Lung function in volunteers before and after exposure to trichloramine in indoor pool environments and asthma in a cohort of pool workers

Gunnar F Nordberg,1 Nils-Goran Lundstrom,1 Bertil Forsberg,1 Annika Hagenbjork-Gustafsson,1 Birgitta J-son Lagerkvist,1 Johan Nilsson,1 Mona Svensson,1 Anders Blomberg,2 Leif Nilsson,1,3 Alfred Bernard,4 Xavier Dumont,4 Helen Bertilsson,1 Kare Eriksson1

ABSTRACT

Objectives: Exposure to trichloramine (NCl3) in indoor swimming-pool environments is known to cause mucous membrane irritation, but if it gives rise to changes in lung function or asthma in adults is not known. (1) We determined lung function in volunteers before and after exposure to indoor pool environments. (2) We studied the occurrence of respiratory symptoms and asthma in a cohort of pool workers.

Design/methods/participants: (1) We studied two groups of volunteers, 37 previously non-exposed healthy persons and 14 pool workers, who performed exercise for 2 h in an indoor pool environment. NCl3 in air was measured during pool exposures and in 10 other pool environments. Filtered air exposures were used as controls. Lung function and biomarkers of pulmonary epithelial integrity were measured before and after exposure. (2) We mailed a questionnaire to 1741 persons who indicated in the Swedish census 1990 that they worked at indoor swimming-pools.

Results: (1) In previously non-exposed volunteers, statistically significant decreases in FEV1 (forced expiratory volume) and FEV% (p=0.01 and 0.05, respectively) were found after exposure to pool air (0.23 mg/m3 of NCl3). In pool workers, a statistically significant decrease in FEV% (p=0.003) was seen (but no significant change of FEV1). In the 10 other pool environments the median NCl3 concentration was 0.18 mg/m3. (2) Our nested case/control study in pool workers found an OR for asthma of 2.31 (95% CI 0.79 to 6.74) among those with the highest exposure. Exposure-related acute mucous membrane and respiratory symptoms were also found.

Conclusions: This is the first study in adults showing statistically significant decreases in lung function after exposure to NCl3. An increased OR for asthma among highly exposed pool workers did not reach statistical significance, but the combined evidence supports the notion that current workroom exposures may contribute to asthma development. Further research on sensitive groups is warranted.

ARTICLE SUMMARY

Article focus

- Exposure to trichloramine (NCl3) in swimming-pool air is known to cause mucous membrane and pulmonary effects, but statistically significant changes in lung function among adults have not been reported.
- Epidemiological studies of asthma among pool workers are not available.

Key messages

- In this study we found for the first time, statistically significant decreases in lung function in volunteers after exposure to pool air with commonly occurring levels of NCl3.
- We found a tendency towards a higher OR for asthma in a nested case reference study within a cohort of 1102 pool workers.
- Our findings support the notion that current workroom exposures of NCl3 may contribute to asthma development.

Strengths and limitations of this study

- This is the first study showing small but statistically significant decreases in lung function after exposure to pool air. It reports an OR for asthma of 2.31 (95% CI 0.79 to 6.74) among pool workers with the highest exposure (after correction for heredity), but this finding did not reach statistical significance.

INTRODUCTION AND OBJECTIVES

Monochloramines, dichloramines and trichloramines are formed following a reaction between ammonia (NH3) or other nitrogen-containing substances present in swimming-pool water when hypochlorite is used as a disinfectant. Trichloramine (NCl3) is the most volatile chloramine and is emitted into the air of indoor swimming-pools. Exposure
concentrations of NCl₃ in pool environments were not known in these outbreaks. It is known, however, that acute respiratory and eye symptoms may occur among recreational swimmers in relation to measured levels of NCl₃ in pool environments and NCl₃ is considered to be the causative agent.

Only few and inconclusive studies have been performed on lung function among adults after exposures to measured levels of NCl₃ in pool environments and additional studies are required.

Clara cell protein 16 (CC16) is an epithelial protective protein in peripheral lung tissue and changes in its serum levels are used as a biomarker of epithelial integrity. It has been shown to be decreased in relation to the frequency of pool attendance. However, changes in serum levels of CC16 have not been studied after short-term exposure to NCl₃.

