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by Salkinoja-Salonen MS, Jokela JK

Affiliation: Department of General Microbiology, University of Helsinki, Finland.

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Measurement of organic halogen compounds in urine as an indicator of exposure

by Mirja S Salkinoja-Salonen, PhD, Jouni K Jokela, MSc

SALKINOJA-SALONEN MS, JOKELA JK. Measurement of organic halogen compounds in urine as an indicator of exposure. Scand J Work Environ Health 1991;17:75—8. The report describes the measurement of urinary organic halogen compounds. The method is an application of the adsorbable organic halogen assay which is widely used for the analysis of industrial waste water and drinking water. It was found that this assay can be applied to human urine if the urine is pretreated to hydrolyze the mucins so as to cleave the neuraminic acid residues responsible for the high viscosity of these slimy proteins. The method was found to be sensitive down to 1 µg of organic halogen/100 ml of urine. Fifty to 260 µg of organic halogen was measured in the night urine of healthy, occupationally unexposed volunteers. Since many toxic chemicals to which man may be exposed environmentally or occupationally are, in fact, halogen compounds, this assay may be used to monitor for human exposure.

Key terms: adsorbable organic halogen, excreted, pollutant.

The majority of all hazardous man-made chemicals contain organically bound chlorine or bromine. This group covers most pesticides and herbicides (1—3) and many antimicrobial agents (4—6) and widely utilized industrial solvents (1, 7—10) that frequently cause occupational poisoning (1, 6—10) and pollute ground water (1, 11). Sixteen out of the 18 organic chemicals, for which the maximal allowable concentration recommended by the World Health Organization at the parts-per-billion (micrograms/liter) level in drinking water are also organic chlorine compounds (12).

The extent to which man is actually exposed to chemicals from the environment is poorly documented. Suitable methods are lacking to analyze the diverse chemicals in any large group of persons. Modern analytical methods, like gas chromatography-mass spectrometry, give accurate results but are costly and require personnel with specific skills. This paper describes the measurement of organic halogen compounds in human urine by microcoulometric titration after adsorption to activated carbon and combustion into hydrogen halides. It is not labor intensive and can be applied to large numbers of urine samples. It is group-specific rather than compound-specific and may thus have potential use for monitoring human exposure to a wide spectrum of chemicals in the environment and in the workplace.

Materials and methods

Collection and pretreatment of urine samples

The entire amount of night urine was collected from the subjects. If the urine could not be analyzed on the same day, the pH was adjusted to ≤ 5 with nitric acid.

Before the analysis the urine was pretreated to desialinate the urine-contained mucins (sialic acid containing mucoproteins) either with neuraminidase (0.01 U/50 ml of urine, > 2 h at 37°C, pH 4 to 5, adjusted with nitric acid) or mild acid hydrolysis at pH 1.3 (nitric acid) for 1 h at room temperature. The urine was then diluted to 1000 ml or the desired density (1.024 g/cm³) with pure water [adsorbable organic halogen (AOX) contents < 5 µg/l; see the section on chemicals and glassware].

Analysis of organic halogen

Aliquots of 50 ml of the pretreated (diluted) urine in triplicate were placed in conical flasks of 250 ml together with sodium nitrate (0.85 g in 5 ml of water) and activated carbon (50 mg). The flasks were closed with teflon-lined screw caps, placed in a gyratory shaker for 1 h or longer (up to overnight) and processed further as described in the standard protocols for determining AOX in water or waste water (13—15).

After incubation, the activated carbon was collected on polycarbonate filter and the AOX measured, after removal of the inorganic chloride (13—15), with the use of the microcoulometric analyzer of Euroglass BV (Delft, The Netherlands), equipped with an automated sample feeder.

Chemicals and glassware

To maintain a low reagent background, pure water (with an AOX content of < 5 µg/l) was used for

1 Department of General Microbiology, University of Helsinki, Finland.

Reprint requests to: Dr M Salkinoja-Salonen, Department of General Microbiology, University of Helsinki, Mannerheimintie 172, SF—00300 Helsinki, Finland.
preparing the reagents and (optional) diluting of the urine. Since common laboratory cleaning agents may contain halogenated disinfectant, only disposable glassware should be used, or the glassware should be washed separately from other laboratory routines. We cleaned with ethanolic potassium hydroxide and rinsed with dilute nitric acid, ethanol, and pure water.

Activated carbon was obtained from Euroglass BV, the polycarbonate filters (0.2 μm) from Nuclepore, and the neuraminidase (EC 3.2.1.18) from Sigma.

The pesticides and their parent compounds used in this study were as follows: aldrin (HHDN, 1,2,3,4, 10,10-hexachloro-6-7-epoxy-1,4,4a,5,6,7,8,8a-octahydroendo-1,4,exo-5,8-dimethanophthalene, Riedel-de Haen AG D-3016, Selze, Germany), 3,5-dichloroaniline (Ishihara Sangyo Kaisha Ltd, Japan), quinotzone (pentachloronitrobenzenzene, Kasei Organic Chemicals, Tokyo), trichloroacetic acid (E Merck, Darmstadt, Germany), 2,4,5-trichlorophenol and pentachlorophenol (Fluka AG, Buchs, Switzerland). The disinfectants were dichlorophene (2,2'-methylene-bis-4-chlorophenol, Givaudan Corporation, Clifton, New Jersey, United States) and Irgasan DP—300 (2,4,4'-trichloro-2'-hydroxydiphenyl ether, Bayer AG, Leverkusen, Germany). The industrial solvents were 1,2-dichlorobenzene (Fluka AG) and 1,1,1-trichloroethane (Rathburn Chemicals Ltd, Walkerburn, United Kingdom). The following pesticides and slimicides were obtained as technical products from the suppliers to the local industry: 2,4-D [(2,4-dichlorophenoxy) acetic acid], DBNPA (2,2-dibromo-3-nitrilopropionamide), and 1,4-bis-bromoacetoxy-2-butene.

