/l CaCO₃           191         Hardness, total HR T         Calibration with basic test 200 Hardness, total           191         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2 – 0.4 mg/l N₂H₄           206         Hydrazine E         0.2 – 0.4 mg/l N₂H₄           207         Hydrazine C   | No. | Method             | Recommended range for user calibration          |
|--|-----|--------------------|---|
| 131         COD MR         50/ mg/l O₂ = 5000 mg/lO₂           204         Colour         operating range           150         Copper T         0.5 – 1.5 mg/l Cu           151         Copper L         2 – 3 mg/l Cu           153         Copper PP         0.5 – 1.5 mg/l Cu           157         Cyanide         0.1 – 0.3 mg/l CN           160         CyA-TEST         30 – 60 mg/l CyA           165         DEHA T         200 – 400 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test 100 Chlorine free           210         H₂O₂ LR L         20-30 mg/l H₂O₂           214         H₂O₂ LR L         200-300 mg/l H₂O₂           219         Hardness, Calcium         100-200 mg/l CaCO₃           191         Hardness, total T         15-25 mg/l CaCO₃           201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           202         Hydrazine P         0.2-0.4 mg/l N₂H₂  | 125 | Chromium           | 1 mg/l Cr                                       |
| 132         COD HR         5 g/l O₂ = 5000 mg/lO₂           204         Colour         operating range           150         Copper T         0.5-1.5 mg/l Cu           151         Copper L         2-3 mg/l Cu           153         Copper PP         0.5-1.5 mg/l Cu           157         Cyanide         0.1-0.3 mg/l CN           160         CyA-TEST         30-60 mg/l CyA           165         DEHA T         200-400 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test100 Chlorine free           213         H₂O₂ LR L         20-30 mg/l H₂O₂           214         H₂O₂ HR L         200-300 mg/l H₂O₂           219         Hardness, Calcium         100-200 mg/l CaCO₃           191         Hardness, Calcium         100-200 mg/l CaCO₃           201         Hardness, total T         15-25 mg/l CaCO₃           201         Hydrazine P         0.2-0.4 mg/l N₂H₄           205         Hydrazine P         0.2-0.4 mg/l N₂H₄           206         Hydrazine C         0.2-0.4 mg/l N₂H₄           207         Hydrazine C         0.2-0.7 mg/l Fe           215         Iron Fe  | 130 | COD LR             | 100 mg/l O <sub>2</sub>                         |
| 132         COD HR         5 g/l O₂ = 5000 mg/lO₂           204         Colour         operating range           150         Copper T         0.5-1.5 mg/l Cu           151         Copper L         2-3 mg/l Cu           153         Copper PP         0.5-1.5 mg/l Cu           157         Cyanide         0.1-0.3 mg/l CN           160         CyA-TEST         30-60 mg/l CyA           165         DEHA T         200-400 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test100 Chlorine free           213         H₂O₂ LR L         20-30 mg/l H₂O₂           214         H₂O₂ HR L         200-300 mg/l H₂O₂           219         Hardness, Calcium         100-200 mg/l CaCO₃           191         Hardness, Calcium         100-200 mg/l CaCO₃           201         Hardness, total T         15-25 mg/l CaCO₃           201         Hydrazine P         0.2-0.4 mg/l N₂H₄           205         Hydrazine P         0.2-0.4 mg/l N₂H₄           206         Hydrazine C         0.2-0.4 mg/l N₂H₄           207         Hydrazine C         0.2-0.7 mg/l Fe           215         Iron Fe  | 131 | COD MR             | 500 mg/l O <sub>2</sub>                         |
| 204         Colour         operating range           150         Copper T         0.5–1.5 mg/l Cu           151         Copper L         2–3 mg/l Cu           153         Copper PP         0.5–1.5 mg/l Cu           157         Cyanide         0.1–0.3 mg/l CN           160         CyA-TEST         30–60 mg/l CyA           165         DEHA T         200–400 μg/l DEHA           167         DEHA PP         200 μg/l DEHA           168         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test 100 Chlorine free           210         H₂O₂ LR L         20-30 mg/l H₂O₂           214         H₂O₂ LR L         20-30 mg/l H₂O₂           219         Hardness, Calcium         100–200 mg/l CaCO₃           219         Hardness, Calcium         100–200 mg/l CaCO₃           210         Hardness, total T         15–25 mg/l CaCO₃           210         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2–0.4 mg/l N₂H₄           206         Hydrazine C         0.2–0.4 mg/l N₂H₄           207<   | 132 | COD HR             |   |
| 151         Copper L         2–3 mg/l Cu           153         Copper PP         0.5–1.5 mg/l Cu           157         Cyanide         0.1–0.3 mg/l CN           160         CyA-TEST         30–60 mg/l CyA           165         DEHA T         200–400 μg/l DEHA           167         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test100 Chlorine free           213         H₂O₂ LR L         20–300 mg/l H₂O₂           214         H₂O₂ HR L         200–300 mg/l H₂O₂           214         H₂O₂ HR L         200–300 mg/l CaCO₃           215         Hardness, Calcium         100–200 mg/l CaCO₃           216         Hardness, total T         15–25 mg/l CaCO₃           217         Hydrazine P         0.2–0.4 mg/l N₂H₄           208         Hydrazine P         0.2–0.4 mg/l N₂H₄           216         Hydrazine C         0.2–0.4 mg/l N₂H₄           217         Hydrazine C         0.2–0.4 mg/l N₂H₄           218         Iron FP         0.1–2 mg/l Fe           219         Iron (Ferin Mo) PP         0.5–1.5 mg/l Fe           220         Iron HR L </td <td>204</td> <td>Colour</td> <td></td>  | 204 | Colour             |   |
| 153         Copper PP         0.5-1.5 mg/l Cu           157         Cyanide         0.1-0.3 mg/l CN           160         CyA-TEST         30-60 mg/l CyA           165         DEHA T         200-400 μg/l DEHA           167         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test100 Chlorine free           213         H₂O₂ LR L         20-30 mg/l H₂O₂           214         H₂O₂ HR L         20-300 mg/l H₂O₂           214         H₂O₂ LR L         200-300 mg/l H₂O₂           214         H₂O₂ LR L         200-300 mg/l H₂O₂           215         Hardness, Calcium         100-200 mg/l CaCO₃           190         Hardness, Calcium         100-200 mg/l CaCO₃           219         Hardness, Stotal T         15-25 mg/l CaCO₃           200         Hardness, total HR T         Calibration with basic test 200 Hardness, total           201         Hydrazine P         0.2-0.4 mg/l N₂H₄           202         Hydrazine C         0.2-0.4 mg/l N₂H₄           215         Iodine         Calibration with basic test 100 Chlorine free           220         Iron T         0.3-0.7 mg/l F   | 150 | Copper T           | 0.5-1.5 mg/l Cu                                 |
| 157 Cyanide 0.1–0.3 mg/l CN 160 CyA-TEST 30–60 mg/l CyA 165 DEHA T 200–400 μg/l DEHA 167 DEHA PP 200 μg/l DEHA 170 Fluoride Calibration with 0 and 1 mg/l F through Mode 40 170 H <sub>2</sub> O <sub>2</sub> T Calibration with basic test100 Chlorine free 171 H <sub>2</sub> O <sub>2</sub> LR L 20–30 mg/l H <sub>2</sub> O <sub>2</sub> 172 H <sub>2</sub> O <sub>3</sub> HR L 200–300 mg/l H <sub>2</sub> O <sub>2</sub> 173 H <sub>2</sub> O <sub>2</sub> HR L 200–300 mg/l CaCO <sub>3</sub> 174 H <sub>2</sub> O <sub>2</sub> HR L 200–300 mg/l CaCO <sub>3</sub> 175 Hardness, Calcium 100–200 mg/l CaCO <sub>3</sub> 176 Hardness, total T 15–25 mg/l CaCO <sub>3</sub> 177 Hydrazine P 0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 178 Hydrazine L 0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 179 Hydrazine C 0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 179 Hydrazine C 0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 170 Hydrazine C 0.3–0.7 mg/l Fe 170 Iron T 0.3–0.7 mg/l Fe 170 Iron (Fe in Mo) PP 0.5–1.5 mg/l Fe 170 Iron LR L 0.5–1.5 mg/l Fe 170 Iron LR L 0.5–1.5 mg/l Fe 170 Iron HR L 0.5–1.5 mg/l Fe 170 Manganese T 1–2 mg/l Mn 170 Manganese PP 0.1–0.4 mg/l Mn 170 Manganese PP 0.1–0.4 mg/l Mn 170 Manganese L 2–3 mg/l Mn 170 Manganese L 2–3 mg/l Mn 170 Molybdate LR PP 1.5–2.5 mg/l Mo 170 Molybdate HR PP 10–30 mg/l Mo 170 Molybdate HR PP 10–30 mg/l Mo 170 Mitrite T 0.2–0.3 mg/l N  | 151 | Copper L           | 2–3 mg/l Cu                                     |
| 160         CyA-TEST         30–60 mg/l CyA           165         DEHA T         200–400 μg/l DEHA           167         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H <sub>2</sub> O <sub>2</sub> T         Calibration with basic test100 Chlorine free           213         H <sub>2</sub> O <sub>2</sub> LR L         20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l CaCO <sub>3</sub> 190         Hardness, Calcium         100–200 mg/l CaCO <sub>3</sub> 201         Hardness, total T         15–25 mg/l CaCO <sub>3</sub> 201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine C         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215         Iodine         Calibration with basic test 100 Chlorine free           216         Iron T         0.3–0.7 mg/l Fe           217         Iron (Fe in Mo) PP         0.5–1.5 mg/l Fe           222         Iron (Fe in Mo) PP         0.5–1.5 mg/l Fe           225         Iron LR 2 L         0.5–1.5 mg/l Fe           226   | 153 | Copper PP          | 0.5-1.5 mg/l Cu                                 |
