Biomass burning at Cape Grim: exploring photochemistry using multi-scale modelling

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Abstract. We have tested the ability of a high-resolution chemical transport model (CTM) to reproduce biomass burning (BB) plume strikes and ozone (O₃) enhancements observed at Cape Grim in Tasmania, Australia, from the Robbins Island fire. The CTM has also been used to explore the contribution of near-field BB emissions and background sources to O₃ observations under conditions of complex meteorology. Using atmospheric observations, we have tested model sensitivity to meteorology, BB emission factors (EFs) corresponding to low, medium, and high modified combustion efficiency (MCE), and spatial variability. The use of two different meteorological models (TAPM–CTM and CCAM–CTM) varied the first (BB1) plume strike time by up to 15 h and the duration of impact between 12 and 36 h, and it varied the second (BB2) plume duration between 50 and 57 h. Meteorology also had a large impact on simulated O₃, with one model (TAPM–CTM) simulating four periods of O₃ enhancement, while the other model (CCAM) simulating only one period. Varying the BB EFs, which in turn varied the non-methane organic compound (NMOC) / oxides of nitrogen (NOₓ) ratio, had a strongly non-linear impact on simulated O₃ concentration, with either destruction or production of O₃ predicted in different simulations. As shown in previous work (Lawson et al., 2015), minor rainfall events have the potential to significantly alter EF due to changes in combustion processes. Models that assume fixed EF for O₃ precursor species in an environment with temporally or spatially variable EF may be unable to simulate the behaviour of important species such as O₃.

TAPM–CTM is used to further explore the contribution of the Robbins Island fire to the observed O₃ enhancements during BB1 and BB2. Overall, TAPM–CTM suggests that the dominant source of O₃ observed at Cape Grim was aged urban air (age = 2 days), with a contribution of O₃ formed from local BB emissions.

This work shows the importance of assessing model sensitivity to meteorology and EF and the large impact these variables can have in particular on simulated destruction or production of O₃ in regional atmospheric chemistry simulations. This work also shows the importance of using models to elucidate the contribution from different sources to atmospheric composition, where this is difficult using observations alone.

1 Introduction

Biomass burning (BB) makes a major global contribution to atmospheric trace gases and particles, with ramifications for human health, air quality and climate. Directly emitted species include carbon monoxide (CO), carbon dioxide (CO₂), oxides of nitrogen (NOₓ), primary organic aerosol, non-methane organic compounds (NMOCs) and black carbon (BC), while chemical transformations occurring in the plume over time lead to formation of secondary species such as O₃, oxygenated NMOC and secondary aerosol. Depending on a number of factors, including magnitude and duration of fire, plume rise, and meteorology, the impact of BB plumes on human health, air quality and climate may be local, regional or global.
BB plumes from wildfires, prescribed burning, and agricultural and trash burning can have a major impact on air quality in both urban and rural centres (Keywood et al., 2015; Luhar et al., 2008; Reisen et al., 2011; Emmons et al., 2010; Yokelson et al., 2011) and regional-scale climate impacts (Andreae et al., 2002; Keywood et al., 2011b; Artaxo et al., 2013; Anderson et al., 2016). In Australia, BB from wild and prescribed fires impacts air quality in both rural and urban areas (Keywood et al., 2015, 2011a; Reisen et al., 2011; Luhar et al., 2008) as well as indoor air quality (Reisen et al., 2011). More generally, as human population density increases, and as wildfires become more frequent (Flannigan et al., 2009; Keywood et al., 2011b), assessing the impact of BB on air quality and human health becomes more urgent (Keywood et al., 2011b; Reisen et al., 2015). In particular, particles emitted from BB frequently lead to exceedances of air quality standards, and exposure to BB particles has been linked to poor health outcomes including respiratory effects, cardiovascular disease and mortality (Reisen et al., 2015; Reid et al., 2016; Dennekamp et al., 2015). There is also increasing evidence that mixing of BB emissions with urban emissions results in enhanced photochemistry and production of secondary aerosol and O₃, which have a non-linear relationship with emissions. Studies have found that modelled O₃ concentration from BB emissions is highly dependant on a range of factors including (a) meteorology (plume transport and dispersion) in global (Arnold et al., 2015) and high-resolution (Lei et al., 2013) Eulerian grid models, (b) absolute emissions and/or biomass burned (Pacifico et al., 2015; Parrington et al., 2012), (c) model grid size resulting in different degrees of plume dilution (Alvarado et al., 2009), and oxidative photochemical reaction mechanisms in Lagrangian parcel models (Mason et al., 2006).

To be able to accurately predict and assess the impact of BB on human health, air quality and climate, models must be able to realistically simulate the chemical and microphysical processes that occur in a plume as well as plume transport and dispersion. In the case of BB plumes close to an urban centre or other sensitive receptor, models can be used to mitigate risks on community by forecasting where and when a BB plume will impact, the concentrations of toxic trace gases and particles in the plume, and potential impact of the BB plume mixing with other sources. Models also allow investigation of the contributions from BB and other sources to observed air quality when multiple sources are contributing. Understanding the relative importance of different sources is required when formulating policy decisions to improve air quality.

Lagrangian parcel models are often used to investigate photochemical transformations in BB plumes as they are transported and diluted downwind (Jost et al., 2003; Trentmann et al., 2005; Mason et al., 2006; Alvarado and Prinn, 2009; Alvarado et al., 2015), while three-dimensional (3-D) Eulerian grid models have been used to investigate transport and dispersion of plumes, plume age, and contributions from different sources. Three-dimensional Eulerian grid models vary from fine spatial resolution on the order of a few kilometres (Luhar et al., 2008; Keywood et al., 2015; Alvarado et al., 2009; Lei et al., 2013) to a resolution of up to hundreds of kilometres in global models (Arnold et al., 2015; Parrington et al., 2012).

Sensitivity studies have allowed the influence of different model components (emissions, plume rise, transport, chemistry) on model output to be investigated. Such studies are particularly important in formation of secondary species such as O₃, which has a non-linear relationship with emissions. Studies have found that modelled O₃ concentration from BB emissions is highly dependant on a range of factors including (a) meteorology (plume transport and dispersion) in global (Arnold et al., 2015) and high-resolution (Lei et al., 2013) Eulerian grid models, (b) absolute emissions and/or biomass burned (Pacifico et al., 2015; Parrington et al., 2012), (c) model grid size resulting in different degrees of plume dilution (Alvarado et al., 2009), and oxidative photochemical reaction mechanisms in Lagrangian parcel models (Mason et al., 2006).

