Tropospheric ozone: respiratory effects and Australian air quality goals

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Abstract

Objective – To review the health effects of tropospheric ozone and discuss the implications for public health policy.

Design – Literature review and consultation with scientists in Australia and overseas. Papers in English or with English language abstracts were identified by Medline search from the international peer reviewed published reports. Those from the period 1980-93 were read systematically but selected earlier papers were also considered. Reports on ozone exposures were obtained from environmental agencies in the region.

Results – Exposure to ozone at concentrations below the current Australian air quality goal (0-12 ppm averaged over one hour) may cause impaired respiratory function. Inflammatory changes in the small airways and respiratory symptoms result from moderate to heavy exercise in the presence of ozone at levels of 0-08-0-12 ppm. The changes in respiratory function due to ozone are short lived, vary with the duration of exposure, may be modified by levels of other pollutants (such as sulphur dioxide and particulates), and differ appreciably between individuals. Bronchial lavage studies indicate that inflammation and other pathological changes may occur in the airways before reductions in air flow are detectable, and persist after respiratory function has returned to normal. It is not known whether exposures to ozone at low levels (0-08-0-12 ppm) cause lasting damage to the lung or, if such damage does occur, whether it is functionally significant. At present, it is not possible to identify confidently population subgroups with heightened susceptibility to ozone. People with asthma may be more susceptible to the effects of ozone than the general population but the evidence is not consistent. Recent reports suggest that ozone increases airway reactivity on subsequent challenge with allergens and other irritants. Animal studies are consistent with the findings in human populations.

Conclusion – A new one hour air quality ozone goal of 0-08 ppm for Australia, and the introduction of a four hour goal of 0-06 ppm are recommended on health grounds.

Ozone (O₃) is a strong oxidant and respiratory irritant. Ozone in the troposphere, the lowest layer of the atmosphere, extending to 10–20 km above the earth’s surface, is formed principally by the action of sunlight on nitrogen oxides (NOₓ), in the presence of reactive organic compounds (ROC). In urban areas NOₓ and ROC are generated largely by motor vehicles and industrial activities that burn fossil fuels. In Australia background levels of ozone range from 0-0-03 ppm. Much higher levels (in the range 0-1–0-4 ppm) occur when photochemical “smog” forms. This is a complex mixture in which nitrogen dioxide (NO₂) and oxygen (O₂) react in the presence of sunlight and ROC, to form O₃, and other oxidant species. In the future, population growth, increased motor vehicle traffic and expanding industrial activity will increase the emissions of the precursors of ozone, and lead possibly to more widespread pollution with photochemical oxidants.

Internationally, air quality goals for ozone averaged over one hour range from 0-06 ppm (Japan) to 0-12 ppm (United States – currently under review). The World Health Organization proposes a one hour air quality goal between 0-076 ppm and 0-1 ppm. In California, where some of the heaviest photochemical oxidant pollution in the world has occurred, the one hour air quality goal for ozone is 0-09 ppm. The current Australian air quality goal for ozone (0-12 ppm averaged over one hour) was last revised in 1985–6. Since then many new studies on the health effects of ozone have been reported, and it is timely to reconsider the scientific basis for the goal.

Human health effects

SYMPTOMS AND LUNG FUNCTION
Symptoms that have been associated with ozone exposure in the urban setting including cough, throat dryness, increased mucus production, wheeze, chest pain or tightness, lassitude, malaise, and nausea. Chamber studies have reported similar findings, although the exposures required to induce symptoms have been higher than those described in field studies.

The most comprehensive information on ozone and lung function in a natural setting comes from studies of children, many of which were conducted at summer camps in North America. In these studies multiple measurements of pulmonary function and concurrent levels of ozone and other air pollutants were made over several weeks. The findings
are summarised in Table 1, where the effect of ozone is displayed as the mean fall in forced expiratory volume in one second (FEV₁) for a unit increase in ozone concentration (measured as parts per million (ppm) averaged over one hour). In all these studies the responses of individual children to ozone varied widely, as shown by Kinney (figure) 9.

