Review

Temporal Changes in Ozone Concentrations and Their Impact on Vegetation

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Abstract: Tropospheric concentrations of phytotoxic ozone (O₃) have undergone a great increase from preindustrial 10–15 ppbv to a present-day concentration of 35–40 ppbv in large parts of the industrialised world due to increased emissions of O₃ precursors including NOₓ, CO, CH₄ and volatile organic compounds. The rate of increase in O₃ concentration ranges between 1 ppbv per decade in remote locations of the Southern hemisphere and 5 ppbv per decade in the Northern hemisphere, where largest sources of O₃ precursors are located. Molecules of O₃ penetrating into the leaves through the stomatal apertures trigger the formation of reactive oxygen species, leading thus to the damage of the photosynthetic apparatus. Accordingly, it is assumed, that O₃ increase reduces the terrestrial carbon uptake relative to the preindustrial era. Here we summarise the results of previous manipulative experiments in laboratory growth cabinets, field open-top chambers and free-air systems together with O₃ flux measurements under natural growth conditions. In particular, we focus on leaf-level physiological responses in trees, variability in stomatal O₃ flux and changes in carbon fluxes and biomass production in forest stands. As the results reported in the literature are highly variable, ranging from negligible to severe declines in photosynthetic carbon uptake, we also discuss the possible interactions of O₃ with other environmental factors including solar radiation, drought, temperature and nitrogen deposition. Those factors were found to have great potential to modulate stomata openness and O₃ fluxes.

Keywords: carbon uptake; manipulation experiment; interaction effect; tropospheric ozone; stomatal ozone flux

1. Changes in O₃ Concentration

Concentrations of O₃ ([O₃]) have been increasing since the preindustrial era due to an increase of its precursors [1]. As an important secondary phytotoxic air pollutant causing injury to plant tissue and a significant decrease of crop and timber yield, it deserves attention from farmers, scientists and the general public. Although the historical [O₃] are unreliable because of a limited number of observations and disagreements in the retrospective modelling [2], it is considered that [O₃] has increased from the baseline of 10–15 ppbv (parts per billion per volume, volume mixing ratio; [3]) to current concentrations of 35–40 ppbv in large parts of the industrialised world [4,5]). Plant species vary in their sensitivity to [O₃], and it seems that genetically based detoxification processes [6] are significant and certainly sufficient to protect plants against any harmful effect of low pre-industrial [O₃]. The concept of “effective O₃ flux”, defined as a balance between stomatal O₃ flux and detoxifying capacity of the plant, was proposed [7]. However, there is a huge variety of clones and cultivars (poplars, beans, etc.), which are sensitive even to low [O₃], demonstrating a strong genetic basis for plant sensitivity to O₃.
Ozone was discovered in 1839 by Christian Friedrich Schönbein during his experiments with the electrolysis of water. At the start of the modern era, [O$_3$] was measured using classical procedures involving titration. In Europe, one of the first measurements was performed by Albert-Lévy in Paris. He showed [O$_3$] to be $11 \pm 2$ ppbv over the period 1876–1910 [8]. Even in high elevations, at Pic du Midi, France, 3000 m a.s.l., a concentration of only 10 ppbv was measured during 1874–1895 with a peak in spring and a minimum in winter [9]. The oldest continuous measurements started at the Arkona-Zingst site (Germany) in 1956: they showed [O$_3$] in the 1950s–1960s to be in the range of 15–20 ppbv [10].

The first harmful effects of O$_3$ were reported in the San Bernardino Mountains of Southern California, in Pinus ponderosa forest [11]. During the 1970s, in inland valleys around Los Angeles, maximum annual [O$_3$] reaching 300–400 ppbv was common [12]. Air pollution led to an increase in the number of days with [O$_3$] > 95 ppbv from 114 in 1963 to 163 days in 1978 [13]. Elsewhere, an increase of 2–4 ppbv per decade was later reported [4], and an increase of 0.35 ppbv per year was seen in South Korea and Japan in 2000–2014 [14,15]. However, at highly polluted urban sites the increase was rapid, 2.6 ppbv per year in Beijing in 2005–2011 [16] and 2 ppbv per year in the Pearl River Delta region [17]. Recently, interannual and decadal changes are reported elsewhere: in the North China plains, there were increases of 3–5 ppbv (2001–2006), mostly attributed to a change of cloud cover and temperature [18] with only a low impact (1–2%) due to afforestation and increased VOC production [19].

In the southern hemisphere, with much less land area and industry, there is a trend of an increasing [O$_3$] of 0.1 ppbv per year from 1990–2015 ranging from 0.04 at Baring head (New Zealand) to 0.21 at Arrival Heights (Antarctica). Overall there seems to be a concentration increase towards southern latitudes [20]. Similarly, an increase of 0.66 ppbv per decade has been observed in Chile at El Tololo mountain [21]. The increase is attributed to the poleward expansion of the Hadley Circulation, bringing the O$_3$-rich air from the stratosphere [20].

Marked diurnal courses of [O$_3$] have usually been found, particularly in large urban agglomerations [22]. O$_3$ is produced over the day, associated with high UV irradiance which drives the photochemistry, whilst at the same time, O$_3$ is being removed by wet and dry depositions on various surfaces and uptake by plants. Typically, NO$_x$ is transported from urban areas at low elevations to rural forested areas where significant amounts of VOC are being produced as natural plant emissions. Thus the appropriate VOC/NO$_x$ ratio for O$_3$ production, ranging between 4 and 15, is achieved [23]. Such middle-range transport of NO$_x$ is responsible for the enhanced production of O$_3$ in rural areas, often at high elevations, and may result in damage of vegetation. A globally averaged lifetime of tropospheric O$_3$ is approximately 23 days [24]. Therefore, O$_3$ could be transported even at long-range between continents [25]. However, its lifetime inside the boundary layer is much shorter because of the surface deposition and chemical reactions, such as reduction of O$_3$ to oxygen. These processes, as well as the spatio-temporal heterogeneity in [O$_3$], are further modulated by the seasonal variability of microclimatic conditions (Figure 1).

