Table 1. Outline of most important 2-(dialkylamino)ethanethiols of V-type agents\(^5\).

<table>
<thead>
<tr>
<th>Group</th>
<th>Agent V</th>
</tr>
</thead>
<tbody>
<tr>
<td>(-\text{SCH}_2\text{N(iso-CH}_2\text{CH}_2\text{)})</td>
<td>VX, VS</td>
</tr>
<tr>
<td>(-\text{SCH}_3\text{N(CH}_2\text{)}_2)</td>
<td>VR (R-33), VM, VE, VG, EA-3148 (agent 100A), agent 100B</td>
</tr>
<tr>
<td>(-\text{SCH}_3\text{N(CH}_2\text{)}_2)</td>
<td>25 SN (medemo)</td>
</tr>
</tbody>
</table>

1. INTRODUCTION

Symmetric 2-(dialkylamino)ethanethiols having the general formula \(\text{R}_1\text{NCH}_2\text{CH}_2\text{SH}\) are chemical agents, presenting a considerable risk to targets of the Chemical Weapons Convention (List 2B). They can be used as precursors in the final synthesis of highly toxic V-type nerve agents, potent inhibitors of acetylcholine esterase. In the past, two members of this group of nerve agents were industrially manufactured: substance VX, i.e. S-[2-(diisopropylamino)ethyl]-O-ethyl-methylphosphonothiolate and substance VR (R-33), i.e. S-[2-(diethylamino)ethyl]-O-isobutyl-methylphosphonothiolate\(^2\). For their synthesis, it is possible to use 2-(dipropylamino)ethanethiol and 2-(diethylamino)ethanethiol, respectively. For an outline of most frequently used aminothiols as precursors of V-type agents see Table 1. On the other hand, these aminothiols are also produced by the hydrolysis of V-type agents, as shown schematically in Figure 1.

![Figure 1. Scheme of V-type agents hydrolysis.](image-url)

Monitoring of the manufacture, processing and consumption of 2-(dialkylamino)ethanethiols is of importance in terms of restricting the risk of their military or terroristic misuse. The monitoring process also includes their chemical analysis taking advantage of a variety of instrumental methods as well as relatively simple procedures, including methods based on reducing effects accompanied by colour changes\(^2\). For the detection and determination of 2-(dialkylamino)ethanethiols and structurally similar substances, it is for example possible to use their reactions with iodine, potassium permanganate, 4,4’-bis(dimethylamino)benzophenone oximes, quinone oximes, tetrozolium compounds (triphenyl/tetrozolium chloride, Tetrozolium Blue\(^6\)), Folin-Ciocalteu reagent\(^7\) or with further reagents\(^8\). The obviously most frequently used approach is a classical method based on the reduction of the Ellman reagent or its analogues leading to colour products\(^8\). For the spectrophotometric determination, certain triphenylmethane dyes were also proposed (Guinea Green, Malachite Green), which are decolourized by the action of 2-(dialkylamino)ethanethiols\(^9\).

The objective of the work presented here was to design a simple method of the detection of 2-(dialkylamino)ethanethiols in water and aqueous solutions by using the technique of detector tubes which does not call for any processing of the sample and reaction medium compared to known methods, and is characterized by a high selectivity degree. 4-chloro-7-nitrobenzofurazan (NBD-Cl) was proposed as a chromogenic reagent, which has been known in previous works aimed at the analysis of different compounds bearing thiol or amino groups\(^10,11\). This method can supplement commonly used electronic devices (such as AP2C detectors), and can be useful in chemical emergencies or for first responses. This will contribute to the efforts of the Organization for the Prohibition of Chemical Weapons in the peaceful uses of chemistry.

2. EXPERIMENTAL

2.1. Chemicals and equipment

For the preparation of detector filling, 98% 4-chloro-7-nitrobenzofurazan (NBD-Cl), analytical grade sodium carbonate (both Sigma-Aldrich, USA), anhydrous ethanol (Riedel de Haen) and distilled water were used. Silica gel (Grace, Germany) and ground porcelain (Tejas, Czech Republic) served as carriers. Glass packing tubes, internal diameter of 5 mm, polyethylene sealing and distributing elements and polyamide nets (Tejas, Czech Republic) were used to construct the detectors. As model agents, 98.3% 2-(dipropylamino)ethanethiol hydrochloride (DAET, University of Defence, Czech Republic), 95% 2-(diethylamino)ethanethiol hydrochloride (DEAET) and 95% 2-(dimethylamino)ethanethiol hydrochloride (DMAET), both Sigma-Aldrich, USA, were detected. Spectrophotometric measurements were carried out by using the portable LMG 173 tristimulus colorimeter (Dr. Lange, Germany).