| 165         DEHA T         200 – 400 μg/l DEHA           167         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H₂O₂ T         Calibration with basic test 100 Chlorine free           213         H₃O₂ LR L         20-30 mg/l H₂O₂           214         H₃O₂ HR L         200-300 mg/l H₂O₂           190         Hardness, Calcium         100–200 mg/l CaCO₃           191         Hardness, Calcium         100–200 mg/l CaCO₃           200         Hardness, total T         15–25 mg/l CaCO₃           201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2–0.4 mg/l N₂H₄           206         Hydrazine C         0.2–0.4 mg/l N₃H₄           207         Hydrazine C         0.2–0.4 mg/l N₃H₄           215         Iodine         Calibration with basic test 100 Chlorine free           210         Iron T         0.3–0.7 mg/l Fe           221         Iron PP         0.1–2 mg/l Fe           222         Iron PP         0.5–1.5 mg/l Fe           223         Iron LR L         0.5–1.5 mg/l Fe           224         Iron HR L         6–8 mg/l Mn </td <td>157</td> <td>Cyanide</td> <td>0.1–0.3 mg/l CN</td>   | 157 | Cyanide            | 0.1–0.3 mg/l CN                                 |
| 167         DEHA PP         200 μg/l DEHA           170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H <sub>2</sub> O <sub>2</sub> T         Calibration with basic test100 Chlorine free           213         H <sub>2</sub> O <sub>2</sub> LR L         20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 191         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 200         Hardness, total T         15-25 mg/l CaCO <sub>3</sub> 201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215         Iodine         Calibration with basic test 100 Chlorine free           220         Iron T         0.3-0.7 mg/l Fe           221         Iron PP         0.1-2 mg/l Fe           222         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           225         Iron LR L         0.5-1.5 mg/l Fe           226         Iron HR L         6-8 mg/l Fe           227         <  | 160 | CyA-TEST           | 30–60 mg/l CyA                                  |
| 170         Fluoride         Calibration with 0 and 1 mg/l F through Mode 40           210         H <sub>2</sub> O <sub>2</sub> T         Calibration with basic test 100 Chlorine free           213         H <sub>2</sub> O <sub>2</sub> LR L         20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 201         Hardness, total T         15-25 mg/l CaCO <sub>3</sub> 201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215         Iodine         Calibration with basic test 100 Chlorine free           220         Iron T         0.3-0.7 mg/l Fe           221         Iron PP         0.1-2 mg/l Fe           222         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           225         Iron LR L         0.5-1.5 mg/l Fe           226         Iron LR 2 L         1-15 mg/l Fe           227         Iron HR L         6-8 mg/l Fe           240 <td< td=""><td>165</td><td>DEHA T</td><td>200-400 μg/l DEHA</td></td<> | 165 | DEHA T             | 200-400 μg/l DEHA                               |
| 210       H <sub>2</sub> O <sub>2</sub> T       Calibration with basic test100 Chlorine free         213       H <sub>2</sub> O <sub>2</sub> LR L       20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214       H <sub>2</sub> O <sub>2</sub> HR L       200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190       Hardness, Calcium       100-200 mg/l CaCO <sub>3</sub> 191       Hardness, Calcium       100-200 mg/l CaCO <sub>3</sub> 200       Hardness, total T       15-25 mg/l CaCO <sub>3</sub> 201       Hardness, total HR T       Calibration with basic test 200 Hardness, total         205       Hydrazine P       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206       Hydrazine L       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207       Hydrazine C       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 208       Holdine       Calibration with basic test 100 Chlorine free         215       Iodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3-0.7 mg/l Fe         221       Iron PP       0.1-2 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         224       Iron LR 2 L       1-15 mg/l Fe         225       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         241       Manganese HR   | 167 | DEHA PP            | 200 μg/l DEHA                                   |
| 213         H <sub>2</sub> O <sub>2</sub> LR L         20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 191         Hardness, total T         15-25 mg/l CaCO <sub>3</sub> 200         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 208         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 209         Iron T         0.3-0.7 mg/l Fe           210         Iron P         0.1-2 mg/l Fe           211         Iron PP         0.1-2 mg/l Fe           212         Iron PP         0.3-0.7 mg/l Fe           223         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           224         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           225         Iron LR L         1-15 mg/l Fe           226         Iron HR L         6-8 mg/l Fe           227         Iron HR L         6-8 mg/l Mn           240         Manganese PP         0.1-0.4 mg/l Mn           241   | 170 | Fluoride           | Calibration with 0 and 1 mg/l F through Mode 40 |
| 213         H <sub>2</sub> O <sub>2</sub> LR L         20-30 mg/l H <sub>2</sub> O <sub>2</sub> 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 191         Hardness, total T         15-25 mg/l CaCO <sub>3</sub> 200         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 208         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 209         Iron T         0.3-0.7 mg/l Fe           210         Iron P         0.1-2 mg/l Fe           211         Iron PP         0.1-2 mg/l Fe           212         Iron PP         0.3-0.7 mg/l Fe           223         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           224         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           225         Iron LR L         1-15 mg/l Fe           226         Iron HR L         6-8 mg/l Fe           227         Iron HR L         6-8 mg/l Mn           240         Manganese PP         0.1-0.4 mg/l Mn           241   | 210 | H,O, T             |   |
| 214         H <sub>2</sub> O <sub>2</sub> HR L         200-300 mg/l H <sub>2</sub> O <sub>2</sub> 190         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 191         Hardness, Calcium         100-200 mg/l CaCO <sub>3</sub> 200         Hardness, total T         15-25 mg/l CaCO <sub>3</sub> 201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215         Iodine         Calibration with basic test 100 Chlorine free           220         Iron T         0.3-0.7 mg/l Fe           221         Iron PP         0.1-2 mg/l Fe           222         Iron PP         0.1-2 mg/l Fe           223         Iron (Fe in Mo) PP         0.5-1.5 mg/l Fe           224         Iron LR L         0.5-1.5 mg/l Fe           225         Iron LR L         1-2 mg/l Fe           226         Iron LR 2 L         1-2 mg/l Mn           242         Manganese T         1-2 mg/l Mn           243         Manganese HR PP         4-6 mg/l Mn   |     |                    |   |
| 190         Hardness, Calcium         100–200 mg/l CaCO <sub>3</sub> 191         Hardness, Calcium         100–200 mg/l CaCO <sub>3</sub> 200         Hardness, total T         15–25 mg/l CaCO <sub>3</sub> 201         Hardness, total HR T         Calibration with basic test 200 Hardness, total           205         Hydrazine P         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206         Hydrazine L         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207         Hydrazine C         0.2–0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215         Iodine         Calibration with basic test 100 Chlorine free           220         Iron T         0.3–0.7 mg/l Fe           221         Iron PP         0.1–2 mg/l Fe           222         Iron (Fe in Mo) PP         0.5–1.5 mg/l Fe           223         Iron (Fe in Mo) PP         0.5–1.5 mg/l Fe           224         Iron LR L         0.5–1.5 mg/l Fe           225         Iron LR L         1–15 mg/l Fe           226         Iron LR L         1–2 mg/l Mn           240         Manganese T         1–2 mg/l Mn           241         Manganese HR PP         4–6 mg/l Mn           242         Manganese HR PP         4–6 mg/l Mn           243         Molybdate LR PP         1.5–2.5 mg/l Mo  | 214 | H,O, HR L          | 200-300 mg/l H <sub>2</sub> O <sub>2</sub>      |
| 191       Hardness, Calcium       100–200 mg/l CaCO₃         200       Hardness, total T       15–25 mg/l CaCO₃         201       Hardness, total HR T       Calibration with basic test 200 Hardness, total         205       Hydrazine P       0.2–0.4 mg/l N₂H₄         206       Hydrazine L       0.2–0.4 mg/l N₂H₄         207       Hydrazine C       0.2–0.4 mg/l N₂H₄         215       Iodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3–0.7 mg/l Fe         221       Iron PP       0.1–2 mg/l Fe         222       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         223       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         224       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         225       Iron LR L       0.5–1.5 mg/l Fe         226       Iron LR 2 L       1–15 mg/l Fe         227       Iron HR L       6–8 mg/l Fe         240       Manganese T       1–2 mg/l Mn         242       Manganese HR PP       4–6 mg/l Mn         243       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate HR PP       10–30 mg/l Mo         254 <t< td=""><td>190</td><td></td><td></td></t<>  | 190 |                    |   |