Thickett et al reported three cases of occupational asthma among British pool workers exposed to NCl₃. There is a lack of epidemiological studies on asthma among those working in swimming-pool environments.

The objectives of the present study were (1) to perform a controlled human exposure study of lung function and biomarkers of pulmonary epithelial integrity in volunteers before and after exposure to indoor swimming-pool environments. (2) To perform an epidemiological study of self-reported asthma and subjective symptoms in a cohort of indoor swimming-pool workers.

**DESIGN, MATERIALS AND METHODS**

**Air sampling and determination of NCl₃**

Exposure measurements in human exposure study

In the two pool environments where our study of volunteers and pool workers took place hypochlorite was used as disinfectant. Air samples were collected in the breathing zone: one sample for each 2 h exposure, in total 51 samples.

Determination of NCl₃ at other indoor swimming-pools

Additional determinations of NCl₃ were performed 2004–2008 at 10 different pool establishments (7 conventional ones and 3 ‘adventure water lands’) in southern Sweden with totally 30 indoor pools. Hypochlorite was used as disinfectant. At each swimming-pool, air was sampled during 3 h at three to four different locations in close vicinity of the pool. The equipment was mounted on a stand with the filter at a height of approximately 1.5 m. Sampling was performed on three different days during winter and three different days during summer.

Air collection and analysis

One litre/min of air was pumped through a filter (quartz filter QM-A 37 mm Whatman International Ltd, Maidstone, England). The filter was soaked in a solution of sodium carbonate and arsenic trioxide (AsO₃) and dried as presented earlier. When NCl₃ is collected on the filter it is reduced to chloride ion (Cl⁻). After sampling, the filters were extracted with 10 ml of ultrapure water, shaken for 30 min and filtered through a 13 mm syringe filter (IC Acrodisc, PALL). The chlorides were analysed in a suppressed ion chromatography system (Triatron 900 autosampler, Spark, The Netherlands); ICSep AN1, Anion column (CETAC, Omaha, USA); SCX membrane suppressor column (Sequant, Umeå, Sweden); JD-21 conductivity detector (Costech Microanalytical Ltd, Tallin, Estonia). The eluent was 7.5 mM NaOH and the suppressor 5 mM H₂SO₄. Control samples of two known chloride concentrations (0.5 and 5 mg/l) and at least two blanks were run together with the samples in each run. The chloride concentrations in the blanks were subtracted from the concentration in the samples. The detection limits of NCl₃ (1.78 and 1.18 µg/m³ for 2 and 3 h samplings, respectively) were determined as three times the mean SD of the amount collected on filters of 10 blanks. The limits of quantification (5.9 and 3.9 µg/m³ for 2 and 3 h samplings, respectively) were determined as 10 times the mean SD for the same blanks.

**Human exposure study**

**Study groups**

Group A: 37 healthy subjects (20 men and 17 women, mean age 24.5 years). They were not regular swimming-pool visitors and they had not visited a swimming-pool within 4 weeks before study start.

Group B: 14 workers at swimming-pools (5 men, 9 women, mean age 39.9 years).

All participants were non-smokers with normal lung function and had no history of allergy or pre-existing lung disease. Subjects were free of airway infection for ≥ 4 weeks prior to the first exposure and throughout the remainder of the study.

**Study design**

The study was conducted in a crossover control manner. Each volunteer was exposed to filtered air in an exposure chamber and on another occasion to an indoor swimming-pool environment. In the exposure chamber, located in a separate building away from swimming-pools, incoming air was adjusted to room temperature and filtered through a particle filter. The exposures were performed in random order. Successive exposures were separated by ≥ 2 weeks. The exposures were performed either between 8:00 h and 10:00 h or between 10:00 h and 12:00 h. All exposures (pool environment or filtered air) lasted for 2 h. The study subject was exercising on a bicycle ergometer with moderate exercise (minute ventilation 20 l/min/m²), during 15 min followed by 15 min of rest, that is, four periods of exercise and four periods of rest.