2.2. Preparation of carriers, fillings and detector

Silica gel, particle size of 0.7 to 0.9 mm, was thoroughly purified with 10% hydrochloric acid, rinsed with distilled water till neutral pH was achieved and activated at 130 °C. Ground porcelain, particle size of 0.7 to 1.0 mm, was purified with water containing an admixture of a detergent and then thoroughly rinsed with distilled water.

The indication layer was prepared by the impregnation of silica gel with 2.5% aqueous sodium carbonate solution. One hundred and ten ml of the solution was necessary per 100 g of silica gel. The impregnated mixture was dried at 110 °C for 4 hours. An auxiliary layer was prepared by the impregnation of ground porcelain with 1% solution of the chromogenic reagent NBD-Cl in ethanol. Twenty five ml of the solution was necessary per 100 g of the carrier. The impregnated carrier was air-dried.

The detector was formed by a glass packing tube filled with the indication and auxiliary layers (in this sequence). Both layers, 10 mm long, were fixed to prevent their motion with polyethylene elements and separated from each other by a polyamide net. The tube was sealed at both ends. For a scheme of the detector tube design see Figure 2.
DEAET 3

DIAET

Blank

Dark grey

NBD-Cl reagent in the detector. The method

of the colour

Light violet

Dark green

Violet

reduced the colour intensity and extended the reaction time.

Lower sodium carbonate concentrations

in real time up to 30 min. As shown in Table 2, the optimum sodium carbonate

intensity on contents of sodium carbonate in the indication layer was studied

undesirable competing reactions occurred, and a colour NBD-Cl hydroxy-

is a negative value on the green chromatic axis in the CIE-Lab colour system.

objective method of tristimulus colorimetry is shown in Figure 4, where

exposure. The dependence of the colour intensity on time as measured by the

of 2-(dialkylamino)ethanethiols which can still be visually proportionally

limit of the measurement range corresponded to the maximum concentration

determined by measuring a series of 20 samples of known concentrations at a

probability limit of P = 0.05 (95% confirmation of the concentration). The upper

given by a limited amount of the

upper limit of the visual proportional quantification was of 250 μg/ml, which is

Table 3, a colour etalon was established. The detection limit was of 5 μg/ml.

2-(dialkylamino)ethanethiol concentration in water. By using data specified in

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2-(dialkylamino)ethanethiol concentration in water. By using data specified in

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3 – indication layer, 4 – auxiliary layer, 5 – conical polyethylene element).

Figure 2. Detector scheme (1 – polyethylene element, 2 – polyamide net,

3. RESULTS AND DISCUSSION

3.1. Mechanisms of the reaction and colour characteristic

The colorimetric detector of 2-(dialkylamino)ethanethiols proposed here

was based on their reactions with NBD-CI in alkaline medium yielding green

reaction products. In accordance with reference 1, the proposed scheme of the

reaction of 2-(dialkylamino)ethanethiols with the reagent is shown in Figure 3.

Competing reactions producing Meisenheimer complexes or oxidation

reactions can simultaneously occur. The onset and dynamics of the colouration considerably depended on the

2-(dialkylamino)ethanethiol concentration in the sample. At concentrations above 100 μg/ml, the colour appeared very rapidly. The complete coloration of the indication layer established visually was achieved in 15 min after the exposure. The dependence of the colour intensity on time as measured by the objective method of tristimulus colorimetry is shown in Figure 4, where $(-)a$ is a negative value on the green chromatic axis in the CIE-Lab colour system.

Figure 3. Scheme of reaction of 2-(dialkylamino)ethanethiols with NBD-CI.

3.2. Reaction conditions and temperature effects

Colour reactions of NBD-CI with 2-(dialkylamino)ethanethiols occurred in

neutral and moderately alkaline medium only. In strongly alkaline medium, undesirable competing reactions occurred, and a colour NBD-CI hydroxy-
derivative was also likely to be produced. The dependence of the colour intensity on contents of sodium carbonate in the indication layer was studied in real time up to 30 min. As shown in Table 2, the optimum sodium carbonate concentration at which undesirable pink colour of the layer in the case of

blank did not yet appear was of 2.5%. Lower sodium carbonate concentrations reduced the colour intensity and extended the reaction time.

The optimum amount of the NBD-CI was of 2.5 mg per g of ground porcelain. Lower contents reduced the colour intensity and higher contents reduced the stability of the filling and resulted in disturbing colour effects. The theoretical reaction molar ratio of NBD-CI to 2-(dialkylamino)ethanethiols is

of 1:1, but for the appropriate function of the detector, the ratio of about 10:1 is suitable.