Broadly speaking, models used for simulating BB plumes comprise (a) a description of the emission source, (b) a determination of plume rise, (c) treatment of the vertical transport and dispersion and (d) a mechanism for simulating chemical transformations in the plume (Goodrick et al., 2013). There are challenges associated with accurately representing each of these components in BB modelling. The description of the emission sources includes a spatial and temporal description of the area burnt, the fuel load, combustion completeness, and trace gas and aerosol emission factors (EFs) (mass of species emitted per mass of fuel burned). The area burned is often determined by a combination of hotspot and fire scar data, determined from retrievals from satellite (Kaiser et al., 2012; Reid et al., 2009; Giglio et al., 2013). Cloud cover may lead to difficulties in obtaining area burnt data, while scars from small fires may be difficult to discern against complex terrain, and low intensity fires may not correspond with a detectable hotspot (Meyer et al., 2008). Emission factors are determined experimentally using either field or laboratory measurements and are typically grouped by biome type. In some regions, such as SE Australia, biomes have been sparsely characterised (Lawson et al., 2015). Furthermore, models use biome-averaged EFs, which do not account for complex intra-biome variation in EFs as a result of temporal and spatial differences in environmental variables. This includes factors such as impact of vegetation structure, monthly average rainfall (van Leeuwen and van der Werf, 2011) and the influence of short-term rainfall events (Lawson et al., 2015). For example, EFs have been shown to vary significantly with fuel moisture, which can vary seasonally (Korontzi et al., 2003; Urbanski, 2013). There may be significant spatial variability in EFs within a biome (Castellanos et al., 2014); taken along with temporal variability, this has been shown to have a large impact on simulated concentrations of BB species in global-scale modelling (van Leeuwen et al., 2013).

Finally, the very complex mixture of trace gases and aerosols in BB plumes creates analytical challenges in quantifying EFs, especially for semi-volatile and low-volatility organics, which are challenging to measure and identify but contribute significantly to secondary aerosol formation and
photochemistry within the plume (Alvarado and Prinn, 2009; Alvarado et al., 2015; Ortega et al., 2013).

Plume rise is a description of how high the buoyant smoke plume rises above the fire and consequently the initial vertical distribution of trace gases and aerosols in the plume (Freitas et al., 2007). This is still a large area of uncertainty in BB models, typically using a generalised plume rise approach, which may include either homogenous mixing, prescribed fractions of emissions distributed according to mixing height, use of parametisations, and finally plume rise calculated according to atmospheric dynamics. A key driver of this uncertainty is the complexity of fire behaviour, resulting in high spatial and temporal variability in pollutant and heat release, which drives variability in plume rise behaviour, such as multiple updraft cores (Goodrick et al., 2013).

Transport and dilution in models is driven by meteorology, particularly wind speed and direction, wind shear and atmospheric stability. Meteorology has a large impact on the ability of models to simulate the timing and magnitude and even composition of BB plume impacts in both local- and regional-scale models (Lei et al., 2013; Luhar et al., 2008; Arnold et al., 2015). For example, too-high wind speeds can lead to modelled pollutant levels that are lower than observed (e.g. Lei et al., 2013), while small deviations in wind direction lead to large concentration differences between modelled and observed levels, particularly when modelling emissions of multiple spatially diverse fires (Luhar et al., 2008). Dilution of BB emissions in large grid boxes in global models may also lead to discrepancies between modelled and observed NO\textsubscript{x}, O\textsubscript{3} and aerosols (Alvarado et al., 2009).

Finally, models use a variety of gas-phase and aerosol-phase physical and chemical schemes, which vary in their ability to accurately represent chemical transformations, including formation of O\textsubscript{3} and organic aerosol (Alvarado and Prinn, 2009; Alvarado et al., 2015). Validating and constraining chemical transformations in models requires high quality, high-time-resolution BB observations of a wide range of trace gas and aerosol species, including important but infrequently measured species such as OH and semi-volatile and low-volatility NMOCs. Field observations, whilst often temporally and spatially scarce, are particularly valuable because the processes and products of BB plume processing are dependent on long-range transport, cloud processing, varying meteorological conditions and heterogeneous reactions.

In this work we test the ability of CSIRO’s high-resolution 3-D Eulerian grid chemical transport model (CTM) to reproduce BB plume observations of the Robbins Island fire reported in Lawson et al. (2015), with a focus on CO, BC and O\textsubscript{3}. We undertake sensitivity studies using varying emission factors associated with a low, medium and high modified combustion efficiency (MCE), which in turn changes the NMOC/NO\textsubscript{x} ratio, in contrast to other sensitivity studies that typically scale emissions of all species by a constant factor (Pacifico et al., 2015; Lei et al., 2013). We also test sensitivity to meteorology by coupling the CTM with two different meteorological models, The Air Pollution Model (TAPM) and CSIRO’s Conformal Cubic Atmospheric Model (CCAM). The fire and fixed observation sites (Cape Grim) were only 20 km apart, and so simulation of the plume strikes is a stringent test of TAPM and CCAM’s ability to reproduce wind speed and direction. Plume rise and chemical mechanism are held constant. Finally, we use TAPM–CTM to separate the contribution of the Robbins Island fire emissions and urban emissions to the observed O\textsubscript{3} enhancements at Cape Grim reported in Lawson et al. (2015) and to determine the age of the O\textsubscript{3}-enhanced air parcels.

2 Methods

2.1 Fire and measurement details

Details of the fire and measurements are given in Lawson et al. (2015). Briefly, BB plumes were measured at the Cape Grim Baseline Air Pollution Station during the 2006 Precursors to Particles campaign, when emissions from a fire on nearby Robbins Island impacted the station. Fire burned through native heathland and pasture grass on Robbins Island some 20 km to the east of Cape Grim for 2 weeks in February 2006. On two occasions an easterly wind advected the BB plume directly to the Cape Grim Station. The first plume strike (BB1) occurred from 02:00 to 06:00 (Australian eastern standard time – AEST) on 16 February, with light easterly winds of 3 m s\textsuperscript{-1}, a temperature of 13 °C and a relative humidity (RH) of 96 %. The second, more prolonged plume strike (BB2) occurred from 23:00 on 23 February to 05:00 on 25 February, with strong easterly winds ranging from 10 to 16 m s\textsuperscript{-1}, temperatures of 16–22 °C and RH in the range of 75–95 %. Under a northerly wind direction, urban air from the city of Melbourne (population 4.2 million) some 300 km away is transported across the ocean (Bass Strait) to Cape Grim.

A wide variety of trace gas and aerosol measurements were made during the fire event (Lawson et al., 2015). In this work, measurements of black carbon (BC), carbon monoxide (CO) and ozone (O\textsubscript{3}) are compared with model output. BC measurements were made using an Aethalometer (Gras, 2007), CO measurements were made using an AGAGE gas chromatography system with a multi-detector (Krummel et al., 2007) and O\textsubscript{3} measurements were made using a TECO analyser (Galbally et al., 2007). For further details see Lawson et al. (2015).