Controlled exposure studies carried out with volunteers in chambers show that short term exposures (less than one hour) to ozone concentrations greater than 0-12 ppm lead to reductions in FEV₁ of up to 10% (table 2). Studies which have reported effects on respiratory function at ozone levels less than 0-12 ppm have generally employed exposures exceeding 1–2 hours, in conjunction with moderate to heavy levels of exercise.10,11 McDonnell11 reported a mean decline in FEV₁ of more than 8% after 0-08 ppm ozone for 6-6 hours. Extrapolation of these data at an eight hour work day of heavy manual labour suggests that ozone concentrations of 0-08, 0-10, and 0-12 ppm would decrease FEV₁ by 9, 15, and 20% respectively.11

The chamber studies also show large intersubject variability in the magnitude of responses, although individual ozone responsiveness is moderately reproducible.14 With longer exposures, effects are observed at lower ozone concentrations, indicating that both peak exposure and cumulative exposure may be important determinants of the lung response.15 Most expected spirometric variables are strongly correlated, suggesting that there is a common functional impairment, probably a restriction of inspiration.

The dose-response relationship between ozone exposure and respiratory impairment varies with the level of activity, duration of exposure, and individual susceptibility. In one controlled exposure study, the changes in FEV₁ and forced vital capacity (FVC) increased threefold between light and very heavy exercise for the same change in ozone concentration.15 In the same study a mean 5% fall in FEV₁ was produced by 0-29 ppm ozone over one hour with moderate exercise, but by only 0-07 ppm when exposure was for 6-6 hours. Within the range of the laboratory data, there seems to be a linear or curvilinear relation between ozone exposure and reduction in respiratory function.16 It is not known whether, at lower concentrations, there is a threshold level below which respiratory function is unaffected by changes in exposure to ozone.

As with symptoms, the respiratory effects on lung function reported in field studies tend to be greater than would be expected on the basis of the chamber experiments alone.16,17 It may be that the effect of ozone on the lung is amplified by the presence of other air pollutants such as acid particulates and NO₂. However, chamber experiments have not shown this to occur with NO₂,10 or sulphuric acid.19 Another possibility is that the ozone levels recorded in field studies systematically underestimate personal exposures. There is some evidence that individual doses recorded by personal monitors tend to exceed estimates from stationary monitors.20

Spirometric tests are relatively insensitive guides to the existence of pathology in the airways. Bronchial lavage studies show that inflammation in the walls of the small airways occurs after exposure to ozone levels of 0-08-0-12 ppm over six hours (with moderate exercise), and this inflammation may be observed up to 18 hours after respiratory function has returned to pre-exposure levels.21 The clinical importance of such changes in the small airways is not known.

**Asthma**

In Australia in recent decades there has been a concurrent rise in mortality from asthma22 and output of vehicle emissions and other ozone precursors. There is little information, however, relating ozone to the incidence of asthma, in Australia or overseas. In the United States six cities study23 of 10–12 year old children, particulates were associated more strongly than...
ozone with bronchitis, chronic cough, and chest illness, while ozone was the only measured pollutant associated with asthma. Holguin et al. studied 42 non-smokers, with stable, well-defined asthma but no other pulmonary disease, during periods of high photochemical oxidant pollution. This is one of the most informative studies of asthma that is available, since it includes measurements of exposure and disease for individuals. Ambient levels of ozone, nitrogen dioxide, pollen, temperature, and relative humidity were determined by air monitoring stations located near the dwellings of all subjects. The average ozone level in the 12 hour study periods was less than 0·05 ppm in all instances; the maximum 12 hour ozone exposure was less than 0·12 ppm for 39 of the 42 subjects. Asthma was more common at higher ozone levels: the probability of an episode increased by 0·025 for an increase of 0·04 ppm ozone.

There is some evidence that ambient ozone levels may be weakly related to the occurrence of severe asthma resulting in hospital admission. A recent study in Melbourne, however, reported that “ozone days”, defined by a one hour ozone value ≥0·09 ppm, were not associated with the frequency of hospital presentations for childhood asthma.

Non-specific bronchial responsiveness is increased by exposure to ozone. Horstman exposed healthy volunteers to graded levels of ozone for 6·6 hours with exercise and found that the dose of methacholine which increased airways resistance by 100% decreased with higher levels of ozone. Folinsbee reported that airway reactivity to inhaled methacholine was approximately doubled after exposure to 0·12 ppm ozone with exercise over 6·6 hours. Moreover, changes in non-specific bronchial hyper-responsiveness induced by ozone may predispose individuals to bronchospasm from other environmental agents such as SO₂, or allergens.

OTHER RESPIRATORY ILLNESSES

There is no consistent evidence linking ozone with respiratory illnesses other than asthma. The Ontario air pollution study showed that hospital admissions due to all respiratory illness were 7% greater on “high ozone days” (0·08–0·2 ppm) than on “low ozone days” (0·01–0·06 ppm). The authors noted unusual peaks of acid aerosol of small particle size in this region in the summer, concomitant with raised ozone and sulphate levels. This increase in hospital admissions was interpreted cautiously as an “acid summer haze effect”, rather than an effect of ozone specifically.