For testing the recovery, these chemicals were dissolved in urine to 5—20 μg of organic halogen (stock solution on ethanol or tetrahydrofuran) per 50 ml.

**Results**

We used the microcoulometric assay to measure the contents of organically bound halogen in human urine. The filterability of human urine through the polycarbonate was poor, and therefore the removal of inorganic halides was incomplete at the nitrate wash, and thus the results were unreliable (too high).

The filterability of urine was dramatically improved upon preincubation of the urine with neuraminidase. The filtration time required for 50 ml of urine was decreased by this treatment from 2—5 h to 10 min or less (ie, a time comparable to that for clear water samples). The same effect was achieved by mild acid hydrolysis of the urine (pH ≤ 1.5).

We tested the halogen recovery of a variety of different organic halogen compounds in urine. The results, recorded in figure 1, showed that, of the 15 chemicals tested, 12 were measured with good recovery (>70% for concentrations of 0.2 to 10 μM) in

**Figure 1.** Measurement of organic halogen compounds in urine by the adsorbable organic halogen (AOX) assay. The recovery of halogen from 15 selected pesticides, disinfecting agents, industrial solvents, and slimicides, present in concentrations of 0.2—10 μM, are shown in the figure. [MCPA = (4-chloro-2-methyl)-phenoxyacetic acid; 2,4-D = 2,4-dichlorophenoxy acetic acid; PCP = pentachlorophenol; 2,4,5-TCP = 2,4,5-trichlorophenol; 3,5-DCA = 3,5-dichloroaniline; TCA = trichloroacetic acid; 1,2-DCB = 1,2-dichlorobenzene; 1,1,1-TCE = 1,1,1-trichloroethane; Br-acetoxy butene = 1,4-bis-bromoacetoxy-2-butene; Kathon 886MW = 5-chloro-2-methyl-3(2H)isothiazolone; DBNPA = 2,2-dibromo-3-nitrilopropionamide]
urine when a hydrolysis or neuraminidase step was included in the standard AOX protocol. The results showed that other organic compounds (urea, uric acid, creatinine, bilirubins) in the urine did not seriously interfere with the assay of the organic halogen compounds.

The AOX content of urine was measured from occupationally unexposed volunteers (N = 51, aged 2—79 years) in Finland. No clear-cut correlation of the urine AOX content with age, sex, or body weight was found. Unexpectedly, our results suggested regional variation. Figure 2 shows examples of results obtained from residents of the Helsinki metropolitan area and those from a small, heavily industrialized town 300 km east of Helsinki. The individual variation of the AOX contents of night urine from the residents of Helsinki ranged from 50 to 90 μg (average: males 71 μg, females 62 μg) and at Imatra from 60 to 190 μg (average: males 132 μg, females 105 μg).

Since one very obvious exposure route of humans to organic halogen compounds is drinking water, the AOX contents of both urine and tap water were measured at six different localities in Finland. The results, depicted in figure 3, indicated a positive correlation between the urinary AOX and the AOX in tap water.

Discussion

The measurement of AOX has been shown to be a valuable tool for monitoring potable water quality (13, 15—17) and undesirable waste-water discharges from industry (14). In this paper we show that it is possible to adapt this method to the monitoring of AOX in human urine.

The filterability of untreated urine was poor, causing the AOX assay to fail, unless modified to remove viscosity caused by mucins excreted by the mucous membranes of the urinary tract. The desired effect was obtained with either neuraminidase or mild acidolysis. Both treatments are known to remove neuraminic acid residues from the mucins (18, 19). Mucins are extremely viscous, rod-shaped molecules, but lose their viscosity when the terminal neuraminic acid residues of their polysaccharide moieties are removed (18).

We found satisfactory recovery for many different randomly selected pesticides, disinfectants, industrial solvents, slimicides, and fungicides (figure 1), chosen as representatives of chemicals to which humans may be exposed occupationally or environmentally. The AOX assay was sufficiently sensitive to detect these chemicals in the urine at the same concentration recommended by the World Health Organization as the acceptable limit in drinking water (1 μg of organic halogen/100 ml) (12). The preliminary results of this work have been reported elsewhere (20).

There was a positive correlation between the urinary AOX contents and those of the residential drinking water (figure 3). This finding may indicate that a significant share of the AOX contained in the drinking water was actually resorbed by the gastrointestinal tract and then renally excreted by the studied individuals. Organic halogen compounds in drinking water disinfected with chlorine have been shown to be mutagenic and are suspected of being carcinogenic (21—25). In Finland the AOX content of drinking water is very high because of the chlorinated disinfection used for humic raw water (24, 25).

Human metabolism is not known to synthesize organic halogen compounds, apart from the iodine-containing hormone thyroxine. If the data in figure 3 are extrapolated to 0 μg of AOX/l of drinking water, there remains a residual AOX in the urine of around 70 μg. It remains to be seen how much of this amount originates from thyroxine turnover and what is the role of airborne exposure and food.

In our opinion, the analysis of urinary AOX will be a useful tool for monitoring human exposure to chem-
icals at the workplace and in the environment. It only takes 10 min to perform, after 1 h of mucin hydrolysis of urine and 1 h for adsorption onto activated carbon has been allowed for. The analysis procedure is simple, requires no costly reagents, and one laboratory person can handle 20 to 30 samples per workday. The method may thus be applied to large populations.

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