**Lung function**

Forced vital capacity (FVC) and forced expiratory volume in 1 s (FEV₁) was determined using a portable
spirometer connected to a computer (KoKo Spirometer and KoKo DigiDoser; Pulmonary Data Service Instrumentation, Inc, Louisville, Kentucky, USA), calibrated in the morning and after every 10th measurement. FEV\textsubscript{1\%} was calculated as a percentage of FVC (FEV\textsubscript{1\%}=FEV\textsubscript{1}×100/FVC). Lung function was measured immediately before and after exercise in a room with non-detectable levels of NCl\textsubscript{3} (<0.002 mg NCl\textsubscript{3}/m\textsuperscript{3}) or in a room adjacent to the exposure chamber.

Blood sampling and determination of biomarkers

We obtained blood samples from the antecubital vein at 0 and 2 h, that is, before and after exposure, and at 4, 6 and 8 h. Peripheral blood was collected into BD Vacutainer tubes (BD, Plymouth, UK). Each sample was allowed to clot for 1–2 h at room temperature, centrifuged at 3000×g and serum was transferred to cryotubes and frozen at −80°C. These samples were sent to the Industrial Toxicology Unit at the Catholic University of Louvain in Brussels (IUTUCL), Belgium for determination of Clara Cell protein 16 (CC16) and surfactant protein D (SPD). CC16 was determined by latex immuno-assay using a rabbit anti-CC16 antibody (Dakopatts, Glostrup, Denmark) and CC16 purified at (IUTUCL) as standards. All samples were run in duplicate at two different dilutions. The between-run and within-run coefficients of variation range 5–10% and results are comparable with ELISA methods. SPD determinations were performed using the Biovendor ELISA kit (Biovendor, Heidelberg, Germany). Analyses were done in duplicate as recommended by the manufacturer.

Total IgE was determined in human serum by a double-antibody sandwich ELISA method (Human IgE ELISA kit, Immunology Consultants Lab; Inc, Newberg, Oregon, USA). The quantity of IgE in the samples was interpolated from a standard curve.

Statistical analyses

All data from CC16 measurements were corrected for diurnal variation according to Helleday et al\textsuperscript{8,9} and recalculated to correspond to 7:00 h. CC16(corr)=CC16 +0.582*T−0.032*T\textsuperscript{2}. T is the time after 7:00 h when the blood sample was taken. Because CC16 values are highest in the morning\textsuperscript{10,11} corrected CC16 values were somewhat greater than measured values.

Statistics: We used repeated measures analyses of variance (Huynh-Feldt corrected) with time and exposure as within-subject factors and group as between-subject factor. Paired t test or Wilcoxon signed rank test was used when comparing exposures to filtered air and pool environment at baseline (0 h) and after exercise (2 h). SPSS V.17.0 was used to perform the statistical analyses. A p value of 0.05 was considered statistically significant.

Epidemiological study

Population

The epidemiological study group included 1741 persons in the Swedish Census of Population and Housing 1990 who had indicated that they worked at swimming-pools. Early 2007 a questionnaire was mailed to them. There was one reminder.

Questionnaire

Questions dealt with: year hired as a pool worker, time periods in various jobs, time spent in swimming-pool environments, various symptoms from the respiratory tract and mucous membranes of the eyes and possible use of medication for asthma 589 women and 513 men, age 30–>80 years (mean age 51.2 years, SD 12) responded (63%). Among 50 non-responders, interviews were performed via telephone. There was a lower prevalence of asthma and respiratory symptoms among the non-responders, not statistically significant.

‘Self reported asthma’ was derived from a positive answer to the following question: “Do you suffer from asthma or have you suffered from asthma?” Whether a person’s asthma started before or after he/she was hired as a pool worker was derived from the combination of questions about year hired as pool worker and when the first symptoms of asthma occurred. Under the general heading ‘Acute symptoms when working in a swimming-pool environment’ there was a question “How large a part of a working day did you usually spend in the swimming-pool environment Hours?”

In a nested case–control study within this cohort, 44 cases of self-reported asthma occurred after the person was hired as a pool worker. In total 128 age-matched and sex-matched controls were selected within the cohort (mean age 50.5 years, SD 10.7).

Exposure assessment

On the basis of information on work titles given by each individual, exposure was classified into three different categories; 0, 1 or 2; where 0 stands for no exposure, 1 for low exposure and 2 for high exposure. The exposure level is not an estimate of the concentration of NCl\textsubscript{3} in air but is based on the average time during a workday the individual spent in the pool area. Those within category 0 did not spend any time in a pool area, for example, a cashier. A person within category 1 did occasionally spend some time in the pool area. A manager of a swimming-pool or a technician belongs to this category. Individuals belonging to category 2 were those spending most of the workday in the pool area, for example, a swimming teacher, or a swimming-pool worker.