There are no epidemiological data on respiratory infections and ozone. Chamber studies have shown, however, that low levels of ozone have the potential to impair immune defence mechanisms. For example, Devlin exposed volunteers to 0·08 or 0·1 ppm for 6·6 hours with moderate exercise (401/min) and reported that alveolar macrophages obtained by lavage had a decreased ability to phagocytise yeast via the complement receptor.

LONG TERM EFFECTS

Recovery of respiratory function after a single exposure to ozone generally occurs within 48 hours although some residual effects may persist for days. In laboratory studies, repeated exposures lead to progressively smaller decreases in respiratory function for a standard level of ozone. In people, long term exposures to ambient ozone seem to produce a functional adaptation which persists for several months. This does not imply tolerance – children in Mexico City who are exposed frequently to high levels of ozone still experience acute ozone-related decrements in lung function when they exercise.

There is little information on the long term effects of ozone on the lung. Detels measured respiratory function of adult never-smokers in two communities in Los Angeles on two occasions, approximately five years apart. In one community, moderate levels of photochemical pollution (mean of 12 month peak hourly values, 1971–82, 0·07 ppm) and low levels of other pollutants were recorded, compared with high levels of photochemical oxidant (mean 0·11 ppm) and relatively high levels of sulphates and particulates in the other setting. A greater decline in respiratory function was observed in the high oxidant setting, but it is not clear how much of this effect should be attributed to ozone rather than other air pollutants. A subsequent paper from the same authors reported more rapid declines in respiratory function in a third community, exposed to high levels of SO₂ and NO₂ but relatively low levels of ozone (mean 1971–82, 0·04 ppm) than in the community with moderate photochemical pollution and low levels of other pollutants.

MORTALITY STUDIES

A study of Los Angeles death records reported that all cause mortality on a given day was correlated with the maximum oxidant concentration on the previous day. A regression model with three variables (NO₂, one day lagged oxidant, and temperature) explained 4% of the short term variation in total mortality. Similar findings were obtained from an analysis of New York mortality data, 1971–76. In one of the most heavily polluted European cities, Athens, peak levels of air pollution (SO₂, ozone and smoke) were associated consistently with a small rise in overall mortality and this association was stronger on days with high temperatures.

Thus, there does seem to be a small, but consistent, association between ozone and daily mortality in some large cities. The increase is small as a fraction of all deaths, and it is difficult to separate the possible effects of ozone from those of other air pollutants.

SUSCEPTIBLE GROUPS

The variability in individual response to ozone is unrelated to the individual’s non-specific airway responsiveness or baseline FEV₁. Smokers may be less sensitive, at least at
concentrations of ozone equivalent to environmental exposure. The lung function response to ozone is modified little by age, but it has been noted that children express symptoms due to ozone less readily than do adults. From the limited data available, it seems that functional responses caused by ozone in the elderly may be less than those in young adults. This may partly reflect reduced physical activity and lower ventilation rates of the elderly, and also lung "stiffening" with age.

It is uncertain whether people with asthma are more susceptible to the effects of ozone, although several recent papers report greater falls in respiratory function after exposure to ozone in those with a history of asthma. On the basis of a small number of studies, there is no evidence of a marked susceptibility to ozone in people with other major respiratory diseases.

Heavy exercise lowers the concentration of ozone at which respiratory effects become apparent and increases the severity of effects, presumably due to the increased dose received by the lung at high levels of ventilation. Athletic performance may be impaired at peak ozone levels around 0.12 ppm. Dose dependent effects are apparent at levels down to 0.08 ppm (over 6-6 hours).

Animal studies
Animal studies suggest that there is no threshold dose for some of the toxic effects of ozone, but little is known about the effects of doses equivalent to those received by humans when ozone concentrations are in the range 0.08-0.12 ppm. In principle, toxicity will occur when the rate of interaction of ozone with its target tissue (or toxicological receptor) exceeds the ability of antioxidant mechanisms to repair that damage. Neither the key toxic events nor the role of the antioxidant mechanisms in these events, however, are well understood at present.

Studies of monkeys exposed to relatively high levels of ozone (>0.25 ppm) suggest that intermittent exposures may produce greater effects than those produced by a continuous exposure regimen, even though the latter results in a higher cumulative dose. This is consistent with findings of controlled exposure studies in humans and suggests that damage results, at least in part, from the repeated attempts to adapt to the irritant challenge as well as to the direct effects of the irritant exposure.