At nightfall [O$_3$] rapidly decreases because of the oxidation of NO to NO$_2$ in the absence of production. Distinct seasonal behaviour patterns have been reported in industrialised and rural areas of Europe and the USA: (1) a broad spring-summer maximum of [O$_3$] in the industrialised parts, but (2) a minimum [O$_3$] in summer and autumn in remote regions [26]. Noticeably, spring [O$_3$] maximum is a northern hemispheric phenomenon, only found in northern and western parts of Europe. In the temperate zone of Central and Eastern Europe, the highest [O$_3$] are observed in summer months when temperatures and irradiances reach their highest values (reviewed in Monks [27]), while these are lowest in winter [28,29]. Moreover, substantially higher [O$_3$] are observed under clear skies than under cloudy skies, but not in winter [29].
Figure 1. Scheme of tropospheric ozone (O₃) formation. Tropospheric ozone is formed in a complex series of photochemical reactions driven by ultraviolet (UV) solar radiation. NO₂ is photolyzed to form NO and an electronically excited oxygen atom, O, which reacts with molecular oxygen in the atmosphere (O₂) to form O₃. However, O₃ may also regenerate NO₂ in the presence of NO, thus keeping a photo-stationary state. Therefore, net O₃ production occurs when O₃ precursors, such as carbon monoxide (CO), methane (CH₄) and volatile organic compounds (VOCs), are present in the atmosphere at appropriate concentrations. This chain of photochemical reactions is catalysed by hydroxide anion (OH⁻), hydroperoxyl radical (HO₂), NO and NO₂. Enhanced O₃ production thus occurs under high levels of UV radiation and when the concentration of precursors reaches critical levels. Transport of precursors and catalysts from urban and industrial lands (CO, NOₓ, NO and VOC) to rural conditions enhances mixing of polluted air plumes with clean rural air and results in a VOC/NOₓ ratio conducive for O₃ formation. Therefore, latitudinal and elevational distribution differences in O₃ concentration are likely caused by the distribution of O₃ precursor sources associated with industrialization development and/or by an elevational increase in UV radiation.

In the Czech Republic, Central Europe, the annual maxima of [O₃] are being shifted towards the later parts of the year. The [O₃] peak has shifted from Day of Year (DOY) 120–170 at the beginning of the millennium towards DOY 160–175 over the following 20 years depending on the locality [30]. The shift is probably caused by the change of meteorological conditions towards warmer and dryer years with consequently more favourable conditions for O₃ formation [30]. However, conflicting results are found in the summer monsoon climate of Beijing: [O₃] maximum is in June, while the lowest values of [O₃] are in December [31]. Similarly, in the Yangtze River Delta, the maximum is found in July with a second maximum in September, followed by a minimum in November [32]. At 38 sites involved in the European Monitoring and Evaluation Programme (EMEP), there was a decrease of [O₃] reported in the 1990s, however later, around 2000, the [O₃] had increased; then, in the 2010s it decreased [33]. Interpretation of the trends and spatial patterns over several past decades has been challenging [34]; however, in Europe, because of the successfully adopted measures to reduce O₃ precursors, O₃ surface concentration decreased by 2% from 2000 to 2014 [35].

In the Arctic, there is no clear trend in Barrow (USA, Alaska, 1981–2010) and Resolute (Canada), although there is an increasing trend in short-term periods [36]. In the southern hemisphere, the strongest increase in [O₃] is reported to be during the austral autumn (March-May) with an increase of 0.14 ppbv per year on average, while in other seasons the increase is only 0.07–0.12 ppbv per year [20]. The exception is South Africa with a sharp increase of 1 ppbv per year over the period 1992–2011 [37]. An overview of [O₃] in different regions of the world, with model predictions for the future, is given by Archibald et al. [5].
Currently, $[O_3]$ and its changes are both measured and modelled, however modelling approaches based on state-of-the-art models may suffer from huge uncertainties [38], and some are unable to track accurately $[O_3]$ from the past.

2. Methodological Approaches

The earliest attempts to examine the response of plants to ozone were conducted in glasshouses, growth rooms or transparent enclosures with air-conditioning [39–41]. These studies were short-term, and often focused on foliar damage, rather than growth or physiology. Over several decades, researchers have moved away from this starting point in an effort to achieve realism and understanding (Figure 2).

![Figure 2](image_url)

**Figure 2.** The scheme suggests a time development of methodological approaches applied to examine the plant responses to $O_3$ in relation to a transition from tissue-level studies, through studies at the level of individuals and stands, up to large-scale modelling.

### 2.1. Open-Top Chambers

Soon, methods for quantifying the long-term growth-responses of plants to atmospheric pollutants (SO$_2$, O$_3$, acid mist) were being developed. From early 1970s researchers began to employ open-top chambers (OTCs), each consisting of a transparent plastic structure, often a cylinder, into which a prepared gas mixture could be injected [42,43]. The ‘open top’ enabled mixing of the chamber air with the ambient air in an attempt to aid mixing and reduce unwanted microclimatological effects, particularly heating, that might occur in a fully closed chamber. OTCs became widely used, not only for pollutants such as O$_3$ and SO$_2$ but also to explore the impact of rising CO$_2$ on plant growth. Features of their designs are illustrated in retrospective reviews by Allen et al. [44] and D’Andrea and Rinaldi [45].

However, it was realised that a distinct microclimate would inevitably develop inside the OTC, with higher temperature and humidity than outside, and a lower wind speed and precipitation. The temperature difference was found to be typically less than two degrees [46]. Differences in humidity were likely to influence stomatal conductance, and the lack of plant movement in the natural wind might influence the thigmo-morphogenic response [47]. For many purposes, small differences in environmental variables may not
matter, as both control and treatment chambers induce the same conditions. On the other hand, plant responses to pollutant levels may well be different when the environmental variables are changed, and extrapolation to present-day ambient conditions may thus be unwise. Another issue is the non-uniform distribution of gas within the chamber. Wind tunnel models were constructed to aid design [48,49], and analysis was made of gaseous diffusion pathways [50]. It was found that the addition of a frustum at the top edge of the chamber prevented large incursions of ambient air in windy conditions, and also economised on the use of the pollutant gas. Attention was given to the points of entry of gas to the chamber. Leaves in the ‘near field’ of the entering gas would receive an unwanted high dosage unless the entry points are well distributed [51]. When the pollutant gas is ozone, this may be a special problem. To distribute the injected gas, many designs featured one or two toroidal (“dough-nut”) perforated gas exchange units made of flexible polythene, and sometimes a fan was fitted.