Statistics

Fisher’s test was used to test differences between proportions. Conditional logistic regression was used for analyses in the nested case–control study and logistic regression for analyses of asthma in relation to years worked in swimming-pool environments. All statistical analyses were performed using the statistical package R, V2.9.0 (www.r-project.org). p Values equal to or less than 0.05 were considered statistically significant.
Ethics
The project was approved by the Regional ethical review board in Umeå, Sweden (Dnr 05-044M) and volunteers provided written informed consent. The study was carried out according to the declaration of Helsinki.

RESULTS
Air sampling
Experimental exposure (human exposure study)
The NCl₃ levels during the experimental exposures were
- Group A: mean 0.23 mg/m³ (SD 0.09)
- Group B: mean 0.15 mg/m³ (SD 0.04)

Other swimming-pools
NCl₃ concentrations in air at the 10 different indoor swimming-pool establishments were between 0.001 and 0.77 mg/m³, median 0.18 mg/m³, arithmetic mean (AM) 0.21 mg/m³ (n=129). The AM concentrations of NCl₃ in each of the 10 different pool establishments were between 0.09 and 0.32 mg/m³. There was no difference in NCl₃ concentrations during summer compared with winter conditions (results not shown).

Human exposure study
Lung function
Group A
Measured FEV₁ volumes among healthy volunteers as well as the difference before and after 2 h of exposure to pool environment or filtered air are summarised in table 1. There was a small, statistically significant decrease (p=0.01) in FEV₁ (mean decrease=0.05 litre) after exposure to swimming-pool air. After exposure to filtered air there was a slight, not statistically significant increase in FEV₁ (mean increase 0.01 litre). When comparing the differences (Δ-values) in FEV₁ before and after exposure to pool environment with the Δ-values for exposure to filtered air in the same individuals, the difference between Δ-values was statistically significant (p=0.01).

FEV% values among healthy volunteers are also given in table 1. After exposure to pool air, there was a small decrease (0.8 FEV%) that was marginally statistically significant (p=0.05). After exposure to filtered air, there was a small (statistically non-significant) increase in FEV% values. When the individual differences (Δ-values) of FEV% before and after exposure to pool air were compared with the corresponding Δ-values in filtered air, a statistically significant difference was demonstrated (p=0.004, paired t test). Airway obstruction is usually defined as FEV% below 70 (www.goldcopd.com). Only one value was below 70 (after exposure) among the healthy volunteers.

Group B
In table 2, FEV₁ values for the swimming-pool workers are summarised. After exposure to pool air there was a small and not statistically significant decrease in FEV₁, 0.01 litre. There was also a small decrease in FEV₁ after

Table 1 Healthy volunteers (n=37): FEV₁ (forced expiratory volume, litre during 1 s) and FEV% (FEV₁×100/forced vital capacity) measured before and after 2 h exercise in filtered air and pool air, respectively. Mean±SD. Mean differences (before-after) within parentheses.

| Expiratory volume | Exposure in filtered air | Exposure in pool air | Difference in changes* |
|-------------------|--------------------------|----------------------|------------------------|
|                   | before                   | after                | Δ-values               | before       | after       | Δ-values       |
| FEV₁              | 4.10±0.85                | 4.11±0.87            | (−0.01)**              | 4.14±0.87    | 4.09±0.86   | (0.05)**     |
|                   |                          |                      |                        | p=0.01       | p=0.01      |               |
| FEV%              | 80.5±5.8                 | 80.9±5.2             | (−0.4)                 | 80.7±5.3     | 79.9±5.3    | (0.8)*       |

**FEV₁ significantly lower after exposure in pool air, p=0.01
*FEV% lower after exposure to pool air, p=0.05.
°Indicates no statistically significant difference.
*Statistical significance of difference between Δ-values in filtered air and in pool air.