| 200       Hardness, total T       15–25 mg/l CaCO₃         201       Hardness, total HR T       Calibration with basic test 200 Hardness, total         205       Hydrazine P       0.2–0.4 mg/l N₂H₄         206       Hydrazine L       0.2–0.4 mg/l N₂H₄         207       Hydrazine C       0.2–0.4 mg/l N₂H₄         215       Iodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3–0.7 mg/l Fe         221       Iron PP       0.1–2 mg/l Fe         222       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         223       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         224       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         225       Iron LR L       0.5–1.5 mg/l Fe         226       Iron LR 2 L       1–15 mg/l Fe         227       Iron HR L       6–8 mg/l Fe         240       Manganese T       1–2 mg/l Mn         242       Manganese HR PP       4–6 mg/l Mn         243       Manganese L       2–3 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate H   | 191 |                    |   |
| 201       Hardness, total HR T       Calibration with basic test 200 Hardness, total         205       Hydrazine P       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 206       Hydrazine L       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 207       Hydrazine C       0.2-0.4 mg/l N <sub>2</sub> H <sub>4</sub> 215       Iodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3-0.7 mg/l Fe         221       Iron PP       0.1-2 mg/l Fe         222       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         223       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         244       Molybdate T       5-15 mg/l Mo         250       Molybdate HR PP       1.5-2.5 mg/l Mo         251       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR P       10-30 mg/l N         265  | 200 | Hardness, total T  |   |
| 205       Hydrazine P       0.2-0.4 mg/l N₂H₄         206       Hydrazine L       0.2-0.4 mg/l N₂H₄         207       Hydrazine C       0.2-0.4 mg/l N₂H₄         215       Iodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3-0.7 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (FPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         243       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate HR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR C       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         265       Nitrate LR       0.5-0.7 mg/l N   | 201 |                    |   |
| 206       Hydrazine L       0.2-0.4 mg/l N₂H₄         207       Hydrazine C       0.2-0.4 mg/l N₂H₄         215       lodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3-0.7 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (TPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         243       Manganese L       2-3 mg/l Mn         245       Manganese L       2-3 mg/l Mo         250       Molybdate T       5-15 mg/l Mo         251       Molybdate HR PP       10-30 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR C       50-70 mg/l No         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N   | 205 |                    |   |
| 207       Hydrazine C       0.2-0.4 mg/l N₂H₄         215       lodine       Calibration with basic test 100 Chlorine free         220       Iron T       0.3-0.7 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (TPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese HR PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR L       50-70 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 206 | Hydrazine L        | - 2 7   |
| 220       Iron T       0.3-0.7 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (TPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N  | 207 | Hydrazine C        |   |
| 220       Iron T       0.3-0.7 mg/l Fe         222       Iron PP       0.1-2 mg/l Fe         223       Iron (TPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N  | 215 | lodine             | Calibration with basic test 100 Chlorine free   |
| 223       Iron (TPTZ) PP       0.3-0.7 mg/l Fe         224       Iron (Fe in Mo) PP       0.5-1.5 mg/l Fe         225       Iron LR L       0.5-1.5 mg/l Fe         226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 220 | Iron T             |   |
| 224       Iron (Fe in Mo) PP       0.5–1.5 mg/l Fe         225       Iron LR L       0.5–1.5 mg/l Fe         226       Iron LR 2 L       1–15 mg/l Fe         227       Iron HR L       6–8 mg/l Fe         240       Manganese T       1–2 mg/l Mn         242       Manganese PP       0.1–0.4 mg/l Mn         243       Manganese HR PP       4–6 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N  | 222 | Iron PP            | 0.1–2 mg/l Fe                                   |
| 225       Iron LR L       0.5–1.5 mg/l Fe         226       Iron LR 2 L       1–15 mg/l Fe         227       Iron HR L       6–8 mg/l Fe         240       Manganese T       1–2 mg/l Mn         242       Manganese PP       0.1–0.4 mg/l Mn         243       Manganese HR PP       4–6 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N   | 223 | Iron (TPTZ) PP     | 0.3–0.7 mg/l Fe                                 |
| 226       Iron LR 2 L       1-15 mg/l Fe         227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 224 | Iron (Fe in Mo) PP | 0.5–1.5 mg/l Fe                                 |
| 227       Iron HR L       6-8 mg/l Fe         240       Manganese T       1-2 mg/l Mn         242       Manganese PP       0.1-0.4 mg/l Mn         243       Manganese HR PP       4-6 mg/l Mn         245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N  | 225 | Iron LR L          | 0.5–1.5 mg/l Fe                                 |
| 240       Manganese T       1–2 mg/l Mn         242       Manganese PP       0.1–0.4 mg/l Mn         243       Manganese HR PP       4–6 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N  | 226 | Iron LR 2 L        | 1–15 mg/l Fe                                    |
| 242       Manganese PP       0.1–0.4 mg/l Mn         243       Manganese HR PP       4–6 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N  | 227 | Iron HR L          | 6–8 mg/l Fe                                     |
| 243       Manganese HR PP       4–6 mg/l Mn         245       Manganese L       2–3 mg/l Mn         250       Molybdate T       5–15 mg/l Mo         251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N   | 240 | Manganese T        | 1–2 mg/l Mn                                     |
| 245       Manganese L       2-3 mg/l Mn         250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 242 | Manganese PP       | 0.1-0.4 mg/l Mn                                 |
| 250       Molybdate T       5-15 mg/l Mo         251       Molybdate LR PP       1.5-2.5 mg/l Mo         252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 243 | Manganese HR PP    | 4–6 mg/l Mn                                     |
| 251       Molybdate LR PP       1.5–2.5 mg/l Mo         252       Molybdate HR PP       10–30 mg/l Mo         254       Molybdate HR L       50–70 mg/l Mo         257       Nickel T       6–8 mg/l Ni         260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N  | 245 | Manganese L        | 2–3 mg/l Mn                                     |
| 252       Molybdate HR PP       10-30 mg/l Mo         254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N  | 250 | Molybdate T        | 5–15 mg/l Mo                                    |
| 254       Molybdate HR L       50-70 mg/l Mo         257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N  | 251 | Molybdate LR PP    | 1.5–2.5 mg/l Mo                                 |
| 257       Nickel T       6-8 mg/l Ni         260       Nitrate LR       0.5-0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2-0.3 mg/l N   | 252 | Molybdate HR PP    | 10-30 mg/l Mo                                   |
| 260       Nitrate LR       0.5–0.7 mg/l N         265       Nitrate TT       10 mg/l N         270       Nitrite T       0.2–0.3 mg/l N  | 254 | Molybdate HR L     | 50-70 mg/l Mo                                   |
| 265 Nitrate TT 10 mg/l N 270 Nitrite T 0.2-0.3 mg/l N  | 257 | Nickel T           | 6–8 mg/l Ni                                     |
| 270 Nitrite T 0.2–0.3 mg/l N   | 260 | Nitrate LR         | 0.5–0.7 mg/l N                                  |
| y .  | 265 | Nitrate TT         | 10 mg/l N                                       |
| 272 Nitrite LR PP 0.1–0.2 mg/l N   | 270 | Nitrite T          | 0.2-0.3 mg/l N                                  |
|  | 272 | Nitrite LR PP      | 0.1-0.2 mg/l N                                  |
| 280 Nitrogen, total LR 10 mg/l N   | 280 | Nitrogen, total LR | 10 mg/l N                                       |
| 281 Nitrogen, total HR 50–100 mg/l N   | 281 | Nitrogen, total HR | 50–100 mg/l N                                   |
| 300 Ozone (DPD) Calibration with basic test 100 Chlorine free  | 300 | Ozone (DPD)        | Calibration with basic test 100 Chlorine free   |