2.2 Chemical transport models

Simulations were undertaken with CSIRO’s CTM, coupled offline with two meteorological models (see below). The CSIRO CTM is a three-dimensional Eulerian CTM with the capability of modelling the emission, transport, chemical transformation, wet and dry deposition of a coupled gas- and aerosol-phase atmospheric system. The CTM was initially
developed for air quality forecasting (Cope et al., 2004) and has had extensive use with shipping emission simulations (Broome et al., 2016), urban air quality (Cope et al., 2014; Galbally et al., 2008), and biogenic (Emmerson et al., 2016) and biomass burning studies (Keywood et al., 2015; Meyer et al., 2008; Luhar et al., 2008).

The chemical transformation of gas-phase species was modelled using an extended version of the Carbon Bond 5 mechanism (Sarwar et al., 2008) with updated toluene chemistry (Sarwar et al., 2011). The mechanism was also extended to include the gas-phase precursors for secondary (gas and aqueous phase) inorganic and organic aerosols. Secondary inorganic aerosols were assumed to exist in thermodynamic equilibrium with gas-phase precursors and were modelled using the ISORROPIA-II model (Fountoukis and Nenes, 2007). Secondary organic aerosol was modelled using the volatility basis set (VBS) approach (Donahue et al., 2006). The VBS configuration is similar to that described in Tsipmidis et al. (2010). The production of S-VI in cloud water was modelled using the approach described in Seinfeld and Pandis (1998). The boundary concentrations in the models for different wind directions were informed by Cape Grim observations of atmospheric constituents during non-BB periods (Lawson et al., 2015). In this work the modelled elemental carbon (EC) output was considered equivalent to the BC measured with an Aethalometer at Cape Grim.

Horizontal diffusion is simulated according to equations detailed in Cope et al. (2009) according to the principles of Smagorinsky et al. (1963) and Hess (1989). Vertical diffusion is simulated according to equations detailed in Cope et al. (2009) according to the principles of Draxler and Hess (1997). Horizontal and vertical advection use the approach of Walcek et al. (2000).

2.2.1 Meteorological models

Prognostic meteorological modelling was used for the prediction of meteorological fields including wind velocity, temperature, water vapour mixing ratio and clouds, radiation, and turbulence. The meteorological fields force key components of the emissions and the chemical transport model. Two meteorological models were used in this work. CSIRO’s TAPM (Hurley, 2008a, b), a limited area, nestable, three-dimensional Eulerian numerical weather and air quality prediction system, and CSIRO’s CCAM, a global stretched-grid atmospheric simulation model (McGregor, 2015, and references therein). The models represent two unique (and independent) approaches for generating the meteorological fields required by the CTM.

For CCAM, 20 km spaced simulations over Australia were used by the CTM (with the same grid spacing) to model large-scale processes on the continent, including the emission and transport of windblown dust, sea salt aerosol and smoke from wildfires. Note that the governing equations for TAPM do not enable this model to simulate spatial scales greater than 1000 km in the horizontal and thus only the CCAM meteorology was available for the continental-scale simulations. TAPM and CCAM 12 km spaced simulations were then used to model the transport of the Melbourne plume to Cape Grim using the CTM (at 12 km grid spacing) with boundary conditions provided by the continental simulation. Nested grid simulations by the CTM at 3 and 1 km grid spacing utilised TAPM and CCAM meteorology simulated at matching grid spacing. The 1 km spaced meteorological fields were also used to drive a 400 m spaced CTM domain that encompassed Robbins Island and Cape Grim. This domain was included in the nested grid system because we wanted to better numerically resolve the spatial extent of the fire and the process of plume advection between Robbins Island and Cape Grim. Figure 1 shows the five nested computational domains used in TAPM–CTM and CCAM–CTM.

In this work the CTM coupled with the CCAM meteorological model is referred to as CCAM–CTM, while the CTM coupled with the TAPM meteorological model is referred to as TAPM–CTM.

Figure 1. The five nested computational domains used in TAPM–CTM and CCAM–CTM, which had cell spacings of 20, 12, 3, and 1 km and 400 m.
2.2.2 Emission inventories

**Anthropogenic emissions**

Anthropogenic emissions for Victoria were based on the work of Delaney et al. (2011). No anthropogenic emissions were included for Tasmania. The north-west section of Tasmania has limited habitation and is mainly farmland, and thus the influence of Tasmanian anthropogenic emissions on Cape Grim are expected to be negligible.

**Natural and biogenic emissions**

The modelling framework includes methodologies for estimating emissions of sea salt aerosol (Gong, 2003), emissions of windblown dust (Lu and Shao, 1999), gaseous and aerosol emissions from managed and unmanaged wild fires (Meyer et al., 2008), emissions of NMOCs from vegetation (Auzzi et al., 2012), and emissions of nitric oxide and ammonia from vegetation and soils. Emissions from all but the wildfires are calculated inline in the CTM at each time step using the current meteorological fields. There were no other major fires burning in Victoria and Tasmania during the study period.

**Emissions – Robbins Island fire**

The area burnt by the fire was determined from hotspots from the Sentinel product (Geosciences Australia), which were derived from MODIS imagery. The hotspots were buffered to give polygon spots at a resolution of 400 ha spot$^{-1}$, then merged into a single polygon for each fire day (Meyer et al., 2008). The fire burnt 2000 ha over the 2-week period, and the direction of fire spread was unknown. As such, the fire scar was divided up into 250 m grids and the hourly areas burnt were calculated using a normalised version of the McArthur Forest Fire Danger Index (FFDI) (Meyer et al., 2008). The models assumed that an equal proportion of each grid burned simultaneously over the 2-week period. The fuel density used was estimated to be 18.7 tC ha$^{-1}$, based on mean mass loads of coarse and fine fuels taken from the biogeochemical production model (VAST 1.2; Barrett, 2002) and converted into carbon mass (Meyer et al., 2008).

The hourly diurnal emissions of all gases and particles from the fire were calculated using the FFDI in which the presence of strong winds will result in faster fire spread and enhanced emissions, compared to periods of lower wind speeds. The effect of wind speed on the fire behaviour and emissions is particularly important during BB2 in which the winds ranged from 10 to 15 m s$^{-1}$. This is evident from Fig. 2 in which hourly emission profiles based on an average diurnal FFDI calculated by Meyer et al. (2008) (which peaks early afternoon) is compared with profiles based on hourly FFDIs generated by TAPM and CCAM meteorology. It can be seen that the use of the dynamic FFDI approach during the BB2 period increases the base emissions by 70% for TAPM meteorology and by 45% for the CCAM meteorology. It is also notable that the use of the dynamic approach with TAPM meteorology leads to the peak emissions occurring overnight on 24 February, which is when the base emissions are at a minimum.