Table 2 Swimming-pool workers (n=14): FEV₁ (forced expiratory volume, litre during 1 s) and FEV% (FEV₁×100/forced vital capacity) measured before and after 2 h exercise in filtered air and pool air, respectively. Mean±SD. Mean differences (before-after) within parentheses.

| Expiratory volume | Exposure in filtered air | Exposure in pool air | Difference in changes* |
|-------------------|--------------------------|----------------------|------------------------|
|                   | before                   | after                | Δ-values               | before       | after       | Δ-values       |
| FEV₁              | 3.56±0.99                | 3.51±0.91            | (0.05)*                | 3.59±0.93    | 3.57±0.92   | (0.014)*     |
|                   |                          |                      |                        | Non-significant | Non-significant |
| FEV%              | 78.86±6.3                | 78.43±5.42           | (0.43)*                | 79.1±4.1     | 77.8±5.1    | (1.36)*       |

*FEV% lower after exposure to pool air, p=0.003 (Wilcoxon signed rank test).
°Indicates no statistically significant difference.
*Statistical significance of difference between Δ-values in filtered air and in pool air.
exposure to filtered air (0.05 litre, p=0.054). When considering the FEV₁ values for the workers (table 2) before and after exposure to pool air, there was a statistically significant decrease of 1.36% (p=0.003). After exposure to filtered air the small decrease in FEV₁ of 0.43% was not statistically significant. Only two FEV% values among the pool workers (one before and one after exposure) were below 70. When comparing the Δ-values in filtered air with those in pool air no statistically significant differences were found. The lack of such differences may be partly related to the lower exposure level in group B compared to Group A.

Biomarkers of pulmonary epithelial integrity: CC16, Group A
Mean CC16corr values and related SDs in previously unexposed healthy volunteers, are shown in figure 1 for 33 of the participants in group A. For the remaining four persons, values were missing and they were therefore excluded from analysis.

At baseline (0 h), mean CC16corr=12.6 µg/l before pool exp (0 h) and 10.3 µg/l immediately before (0 h) exposure to filtered air. This difference (p=0.018, paired t test) is difficult to explain because the same volunteers were exposed to both pool environment and filtered air and they were randomly assigned to either exposure.

CC16, Group B
Results are shown in figure 1. The mean CC16corr was 6.5 µg/l before both pool and filtered air exposures.

The difference between groups A and B persisted during and after exposure (0–8 h) and is statistically significant (p<0.001 repeated measures analysis of variance on log transformed data). There is also a different change with time. Group A decreases with time and group B increases with time. The difference in trend is statistically significant p=0.038.

The decrease with time in group A during and after exposure to pool environment as well as filtered air is statistically significant (p<0.05, GLM repeated analysis model). In groups A and B there is no statistically significant difference in change with time between pool exposure and filtered air. For improved analysis, values were converted to their natural logarithms, SDs decreased,
providing improved statistical conditions, but no statistically significant effect of exposure could be shown (data not shown).

**SPD values**, shown in figure 2, also display a change with time, with lower values with increasing time intervals from initiation of exposure. Considering the log-transformed SPD variable, there was a difference (p<0.05) before and after exposure (ie, SPD values were higher at 0 than at 2 h) and there was a further decrease (p<0.01) with time at 2–8 h (figure 2). This decrease was similar for exposure to pool air and filtered air. In groups A and B we found no statistically significant changes in SPD values in relation to exposure.

**IgE**
The median IgE value was low 1 mg/l in group A and 0.0 in group B.

**Epidemiological study**
There was a statistically significant relationship between the number of hours, during an average day, spent in the swimming-pool environment and the percentage of workers reporting acute symptoms during work (p<0.01; logistic regression). Frequent symptoms were: dyspnoea (13%), cough (23%), nose irritation (29%), throat irritation (24%) and eye irritation (37%).

In the nested case–control study, the OR for asthma was 2.53 (95% CI 0.89 to 7.19) for persons with exposure level 2 (114 controls and 42 cases) compared with persons exposed to level 0 or 1 (14 controls and 2 cases). After correction for heredity, the corresponding numbers were: OR 2.31 (95% CI 0.79 to 6.74).

These values refer to cases of self-reported asthma occurring after they started pool work, compared with controls without asthma.

Among individuals who worked more than 1 year, there was a tendency to a reduced risk of developing asthma in relation to the number of years of work in swimming-pool environments. Only asthma cases that occurred after they started to work as pool workers were considered. This tendency was, however not, statistically significant p=0.07.