| No.        | Method                           | <b>Recommended range for user calibration</b> Calibration with basic test 100 Chlorine free |
|------------|----------------------------------|---|
| 290<br>292 | Oxygen, active Oxygen, dissolved | possible against meter for dissolved oxygen   |
| 329        | pH-Value LR                      | 6.0–6.6   |
| 330        | pH-Value T                       | 7.6–8.0   |
| 331        | pH-Value L                       | 7.6–8.0   |
| 332        | pH-Value HR                      | 8.6–9.0   |
| 70         | PHMB                             | 15–30 mg/l  |
| 320        | Phosphate LR T                   | 1–3 mg/l PO <sub>4</sub>  |
| 321        | Phosphate HR T                   | 30–50 mg/l PO <sub>4</sub>  |
| 323        | Phosphate, ortho PP              | 0.1–2 mg/l PO <sub>4</sub>  |
| 324        | Phosphate, ortho TT              | 3 mg/l PO <sub>4</sub>  |
| 327        | Phosphate 1, ortho C             | 20–30 mg/l PO <sub>4</sub>  |
| 328        | Phosphate 2, ortho C             | 1–3 mg/l PO <sub>4</sub>  |
| 325        | Phosphate, total TT              | 0.3–6 mg/l P  |
| 326        | Phosphate, hydr. TT              | 0.3–0.6 mg/L P  |
| 334        | Phosphate LR L                   | 5–7 mg/L PO <sub>4</sub>  |
| 335        | Phosphate HR L                   | 30–50 mg/L PO <sub>4</sub>  |
| 316        | Phosphonate                      | 1–2 mg/l PO <sub>4</sub>  |
| 338        | Polyacrylate L                   | 15–20 mg/l Polyacrylic Acid 2′100 sodium salt   |
| 340        | Potassium                        | 3 mg/l K  |
| 350        | Silica                           | 0.5–1.5 mg/l SiO <sub>2</sub>   |
| 351        | Silica LR PP                     | 1 mg/l SiO <sub>2</sub>   |
| 352        | Silica HR PP                     | 50 mg/l SiO <sub>2</sub>  |
| 353        | Silica L                         | 4–6 mg/l SiO <sub>2</sub>   |
| 212        | Sodium hypochlorite              | 8 %   |
| 360        | Sulfate PP                       | 50 mg/l SO <sub>4</sub>   |
| 355        | Sulfate T                        | 50 mg/l SO <sub>4</sub>   |
| 365        | Sulfide                          | 0.2-0.4 mg/l S  |
| 370        | Sulfite                          | $3-4 \text{ mg/l SO}_3$   |
| 384        | Suspended Solids                 | operating range   |
| 386        | Turbidity                        | operating range   |
| 388        | Triazole PP                      | 6 mg/ Benzotriazole   |
| 390        | Urea                             | 1–2 mg/l CH <sub>4</sub> N <sub>2</sub> O   |
| 400        | Zinc                             | 0.2-0.4 mg/L Zn   |
| 405        | Zinc                             | 1–1.5 mg/L Zn   |