Three different sets of fire EFs, corresponding to low, medium and high MCE, were used to test the sensitivity of the models, where MCE $= \Delta$CO$_2$/\DeltaCO + \DeltaCO$_2$ (Ferek et al., 1998). We used published EFs of CO and CO$_2$ from temperate forests (Akagi et al., 2011) to calculate a typical range of MCEs for temperate fires, including an average (best estimate) of 0.92 and a lower (0.89) and upper estimate (0.95). Fires with MCEs of approximately 0.90 consume biomass with approximately equal amounts of smouldering and flaming, while MCEs of 0.99 indicate complete flaming combustion (Akagi et al., 2011). Therefore, the calculated range of MCEs (0.89–0.95) corresponds to fires in which both smouldering and flaming occurs, with a tendency for more flaming combustion in the upper estimate (0.95) compared to a tendency for more smouldering in the lower estimate (0.89).

In previous smoke modelling work, CCAM–CTM and TAPM–CTM used savannah EFs from Andreae and Merlet (2001). However, as Robbins Island is in a temperate region, the Andreae and Merlet (2001) savannah EFs used in the models were adjusted to reflect temperate EFs based on the following methodology. Minimum, mean and maximum CO EFs for temperate forests from Agaki et al. (2011) were used for lower, best-estimate and upper MCEs. For all other species, savannah EFs (corresponding to an MCE of 0.94) were adjusted to EFs for the lower, best-estimate and upper MCEs using published relationships between MCE and EF (Meyer et al., 2012; Yokelson et al., 2007, 2003, 2011).

For example, to adjust the Andreae and Merlet (2001) savannah EFs (corresponding to an MCE of 0.94) to our temperate best-estimate EF (corresponding to an MCE of 0.92) the Andreae and Merlet (2001) NO EF was reduced by 30%, the NMOC EFs were increased by 30%, the BC EF was reduced by 30% and the OC EF was increased by 20%. Table 1 gives EFs for the original savannah EF (Andreae and Merlet, 2001) and the adjusted EF used in this work. The NO$_x$ / NMOC ratios used are also shown and vary by a factor of 3 between the low- and high-MCE scenarios, mainly driven by the variability in NO emissions with MCE. The EF calculated from observations for this fire are shown for comparison (Lawson et al., 2015).

We recognise that calculating EF in this way is approximate; however, the purpose of including a range of EFs was to explore the model’s sensitivity to EF. While EFs were calculated for the Robbins Island fire for several species (Lawson et al., 2015), these are only available for a subset of species required by the CB05 chemical mechanism. The adjustment of the Andreae and Merlet (2001) savannah EFs to a lower MCE (0.89) resulted in good (±20%) agreement with the calculated EFs for CO, BC and several NMOCs from Lawson et al. (2015), in which the MCE was calcu-
With respect to plume rise, the Robbins Island fire was a relatively low-energy burn (Lawson et al., 2015), and as noted by Paugam et al. (2016), the smoke from such fires is largely contained within the planetary boundary layer (PBL). Given that ground-based images of the Robbins Island smoke plume support this hypothesis, in this work we adopted a simple approach of mixing the emitted smoke uniformly into the model’s layers contained within the PBL. The plume was well mixed between the maximum of the PBL height and 200 m above the ground, with the latter included to account for some vertical mixing of the buoyant smoke plume even under conditions of very low PBL height. The high wind speeds, particularly during the second BB event, also suggest that the plume was not likely to be sufficiently buoyant to penetrate the PBL.

3 Results and discussion

3.1 Modelling sensitivity study

The ability of the models to reproduce the two plume strikes (BB1 and BB2; described in Lawson et al., 2015) was tested. The period examined was 13 February 2006 to 28 February 2006. The sensitivity of the models to meteorology, EFs and spatial variability was also investigated and is discussed below. Observational and model data shown are hourly averages. Table 2 summarises the main findings of the model sensitivity study. A MODIS True Color Aqua image of the Robbins Island fire plume is shown in Fig. 3 from 23 February 2006, with the modelled plume during the same period.

3.1.1 Sensitivity of modelled BB species to meteorology

Qualitative and quantitative assessments of model performance for meteorological parameters were undertaken for both TAPM and CCAM. Hourly observed and modelled winds, temperature, humidity and PBL are compared and discussed in the Supplement (Figs. S2–S8). Briefly, both TAPM
Table 1. EF used in model sensitivity studies, corresponding to low (MCE = 0.89), medium (MCE = 0.92) and high (MCE = 0.95) MCEs. A subset of the total species included in the CB05 lumped chemical mechanism are shown. Also shown are savannah EFs from Andreae and Merlet (2001) (A&M) and EFs calculated from BB2 in previous work (Lawson et al., 2015). NO: nitric oxide, CO: carbon monoxide, PAR: paraffin carbon bond, OLE: terminal olefin carbon bond, TOL: toluene and other monoalkyl aromatics, XYL: xylene and other polyalkyl aromatics, BNZ: benzene, FORM: formaldehyde, ALD2: acetaldehyde, EC: elemental carbon < 10 µm and OC: primary organic carbon < 10 µm.

<table>
<thead>
<tr>
<th>Species</th>
<th>A&amp;M (2001) MCE 0.94</th>
<th>Lawson et al. (2015) MCE 0.88</th>
<th>Used in this work MCE 0.89</th>
<th>MCE 0.92</th>
<th>MCE 0.95</th>
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<td>NO</td>
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<td>NA</td>
<td>0.8</td>
<td>2.7</td>
<td>4.7</td>
</tr>
<tr>
<td>CO</td>
<td>65</td>
<td>127</td>
<td>121</td>
<td>89</td>
<td>57</td>
</tr>
<tr>
<td>PAR</td>
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<td>2.02</td>
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</tr>
<tr>
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<td>0.06</td>
<td>0.04</td>
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<td>11.99</td>
<td>2.97</td>
<td>1.20</td>
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NA: not available.

Table 2. Summary of sensitivity study results, including meteorology, emission factors and spatial variability.