Despite these difficulties, OTCs have been widely used. They have the great advantage of being relatively inexpensive, and therefore a number of them can be deployed in an experimental design which includes appropriate replication. Plants may be rooted in native soil, thus minimising the unwanted effects of root restriction usually seen in pot-grown plants in glasshouses or growth-rooms. Moreover, the environment inside the OTC more-or-less follows the outside environment and thus plants experience almost natural photon irradiance, photoperiods and seasonality. Further, they have been used successfully in many outdoor settings: for field crops [52], for trees [53] and for natural ecosystems [54]. As knowledge progresses, researchers have been able to use OTCs in new and imaginative ways, for example, to investigate the important interactions of \( \text{O}_3 \) pollution with nitrogen supply [55], to investigate competition between species in different \( \text{O}_3 \) scenarios [56,57] and to study the impact of \( \text{O}_3 \) on root growth and development [58].

2.2. Free-Air Systems

The next step was the development of fumigation systems without chambers [59,60]. In most of the early work, the main interest was \( \text{SO}_2 \) and the system was known as the free-air controlled enrichment system (FACE). Later, when used by the \( \text{CO}_2 \) research community this became free-air \( \text{CO}_2 \) enrichment (also FACE). There are no walls or barriers; the gas is distributed to the open air from a circle of towers and the rate is controlled so that the concentrations at the canopy level are constant [61]. The installation and running costs are relatively high, and the degree of replication is usually limited by the funds or area of land available for the project. FACE experiments using \( \text{O}_3 \) have now been used many times, particularly on crop species and young trees [62–64]. In some of these experiments, more than one gas may be used to examine the interactions, particularly between \( \text{O}_3 \) and \( \text{CO}_2 \) [62,65,66].

FACE rings may be quite small (diameter just a few metres) for use with short vegetation [56] or huge (diameter 35 m) for mature forests [67]. To obtain meaningful results, experiments may be conducted for just one growing season in the case of annual crops, or they may be extended for many years in the case of forests [68]. Where such major facilities are established, it becomes desirable and highly effective to include eco-physiological experiments, including both above- and below-ground components [69].

For trees, there has been a preponderance of work on seedlings and saplings. For mature trees, the scale of construction, both for OTCs and FACE rings, is often beyond the funding capabilities. Where two gases are involved, 12 units would seem to be the minimum (see the experimental design of Isebrands et al. [62]). Seedlings and saplings inevitably have different growth responses from mature trees and cannot ever be a realistic surrogate for them [70,71]. Likewise, in studies on trees, there is a tendency to work on fast-growing species, which may not be representative of the world’s forests.
2.3. Micrometeorological Approaches

With the rapid adoption of micrometeorological techniques, particularly eddy-covariance, it became possible to measure directly the flux of trace gases to vegetation. Eddy-covariance requires sensors that respond quickly to concentration changes with a frequency of several Hz. For the case of O\textsubscript{3}, the gas could be measured by chemiluminescence and such sensors were developed very early [72,73]. Fast sensors for CO\textsubscript{2} and H\textsubscript{2}O, and their use to measure CO\textsubscript{2} fluxes came about a decade later [74]. This opened up new possibilities to utilise the naturally occurring [O\textsubscript{3}] to measure and model the photosynthetic response (as CO\textsubscript{2} flux) of mature vegetation to episodes of pollution. This line of research is expected to provide an understanding of exactly how the fundamental ecophysiology of the plant responds to the wildly varying [O\textsubscript{3}], and how this response depends on hourly, daily or seasonal changes in natural conditions by interrogating a long time-series (preferably over several years).

As there are several hundred long-established and well-maintained CO\textsubscript{2} flux stations in the world [75,76], it would seem to be a simple matter to add O\textsubscript{3} sensors to those stations that measure trace gases and the flux of water vapor and heat [77–79]. Usually, the water vapor fluxes are used in conjunction with other variables to estimate the stomatal conductance of the canopy, and thus it has been possible to partition the overall flux of O\textsubscript{3} into stomatal and non-stomatal components. The non-stomatal components of the flux are unlikely to influence the plant’s physiology directly: they include deposition to inert surfaces and reaction with gaseous compounds in the canopy air-space.

It has been found that O\textsubscript{3} uptake by plant canopies is greatest when the CO\textsubscript{2} uptake and stomatal conductance to O\textsubscript{3} are maximal [78,80–83]. Moreover, the photosynthetic uptake of CO\textsubscript{2} is more strongly related to the stomatal conductance to ozone than simply to [O\textsubscript{3}] [28,70,84,85]. In a more recent paper, Jurán et al. [29] show how diffuse sky conditions are associated with higher stomatal uptake of O\textsubscript{3} even though the concentrations of ozone are higher in clear sky conditions. The likely explanation of the diffuse-sky effect is that meteorological conditions are more humid under diffuse skies and thus stomata are more open because of the well-established stomatal response to vapor pressure deficit (VPD; [84]).

A full analysis of the effect of O\textsubscript{3} on CO\textsubscript{2} flux from micrometeorological data is however beset with difficulties because the climatological variables are strongly intercorrelated. In particular, solar radiation is responsible for the photochemical production of O\textsubscript{3} as well as for photosynthesis. Fares et al. [85] tried to overcome the confounding effect by first fitting a simple relationship between solar radiation and CO\textsubscript{2} flux, and thereafter examining the effect of O\textsubscript{3} on the residuals of this relationship. There is scope for the development of a standardised suite of statistical tools to identify the optimal methodology for data-mining in this important area.

2.4. Modelling Approaches

How should this wealth of research data, experimental and observational, be modelled to enable upscaling? The first approach is to develop mechanistic models. Modelling the uptake of gases by leaf canopies using the basic principle of an Ohm’s Law resistance network was developed by Monteith and others in the 1960s in relation to CO\textsubscript{2} and water vapor and is widely applied at scales from leaf to landscape [86,87]. The same type of model, with certain modifications, might be used to simulate the effect of O\textsubscript{3} on CO\textsubscript{2} uptake. The resistance network needs to allow for the uptake through stomatal and non-stomatal pathways [88], and a robust stomatal model is required that can respond to environmental variables [89]. The important and more difficult step is to go beyond modelling uptake, towards modelling the effective dose of O\textsubscript{3}. Ideally, this would require knowledge of the species-dependent internal defences (detoxification) of the leaf tissues. The effective ozone dose is then the difference between the integrated uptake by the stomata and the integrated detoxification rate.
The second modelling approach is the application of machine-learning methods with artificial neural networks, ANN [90]. Prediction using ANN is widely used in industry and has been successfully applied to model CO$_2$ fluxes [91–95]. It requires large data sets, so that all combinations of environmental variables are represented. Savi et al. [90] demonstrated the efficacy of the approach applied to model O$_3$ uptake by analysis of flux data from 11 sites of diverse ecosystem types and locations from the north to the south of Europe. They found the reduction in net ecosystem productivity caused by O$_3$ was highly site-specific and varied 0 to 2.6 percent.