#### Store user calibration

100 Chlorine T 0.02-6 mg/l Cl2 0.90 mg/l free Cl2 Perform the required method as described in the manual using a standard of known concentration instead of the water sample.





If the test result is displayed press [MODE], [Shift] + [4] [5] keys and confirm with  $[ _{\bullet} ]$  key.

<user calibration>
100 Chlorine T
0.02-6 mg/l Cl2
0.90 mg/l free Cl2
up: ↑, down: ↓
save: 」

The display shows:

Pressing the arrow key  $[\blacktriangle]$  once increases the displayed result.

Pressing the arrow key  $[\ensuremath{\blacktriangledown}]$  once decreases the displayed result.

Press keys till the displayed result corresponds to the value of the standard.



Confirm with [ع] key to store the new calibration factor.

Cancel user calibration by pressing [ESC] key.

#### Jus Factor saved

The display shows:

100 Chlorine T 0.02-6 mg/l Cl2 1.00 mg/l free Cl2

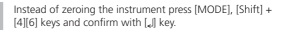
Now the method name is displayed inverse and the test result is calculated with the new calibration factor.

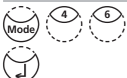
#### **Delete user calibration**

This chapter only applies for methods which can be user calibrated.

100 Chlorine T 0.02-6 mg/l Cl2 Select the required method.







<user calibration>
100 Chlorine T
0.02-6 mg/l Cl2
clear user
calibration?
YES: 1, NO: 0

The display shows:



Press [Shift] + [1] keys to delete user calibration.

Press [Shift] + [0] keys to keep the valid user calibration.

The instrument goes back to Zero-query automatically.

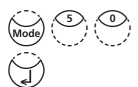
#### 2.4.6 Lab function

#### Reduced operator guidance => "Profi-Mode"

This function may be used for routine analyses with many samples of one method. The following information is always stored in the methods:

- a) Method
- b) Range
- c) Date and time
- d) Differentiation of results
- e) Detailed operator instruction
- f) Compliance with reaction periods

If the Profi-Mode is active, the photometer provides only a minimum of operator instructions. The criteria specified above in d, e, f are no longer included.



Press [MODE], [Shift] + [5][0] keys in succession.

Confirm with [ع] key.

<Profi-Mode>
ON:1 OFF:0

The display shows:



- Press [Shift] + [0] keys to switch the Profi-Mode off.
- Press [Shift] + [1] keys to switch the Profi-Mode on.

switched off

The display shows:

or

switched on



Confirm with [4] key.

#### Note:

- Storage of test results is possible. When results are stored the display also shows "Profi-Mode".
- 2. The selected settings are kept by the photometer even whein it is switched off. To change photometer setting a new setting is required.

#### One Time Zero (OTZ)

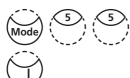
OneTimeZero is available for all methods where Zero is performed in a 24 mm  $\emptyset$  round vial with sample water (see chapter 1.1 Table of Methods).

OneTimeZero can be used for different tests providing the tests are performed with the same sample water and under the same test conditions. When changing the method, it is not necessary to perform a new Zero. The test can be carried out straight away.

When the instrument is first being used for an OTZ compatible method and OneTimeZero is activated, the instrument will request a new Zero with "prepare OT-Zero". Perform Zero as described in the method. This Zero will be stored and used for all methods with OTZ function until the instrument is switched off.

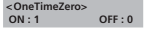
If necessary, a new Zero can be performed by pressing [Zero] key at any time.

#### Switching the "OTZ-Function" on and off:



Press [MODE], [Shift] + [5][5] keys.

Confirm with  $[\c ]$  key.



The display shows:



- Press [Shift] + [0] keys to switch the OTZ off.

switched on

• Press [Shift] + [1] keys to switch the OTZ on.

switched off or

The display shows:



Confirm with [4] key.

The instrument goes back to mode menu automatically.

#### Note:

1. The specified accuracy is valid for all test results when Zero is performed for each test (OneTimeZero function is switched off).

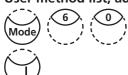
#### 2.4.7 User operations

#### User method list

After switching on the instrument a scroll list of all available methods is automatically shown in the display. To shorten this list according to the requirements of the user it is possible to create a user defined scroll list.

The program structure requires that this list must have at least one active (switched on) method. For this reason it is necessary to activate first all required methods and then to switch off the automatically activated one if this method is not required.

#### User-method list, adaptation



Press [MODE], [Shift] + [6][0] keys.

Confirm with  $[\n ]$  key.

The display shows:

Start with [4] key.

<Method list>
>> 30 • Alkalinity-tot
40 • Aluminium
50 • Ammonium

The complete method list is displayed.

Methods with a point [•] behind the method number will be displayed in the method selection list. Methods without a point will not be displayed in the method selection list.

>> 30 • Alkalinity-tot



Press key  $[\blacktriangle]$  or  $[\blacktriangledown]$  to select the required method from the displayed list.

>> 30 Alkalinity-tot



Switch with [F2] key between "active"  $[\bullet]$  and "inactive"  $[\cdot]$ .

>> 30 • Alkalinity-tot

Select next method, activate or inactivate it and continue.



Confirm with [4] key.

Cancel without storing by pressing [ESC] key.

#### Recommendation:

If only a few methods are required it is recommended to perform Mode 62 first, followed by Mode 60.

All user Polynomials (1-25) and Concentrations (1-10) are displayed in the method list, although they are not programmed by the user. Non-programmed user methods can't be activated!

#### User method list, switch all methods on

This mode function activates all methods. After switching on the instrument a scroll list of all available methods is automatically shown in the display.



Press [MODE], [Shift] + [6][1] keys.



Confirm with [4] key.

<Mlist all on> switch on all methods YES: 1, NO: 0

The display shows:



- Press [Shift] + [1] keys to display all methods in the method selection list
- Press [Shift] + [0] keys to keep the valid method selection list.

The instrument goes back to mode menu automatically.

#### User method list, switch all methods off

The program structure requires that the method list must have at least one active (switched on) method. For this reason the instrument activates one method automatically.



Press [MODE], [Shift] + [6][2] keys.



Confirm with [4] key.

<Mlist all off> switch off all methods YES: 1, NO: 0

The display shows:



- Press [Shift] + [1] keys to display only one method in the method selection list
- Press [Shift] + [0] keys to keep the valid method selection list.

The instrument goes back to mode menu automatically.

#### User Concentration Methods

It is possible to enter and store up to 10 User Concentration Methods.

Therefor you need 2 to 14 standards of known concentration and one blank (deionised water or reagent blank value). The Standards should be measured with increasing concentrations and from the brightest to the darkest colouration.

The measuring range for "Underrange" and "Overrange" is defined with –2600 mAbs\* and +2600 mAbs\*. After selection of a method the concentration of the lowest and highest used standard is displayed as measuring range. The operation range should be within this range to achieve best results

\*1000 mAbs = 1 Abs = 1 E (displayed)

#### **Entering a User Concentration:**



Press [MODE], [Shift] + [6][4] kevs.

Confirm with [4] key.

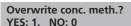
# **Entry Procedure:**

The display shows:

< User concentr.> choose no.: (850-859)

Enter a method number in the range from 850 to 859, e.g.: [Shift] + [8][5][0]

Confirm with [4] key.



#### Note:

wavelength: 1: 530 nm 4: 430 nm 2: 560 nm 5: 580 nm 3: 610 nm 6: 660 nm if the entered number has already been used to save a concentration the display shows the query:

- Press [Shift] + [0] or [ESC] keys to go back to method no. query.
- Press [Shift] + [1] keys to start entry mode.

Enter the required wavelength, e.g.: [Shift] + [2] for 560 nm.



choose unit:

Press [▲] or [▼] keys to select the required unit.



%

Confirm with [4] key.