<table>
<thead>
<tr>
<th>Sensitivity study</th>
<th>Species</th>
<th>TAPM–CTM simulation</th>
<th>CCAM–CTM simulation</th>
<th>Comments/drivers of model outputs</th>
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<tr>
<td>Meteorology</td>
<td>BC and CO</td>
<td>BB1 plume strike +3 h Duration 12 h (actual 5 h)</td>
<td>BB1 plume strike −12 h Duration 36 h intermittent (actual 5 h)</td>
<td>Narrow BB plume. Differences in plume strike due to timing of wind direction change; wind speeds; direct or indirect advection of plume over Cape Grim</td>
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<td></td>
<td>BB2 plume strike −26 h Duration 50 h (actual 29 h)</td>
<td>BB2 plume strike −26 h Duration 57 h (actual 29 h)</td>
<td>Wind direction differences driven by gravity wave oscillations; timing of wind direction change; different wind speeds driving absolute BB emissions and plume dispersion</td>
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<tr>
<td>O3</td>
<td></td>
<td>Four O3 peaks simulated (two observed, two not)</td>
<td>One O3 peak simulated (observed)</td>
<td>Differences in simulated wind speed and direction (and EF – see below)</td>
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<tr>
<td>Emission factors</td>
<td>BC and CO</td>
<td>BC peak magnitude varies by factor 3, CO factor 2 with different EF runs</td>
<td>As for TAPM -CTM</td>
<td>Concentrations vary according to EF input ratios.</td>
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<tr>
<td></td>
<td>O3</td>
<td>Two peaks with high EF sensitivity, two peaks with no EF sensitivity</td>
<td>One peak with no EF sensitivity</td>
<td>Different NMOC / NOx emission ratios (varies with MCE) drive destruction or production of O3 in fire-related peaks. MCE 0.89 TAPM–CTM simulation gives best agreement with observations</td>
</tr>
<tr>
<td>Spatial variability</td>
<td>CO</td>
<td>Differences of up to &gt; 500 ppb in grid points 1 km apart (BB2)</td>
<td>NA</td>
<td>Narrow BB plume</td>
</tr>
<tr>
<td></td>
<td>O3</td>
<td>Differences of up to 15 ppb in grid points 1 km apart (BB1)</td>
<td>n/a</td>
<td>Narrow ozone plume generated downwind of fire</td>
</tr>
</tbody>
</table>
Figure 4. Model concentration isopleth of BC for TAPM–CTM (left panels) and CCAM–CTM (right panels). Panels show 12 h time intervals during BB1, including the BB plume intermittently striking Cape Grim, followed by the change in plume direction with wind direction change. Arrows are wind vectors. The time series of observed and modelled wind direction for BB1 are shown above with golden bands highlighting the periods corresponding to the panels.
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Figure 5. Model concentration isopleth of BC for TAPM–CTM and CCAM–CTM at 05:00 on 24 February during BB2. Arrows are wind vectors. The time series of observed and modelled wind direction for BB2 is shown above with a golden band highlighting the period corresponding to the panels.

and CCAM demonstrated reasonable skill in modelling the meteorological conditions, with the TAPM simulations being slightly better than the CCAM simulations with respect to the low-level wind, temperatures and RH and CCAM simulations being slightly better in terms of PBL height.

Primary species – CO and BC

Figures 4 and 5 show concentration isopleths for BC generated by TAPM–CTM and CCAM–CTM for BB1 and BB2 respectively. The simulated and observed time series concentrations of CO and BC for the two different models (TAPM–CTM and CCAM–CTM) and for three different sets of EFs (discussed in Sect. 3.1.2) are shown in Fig. 6. TAPM–CTM and CCAM–CTM both reproduce the observed plume strikes (BB1 and BB2). The impact of meteorology on the plume strike timing and duration is discussed below.

Both models overestimate the duration of BB2 and simulate the plume strike occurring earlier than observed. TAPM–CTM predicts that BB2 is 26 h earlier than observed and that BB2 persists for 50 h (observed duration 29 h). CCAM–CTM predicts that BB2 is 26 h earlier than observed and that BB2 persists for 57 h. It should be noted that there is a brief observed enhancement of BB species that corresponds with the beginning of the modelled BB2 plume strike, some 24 h prior to the prolonged observed event. This was likely due to the edge of the plume impacting the station briefly.

In both observed BB1 and BB2 the plume strike at Cape Grim occurred just prior to a wind direction change from easterly (fire direction) to south-westerly. The timing of the wind direction change in the models is therefore crucial to correctly predicting plume strike time and duration. In BB1 CCAM predicts an earlier wind direction change with higher wind speeds, which advects the plume directly over Cape Grim, while TAPM predicts a later wind change, lower wind speeds and advection of only the edge of the plume over Cape Grim. The higher concentrations CO and BC in BB1 predicted by CCAM–CTM are likely due to the direct advection of the plume over the site compared to only the plume edge in TAPM–CTM.

In BB2, both TAPM–CTM and CCAM–CTM predict direct strikes of the Robbins Island smoke plume on Cape Grim because the wind direction is modelled to be predominantly easterly for the duration of the event (Fig. 5). Both models simulate some backing and veering of the wind direction
for the duration of BB2 due to gravity wave processes that lead to intermittent strikes on Cape Grim as the Robbins Island smoke plume sweeps to the north and south of Cape Grim. The gravity wave oscillations are more pronounced in CCAM–CTM than TAPM–CTM (and thus the plume strikes are more pronounced from the former) due to differences in how the models are coupled to large-scale synoptic forcing. The event is eventually curtailed by the passage of a south-westerly change.

Figure 5 shows that TAPM–CTM predicts the onset of the change to occur about 6 h ahead of the observed change and thus the BB2 event ends too early for this meteorological simulation. CCAM–CTM models the south-westerly change to occur 1 h after observed, leading to the modelled BB2 event extending beyond the observed duration for this meteorological simulation.

Differences in the magnitude of the modelled CO and BC peaks for TAPM–CTM and CCAM–CTM have two principal causes: (a) the coupling of the smoke emissions to the TAPM and CCAM meteorology via the FFDI scaling leads to approximately 20 % higher emissions in the case of the TAPM–CTM simulations; (b) the CCAM wind speeds are 20–50 % higher than the TAPM wind speeds during BB2, which in combination with the emission differences, leads to TAPM–CTM generating near-surface smoke concentrations that are up to 80 % higher than CCAM–CTM. Mixing depth can also play an important role in plume dispersion; however, the PBL heights generated by both models are similar and generally low during BB2 due to the easterly wind direction and the mainly maritime upwind fetch.

Secondary species – O3

Figure 6e and f show the simulated and actual O3 concentration time series for TAPM–CTM and CCAM–CTM for three different sets of EFs (discussed in Sect. 3.1.2). The two observed O3 peaks that followed BB1 and BB2 can clearly be seen in the time series of observations. Figure 7 shows the TAPM–CTM and CCAM–CTM concentration isopleths of O3 enhancement downwind of the fire during BB1 at 11:00 and 13:00 on 16 February.

Again the simulated meteorology has a major impact on the ability of the models to reproduce the magnitude and timing of the observed O3 peaks. TAPM–CTM reproduces the major O3 peak observed following BB2 and captures part of the O3 peak following BB1. For the peak following BB1 it underpredicts the peak duration and fails to capture the subsequent observed peaks on 18 and 19 February. TAPM–CTM also shows two additional O3 peaks, which were not observed, about 24 h prior to the BB1 and BB2 peaks. The magnitude of these additional peaks shows a strong dependency on the EF, suggesting an influence of fire emissions. This is discussed further below and in Sect. 3.2.1. Compared to TAPM–CTM, CCAM–CTM predicts fewer distinct peaks of ozone above the background (where background is 15–17 ppb) throughout the entire period. Both TAPM–CTM and CCAM–CTM show depletion of O3 below background levels, which was not observed, and this is discussed further in Sect. 3.1.2.