To address the policy issues regarding pollution control, models need to be at regional or even global scale (“upscaling”). This has been tried by [89,94,95]. For example, Oliver et al. [89] adapted the UK’s JULES model by incorporating data-based assumptions about the threshold ozone uptake required to reduce photosynthesis. They found rather little data on dose-response relationships; however, they made reasonable assumptions about the different sensitivity of different plant functional types (PFTs). The model was run for the whole of Europe in the period 1901–2050. O$_3$ was found to reduce the gross primary productivity (GPP) by 4–9 percent and carbon was lost from the soil. Boreal regions suffered less than temperate regions.

3. Effect on Carbon Uptake from Leaf to Ecosystem Level

In the atmosphere, O$_3$ is known to react with double bonds between carbon atoms to produce aldehydes, ketones or higher oxidised molecules— that has been known since 1840 when O$_3$ was discovered. The mechanism is the same in plants, where, after penetrating through the stomatal apertures, ozone molecules oxidise the fatty acids of cell/organelle membranes; this leads to the formation of reactive oxygen species (ROS) causing damage of tissues (Figure 3). Such damage to photosynthetic membranes, despite the plant’s increased defensive production of ROS scavenging enzymes ascorbate [96] and compounds with antioxidative capacity (carotenoids; [97,98]), inevitably leads to local necrotic cell death or early senescence [99]. Among others, Luwe and Heber [6] have shown that elevated [O$_3$] increases concentrations of reduced and oxidised forms of ascorbate in the apoplast of leaves of different plant species. These transient increases are, however, often insufficient to protect leaf tissues. The yellowish mottling occurs particularly close to stomata and appears more often in older than young leaves [100]. Microscopic studies identified enlargement of intercellular space and chloroplast injuries, including thylakoid swellings and membrane disruption, as typical symptoms of O$_3$ impact [97]. Such reduced photosynthetically active leaf area leads to a reduced carbon uptake [101].

Figure 3. Damaging O$_3$ effects at the cellular and leaf levels influence the carbon allocation at tree and ecosystem level.
However, O₃ may affect carbon uptake at various physiological levels. Exposure to chronic [O₃] closes stomatal pores leading thus to a reduced stomatal conductance to CO₂ diffusion and consequently to a reduced photosynthetic CO₂ assimilation [102]. Moreover, O₃ reduces photosynthetic CO₂ uptake via reduced Rubisco (ribulose-1,5-bisphosphate carboxylase/oxygenase) content [103,104]. These effects need to be related to growth and carbon economy at the ecosystem level. In a six-year free-air fumigation at a German forest, Matissek et al. [105] reported a 44% decline in stem productivity in Fagus sylvatica exposed to twice-ambient [O₃]. Reductions in biomass accumulation have been associated with a modified carbon allocation to plant organs. Based on the meta-analysis of temperate and boreal forests of the northern hemisphere, Wittig et al. [106] reported a significant decrease of the root-to-shoot ratio under elevated [O₃] indicating greater sensitivity of root biomass to [O₃]. O₃-induced reduction in root surface area per soil volume unit [107] can result in decreases of water and nutrition uptakes. Investigation of carbon pools revealed faster O₃-induced turnover of leaves/needles, reduction of canopy carbon pools and a substantial increase in carbon deposited to the forest floor [108].

Several metrics have been developed to assess the effect of O₃ on plants and to relate threshold [O₃] to relative yield loss. For example, the index AOT40 (accumulated dose of ozone over a threshold of 40 ppbv), which has to be interpreted with regional and meteorological aspect, has been established. This index is calculated over the sunlight hours and whole growing season, which is being prolonged towards a larger number of days in line with earlier phenological phase occurrence [109,110]. The highest and lowest AOT40 values are reported from Mediterranean regions (38,359 ppb h) and Northern Europe (5094 ppb h), respectively. In Continental Central Europe, AOT40 ranges between 13,636 and 23,515 ppb h, while it is 8207–13,751 ppb h in Atlantic Central Europe [35]. However, this AOT40 index takes into account only of O₃ exposure, but not the physiological properties enabling O₃ diffusion to plant tissues, which is directly responsible for the damage. Therefore, an alternative index based on stomatal O₃ uptake, POD₃ (phytotoxic O₃ dose above a flux threshold of Y nmol O₃ m⁻² s⁻¹) has also been advanced. The threshold is species-specific and depends on the detoxifying capacity of the plant (e.g., [111]). The value of Y ranges from 7 in Alnus glutinosa to 0–1 nmol O₃ m⁻² s⁻¹ in Fagus sylvatica. The minimum values of POD₀ were found in Northern Europe (14 mmol m⁻² year⁻¹), while maximum values of 29.7–32.1 mmol m⁻² year⁻¹ were observed in Mediterranean and Atlantic regions of Europe [35].

While POD₃ is mainly used in scientific and modelling studies, AOT40 still prevails in legislation (European Council Directive 2008/50/EC and monitoring activities [112]). Protection of vegetation recommended by UNECE [113] sets an exposure-based critical level of AOT40 as 5000 ppbv h. Attitudes may change, and POD₃ is now being discussed as a potential integral part of new legislation in Europe [114]. While AOT40 decreased and POD₀ increase in Lithuanian forests over the period 2007–2014 [115], Klingberg et al. [116] reported a reduction of both indices in Picea abies at EMEP sites. More recently, Karlsson et al. [117] confirmed a reduction in AOT40 but did not find a change in POD₁ for the same tree species. Between 2000 and 2014, AOT40 decreased in most of the European countries (except rural northern areas of Iceland, Svalbard and Sweden), while POD₀ increased from 0.03 to 1.06 mmol O₃ m⁻² year⁻¹ across Europe [35]. However, in warm and dry years, AOT40 increased [118] and POD₁ decreased [119] when compared to wet seasons. To correctly determine the long-term trends in the development of these indices, continual time series over several decades are, therefore, very much needed.