#### choose resolution

1: 1

2: 0.1

3: 0.01

4: 0.001



Press the appropriate numerical key to select the required resolution, e.g.: [Shift] + [3] for 0.01.

#### Note:

Please enter the required resolution according to the instrument pre-sets:

| range       | max. resolutions |
|-------------|------------------|
| 0.0009.999  | 0.001            |
| 10.0099.99  | 0.01             |
| 100.0 999.9 | 0.1              |
| 10009999    | 1                |

# Measurement procedure with standards of known concentration:

The display shows:

Prepare Zero and press [Zero] key.

#### Note:

Use deionised water or reagent blank value.

The display shows:

Enter the concentration of the first standard; e.g.: [Shift] + [0][.][0][5]

- One step back with [ESC].
- Press [F1] key to reset numerical input.

Confirm with [4] key.

The display shows:

Prepare the first standard and press [Test] key.

The display shows the input value and the measured absorption value. Confirm with  $[\cline{L}]$  key.

Enter the concentration of the second standard; e.g.: [Shift] + [0][.][1]

- One step back with [ESC].
- Press [F1] key to reset numerical input.

Confirm with [4] key.

< User concentr.> prepare Zero press ZERO



< User concentr.> Zero accepted

S1: +



<User concentr.> S1: 0.05 mg/l prepare press TEST



S1: 0.05 mg/l mAbs: 12

S1 accepted S2: +\_\_\_\_

↓ | ESC | F1



S2: 0.10 mg/l prepare press TEST S2: 0.10 mg/l

mAbs: 150 🚽

S2 accepted
S3: +\_\_\_\_
\_ | ESC | F1 | Store



stored!

Prepare the second standard and press [Test] key.

The display shows the input value and the measured absorption value. Confirm with [ ] key.

#### Note:

- Perform as described above to measure further standards
- The minimum of measured standards is 2.
- The maximum of measured standards is 14 (S1 to S14).

If all required standards or the maximum value of 14 standards are measured press [Store] key.

The display shows:

The instrument goes back to the mode menu automatically.

Now the concentration is stored in the instrument and can be recalled by entering its method number or selecting it from the displayed method list.

#### TIP:

Save all your concentration data in a written form because in case of power outage (e.g. changing the battery) all concentration data will be lost and must be entered again.

You might want to use Mode 67 to transfer all concentration data to a PC.

## **User Polynomials**

It is possible to enter and store up to 25 User Polynomials.

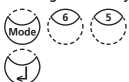
The program allows the user to apply a Polynomial up to the 5th degree:

$$y = A + Bx + Cx^2 + Dx^3 + Ex^4 + Fx^5$$

If only a Polynomial of a lower degree is necessary the other coefficients are specified as zero (0), e.g.: for the 2nd degree is D, E, F = 0.

The values of the coefficients A, B, C, D, E, F must be entered in an academic notation with maximal 6 decimal places, e.g.: 121,35673 = 1,213567E+02

## **Entering a User Polynomial:**



Press [MODE], [Shift] + [6][5] keys.

Confirm with [4] key.

<User polynoms> choose no.: \_\_\_\_ (800-824) The display shows:



Enter a method number in the range from 800 to 824, e.g.: [Shift] + [8][0][0]



Overwrite polynom? YES: 1. NO: 0 Confirm with [4] key.

#### Note:

if the entered number has already been used to save a polynomial the display shows the query:

- Press [Shift] + [0] or [ESC] keys to go back to method no. query.
- Press [Shift] + [1] keys to start entry mode.

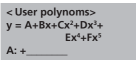
Enter the required wavelength, e.g.: [2] for 560 nm.



wavelength:

1: 530 nm

2: 560 nm



4: 430 nm

5: 580 nm

6: 660 nm



A: 1.32\_\_\_\_ E+\_\_\_



B: +\_\_\_\_

 Press [▲] or [▼] key to change between plus and minus sign

- Enter data of the coefficient A including decimal point,
   e.g.: [Shift] + [1][.][3][2]
- Press [F1] key to reset numerical input.

Confirm with [ ] key.

- Press [▲] or [▼] key to change between plus and minus sign
- Enter the exponent of the coefficient A, e.g.: [Shift] + [3]

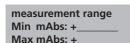
Confirm with [4] key.

Successively the instrument queries the data for the other coefficients (B, C, D, E and F).

#### Note:

If zero [0] is entered for the value of the coefficient, the input of the exponent is omitted automatically.

Confirm every input with [4] key.





Enter measurement ranges from -2600 to +2600 mAbs.

- Press [▲] or [▼] key to change between plus and minus sign.
- Enter the values in Absorbance (mAbs) for the upper limit (Max) and the lower limit (Min).

Confirm every input with [4] key.

# choose unit: >> mg/l g/l mmol/l mAbs µg/l E A

Press [▲] or [▼] keys to select the required unit.



Confirm with [ ] key.

#### choose resolution

1: 1 2: 0.1 3: 0.01 4: 0.001



Press the appropriate numerical key to select the required resolution, e.g.: [Shift] + [3] for 0.01.

#### Note:

Please enter the required resolution according to the instrument pre-sets:

| range       | max. resolutions |
|-------------|------------------|
| 0.0009.999  | 0.001            |
| 10.0099.99  | 0.01             |
| 100.0 999.9 | 0.1              |
| 10009999    | 1                |

#### stored!

The display shows:

The instrument goes back to the mode menu automatically.

Now the polynomial is stored in the instrument and can be recalled by entering its method number or selecting it from the displayed method list.

#### TIP:

- 1. Save all your polynomial data in a written form because in case of power outage (e.g. changing the battery) all polynomial data will be lost and must be entered again.
- 2. You might want to use Mode 67 to transfer all polynomial data to a PC.

#### **Delete User Methods (Polynomial or Concentration)**

In principle a valid user method can be overwritten.

An existing user method (Polynomial or Concentration) can be totally deleted as well and is removed out of the method selection list:



Press [MODE], [Shift] + [6][6] keys.



Confirm with [4] key.

<User m. clear> choose no.: (800-824), (850-859) The display shows:



Enter the number of the User Method you want to delete (in the range from 800 to 824 or 850 to 859), e.g.: [Shift] + [8][0][0]



Confirm with [4] key.

M800 delete? YES: 1, NO: 0 The query is displayed:



Press [Shift] + [1] keys to delete the selected User Method.



Press [Shift] + [0] keys to keep the valid User Method.

The instrument goes back to mode menu automatically.

# **Print Data of User Methods (Polynomials & Concentration)**

With this Mode function all data (e.g. wavelength, unit ...) of stored user polynomials and concentration methods can be printed out or transferred with HyperTerminal to a PC.





Press [MODE], [Shift] + [6][7] keys.



Confirm with [4] key.

The display shows:



Press [4] key to print out the data (e.g. wavelength, unit, ...) of all stored User Methods.



The display shows e.g.:

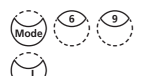
After data transfer the photometer goes back to mode menu automatically.

#### **Initialise User Method System (Polynomials & Concentration)**

Power loss will cause incoherent data. The user method system must be initialised with this mode function to set it to a predefined state.

#### ATTENTION:

All stored user methods (polynomial & concentration) are deleted with initialisation.



Press [MODE], [Shift] + [6][9] keys.

Confirm with [4] key.



The display shows:



Confirm with [4] key.