Exposure of animals to ozone on successive days leads to an attenuation of the decrement of lung function. The biological importance of this functional adaptation is not known. Some studies have shown that persistent damage to lung cells may occur at the same time as so-called "adaptive" changes in respiratory function. It has yet to be determined if adaptation is a means of protecting from ozone toxicity with prolonged exposure. An alternate hypothesis is that the tissue is no longer responding to the stimulus provided by the ozone insult because of structural or functional damage.

Little is known about the effects of ozone on lung development in early life. Generally, the lungs of neonatal animals are more resistant to ozone than those of older animals. However, the influence of animal age on response to ozone depends upon the particular response that is studied.

Animal experiments have shown interactions between ozone and other air pollutants. For example, a study of allergen sensitivity in mice noted synergistic effects between ozone and sulphuric acid. In rats, synergism has been noted between O_3 and NO_2 using parameters such as lung weight and detoxification enzyme activities, or lung lavage protein as an index of damage. These interactive effects have been observed, in animals, at levels of pollutants equivalent to those found in the ambient atmosphere.

The limited data available suggest humans are more sensitive to the effects of ozone than the most commonly studied laboratory animal, the rat. For example, rats exposed to 0.5 ppm ozone for 6-6 hours had functional decrements similar to those seen in humans exposed to 0.12 ppm for the same time. It is not known whether this difference in sensitivity extends to other ozone induced effects (for example, inflammatory responses).

Implications for public health
There are many uncertainties surrounding the estimation of the health effects of ozone. In most instances human studies rely on indirect measures of the effective dose to the lung. Personal exposures are frequently estimated from measurement of ambient ozone at a small number of test sites, although it is well known that ozone levels may vary greatly over a small area depending on air flows and local scavenging. In field studies, individual doses are also influenced by factors such as the time spent outdoors, physical activity patterns, and ventilation rates. The consequences of these uncertainties in measurement of the dose of ozone are not predictable, although in the absence of systematic error, the effects of ozone will generally be underestimated.

In natural settings ozone always occurs in association with other pollutants. These pollutants may themselves cause illness and respiratory impairment, and it is frequently difficult in epidemiological studies to control confounding by factors such as NO_2 and particulates. A separate issue is that of interaction or effect modification. The effect of ozone on the respiratory tract may be amplified when exposure occurs in conjunction with other agents such as SO_2. Interactions of this kind complicate risk assessments, and indeed from the point of view of health policy it would be preferable to frame standards in terms of commonly-encountered mixtures or combinations of pollutants (such as the total of ozone and NO_2).

The objective of air quality guidelines is to set levels of air pollution which will avoid, or minimise, adverse health effects. There is contention over whether the changes that are...
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observed after exposure to ozone in the range 0·08 ppm to 0·12 ppm should be described as adverse health effects. For example, it may be argued that the loss of lung function that follows exposure to low levels of ozone (≤ 0·12 ppm) is reversible, is not disabling, and is not known to cause harm in the long term, and therefore should not be labelled as an “adverse” health effect. Another view, which we support, is that any impairment of lung function is by definition an adverse effect on health, especially when the range of individual responses is very wide, and the effect of exposure (to ozone) may be increased by the presence of other pollutants.

There remains considerable uncertainty about the long term effects of low levels of ozone on human health. Animal studies show possible mechanisms by which ozone may cause irreversible damage to the lung, although fibrosis and serious lung disease have not been observed after chronic exposures, even at high levels. Studies in human populations show greater declines in respiratory function with time in communities with high ozone levels, but it remains unclear how much this is due to ozone and not to other, coexisting pollutants, and the functional significance of the observed effect is not known. There are no data from prospective studies of children linking ozone and lung function, and therefore it is not known whether ozone at levels around 0·12 ppm affects lung growth in children.

Impact of tropospheric ozone on health in Australia

Australian cities vary widely in the magnitude of their problems with photochemical pollution. Over the last decade Melbourne and Sydney have often experienced ozone levels exceeding the current NHMRC one hour guideline of 0·12 ppm. (Ozone levels are measured less comprehensively in other Australian cities. From the available data it seems likely that concentrations about 0·12 ppm rarely occur outside Melbourne and Sydney). The factors that influence ozone levels include population size, area of settlement, motor vehicle use, industrial activity, and topography and weather patterns unique to each city. The relative importance of these factors is not well understood. Despite considerable urban growth in the last decade, in Sydney the number of days each year on which the air quality goal for ozone was exceeded has tended to fall. (New South Wales Environment Protection Agency, unpublished data).