Figure 7 shows that there are differences in wind fields between TAPM–CTM and CCAM–CTM as well as different simulated concentrations of O3 generated from the fire. This is discussed further in Sect. 3.1.2. To summarise, the impact of using two different meteorological models for a primary species such as BC was to vary the modelled time of impact of the BB1 plume strike by up to 15 h (CCAM–CTM −12 h and TAPM–CTM +3 h, with actual plume strike time = 0 h) and to vary the plume duration between 12 and 36 h (actual duration 5 h). For BB2, different meteorological models predicted the same impact time (TAPM–CTM and CCAM–CTM both −26 h, with the actual plume strike time = 0 h) and showed the plume duration to vary between 47 and 60 h (actual duration 29 h).

For O3, the use of different meteorological models led to one model (TAPM–CTM) reproducing both observed peaks plus two additional peaks, while the other model (CCAM–CTM) captured only one defined O3 peak over the time series of 2 weeks.

3.1.2 Sensitivity of modelled BB species to emission factors

Primary species – CO and BC

Figure 6a–d show the simulated and observed concentrations of BC and CO for MCE = 0.89, MCE = 0.92 and MCE = 0.95 (see Sect. 2.2.2). Because CO has a negative relationship with MCE, and BC has a positive relationship with MCE, the modelled BC concentrations are highest for model runs using the highest MCE, while the modelled CO concentrations are highest for model runs using the lowest MCE (Fig. 6).

Changing the EF from low to high MCE varies the modelled BC concentrations during BB1 and BB2 by a factor of ∼ 3 for BC and a factor of ∼ 2 for CO and increases the EF ratio of BC / CO by a factor of ∼ 6, in proportion to the difference in EF input to the models.

Quantile–quantile plots of observed and modelled ratios of BC / CO during BB1 and BB2 for the different EF scenarios are shown in Fig. 8. BC / CO ratios were used to minimise uncertainty resulting from errors in modelling transport and dilution (and mixing height), thus enabling a focus on the impact of EF variability. A period incorporating both the modelled and observed BB1 and BB2 was used for the analysis. The TAPM–CTM simulation with MCE = 0.89 performed best, with greater than 60 % of the model percentiles falling within a factor of 2 of the observed. The CCAM–CTM simulation with MCE = 0.89 was the second best performer with 50 % of the modelled percentiles falling within a factor of 2 of the observed. Overestimates of the EC / CO ratio by up to
Figure 6. Simulated CO using (a) TAPM–CTM and (b) CCAM–CTM, simulated BC using (c) TAPM–CTM and (d) CCAM–CTM, and simulated O\textsubscript{3} using (e) TAPM–CTM and (f) CCAM–CTM. Coloured lines represent different MCE EF simulations; black symbols are observations.

A factor of 8 occur for some percentiles for the MCE = 0.95 scenarios, while the scenarios with no fire significantly underestimated the observed ratio. Plots of mean fractional bias and mean fractional error (Figs. S11 and S12) show that the TAPM–CTM simulation with MCE = 0.89 has the smallest bias and error, followed by the CCAM–CTM simulation with MCE = 0.89. As discussed previously, there is uncertainty in the derivation of EF as a function of MCE, as this was based on relationships from a small number of studies. Nevertheless, the percentile, bias and error analysis indicate that using EFs corresponding to an MCE of 0.89 gives the best agreement with the observations for the BC/CO ratio. This is in agreement with the calculated MCE of 0.88 for this fire (Lawson et al., 2015).

Secondary species – O\textsubscript{3}

For secondary species such as O\textsubscript{3} (Fig. 6e and f), the relationship between EF precursor gases and model output is more complex than for primary species such as CO and BC because the balance between O\textsubscript{3} formation and destruction is dependent on the degree of dilution of the BB emissions and also factors such as the NMOC composition and the NMOC/NO\textsubscript{x} ratio.

TAPM–CTM (Fig. 6e) reproduces the magnitude of both observed peaks following BB1 and BB2 (BB1 maximum observed = 33 ppb, modelled = 31 ppb; BB2 maximum observed = 34 ppb, modelled = 30 ppb). Interestingly, the magnitude of O\textsubscript{3} for these two peaks is the same for different EF inputs of O\textsubscript{3} precursors from the Robbins Island fire, sug-
gesting that the BB emissions are not responsible for these enhancements as demonstrated in Sect. 3.2. In contrast, the two additional peaks modelled but not seen in the observations are heavily dependent on the input EF. For the first additional modelled peak, which was predicted at the time of BB1 observations on 16 February, all EF scenarios result in an O$_3$ peak, with the MCE = 0.92 model scenario resulting in the highest predicted O$_3$. For the second additional modelled peak just prior to the BB2 observations on 23 February, only the MCE = 0.89 scenario results in net O$_3$ production, while MCE = 0.92 and MCE = 0.95 scenarios lead to net O$_3$ destruction.

This differing response to EF for the TAPM–CTM runs suggests the importance of the NO EF for O$_3$ production in BB plumes. Unfortunately, there were no oxides of nitrogen measurements made during the fire to test the models. For the first simulated additional peak prior to BB1, while the medium NO EF (MCE = 0.92) resulted in the highest O$_3$ peak (with corresponding NO of 3.7 ppb and NO$_2$ of 4.5 ppb), the lower NO EF in the 0.89 MCE run perhaps indicates that insufficient NO was present to drive O$_3$ production (corresponding NO of 0.5 ppb and NO$_2$ of 1.5 ppb), which is in line with studies that have shown that BB plumes are generally NO$_x$ limited (Akagi et al., 2013; Jaffe and Wigder, 2012; Wigder et al., 2013). Conversely, the highest input NO EF (MCE = 0.95) led to net destruction of O$_3$ (NO of 9 ppb, NO$_2$ of 7 ppb), which is due to titration of O$_3$ with the larger amounts of NO emitted from the fire in these runs as indicated by excess NO (NO/NO$_2$ ratio > 1) at Cape Grim (where NO has a positive relationship with MCE). For the second additional peak prior to BB2, only the lowest NO EF run (MCE = 0.89) resulted in net production of O$_3$ (NO of 1.5 ppb, NO$_2$ of 2.6 ppb). In the medium- and high-MCE runs, the background O$_3$ concentration is completely titrated (0 ppb), with NO concentrations of 10 and 20 ppb and NO/NO$_2$ ratios of 1.3 and 2.6 respectively.