Initialising? YES: 1, NO: 0 The query is displayed:



• Press [Shift] + [1] keys to start initialisation.

• Press [Shift] + [0] keys to to cancel without initialisation.

The instrument goes back to mode menu automatically.

# 2.4.8 Special functions

#### **Langelier Saturation Index (Water Balance)**

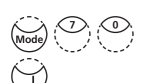
For calculation the following tests are required:

- pH-value
- Temperature
- Calcium hardness
- Total Alkalinity
- TDS (Total Dissolved Solids)

Run each test separately and note the results.

Calculate the Langelier Saturation Index as described:

#### **Calculation of Langelier Saturation Index**



With Mode 71 (see below) it is possible to select between degree Celsius or degree Fahrenheit.

Press [MODE], [Shift] + [7][0] keys.

Confirm with [4] key.

<Langelier> temperature °C: 3°C <=T<=53°C +

The display shows:



Enter the temperature value (T) in the range between 3 and 53°C and confirm with [ $\downarrow$ ] key. If °F was selected, enter the temperature value in the range between 37 and 128°F.

calcium hardness 50<=CH<=1000

The display shows:



Enter the value for Calcium hardness (CH) in the range between 50 and 1000 mg/l CaCO $_3$  and confirm with  $[ \_ ]$  key.

tot. alkalinity 5<=TA<=800

The display shows:



Enter the value for Total Alkalinity (TA) in the range between 5 and 800 mg/l  $CaCO_3$  and confirm with [ $\downarrow$ l] key.

total dissol. solids 0<=TDS<=6000

The display shows:



Enter the value for TDS (Total Dissolved Solids) in the range between 0 and 6000 mg/l and confirm with [4] key.



The display shows:



Enter the pH-value in the range between 0 and 12 and confirm with  $[\t _{=}]$  key.

<Langelier>
Langelier
saturation index
0.00
Esc

The display shows the Langelier Saturation Index.

Press [ ] key to start new calculation.

Return to mode menu by pressing [ESC] key.

#### Operating error:

#### **Examples:**

Values out of defined range:

CH<=1000 mg/l CaCO3!

The entered value is too high.

CH>=50 mg/l CaCO3!

The entered value is too low.



Confirm display message with [A] key and enter a value in the defined range.

# Selection of temperature unit

Entering the temperature value is possible in degree Celsius or degree Fahrenheit. Therefore the following preselection is (once) required.





Press [MODE], [Shift] + [7][1] keys.



Confirm with [4] key.

<temperature>
1: °C 2: °F

The display shows:



Press [Shift] + [1] keys to select degree Celsius.

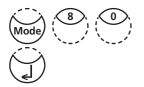
(2)

Press [Shift] + [2] keys to select degree Fahrenheit.

The instrument goes back to mode menu automatically.

# 2.4.9 Instrument basic settings 2

# **Adjusting display contrast**



Press [MODE], [Shift] + [8][0] keys.

Confirm with  $[\c ]$  key.

#### <LCD contrast>

1↑ 1↓

The display shows:



 Press arrow key [A] to increase contrast of the LCD display about one unit.



10 ↑

 Press arrow key [V] to decrease contrast of the LCD display about one unit.



10 ↓

 Press [Store] key to increase contrast of the LCD display about ten units.

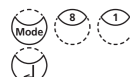


 Press [Test] key to decrease contrast of the LCD display about ten units



Confirm with [4] key.

# **Adjusting display brightness**



Press [MODE] [8] [1] keys.

Confirm with [4] key.

<LCD brightness>

1 1

The display shows:



Press  $[\![ \Delta ]\!]$  key to increase brightness of the display about one unit.



Press  $[\mathbf{V}]$  key to decrease brightness of the display about one unit.

10 ↑ 10 ↓
Store

Press [Store] key to increase brightness of the display about ten units.

Press [Test] key to decrease brightness of the display about ten units.

**0...254 : 200** The display shows:

The brightness can be selected between 0 and 254 units, e.g.: 200.



Confirm with [4] key.

# 2.4.10 Instrument special functions /service

#### **Photometer-Information**



Press [MODE], [Shift] + [9][1] keys.

Confirm with [ع] key.

<System-Info> Software: V201.001.1.001.002 more: ↓, cancel: Esc

This method informs you about the current software version, about the number of performed tests and free memory capacity.



Press arrow key  $[ \mathbf{\nabla} ]$  to display the number of performed tests and free memory capacity.

<System-Info> Number of Tests: 139 free records left 999 cancel: Esc

Finish with [ESC] key.

#### 2.5 Data transfer

To print data or to transmit to a PC the optional IRIM (Infra-Red Interface Module) is required.

# 2.5.1 Data Printing

Besides the IRIM module the following printer is required to print data directly using the USB Interface of the module: HP Deskjet 6940.

#### 2.5.2 Data transfer to a personal computer

Besides the IRIM a transfer program, is required to transmit test results.

Please find detailed information in the IRIM manual or at our homepage in the download-area.

## 2.5.3 Internet Updates

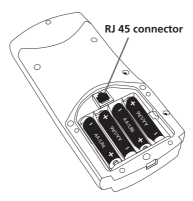
To connect the instrument to the serial interface of a computer the optional connection cable with integrated electronic system is required.

It is possible to update new software applications and additional languages via the internet. Please find detailed information at our homepage in the download-area (as soon as available).

How to open and close the battery compartment cover see chapter 2.1.3!

#### Please Note:

To prevent loss of stored test results store or print them out before performing an Update. If the update procedure is interrupted (eg. interruption of connection, LoBat., etc.) the instrument isn't able to work (no display). The instrument will only work again after completing the data transfer.



# Part 3

**Enclosure** 

# 3.1 Unpacking

Carefully inspect all items to ensure that every part of the list below is present and no visible damage has occurred during shipment. If there is any damage or something is missing, please contact your local distributor immediately.

# 3.2 Delivery contents

Standard contents for MD 600:

| $\bigvee$ |                              |
|-----------|------------------------------|
|           | 1 Photometer in plastic case |
|           | 4 batteries (Type AA/LR 6)   |
|           | 1 Instruction manual         |
|           | 1 Guarantee declaration      |
|           | 1 Certificate of compliance  |
|           | Adapter for 16 mm Ø vials    |
|           | Adapter for 13 mm Ø vials    |

Cleaning brush

Stirring rod, plastic

Reagent sets, IRIM module and connection cable with integrated electronic system are not part of the standard scope of delivery. Please see the General Catalogue for details of available reagent sets.

# 3.3 blank because of technical requirements

Round vials with cap, height 48 mm, Ø 24 mm

Round vials with cap, height 90 mm, Ø 16 mm

#### 3.4 Technical data

Display Graphic Display with backlight

Serial Interface IR interface for data transfer

RJ45 connector for internet updates (see chapter 2.5.3)

Light source light-emitting diode – photosensor – pair arrangement

in a transparent measurement chamber

Wavelength ranges:

 $\lambda 1 = 530$  nm IF  $\Delta \lambda = 5$  nm  $\lambda 2 = 560$  nm IF  $\Delta \lambda = 5$  nm  $\lambda 3 = 610$  nm IF  $\Delta \lambda = 6$  nm  $\lambda 4 = 430$  nm IF  $\Delta \lambda = 5$  nm  $\lambda 5 = 580$  nm IF  $\Delta \lambda = 5$  nm  $\Delta 6 = 660$  nm IF  $\Delta \lambda = 5$  nm IF = Interference filter

Wavelength accuracy  $\pm 1 \text{ nm}$ 

Photometric accuracy\*  $2\% FS (T = 20^{\circ}C - 25^{\circ}C)$ 

Photometric resolution 0.005 A

Measuring range of absorbance -2600 - 2600 mAbs

Protection conforming to IP 68 (1 h, 0.1 m)

Operation Acid and solvent resistant touch-sensitive keyboard with

integral beeper as acoustic indicator.