The variability in annual exceedances shows the very strong effect of weather on concentrations of pollutants in the urban atmosphere. While source strengths define potential maxima, sunlight, weather conditions, including wind speed and direction, inversion heights and circulation patterns over a city will determine whether concentrations will actually achieve that potential.

What would be the savings in Australia, in health terms, if the air quality goal for ozone were lowered? The number of people in Australia exposed each year to levels of ozone around the current air quality goal of 0·12 ppm in one hour is unknown, but is likely to be very large. For example, in Melbourne it is estimated that between 1982 and 1990 approximately 16 exceedances per annum would occur if a goal of 0·08 ppm were introduced in place of 0·12 ppm.22 Typical exceedances occur over areas that may include about 10% of the surface area of the Melbourne statistical division (total population 3 million). The number of people exposed on each occasion when the monitored level of ozone exceeds 0·08 ppm depends on many factors, including population density and activity patterns (such as time spent outdoors), and therefore no precise estimate can be made. However it seems likely that several hundred thousand people in Melbourne are exposed each year on at least one occasion to levels of ozone between 0·08 and 0·12 ppm, averaged over one hour. In Sydney, the numbers exposed to such levels of ozone may be greater still. Between 1990 and 1992, the number of hours per year when levels of 0·08 ppm and higher were recorded in Sydney was 60, compared with 16 in Melbourne. (S McPhail, personal communication). Exceedances between 0·08 and 0·12 ppm are unlikely to cause respiratory embarrassment for most of the population. Some susceptible individuals will experience a temporary loss of function amounting to 10% or more of FEV1, which may be disabling, although the number of these people is not known. Persons undertaking heavy physical activity out of doors during the period when ozone levels are between 0·08 and 0·12 ppm are likely to experience symptoms (such as cough, throat dryness) and reduced peak athletic performance.

Some people with asthma are likely to experience additional asthma attacks as a consequence of exposure to ozone at 0·08 to 0·12 ppm, either directly from irritant effects of ozone or due to sensitisation to subsequent allergens, but it is not known at present how many attacks of asthma may be attributed to ozone. Asthma is common in Australia, affecting approximately 10% of adults24 and 20–25% of children.25 Therefore a small increase in risk of an asthma attack for an individual represents a substantial additional burden of illness, nationally, as a consequence of widespread exposure to increased ozone levels.

Conclusions

In our judgement, there is firm evidence that exposure to ozone at around 0·12 ppm over one hour may lead to a reduction in respiratory function, inflammatory changes in the lung, and increased reactivity of airways. The changes are small, on average, but there is a wide range of individual susceptibility. Little is known about the effects of chronic exposures on health because few good studies have been carried out, but animal studies indicate that exposure to ozone may cause long term damage to the lung. The number of people who may be exposed to high levels of ozone is large and, currently, there is no margin of safety between
the goal and the level at which adverse health effects are observed.

We recommend a one hour goal of 0.08 ppb. This is not a "no effect" level - the current evidence suggests that more sensitive measurements of lung function than are currently available will show deleterious effects of ozone at levels down to, and perhaps including, background. At present, reductions in respiratory function have been observed at 0.08 ppb but only in the presence of heavy exercise sustained for two hours or longer.

We also recommend a new four hour goal of 0.06 ppm. The reasons for this are that the effects of ozone on pulmonary function accumulate over several hours,11 and raised ozone levels in towns are typically spread over a period of 3-6 hours. We suggest that the averaging time for a second goal should be set at four hours. To avoid health effects, the published reports suggest that this level should be lower than 0.08 ppb, and we propose a figure that lies approximately midway between the level that is currently known to induce health effects and background.

The one hour goal may be stringent that a one hour goal, if there are short lived peaks in ambient ozone levels. At present, for example, fewer exceedances of the 0.08 ppm level, averaged over four hours, are recorded in Sydney than exceedances of the current objective (0.12 ppm, measured over one hour). We recommend that a one hour goal be retained to control peak exposures (and to maintain a consistent index of air quality) in addition to the four hour goal.

The margin between the air quality goal for ozone and background levels is not wide, and would be approximately halved by reducing the one hour goal from 0.12 to 0.08 ppm. As the goal approaches background the cost of controlling emissions will increase sharply, and at some point it is expected that society will judge that the cost of further improvement in air quality is too great to bear. Social judgements must be made about acceptable costs and tolerable detriment to health. The next step is to explore the social, economic and environmental consequences of adopting and enforcing more stringent air quality goals for ozone.

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