In contrast, the CCAM–CTM model (Fig. 6f) simulations reproduce only the first observed O$_3$ peak associated with BB1 (modelled = 27 ppb, measured = 34 ppb). This modelled O$_3$ peak does not show an influence of MCE on O$_3$ con-

Figure 7. Model concentration isopleth showing O$_3$ enhancement downwind of the fire during BB1 at 11:00 and 13:00 on 16 February for (a) TAPM–CTM and (b) CCAM–CTM. The spatially variable plume and complex wind fields are shown. Arrows are wind vectors.
concentration, in agreement with TAPM, again suggesting no influence from fire emissions as later demonstrated in Sect. 3.2. The CCAM model runs also show significant titration of O$_3$ during BB1 and BB2 for the medium- and high-MCE model runs, with $\sim$ 24 and $\sim$ 48 h of significant O$_3$ depletion below background concentrations being modelled for each event, which was not observed.

Quantile–quantile plots of modelled and observed concentrations of O$_3$ for all EF scenarios are shown in Figs. 9 and S13. Model performance was assessed for both the BB and the background periods in order to test the ability of the models to reproduce O$_3$ from both the fire and other sources, including urban sources. The modelled O$_3$ concentrations from the TAPM–CTM simulation with MCE = 0.89 are close to the 1 : 1 line with observations for all of the sampled percentiles and demonstrate that this scenario is in best agreement with observations. As stated previously, it is also in agreement with the calculated MCE of 0.88 for BB2 (Lawson et al., 2015). Ozone titration, which was not observed, in the MCE = 0.92 and MCE = 0.95 scenarios is visible as a significant deviation from the 1 : 1 line in Fig. 9. With the exception of these titration events, all of the sampled model concentration percentiles fall within a factor of 2 of the observations. Plots of mean fractional error and mean fractional bias (Figs. S14 and S15) show that the error and bias are very low for all runs and fall within performance guidelines.

The different EF scenarios presented here suggest that varying model EFs have a major impact on whether the models simulate production or destruction of O$_3$, particularly important at a receptor site in close proximity to the BB emissions. In the previous work (Lawson et al., 2015), the MCE for the first 10 h of BB2 was calculated as 0.88. However, later in BB2, a rainfall event led to changes in the NMOC / CO and BC / CO ratios. This suggests that during the course of BB2 the MCE decreased and thus EFs changed. As such, the use of fixed BB EFs in this work and in other models may lead to incorrect prediction of important species such as O$_3$.

3.1.3 Sensitivity of modelled concentrations to spatial variability

The near-field proximity of the Robbins Island fire (20 km) to Cape Grim, the narrowness of the BB plume and the spatial complexity of the modelled wind fields around north Tasmania are likely to result in strong heterogeneity in the modelled concentrations surrounding Cape Grim. We investigated how much model spatial gradients vary by sampling TAPM–CTM output with MCE = 0.89 at four grid points located 1 km to the north, east, south and west of Cape Grim.
Primary species – CO

Figure 10a shows a time series of the modelled CO output of the difference between Cape Grim and each grid point 1 km either side.

The plotted CO concentration is equal to the other location [CO] (N, S, E, W) – Cape Grim [CO].

The figure clearly shows that there are some large differences in the modelled concentrations of CO between grid points for both BB1 and BB2. Particularly large differences were seen for BB2, with the north grid point modelled concentrations in BB2 over 500 ppb lower than at the Cape Grim grid point, while at the southerly grid point the modelled CO was up to 350 ppb higher. Smaller differences of up to 250 ppb between the east and Cape Grim grid points were observed for BB1. This indicates that the plume from the fire was narrow and had a highly variably impact on the area immediately surrounding Cape Grim.

Figure 10b shows the observed cumulative concentration of CO over the 29 h duration of BB2 at Cape Grim, as well as the modelled cumulative concentration at Cape Grim and at the four grid points either side. This figure shows the variability in concentration with both location and time. TAPM–CTM’s underestimation of the observed CO is visible by hour 20. TAPM–CTM begins to show differences in modelled cumulative CO concentrations between the five grid points (including Cape Grim) by hour 10. At the end of BB2, TAPM–CTM predicts that there are differences of 5–30 % between the cumulative modelled CO concentration at Cape Grim and the grid points to the north, east, south and west. This variability modelled between sites that are closely located highlights the challenges with modelling the impact of a near-field fire at a fixed single-point location. This also highlights the high spatial variability that may be missed in similar situations by using a coarser resolution model, which would dilute emissions in a larger grid box.

Ozone (O₃)

Figure 10c shows a time series of the modelled O₃ output of the difference between Cape Grim and each grid point 1 km either side, where plotted O₃ concentration is equal to the other location [O₃] (N, S, E, W) – Cape Grim [O₃].

The modelled TAPM–CTM concentrations are very similar at all grid points when not influenced by BB emissions. The variability increases at the time of BB1 and BB2, with differences mostly within 2–3 ppb, but up to 15 and 10 ppb at the east and west sites for BB1. This largest difference corresponds to the additional modelled O₃ peak that was not observed and showed strong dependency on EF (see Sect. 3.1.2) and provides further evidence that local BB emissions drive this enhancement.

The TAPM–CTM output for O₃ for BB1 (Fig. 7) shows O₃ enhancement downwind of the fire at 11:00 and 13:00 on 16 February. The very localised and narrow O₃ plume is dispersed by the light (2 m s⁻¹) and variable winds, and Cape Grim is on the edge of the O₃ plume for much of this period, explaining the high variability seen in Fig. 6c.

In summary, there is a large amount of spatial variability in TAPM–CTM for primary species such as CO during the BB events, with differences of > 500 ppb in grid points 1 km apart. This is due to the close proximity of the fire to the observation site and narrow-plume non-stationary meteorology. For O₃, there is up to 15 ppb of difference between grid
points for a narrow O$_3$ plume that is formed downwind of the fire.

The highly localised nature of the primary and in some cases secondary species seen here highlights the benefits of assessing spatial variability in situations with a close-proximity point source and a fixed receptor (measurement) site.

3.2 Exploring plume chemistry and contribution from different sources

3.2.1 Drivers of O$_3$ production

In previous work on the Robbins Island fire, it was noted that the increases in O$_3$ observed after both BB1 and BB2 were correlated with increased concentration of HFC-134a (Lawson et al., 2015). This indicated that transport of photochemically processed air from urban areas to Cape Grim was likely the main driver of the O$_3$ observed, rather than BB emissions (Lawson et al., 2015). However, during BB1 in a calm sunny period with minimal urban influence, an increase in O$_3$ was observed alongside a period of particle growth and elevated BC, suggesting possible BB influence. Normalised excess mixing ratios (NEMRs) observed during BB2 were also in the range of those observed elsewhere in young BB plumes (Lawson et al., 2015) (where NEMR is an excess mixing ratio normalised to a non-reactive co-emitted tracer, in this case CO; see Akagi et al., 2011).