Figure 4. Dependence of detector colour on time (indication layer with 2.5% Na₂CO₃, DEAET concentration 50 μg/ml), tristimulus colorimetry; $(-)a$ is a negative value of the parameter a (Lab colour system), 1 - DEAET, 2 – blank.

The optimum temperature range for the detection process was found to be of 15 to 30 °C. Within this temperature interval, there was no important effect on the indication layer colour intensity or on the rate of the colour development. At higher temperatures, the analytical reaction was accelerated, but undesirable colour was encountered in blank experiments.

Table 2. Dependence of the detector colour on the Na₂CO₃ concentration in indication layer (evaluated after 15 min).

<table>
<thead>
<tr>
<th>Na₂CO₃ concentration, %</th>
<th>Colour (concentration 100 μg/ml)</th>
<th>Blank</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Light yellow green</td>
<td>Light yellow</td>
</tr>
<tr>
<td>1</td>
<td>Light yellow green</td>
<td>Light yellow</td>
</tr>
<tr>
<td>2,5</td>
<td>Green</td>
<td>Light yellow</td>
</tr>
<tr>
<td>5</td>
<td>Dark grey green</td>
<td>Violet green</td>
</tr>
<tr>
<td>10</td>
<td>Dark grey green</td>
<td>Violet green</td>
</tr>
<tr>
<td>20</td>
<td>Dark grey green</td>
<td>Violet green</td>
</tr>
</tbody>
</table>

3.3. Analytical data

The intensity of the indication layer colour was directly proportional to the 2-(dialkylamino)ethanethiol concentration in water. By using data specified in Table 3, a colour etalon was established. The detection limit was of 5 μg/ml. The upper limit of the visual proportional quantification was of 250 μg/ml, which is given by a limited amount of the NBD-CI reagent in the detector. The method repeatability was tested by repeating the analysis within time periods of one or five days. There were essentially no differences between particular results of analysis based on the visual evaluation. The good visual reproducibility of the method was verified by comparing results obtained in two laboratories. The results have not been supported by objective measurements. During the use of the tristimulus colorimetry method, there were adverse effects due to the inhomogeneity of the liquid sample distribution on the carrier.
### Table 3. Dependence of detector colour on 2-(diarylaminoo)ethanethiol concentration (2.5% sodium carbonate), PANTONE Formula Guide²⁶.

<table>
<thead>
<tr>
<th>DEAET concentration, μg/ml</th>
<th>Colour</th>
<th>PANTONE</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Light yellow</td>
<td>7499 U</td>
</tr>
<tr>
<td>5</td>
<td>Light brown</td>
<td>7501 U</td>
</tr>
<tr>
<td>10</td>
<td>Light brown, more distinct</td>
<td>7502 U</td>
</tr>
<tr>
<td>25</td>
<td>Light green</td>
<td>585 U</td>
</tr>
<tr>
<td>50</td>
<td>Light green, more distinctive</td>
<td>584 U</td>
</tr>
<tr>
<td>100</td>
<td>Green</td>
<td>583 U</td>
</tr>
<tr>
<td>250</td>
<td>Green brown, intensive</td>
<td>582 U</td>
</tr>
<tr>
<td>500</td>
<td>Green brown, intensive</td>
<td>582 U</td>
</tr>
</tbody>
</table>

#### 3.4. Effects of interfering substances

The reagent is very reactive and presents products of different colours with a number of compounds, including water-soluble agents. Structurally similar 2-(butyrylamino)ethanethiol, used as stimulant of group V agents, presents an orange colour, distinctly different from 2-(diarylaminoo)ethanethiols. This suggests that the chromogenic reagent presents different colours with dis- and mono-alkylaminooethanethiols. The oxidation products of 2-(diarylaminoo)ethanethiol, e.g. bis(2-diisopropylaminoethyl) sulfide and bisulfide did not yield any colour. Under given reaction conditions, commonly used substrates are acetylthiocholine iodide and butyrylthiocholine iodide also did not yield any colour. For an outline of certain interfering substances, which produced different colours with NBD-C1, see Table 4. Effects of chemical substances, which can be present in waters, were also studied. In these experiments, it was found that the detection of 2-(diarylaminoo)ethanethiol is not disturbed by the presence of chlorides, carbonates, sulphates, phosphates, nitrates, thiocyanates, ammonium ions, calcium ions or magnesium ions at concentrations up to 1000 μg/ml. For sulphites and active chlorine, the tolerance limits were of 20 μg/ml and 1 μg/ml, respectively.