Power supply 4 batteries (Type AA/LR 6);

lifetime: approx. 26 hours continuous use or 3500 tests

Auto off 20 minutes after last function,

30 seconds acoustical signal before switch off

Dimensions approx. 210 x 95 x 45 mm (unit)

approx. 395 x 295 x 106 mm (case)

Weight (unit) approx. 450 g

Working condition  $5-40^{\circ}\text{C}$  at max. 30-90% relative humidity

(without condensation)

Language options English, German, French, Spanish, Italian, Portuguese,

Polish; further languages via Internet Update

Storage capacity ca. 1000 data sets

# Subject to technical modification!

To ensure maximum accuracy of test results, always use the reagent systems supplied by the instrument manufacturer.

<sup>\*</sup> measured with standard solutions

# 3.5 Abbreviations

| Abbreviation | Definition   |
|--------------|--|
| °C           | degree Celsius (Centigrade)  |
| °F           | degree Fahrenheit $^{\circ}F = (^{\circ}C \times 1.8) + 32$  |
| °dH          | degree German Hardness   |
| °fH          | degree French hardness   |
| °eH          | degree English Hardness  |
| °aH          | degree American Hardness   |
| Abs          | Absorption unit (△ Extinction E)   |
|              | 1000 mAbs = 1 Abs ≙ 1 A ≙ 1 E  |
| μg/l         | (= ppb) Microgram per litre  |
| mg/l         | (= ppm) Milligram per litre  |
| g/l          | (= ppth) gram per litre  |
| KI           | Potassium iodide   |
| Ks4.3        | Acid demand to pH 4.3 – this method is similar to Total Alkalinity but converted into the unit "mmol/l", as the German DIN 38409 demand. |
| TDS          | Total Dissolved Solids   |
|              |  |
| LR           | Low Range  |
| MR           | Medium Range   |
| HR           | High Range   |
| С            | Reagents from Chemetrics®  |
| L            | Liquid reagent   |
| Р            | Powder (reagent)   |
| PP           | Powder Pack  |
| T            | Tablet   |
| TT           | Tube Test  |
|              |  |
| DEHA         | N,N-Diethylhydroxylamine   |
| DPD          | Diethyl-p-phenylendiamine  |
| DTNB         | Ellmans reagent  |
| PAN          | 1-(2-Pyridylazo)-2-napthol   |
| PDMAB        | Paradimethylaminobenzaldehyde  |
| PPST         | 3-(2-Pyridyl)-5,6-bis(4-phenylsulfonic acid)1,2,4-triazine   |
| TPTZ         | 2,4,6-Tri-(2-Pyridyl)-1,3,5-triazine   |

# 3.6 Troubleshooting

# 3.6.1 Operating messages in the display / error display

| Display                                 | Possible Causes   | Elimination  |
|---|---|--|
| Overrange                               | reading is exceeding the range  | if possible dilute sample or use other measuring range   |
|   | water sample is too cloudy  | filtrate water sample  |
|   | too much light on the photo cell  | seal on the cap?<br>Repeat measurement with seal<br>on the cap of the vial.                                |
| Underrange                              | result is under the detection limit   | indicate result with lower x mg/l x = low end of measuring range; if necessary use other analytical method |
| Storagesystem<br>error<br>use Mode 34   | mains power fails or is not connected   | insert or change battery.<br>Delete data with Mode 34  |
| Battery warning                         | warning signal every 3 minutes<br>warning signal every 12 seconds                                       | capacity of the battery is too low; change the batteries   |
|   | warning signal, the instrument switches itself off  | change the batteries   |
| Jus Overrange<br>E4                     | The user calibration is out of the accepted range   | Please check the standard, reaction time and other possible faults.  |
| Jus Underrange<br>E4                    |   | Repeat the user calibration.   |
| Overrange<br>E1                         | The concentration of the standard is too high/too low, so that during user calibration the limit of the | Perform the test with a standard of higher/lower concentration   |
| Underrange<br>E1                        | range was exceeded  |  |
| E40 user<br>calibration<br>not possible | If the display shows Overrange/<br>Underrange for a test result a<br>user calibration is not possible   | Perform the test with a standard of higher/lower concentration   |
| Zero not accepted                       | Light absorption is too great or too low  | Refer to chapter 2.3.4<br>Performing Zero.<br>Clean sample chamber.<br>Repeat zeroing.                     |

| Display  | Possible Causes   | Elimination  |
|--|---|--|
| ???  | The calculation of a value<br>(e.g. combined Chlorine) is<br>not possible | Test procedure correct?<br>If not – repeat test  |
| Example 1  |   | Example 1:   |
| 0,60 mg/l free Cl<br>??? comb Cl<br>0,59 mg/l total Cl |   | The readings for free and total Chlorine are different, but considering the tolerances of each reading they are the same. For this reason the combined Chlorine is most likely zero.                         |
| Example 2  |   | Example 2:   |
| Underrange<br>??? comb Cl<br>1,59 mg/l total Cl        |   | The reading for free Chlorine is under the detection limit. The instrument is not able to calculate the combined Chlorine. In this case the combined Chlorine is most likely the same as the total Chlorine. |
| Example 3  |   | Example 3:   |
| 0,60 mg/l free Cl<br>??? comb Cl<br>Overrange          |   | The reading for total Chlorine is exceeding the range. The instrument is not able to calculate the combined Chlorine. The test should be repeated with a diluted sample.                                     |
| Error absorbance<br>e.g.: T2>T1                        | Fluoride calibration was not correct                                      | Repeat calibration   |

# 3.6.2 General

| Finding   | Possible Causes  | Elimination   |
|---|--|---|
| Test result deviates from the expected.   | Chemical species not as required.                              | Press arrow keys to select the required chemical species.                           |
| No differentiation:<br>e.g. for the Chlorine test<br>there is no selection<br>between differentiated,<br>free or total. | Profi-Mode is switched on.                                     | Switch Profi-Mode off with Mode 50.   |
| The pre-programmed countdown is not displayed.  | Countdown is not activated and/or the Profi-Mode is activated. | Switch the countdown on with Mode 13 and/or switch the Profi-Mode off with Mode 50. |
| It seems that a method is not available.  | Method is not activated in the user method list.               | Activate the required method in the user method list with Mode 60.                  |

#### 3.7 **Declaration of CE-Conformity**

Manufacturer

Name of manufacturer: **Tintometer GmbH** 

> Schleefstraße 8-12 44287 Dortmund

Germany

declares that this product

Product name: AL400

meets the requirements of the following guidelines:

Declaration of EC-Conformity according to DIRECTIVE 2004/108/EC OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 2004. December the 15th and DIRECTIVE 2011/65/EU OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 2011, June the 8th

Applied standard

DIN EN 61326-1:2006

Immunity test requirements for equipment intended for use in industrial locations (Table 2)

Emission according to the requirements for class B equipment

Dortmund, 8th October 2014

Cay-Peter Voss, Managing Director