Recently, a new flux-based index combining stomatal exposure and crown defoliation has been determined to define critical levels (CLₑf) for forest protection against O₃-induced visible injuries. Sicard et al. [120] recommended CLₑf to be less than 5 mmol m⁻² year⁻¹ POD₁ for broadleaved species and less than 12 mmol m⁻² year⁻¹ POD₁ for conifers. CLₑf representing ≥25% of crown defoliation is recommended to be maximal 17,000 and 19,000 ppbv h of AOT40 for conifers and broadleaved species, respectively. It is obvious that those new indices are inevitably linked to POD₃ and AOT40 and only new limits are set.
As the injuries induced by O$_3$ deposition on cuticle are usually small [82], the negative effect of O$_3$ uptake is connected mainly to stomatal O$_3$ flux. The total flux of ozone to vegetation may be thought of as two components: stomatal flux (uptake through the stomatal pores) and non-stomatal flux (deposition to other surfaces in the canopy and also reaction with gaseous compounds in the canopy air-space). The ratio between stomatal and total O$_3$ flux depends on actual microclimatic conditions and differs in various ecosystems (Table 1). The highest seasonal maxima of total O$_3$ flux were recorded in Quercus ilex forest [119] followed by Populus grandidentata [121], Larix decidua and Pinus halepensis [122,123]. Daily mean values range from 0.8 nmol m$^{-2}$ s$^{-1}$ in Pinus sylvestris forest in Belgium [92] to 8.6 nmol m$^{-2}$ s$^{-1}$ in Q. ilex forest in Italy. See Table 1 for more details.

Stomatal flux is determined by [O$_3$] and two resistances connected in series (leaf boundary layer resistance and the stomatal resistance). While boundary layer resistance depends on wind speed and heat flux, stomatal resistance is primarily influenced by irradiance and VPD [86]. Stomatal O$_3$ flux was found to be 37% of total O$_3$ flux in a northern mixed hardwood forest [121], but it was 21% in semi-arid regions of Israel [122], and only 15% in Larix decidua, Alps, Italy [123]. However, in subalpine coniferous forest dominated by Engelmann spruce (Picea engelmannii) and subalpine fir (Abies lasiocarpa) in southern Wyoming, USA, 59% of stomatal O$_3$ flux was found as an annual average [81]. Similarly, stomatal O$_3$ flux dominated in Czech P. abies mountainous forest under moderately cool and humid climate [124]. Juráň et al. [29] found that stomatal flux represents 53.5% of total O$_3$ flux during summer days with partly-cloudy conditions, but it decreases to 43.5% during sunny days. Moreover, a fraction of stomatal O$_3$ flux could be further modulated by forest age [124,125]. Comparison of modelled and measured fluxes could be found elsewhere [29]. See Table 1 for more details.

Non-stomatal O$_3$ flux includes deposition onto the soil, stems and branches, cuticles and any external surface present. Chemical O$_3$ sink also contributes to non-stomatal flux involving the reactions of O$_3$ molecules with VOCs, NO and aerosols. Non-stomatal flux dominates in spring and summer because of the exponential increase of VOC concentration with increasing air temperature and solar radiation [126,127]. It is the period, when [O$_3$] is usually the highest. Contrary to that, non-stomatal fluxes are negligible over the winter in temperate forests due to low VOC emissions to the atmosphere [128]. They were significant even at a moorland site in Scotland, without the complications of complex forest canopies: the non-stomatal O$_3$ flux was up to 70% of the total flux [77]. These findings suggest that most of O$_3$ deposits on leaf cuticles and/or wet layer of the moss, below the sparse herbaceous canopy.

Here we summarise the effects of O$_3$ on NEP (net ecosystem productivity) and GPP (gross primary productivity) in several forest ecosystems estimated by eddy-covariance technique and modelling approaches (Table 2). The impacts of O$_3$ are very diverse. There is no effect in mature Belgian Scots pine forest on GPP measured over 15 years, although critical levels of AOT40 and POD$_1$ were exceeded in each year of measurement [85]. Similarly, in a poplar stand, Belgium, no effect on NEP was reported [129], even though stomatal O$_3$ flux amounted up to 59% of the total O$_3$ flux. On the other hand, a reduction of NEP was reported in Czech [124] and Swiss forests [130], particularly in Norway spruce and European beech stands. After 20 years of monitoring, the only mild effects of O$_3$ on GPP and photosynthesis were observed in a broad-leaf Harvard forest. These findings were attributed to the fact that 40% of photosynthesis occurs lower in the canopy, in shade, where stomatal conductance and [O$_3$] are lower [131]. So the canopy structure can also modulate the effect of O$_3$.
Table 1. Examples of fractions of stomatal O$_3$ fluxes to total O$_3$ fluxes from different forest ecosystems. Notes: EC—eddy-covariance; *—Total deposition flux in µg m$^{-2}$ s$^{-1}$; **—value not specified.

| Forest Type                  | Species                  | Country       | Total Deposition Flux (nmol m$^{-2}$ s$^{-1}$) | Stomatal Flux (% of Total) | Approach    | Reference |
|------------------------------|--------------------------|---------------|-----------------------------------------------|----------------------------|-------------|-----------|
| Subalpine coniferous         | *Picea engelmannii* and *Abies lasiocarpa* | Wyoming, USA  | 0.5–0.6 * (summer max)                        | 59                         | EC          | [81]      |
| Mountainous                  | *Picea abies*            | Czech Republic| 7.09 (daily mean)                             | dominant **                | modelling   | [124]     |
| Mountainous                  | *Picea abies*            | Czech Republic| 14 (summer max)                               |                           | EC          | [29]      |
| Northern mixed hardwood      | *Populus grandidentata*  | Michigan, USA | 27.7 (seasonal max)                           | 37                         | EC          | [121]     |
| Evergreen Mediterranean      | *Quercus ilex*           | Italy         | 6.9–8.6 (daily average)                       | 34.4                       | EC          | [119]     |
| Coniferous                   | *Pinus sylvestris*       | Belgium       | 0.8–5.8 (daily mean)                          | 26                         | modelling   | [92]      |
| Coniferous                   | *Picea abies*            | Denmark       | 0.5 * (5-years mean)                           | 21                         | modelling   | [125]     |
| Coniferous                   | *Pinus halepensis*       | Israel        | 5–10 (seasonal range)                         | 21                         | EC          | [122]     |
| Alpine                       | *Larix decidua*          | Italy         | 40 (summer daily max)                         | 15                         | EC          | [123]     |
Generally, a strong correlation of GPP to AOT40 index was shown [132]. Among others, a tight linear decrease of whole-plant dry mass with increasing daylight AOT40 was found in Japanese larch (Larix kaempferi) and beech (Fagus crenata) seedlings [133]. Comparing to the preindustrial era, a reduction of 1–16% of GPP was reported for USA vegetation covers [134]. Similarly, Karlsson [135] reported a reduction of living biomass stock due to O₃ to be 2% in northern European countries but up to 32% in central European countries [136]. One of the highest, as much as 24.8%, reduction rates of NEP were for a Norway spruce forest [28] with the highest decrease in July, the warmest month of the year.