**DISCUSSION**
Our observations of statistically significant decreases in FEV₁ and FEV₁% in previously non-exposed volunteers and in FEV₁% in pool workers after exposure to pool air...
are the first such observations in adults. Carbonelle et al. reported an increase in FEV₁/VC among children and a non-statistically significant decrease in adults (n=13) after they had attended a chlorinated pool. Carbonelle et al. found FEV₁/VC to be unchanged in 11 young adults after swimming in a non-chlorinated pool and slightly, but not statistically significantly decreased after swimming in a chlorinated pool. The lack of statistically significant decrease may be related to the fact that only 11 adults were studied, while the statistically significant decrease in our study was based on 37 previously unexposed healthy volunteers. The findings in volunteers were further supported by statistically significant differences in Δ-values. In the 14-pool workers, only one measurement of lung function (FEV₁) was statistically significantly decreased and no statistically significant difference was seen when Δ-values were compared. A possible effect in pool workers at the exposure level of our study (0.15 mg/m³) may be considered uncertain. Very few FEV₁ values were below 70 (indicating no clinically significant airway obstruction within the study group). The reduction in FEV₁% seen after exposure in pool air here, albeit small, may be a sign of an obstructive airway effect. In children, Bernard et al. found a statistically highly significant relationship between cumulative pool attendance during kindergarten and PEF 15 (post-exercise reduction of peak expiratory flow by 15%), providing supportive evidence of airway effects of exposure to chlorinated pool environments.

CC16 levels in serum increase when lung epithelium permeability is adversely affected by air pollutants or other lung toxicants. On the contrary, reduced levels of CC16 in lung lavage fluid occur in several lung disorders, probably due to a decrease in the production of CC16 as a consequence of a depletion of Clara cells. We found a statistically significant difference in the serum level of CC16 between pool workers compared to volunteers. This finding is consistent with our previous finding of a lower CC16 value in school children frequently attending indoor swimming pools than in those with a low attendance at such pools. The difference between workers and previously unexposed healthy volunteers may be due to the older age of the workers but is more likely due to repeated exposures because a similar difference occurred among school children and all these differences may be due to a depletion of Clara cells. We did not find any statistically significant exposure-related changes in concentrations of the biomarkers of pulmonary epithelial integrity (CC16 and SPD) after exposure to pool air for 2 h. The lack of such an exposure-related change was probably due to the relatively short exposure duration and low exposure level of NCl₃. Another possible explanation is that NCl₃ acts preferentially in the more proximal parts of the respiratory tract, inducing a mild constriction of the central airways, but with less interference in the terminal bronchioles, where the Clara cells are located. In previous studies of volunteers exposed to ozone, we found both a decrease in FEV₁ and an increase in serum CC16 concentrations after exposure.

deally, all exposures should have been performed at the same hour, because it is known that CC16 has diurnal variation. However, for practical reasons exposures were started at somewhat different times during the day and all CC16 values in the present study were corrected for diurnal variation. Such correction is essential, but introduces a certain element of uncertainty. In spite of such correction, there was a statistically significant decrease with time of experiment from 0 to 8 h in group A (regardless of exposure to NCl₃). This indicates that the real diurnal variation exceeded the one assumed in the employed correction calculation. For group B there is an opposite trend with time, possibly related to an inadequate correction of the values in this group. The pool workers were older and had been more exposed to NCl₃ during many years of work in pool environments. Our data on SPD, with a statistically significant decrease with time between 0 and 8 h, confirm previously reported diurnal variation.

The absence of exposure-related effects (after 2 h exposure) on serum concentrations of CC16 and SPD in combination with small, statistically significant decreases in FEV₁ and FEV₁% show that the 2 h exposure level in this experiment can be regarded as the Lowest-Observed-Adverse-Effect-Level on the lung for this group of volunteers. It should be borne in mind that individuals with increased sensitivity to adverse respiratory effects, like those with pre-existing asthma, were not included in the present study. Our observation may be of use in relation to administrative action in setting exposure limits for NCl₃. To our knowledge, no health-based limit values for occupational or environmental exposures have yet been set for NCl₃. A technical value of 0.2 mg/m³ was recently recommended in Germany.