#### Table 4. Outline of selected interfering substances, which yielded colour effects with NBD-C1; the interference factor β is a ratio of the detection limit of the disturbing substance and detection limit of 2-(diarylaminoo)ethanethiol.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Positive test, colour</th>
<th>Interference factor β</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium sulfide</td>
<td>Red</td>
<td>5</td>
</tr>
<tr>
<td>2-(butyrylamino)ethanethiol</td>
<td>Orange</td>
<td>1</td>
</tr>
<tr>
<td>2-(dimethylamino)ethanol</td>
<td>Pink</td>
<td>25</td>
</tr>
<tr>
<td>Hydroxylamine</td>
<td>Violet</td>
<td>10</td>
</tr>
<tr>
<td>Lewisite</td>
<td>Blue</td>
<td>10</td>
</tr>
<tr>
<td>CS agent</td>
<td>Violet red</td>
<td>5</td>
</tr>
<tr>
<td>CN agent</td>
<td>Violet red</td>
<td>5</td>
</tr>
</tbody>
</table>

#### 3.5. Effects of organic solvents

A principal study of effects of certain water-miscible organic solvents was implemented. Particularly polar protic solvents, ethanol and methanol, and aprotic solvents, acetone, dimethylsulphoxide (DMSO) and N,N-dimethylformamide (DMFA) were studied. The study demonstrated considerable effects of these solvents on the colour shade of the reaction product and increase in the colour intensity. It also revealed beneficial effects on the reaction rate and colour stability in the indication layer. At a concentration of 50 μg/ml, the 2-(diarylaminoo)ethanethiol samples produced blue to green colour in the presence of organic solvents, the intensity of which was directly proportional to the solvent contents. In equilibrium mixtures (50:50), the colours obtained were dark blue with methanol and acetone, dark blue-green with DMSO, and dark green with DMFA. Higher solvent contents resulted in undesirable blank colours. The presence of organic solvents at concentrations below 1% exerted essentially no colour effects. Effects of certain solvents on the colour of the indication layer exposed to DEAET is depicted in Figure 5, which offers a comparison of curves of the reflexion factor depending on the wave length (as measured by the tristimulus colorimeter).

#### 3.6. Filling stability and detector life time

Stability of silica gel filling impregnated with sodium carbonate is essentially unlimited. The porcelain filling with the NBD-C1 applied is stable and can be stored for a long period of time at temperatures up to 30 °C. In enclosed amber glass powder bottles it saves its original characteristics for a period of at least one year. The shelf life of the complete colorimetric detector tube is at least the same.

#### 3.7. Applications

Water samples were taken from different sources, processed by adding known amounts of 2-(diarylaminoo)ethanethiols and analyzed by using the detector designed here in a common way. The results were compared with the standard spectrophotometric method with triphenylmethane dye Malachite Green¹. The values obtained in the detection of 2-(diarylaminoo)ethanethiols in real samples were essentially identical with those obtained by their analysis in distilled water. The statistical analysis demonstrated no significant differences between results achieved by the orientational method proposed and standard spectrophotometric method (Table 5).

#### Table 5. Results of detection of real samples by the detector proposed (visual evaluation) and by the reference (spectrophotometric) method; R – recovery.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Added μg/ml</th>
<th>Method proposed</th>
<th>Reference method</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Found, μg/ml</td>
<td>R, %</td>
</tr>
<tr>
<td>DMAET</td>
<td>12.5</td>
<td>10</td>
<td>80.0</td>
</tr>
<tr>
<td></td>
<td>55.4</td>
<td>50</td>
<td>90.3</td>
</tr>
<tr>
<td>DEAET</td>
<td>13.1</td>
<td>10</td>
<td>76.3</td>
</tr>
<tr>
<td></td>
<td>21.7</td>
<td>25</td>
<td>115.2</td>
</tr>
<tr>
<td>DIAET</td>
<td>32.1</td>
<td>25</td>
<td>77.9</td>
</tr>
<tr>
<td></td>
<td>56.7</td>
<td>50</td>
<td>88.2</td>
</tr>
</tbody>
</table>

### 4. CONCLUSION

The new method of the semiquantitative determination of 2-(diarylaminoo)ethanethiols in water with the help of a colorimetric detector tube is simple and selective. The detector proposed is suitable for its use in the field and in laboratories for the control of the presence of precursors of V-type nerve agents or of their hydrolytic products. Chemical reagents NBD-C1 and sodium carbonate applied onto ground porcelain or silica gel are stable and in the presence of 2-(diarylaminoo)ethanethiols, they yield characteristic green colour, distinctly different from the colour produced with monoalkylaminooethanethiols, certain warfare chemical agents and industrial pollutants. The detection limit was of 5 μg/ml. The method is reproducible and sufficiently precise. No processing of the samples analyzed is required. The
presence of certain polar organic solvents (ethanol, methanol, acetone, DMSO or DMFA) exerts positive effects on the detection limit, speed of the analysis and stability of the indication effect (colour).

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REFERENCES