Wang et al. [137] modelled biomass carbon stock over 500 years involving a successional series of the temperate deciduous forests. No change of carbon stock was reported due to the change of forest species over the time period with the increasing dominance of isoprene-emitting species. Isoprene acts as a shielding agent preventing O₃ to enter the stomatal aperture—isoprene outside of the leaf reacts with O₃ [138]. It is clear, that O₃ was not an issue for half a millennium, however, it shows a possible direction of how natural ecosystems might possibly evolve and adapt in a O₃-rich world by a mechanism incorporating successional dynamics.

Table 2. Effects of O₃ on various carbon-related criterion. GPP—gross primary production, NEE—net ecosystem exchange, NEP—net ecosystem productivity.

| Type of Ecosystem              | Dominant Plant          | O₃ Effect          | Country      | Criterion | Reference |
|--------------------------------|-------------------------|--------------------|--------------|-----------|-----------|
| Mature stand Scots pine        | neutral                 | Belgium            | GPP          | [92]      |
| Plantation mix of poplars      | neutral                 | Belgium            | NEE          | [129]     |
| Mature stand Stone pine        | neutral                 | Italy              | GPP          | [85]      |
| Mixed hardwood/conifer forests | Red oak, Red maple      | negligible         | USA          | GPP       | [131]     |
| USA vegetation                 | -                       | reduction 1–16%     | USA          | GPP       | [135]     |
| Young stand Norway spruce      | reduction               | Czech Republic     | NEP          | [124]     |
| Young stand Norway spruce      | reduction 24.8%         | Czech Republic     | NEP          | [28]      |
| Young stand Ponderosa pine     | reduction 12%           | USA                | GPP          | [85]      |
| Orchard Orange orchard         | reduction 19%           | USA                | GPP          | [85]      |
| Flux sites in Europe and USA   | reduction 6–29% deciduous forest reduction 4–20% evergreen needle leaf forest | Europe, USA | biomass | [136] |

4. Interactive Effects of O₃

As mentioned above, O₃ triggers the generation and accumulation of H₂O₂ and/or superoxide, i.e., ROS inducing cell death, depending on the species, accession and cultivar [139]. However, the importance of constitutive (i.e., genetic) protection of plants against O₃ remains unclear [140] and is thought to be relatively small as compared to the environmentally induced O₃ protection and acclimation adjustment. Accordingly, the high variability of observed plant/ecosystem responses to O₃ reported in Table 2 is likely to be due to interactive effects of other co-occurring environmental factors. Here we summarise the potential interactive effects of enhanced [O₃] with solar radiation, temperature, drought and nitrogen supply (Figure 4), particularly on tree species and forest canopies. These factors are among those found to have the greatest potential to modulate O₃ fluxes [141]. However, we do not address the issue of the interactive effect of elevated [O₃] and [CO₂] concentrations in this review. This interaction is important and deserves a detailed analysis [89] but it is beyond the scope of the present review.
Figure 4. The scheme of potential interactive effects of O$_3$ and other environmental factors. Thin open arrows indicate an amelioration of negative O$_3$ effects; thick dark arrows indicate an enhancement of O$_3$ effects on plants/trees.

4.1. Interactive Effects of Solar Radiation and O$_3$

Because of the light attenuation, leaves develop differently in upper and lower regions of dense canopies. Sun- and shade-acclimated leaves are formed. Tjoelker et al. [142] reported higher O$_3$ sensitivity of shade leaves compared with sun leaves of sugar maple (Acer saccharum) exposed to twice-ambient [O$_3$]. Shade leaves, compared to sun leaves, had greater proportional reductions in light-saturated rate of CO$_2$ uptake associated with strong decreases in chlorophyll content and apparent quantum efficiency. Similarly, high [O$_3$] reduced height growth of tomato plants (Lycopersicon esculentum) grown at low irradiances, whereas it had negligible, or a stimulating, effect at high irradiances [143]. It is particularly true at high relative air humidity, typical for deep inside the canopy, stimulating the negative effects of O$_3$. Such variability of ozone sensitivity along the vertical canopy gradients has substantial consequences for the modelling of carbon gain [142,144].

Several studies have also investigated the effect of light spectral composition, particularly the combined effect of shortwave ultraviolet (UV) radiation and high [O$_3$] typical for high elevations (Figure 1). Generally, it has been shown that low, ecologically relevant doses of UV-B radiation ameliorate the detrimental effects of O$_3$. Studies of evergreen tree species, Norway spruce and Scots pine, under twice-ambient [O$_3$] showed a development of chlorotic mottling with a reduction of photosynthetic capacity and carboxylation efficiency at near zero UV-B, but not at natural UV-B intensities [145]. Such findings indicate an important role of UV radiation in the activation of numerous protective mechanisms and enhancing plant resistance [146] including an accumulation of antioxidants [147] and flavonoids in plant tissues [148] among others.

4.2. Interactive Effects of Temperature and O$_3$

In the previous experiments, combined effects of O$_3$ with enhanced air temperatures above the ambient as well as chilling temperatures were studied (Figure 4). The experiments with silver birch (Betula pendula) under boreal conditions have shown that temperature-stimulated growth and carbon allocation in tree biomass is counteracted by simultaneous harmful effects of O$_3$ [149]. While elevated [O$_3$] (1.2 × ambient) accelerated leaf senescence, it was delayed by a temperature increase (ambient + 1.2 °C). However, under the combination treatment O$_3$ substantially reduced the temperature effect. Such responses led to an increase of photosynthetic (foliage biomass) to non-photosynthetic (woody biomass) tissue ratio under elevated temperature, but it decreases under O$_3$ and combined treatments. An antagonistic interactive effect of O$_3$ and temperature was also
found on the thickness of abaxial epidermis of European aspen (*Populus tremula*). While leaves exposed to elevated [O$_3$] had significantly thicker adaxial epidermis, elevated temperature reduced the impact of O$_3$ [150].