Bernard et al. showed that serum total IgE was a factor determining the risk of adverse pulmonary effects after exposure to pool environments. Serum levels of total IgE in the volunteers and workers of our study were low. The absence of an increased level of total serum IgE among the present volunteers indicates that individuals with possibly increased sensitivity due to increased IgE had been successfully excluded. Further studies on persons with elevated serum IgE would be of interest. Another group that may suffer respiratory effects at lower air concentrations of NCl₃ are competitive swimmers because their breathing volumes exceed those of the volunteers in the present study. Helenius et al. found increased respiratory symptoms and bronchial responsiveness in elite swimmers.

Our study indicates that employees in Swedish indoor pools are exposed to approximately the same level of NCl₃ as employees in France and Belgium. We found median NCl₃ concentrations of 0.18 mg/m³ (mean 0.21 mg/m³) in 10 different premises, while Hery et al. reported 0.14–0.91 mg/m³ and Massin et al. reported a mean of 0.24 mg/m³ in public pool environments and 0.67 mg/m³ in establishments with private owners. There are no previous published data on NCl₃ exposure in Swedish indoor pools. The work environment, that is, ventilation and the use of sodium hypochlorite as
disinfectant has probably not changed during the past few decades. This makes it reasonable to estimate that pool workers have been exposed to NCl₃ at approximately the same levels as reported in this study.

In the epidemiological part of the present study, we found a statistically significant relationship between the number of hours spent in swimming-pool environments and the percentage of workers reporting acute symptoms when working. The percentage varied from 13% for dyspnoea to 37% for eye irritation. These findings are in accordance with previous observations in France³ and Holland.¹ These are subjective symptoms reported in a questionnaire also collecting exposure information and there is a possibility for recall bias. However, similar clear outcomes have been reported also in other studies.²³

Our nested case-referent study found an OR for asthma of 2.53 (95% CI 0.89 to 7.19) for workers with more extensive exposure in pool areas (exposure level 2 compared to persons with exposure level 0 or 1). After correction for heredity the corresponding numbers were: OR 2.31 (95% CI 0.79 to 6.74). These values refer to cases of self-reported asthma occurring after they started to work in swimming-pool environments, compared to controls without asthma.

Cases of asthma in pool workers have been reported in the UK,⁸ but no epidemiological evidence has been reported. The findings of the present study did not reach statistical significance and provide only limited support for a causal relationship between asthma and work at indoor swimming–pools. Individuals who are fit for these types of jobs tend to exercise more regularly and may notice respiratory symptoms; this may contribute to confounding. The fact that there was a tendency towards a decreasing risk of asthma in workers with longer work history may indicate a healthy worker effect due to the irritating properties of NCl₃ in pool environments. A recent study,²¹ reported a higher prevalence (4.5%) of new-onset asthma among recreational swimmers with >320 h of cumulative pool attendance compared to 0.4% among swimmers with <320 h of pool attendance, thus supporting a role for exposure at chlorinated pools for development of asthma. In children engaged in recreational swimming, a statistically significant relationship was shown between cumulative attendance at indoor swimming-pools and the probability of developing asthma in those with increased total IgE in serum.¹³¹⁹ Attendance at chlorinated pools before the age of 2 yrs increased the risk of bronchiolitis and asthma.²²

The present findings support the previously advanced hypothesis⁷¹³¹⁹²¹ that exposures to NCl₃ levels commonly occurring in indoor swimming-pool environments can cause acute airway and mucosal symptoms as well as changes in lung function and deterioration of asthma.

CONCLUSIONS
For the first time in adults, statistically significant but small decreases in lung function were found in previously unexposed subjects after exposure to pool air containing 0.23 mg/m³ of NCl₃ compared to filtered air. The changes in lung function occurred in adults without any signs of allergy and with low IgE values. In a cohort of pool workers we found exposure-related acute mucous membrane and respiratory symptoms. An increased OR for asthma (OR 2.31, 95% CI 0.79 to 6.74) was indicated in workers in the highest exposure category compared to lower exposures. Our observations give support to a previously advanced hypothesis that current exposures to NCl₃ can cause adverse effects on mucous membranes and lungs of humans and contribute to the development of asthma. Further research in sensitive groups is warranted.