To explore this further, TAPM–CTM was used to determine the degree to which the local fire emissions, and urban emissions from mainland Australia, were driving the observed O$_3$ enhancements. The scenario with EF corresponding to MCE = 0.89 was used, as discussed previously.

Figure 11 shows the simulated ozone for all sources (“with BB”) and all sources excluding the Robbins Island fire (“no BB”). There are two additional distinct ozone peaks in the with BB simulation (Fig. 11). These occurred during or close to the plume strikes and are short-lived (3 and 5 h) events. These same two peaks showed a strong dependence on model EF in Sect. 3.1.2. In contrast, the two peaks attributed to transport of air from mainland Australia are of longer duration and occur after the plume strikes.

Of the two modelled fire-derived O$_3$ peaks, the first modelled peak (33 ppb) corresponds with a small (21 ppb) observed peak during BB1 (Period B in Lawson et al., 2015), but the second modelled fire-derived O$_3$ peak is not observed. As shown in Fig. 7 and discussed in Sect. 3.1.3, according to TAPM–CTM the O$_3$ plumes generated from fire emissions were narrow and showed a strong spatial variability. Given this, it is challenging for TAPM–CTM to predict the exact timing and magnitude of these highly variable BB-generated O$_3$ peaks impacting Cape Grim. This is likely why there is good agreement in timing and magnitude between model and observations for the large-scale, spatially homogeneous O$_3$ plumes transported from mainland Australia but a lesser agreement for the locally formed, spatially variable O$_3$ formed from local fire emissions.

In summary, TAPM–CTM suggests that the two largest observed O$_3$ peaks following BB1 and BB2 were urban air transported from mainland Australia and some O$_3$ formation was driven by emissions from the local fire event. TAPM–CTM captures the magnitude and timing of the larger-scale urban-derived peaks well but is challenged by the timing and magnitude of O$_3$ from local BB emissions.

3.2.2 Plume age

TAPM–CTM was used to estimate the physical age of air parcels reaching Cape Grim over the 2-week period of the Robbins Island fire. The method is similar to the Eulerian effective physical age of emission metric, accounting for mix-
ing and chemical decay from Finch et al. (2014) and has been previously described in Keywood et al. (2015). Briefly, two model simulations were run for scenarios that included all sources of nitric oxide (NO) in Australia; the first treated NO as an unreactive tracer, the second with NO decaying at a constant first-order rate. The relative fraction of the emitted NO molecules remaining after 96 h was then inverted to give a molar-weighted plume age. As urban emissions are a larger NO source than BB, this approach would weight the age in the favour of the urban emissions if air masses from these two sources were mixed. However, as shown in Fig. 11, there are distinct periods when BB or urban sources dominate. As there is little mixing of air from the two sources, there are unlikely to be issues with the calculated age being weighted towards one source. The modelled NO tracer (decayed version) modelled plume age (hours) and the observed O$_3$. Direct BB1 and BB2 plume strikes can be clearly seen with increases in NO corresponding with a plume age of 0–2 h. The plume age then gradually increases over 24 h in both cases, peaking at 15:00 on 17 February during BB1 (age of plume is 40 h) and peaking at 17:00 on 25 February during BB2 (age of plume is 49 h). The peak observed O$_3$ enhancements correspond with the simulated plume age in both BB1 and BB2 (with an offset of 2 h for BB1) and the observed HFC-134a, suggesting that the plume that transported O$_3$ from Melbourne to Cape Grim was approximately 2 days old. TAPM–CTM also simulates a smaller NO peak alongside the maximum plume age, indicating transport of decayed NO from the mainland to Cape Grim.

As reported in Lawson et al. (2015), during BB2 NEMRs of $\Delta$O$_3$/ΔCO ranged from 0.001 to 0.074, in agreement with O$_3$ enhancements observed in young BB plumes elsewhere (Yokelson et al., 2003, 2009). However, the modelling reported here suggests that almost all of the O$_3$ observed during BB2 was of urban not BB origin. This suggests that NEMRs should not be used in isolation to identify the source of observed O$_3$ enhancements and highlights the value of utilising air mass back trajectories and modelling to interpret the source of O$_3$ enhancements where there are multiple emission sources.

4 Summary and conclusions

In this work we have used a unique set of opportunistic BB observations at Cape Grim Baseline Air Pollution Station to test the ability of CSIRO’s high-resolution (400 m grid cell) CTM to reproduce primary (CO, BC) and secondary (O$_3$) BB species in challenging non-stationary, inhomogeneous and near-field conditions. We tested the sensitivity of the CTM to three different parameters (meteorology, MCE and spatial variability) while holding the plume rise and the chemical mechanisms constant.

We found that meteorology, EF and spatial variability have a large influence on the modelled output mainly due to the close proximity of the fire to the receptor site (Cape Grim). The lower MCE (MCE = 0.89) TAPM–CTM model simulation provided the best agreement with the observed concentrations, in agreement with the MCE calculated from observations of 0.88 (Lawson et al., 2015). The changing EFs, in particular NO dependency on MCE, had a major influence on the simulated O$_3$ concentrations, with a tendency of the models in some configurations to fail to simulate observed O$_3$ peaks and to simulate complete titration of O$_3$ that was not observed. As shown in the previous work (Lawson et al., 2015), minor rainfall events have the potential to significantly alter EF due to changes in combustion processes. This work suggests that varying model EF has a major impact on whether the models predict production or destruction of O$_3$, which is particularly important at a receptor site in close proximity to the BB emissions. Models that assume a fixed EF for O$_3$ precursor species in an environment with temporarily and spatially variable EF may therefore be challenged to correctly predict the behaviour of important species such as O$_3$.

There were significant differences in model output between Cape Grim and grid points 1 km away, highlighting the narrowness of the plume and the challenge of predicting when the plume would impact the station. This also highlights the high spatial variability, which may be missed in similar situations by using a coarser-resolution model that would dilute emissions in a larger grid box.

TAPM–CTM was used to distinguish the influence of the two sources on the observed O$_3$ enhancements that followed BB1 and BB2. Transport of a 2-day-old urban plume some 300 km away from Melbourne was the main source of the O$_3$ enhancement observed at Cape Grim over the 2-week period of the fire. Despite NEMRs of $\Delta$O$_3$/ΔCO during BB2 being similar to those observed in young BB plumes elsewhere, this work suggests that NEMRs should not be used in isolation to identify the source of observed O$_3$ enhancements and highlights the value of utilising air mass back trajectories and modelling to interpret the source of O$_3$ enhancements where there are multiple emission sources.

Data availability. The underlying research data can be accessed upon request to the corresponding author (Sarah Lawson; sarah.lawson@csiro.au).

The Supplement related to this article is available online at https://doi.org/10.5194/acp-17-11707-2017-supplement.

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