At the leaf level, in silver birch (*Betula pendula*) and European aspen (*Populus tremula*), a negative O$_3$ effect on CO$_2$ assimilation rate was partly compensated by the elevated temperature demonstrating an interactive effect of both the treatments [151]. Similarly, an interactive effect on intrinsic water use efficiency (iWUE), defined as the ratio between CO$_2$ assimilation rate and stomatal conductance, was observed when elevated [O$_3$] reduced the stimulatory effect of elevated temperature on iWUE. Such interaction was statistically significant in *B. pendula*, but not in *P. tremula*.

Metabolomic analyses did not reveal any interactive effects of elevated temperature and [O$_3$] on most metabolites and metabolic pathways [152]. Nevertheless, combined treatment of these factors led to an enhanced content of amino acids, intermediates of citric acid cycle and some carbohydrates as well as γ-aminobutyric acid (GABA) compared to individual O$_3$ treatment. These compounds are often associated with plant stress response and/or play a role of signalling molecules under environmental stresses [153]. Such plasticity of the plant metabolome ensures no alteration in the seedlings’ sensitivity to autumn and spring frosts as demonstrated by the temperature required for 50% lethality [152].

Under natural conditions, enhanced temperatures are often associated with enhanced VPD values having a substantial effect on the openness of stomata [154,155] and consequently on stomatal O$_3$ flux [28,29]. Accordingly, lower stomatal O$_3$ fluxes and lower O$_3$-induced injuries may be observed in warm Mediterranean conditions than in the cooler boreal and temperate forest ecosystems [156–158].

Although winter O$_3$ fluxes into the forest ecosystems are relatively small compared to the summer fluxes [29], evergreen urban trees/forests in particular may be subjected to elevated [O$_3$] episodes in parallel to chilling, freezing and/or winter desiccation. Barnes et al. [159] investigated this topic using pea (*Pisum sativum*) as a model plant species. They found an amplification of freezing injury under elevated [O$_3$], including greater electrolyte leakage, that resulted in the deactivation of photochemical reactions of photosynthesis. Although these results are difficult to apply to woody plants that are well-adapted to winter conditions, these experiments revealed greater damage in pea cultivars with a slow stomatal closure under elevated [O$_3$] and having thus a greater O$_3$ flux in the internal leaf tissues.

Later research focused particularly on the interactive effects of elevated [O$_3$] and spring frost events [160,161]. While ozone (1.3 × ambient) exacerbated the effect of frost (applied after bud burst) only on diameter increment of Silver birch (*Betula pendula*), interactive effects on a number of physiological variables, including photochemical quantum yield of photosystem II, photochemical quenching of absorbed light energy and chlorophyll and carotenoid contents, were observed [149]. These findings lead to the conclusion that combined exposure to elevated [O$_3$] and frost appears to damage tree saplings more than a single stress. In contrary, no increase in freeze-induced damage of overwintering birch buds was reported under elevated [O$_3$] [161]. This is likely to be associated with an increased ratio of raffinose to sucrose and the transcript levels of the dehydrin gene. On the other hand, alterations in carbohydrate metabolism of buds is highly genome-specific and more work is needed [161].

4.3. Interactive Effect of Drought and O$_3$

Periods with high tropospheric [O$_3$] are often associated with hot and dry summers. Increased frequency, intensity and duration of such conditions are predicted in the future [162]. Accordingly, it is very likely that many tree species and forest stands are, and will be, simultaneously exposed to enhanced [O$_3$] and insufficient water supply. Some experiments have fortuitously coincided with naturally occurring drought. Matyssek et al. [70] describe a highly comprehensive free air experiment on adult beech (*Fagus sylvatica*) lasting five years, which included the extreme drought of 2003. Drought-induced stomatal closure decoupled the uptake of O$_3$ from [O$_3$], i.e., stomatal closure conferred a protective effect.
No significant interaction of O\textsubscript{3} and drought was detected in the growth and leaf traits of Japanese beech (Fagus crenata) during the first growing season, but significant interactive effects on photosynthesis were found in the second growing season\cite{163,164} indicating thus the importance of the stress duration. Limited water supply induced amelioration of the negative O\textsubscript{3} effects due to a reduced stomatal O\textsubscript{3} flux caused by reduced leaf water potential and stomatal conductance under the drought conditions\cite{165}. Similarly, reduced stomatal O\textsubscript{3} fluxes were observed in coniferous Norway spruce (Picea abies) under the periods of low soil humidity\cite{29,166}.

Metabolomic studies have shown that drought activates the pathways from glucose to xylulose, xylose, inositol and several polyols and sugar alcohols\cite{153}. Polyols, such as sorbitol and mannitol, act as osmoprotectants and protectants of thiol-regulated enzymes involved in deactivation of free radicals, such as hydroxyl radicals. These biochemical adjustments, together with an accumulation of glutathione antioxidant under limited soil water content\cite{164}, contribute to plant cross-tolerance against other oxidative stresses. Soil drought, and even more in combination with ozone, enhanced pools of ascorbate, lutein and xanthophylls in Norway spruce needles\cite{167}. Together with an increased epoxidation status of the xanthophyll cycle, an increased antioxidant capacity in plants is suggested. Accordingly, antagonistic effects of O\textsubscript{3} and drought on leaf photosynthetic traits are mostly reported in both broadleaved\cite{163,168} and coniferous\cite{169} tree species. However, antagonistic effects of O\textsubscript{3} and water stress on the growth are rare\cite{170} and are mostly additive\cite{171,172}. In conclusion, drought stress reduces but does not protect tree species from injurious O\textsubscript{3} impacts on growth and physiological processes associated with carbon assimilation\cite{173,174}. Most recently, Xu et al.\cite{175} have shown that limited water availability does not protect poplar saplings from O\textsubscript{3}-induced reduction of water use efficiency.