REFERENCES
1. Jacobs JH, Spaan S, van Rooy GB, et al. Exposure to trichloramine and respiratory symptoms in indoor swimming pool workers. Eur Respir J 2007;29:690–8.
2. Bowen AB, Kile JC, Otto C, et al. Outbreaks of short-incubation ocular and respiratory illness following exposure to indoor swimming pools. Environ Health Perspect 2007;115:267–71.
3. Massin N, Bohadana AB, Wild P, et al. Respiratory symptoms and bronchial responsiveness in lifeguards exposed to nitrogen trichloride in indoor swimming pools. Occup Environ Med 1998;55:258–63.
4. Carbonnelle S, Francaux M, Doyle I, et al. Changes in serum pneumoproteins caused by short-term exposures to nitrogen trichloride in indoor chlorinated swimming pools. Biomarkers 2002;7:464–78.
5. Carbonnelle S, Bernard A, Doyle IR, et al. Fractional exhaled NO and serum pneumo-proteins after swimming in a chlorinated pool. Med Sci Sports Exerc 2008;40:1472–6.
6. Blomberg A, Mudway I, Svennson M, et al. Clara cell protein as a biomarker for ozone-induced lung injury in humans. Eur Respir J 2003;22:983–8.
7. Lagerkvist BJ, Bernard A, Blomberg A, et al. Pulmonary epithelial integrity in children: relationship to ambient ozone exposure and swimming pool attendance. Environ Health Perspect 2004;112:1769–71.
8. Thickett KM, McCord JS, Gerber JM, et al. Occupational asthma caused by chloramines in indoor swimming-pool air. Eur Respir J 2002;19:827–32.
9. Hery M, Hecht G, Gerber JM, et al. Exposure to chloramines in the atmosphere of indoor swimming pools. *Ann Occup Hyg* 1995;39:427–39.

10. Bernard A, Marchandise FX, Depelchin S, et al. Clara cell protein in serum and bronchalveolar lavage. *Eur Resp J* 1992;5:1231–8.

11. Hermans C, Aly O, Nyberg BI, et al. Determinants of Clara cell protein (CC16) concentration in serum: a reassessment with two different immunoassays. *Clin Chim Acta* 1998;272:101–10.

12. Helleday R, Segerstedt B, Forsberg B, et al. Exploring the time dependence of serum Clara cell protein as a biomarker of pulmonary injury in humans. *Chest* 2006;130:672–5.

13. Bernard A, Carbonelle S, Michel O, et al. Lung hyperpermeability and asthma prevalence in schoolchildren: unexpected associations with the attendance at indoor chlorinated swimming pools. *Occup Environ Med* 2003;60:385–94.

14. Hermans C, Bernard A. Clara cell protein: characteristics and potential applications as marker of lung toxicity. *Biomarkers* 1999;1:3–8.

15. Broeckaert F, Arsalane K, Hermans C, et al. Lung epithelial damage at low concentrations of ambient ozone. *Lancet* 1999;353:900–1.

16. Hermans C, Bernard A. State of the art. Lung epithelium-specific proteins. Characteristics and potential applications as markers. *Am J Respir Crit Care Med* 1999;159:646–78.

17. Hoegh SV, Sorensen GL, Torne I, et al. Long-term stability and circadian variation in circulating levels of surfactant protein D. *Immunobiology* 2010;215:314–20.

18. German Working Group on Indoor Guide Values of the Federal Environment Agency. Risk assessment of trichloramine in the air of indoor swimming pools. *Bundesgesundheitsbl* 2011;54:997–1004 (in German with abstract in English)

19. Bernard A, Carbonnelle S, De Burbure C, et al. Chlorinated pool attendance, atopy, and the risk of asthma during childhood. *Environ Health Perspect* 2006;114:1567–73.

20. Helenius L, Tiikanen HO, Sama S, et al. Asthma and increased bronchial responsiveness in elite athletes: atopy and sports event as risk factors. *J Allergy Clin Immunol* 1998;101:646–52.

21. Ferrari M, Schenk K, Mantovani W, et al. Attendance at chlorinated indoor pools and risk of asthma in adult recreational swimmers. *J Sci Med Sport* 2011;14:184–9.

22. Viosin C, Sardella A, Marcucci F, et al. Infant swimming in chlorinated pools and the risk of bronchiolitis, asthma and allergy. *Eur Resp J* 2010;36 41–7.