Modelling studies across diverse environmental conditions in Europe\cite{156,158} have shown that the arid conditions of South-European regions result in reduced stomatal conductance and consequent lower stomatal O\textsubscript{3} fluxes irrespective of the high AOT40s that prevail in these regions. Together with an enhanced stimulation of protective mechanisms against ROS\cite{153} injuries of Mediterranean tree species are less than those in trees grown under the conditions of temperate and/or boreal zones\cite{144,158}.

4.4. Interactive Effect of Nitrogen Supply and O\textsubscript{3}

Nitrogen availability is a limiting factor of forest production, particularly in N-poor boreal ecosystems (reviewed in Högberg et al.\cite{176}); however, excessive nitrogen deposition may lead to the imbalance between carbon and nitrogen in plant tissues having consequent negative effects on total content of phenolic compounds\cite{177}. A meta-analysis study\cite{178} revealed increased concentrations of phenolics and terpenes in response to elevated [O\textsubscript{3}]. Angiosperms were found to be more responsive than gymnosperms. Such acclimation consequently resulted in reduced insect herbivore activity. Insufficient amounts of phenolics under excessive nitrogen availability may lead, in contrast, to increased sensitivity of plants to abiotic and biotic stresses and consequently to reduced plant growth and development as shown in the case of Scots pine (Pinus sylvestris;\cite{179}). On the other hand, reduced nitrogen availability may increase plant sensitivity to O\textsubscript{3} because of the insufficient catalytic activities of enzymes involved in defence processes\cite{153,180}. However, the reported findings on interactive effects of elevated [O\textsubscript{3}] and nitrogen supply are inconsistent.

No interactive effect of nitrogen availability on stomatal O\textsubscript{3} flux and biomass production was reported in birch (Betula pendula) grown under a boreal environment\cite{181}, evergreen broadleaf camphorwood (Cinnamomum camphora) grown under sub-tropical regions\cite{182} or Quercus serrata, C. sieboldii, P. densiflora and C. japonica seedlings grown under temperate conditions (summarised in Watanabe et al.\cite{164}). On the other hand, significant interactions of O\textsubscript{3} and nitrogen load on dry-biomass increments were found in F. crenata and L. kaempferi\cite{164} enhancing the O\textsubscript{3} sensitivity of trees with increasing nitrogen supply. Similarly, Marzuoli et al.\cite{183} showed that increased wet N-deposition
triggers negative O$_3$ effects on biomass formation of young European hornbeam trees (*Carpinus betulus*). Also, Handley and Grulke [184] reported greater decline in photosynthetic capacity, defined as the CO$_2$ assimilation rate at saturating light intensity and [CO$_2$], in fertilised California black oak (*Quercus kelloggii*). These findings led to the conclusion that N-fertilisation tend to increase plant sensitivity to O$_3$.

It has been proven that reduced CO$_2$ assimilation rate under conditions of high [O$_3$] and nitrogen supply is rather caused by reduced allocation of nitrogen to soluble proteins, including the Rubisco enzyme, while nitrogen content per unit leaf area, stomatal conductance as well as mesophyll conductance to CO$_2$ diffusion remain unchanged [185,186].

The species-specificity in response to the combined effect of elevated [O$_3$] and nitrogen supply was reported by Fusaro et al. [187] for two Mediterranean tree species. Nitrogen supply ameliorated the harmful effects of O$_3$ in deciduous manna ash (*Fraxinus ornus*), but not in evergreen holm oak (*Quercus ilex*). It seems that nitrogen addition particularly increased activities of SOD (superoxide dismutase), CAT (catalase) and GSH (glutathione peroxidase) antioxidative enzymes involved in O$_2^-$ and H$_2$O$_2$ scavenging in *F. ornus*, while an alternative antioxidant response such as the emission of volatile organic compounds were preferentially activated in *Q. ilex*. Nevertheless, the activities of these antioxidative enzymes were not significantly influenced in spring wheat plants exposed only to elevated [O$_3$] [188].

5. Conclusions

What have we learned from fifty years of research? Experimental exposure of plants to high concentrations of ozone, or the exposure of plants to charcoal-filtered air, tell us that almost all species so far examined are damaged to some extent by ozone at the concentrations that often prevail in all regions of the world. This inevitably has a huge economic consequence for horticulture, agriculture and forestry, perhaps greater than that of any other pollutant gas [189]. Not all species are equally sensitive, and so the balance of competition between weeds and crops is likely to be altered.

The question of ‘how much damage’ is more difficult to answer, as the plant response depends not only on the species concerned but also on other stress factors. Drought and the supply of nitrogen are particularly variable across years and sites, and experiments have shown how these can interact with ozone to cause damage levels which can be hard to predict. The first attempts to evaluate the impact of crop losses were the NCLAN (National Crop Loss Assessment Network) in the USA [190] and the European Open Top Chamber Programme [191]. There were many further experiments in Asia. Feng and Kobashi [192] performed a meta-analysis of available data for key crops. The percentage crop losses were: potatoes 5.3, barley 8.9, wheat 9.7, rice 17.5, beans 19 and soybeans 7.7. These are probably still the best overall assessments we have of the overall ozone damage to crops. Ozone may well become a threat to food security as the O$_3$ precursors NO$_x$ and VOCs increase further due to human activities.

As we have seen in Section 4, the interaction of ozone with other environmental variables is often highly significant but varies with species and climatology; thus it is hard to predict and quite challenging to build a framework of regulations based on the concept of dose. In brief, high (non-excessive) irradiances, mild temperature increases and particularly drought tend to ameliorate negative effects of O$_3$ on plant physiology and growth, whereas freezing temperatures and high nitrogen deposition may increase plant sensitivity to O$_3$. Most recent studies discovered also the potential of O$_3$ to modulate the colonisation of roots by arbuscular mycorrhizal fungi [193] and herbivores attacks [194]—topics worth to be further investigate in trees.

For forests, the question of ‘how much damage?’ is harder to answer as the older work has been done on seedlings which may not be good surrogates for mature trees [71,106,195,196]. In some cases, tree seedlings and saplings have shown up to 40% less growth in ambient O$_3$ than when in charcoal filtered air [106]. FACE experiments have enabled limited enrichment experimentations on mature trees but the number of experimental units and the number of trees is inevitably rather small. Consequently, the statistical power of such experiments is low.
Long-term eddy-covariance, coupled to models, may be the way forward (see Section 2.3). The technology is well-developed and the limitations of the technique have been widely discussed [198]. Forest ecosystems are of special concern, because of the risk to the global carbon sink and the biodiversity that forests contain.

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