1 On the relationship between the mesospheric sodium layer and the meteoric input function

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# 10 Abstract

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- 12 This study examines the relationship between the concentration of atmospheric sodium and its
- 13 Meteoric Input Function (MIF). We use the measurements from the Colorado State University (CSU)
- 14 Lidar and the Andes Lidar Observatory (ALO) with a new numerical model that includes sodium
- 15 chemistry in the mesosphere and lower thermosphere (MLT) region. The model is based on the
- 16 continuity equation to treat all sodium-bearing species and runs at a high temporal resolution. The
- 17 model simulation employs data assimilation to compare the MIF inferred from the meteor radiant
- 18 distribution and the MIF derived from the new sodium chemistry model. The simulation captures the
- 19 seasonal variability of sodium number density compared with lidar observations over the CSU site.
- 20 However, there were discrepancies for the ALO site, which is close to the South Atlantic Anomaly (SAA)
- region, indicating it is challenging for the model to capture the observed sodium over ALO. The CSU site
- had significantly more lidar observations (27,930 hours) than the ALO sites (1872 hours). The simulation
- revealed that the uptake of the sodium species on meteoric smoke particles was a critical factor in
- determining the sodium concentration in MLT, with the sodium removal rate by uptake found to be
- approximately three times that of the NaHCO<sub>3</sub> dimerization. Overall, the study's findings provide
   valuable information on the correlation between MIF and sodium concentration in the MLT region,
- 27 contributing to a better understanding of the complex dynamics in this region. This knowledge can
- inform future research and guide the development of more accurate models to enhance our
- 29 comprehension of the MLT region's behavior.
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- 31 **Keywords:** Sodium layer, sodium chemistry, meteor radiant distribution, meteoric input function
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- 33 Key points:
  - A high-time resolution, time-dependent Na chemistry model is developed.
  - Ablated global meteoroid material inputs inferred from ALO and CSU observations are about 83±28 t d<sup>-1</sup> and 53±23 t d<sup>-1</sup>, respectively.
- The frequency of meteor occurrences might not provide a precise reflection of the mass of
   meteoroid material input.
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#### 41 1. Introduction

- 42 Micro-meteoroids enter the Earth's atmosphere day and night, depositing their constituents into the
- 43 atmosphere via ablation, creating a region that hosts various metal species, for example, Fe, K, Si, Mg,
- 44 Ca, and Na, in both neutral and ion form (Plane et al., 2015; Plane et al., 2021; and references therein).
- 45 The region is commonly referred to as the mesosphere and lower thermosphere (MLT), located between
- 46 75 and 110 km altitude. The metal layers in the MLT often serve as the tracers that facilitate the
- 47 investigation of the dynamical and chemical processes within the region (Takahashi et al., 2014; Qiu et
- 48 al., 2021). Quantitative measurements of metal atoms have been made since the 1950s (Hunten, 1967)
- 49 through a variety of ground or space-borne technologies (Koch et al., 2021; Koch et al., 2022). The large
- 50 resonant scattering cross-section (Bowman et al., 1969) and the substantial presence of the sodium
- atom in the MLT make it one of the most researched metal layers in the atmosphere (Yu et al., 2022).
- 52 The sodium layer is usually studied via observations carried out by resonance lidars, satellites, and
- 53 through Na D-line emission at 589.0nm and 589.6nm (Plane, 2010; Plane et al., 2012; Hedin and
- 54 Gumbel, 2011; Langowski et al., 2017; Andrioli et al., 2019; Li et al., 2020a). The sodium vertical profiles
- retrieved by lidars have been commonly used as a tracer to study atmospheric dynamics, e.g., gravity
- 56 waves, wind shear, etc. The long-term seasonal and short-term diurnal variability of metallic species
- 57 have been investigated by several studies (Feng et al., 2013; Marsh et al., 2013; Cai et al., 2019a, b; Yu et
- al., 2022; She et al., 2023). A typical sodium chemistry scheme consists of neutral chemistry, ion
- 59 chemistry, and photolysis. The sodium chemistry research in recent years has primarily been based on
- 60 the sodium chemistry model by Plane (2004), which has been cited in various subsequent works,
- 61 including Bag et al. (2015) and references therein.

62 As meteoroids are the primary source of metal layers in the atmosphere, including the sodium layer, 63 the Meteoric Input Function (MIF) plays a crucial role in the modeling of metallic layers in the 64 atmosphere. The MIF is a function designed to model the temporal and spatial variability of the 65 meteoroid on the atmosphere (Pifko et al., 2013). Sporadic meteors are estimated to make up more 66 than 95% of the total meteoroid population by comparing the number of meteors originating in sporadic 67 sources to those originating in known shower meteor sources (Chau and Galindo, 2008). This highlights 68 the importance of incorporating sporadic meteor data in the MIF to accurately understand sodium 69 concentration in the mesosphere and lower thermosphere (MLT) region and its correlation with 70 meteoroid material input. It is well established that there are six apparent sources of sporadic meteors, 71 namely North and South Apex (NA and SA); North and South Toroidal (NT and ST); and Helion and Anti-72 Helion sources (H and AH) (Campbell-Brown, 2008; Kero et al., 2012; Li et al., 2022). However, the 73 relative strength of these meteor radiant sources varies among the studies. For example, the NA and SA 74 sources are found to be much stronger than other sources in results obtained with High Power Large 75 Aperture (HPLA) radars (Chau et al., 2007; Kero et al., 2012; Li and Zhou, 2019), while specular meteor 76 radars found the difference to be much smaller (Campbell-Brown and Jones, 2005; Campbell-Brown, 77 2008). The detection sensitivity varies significantly among different facilities. For instance, the Arecibo 78 Observatory (AO) at 18° N, 66° W detects approximately 20 times more meteors per unit area per unit 79 time than the Jicamarca Radio Observatory (JRO) at 12° S, 77° W, and at least 800 times more meteors 80 than the Resolute Bay Incoherent Scatter North (RISR-N) radar at 75° N, 95° W, despite all being HPLA 81 facilities (Li et al., 2020, 2023a; Hedges et al., 2022). While meteor flux does exhibit variations based on 82 time and latitude, these fluctuations alone cannot explain the magnitude of the observed difference.

- 83 Consequently, the total mass of the meteors that enter the Earth's atmosphere is subject to significant
- 84 uncertainties. In the existing Whole Atmosphere Community Climate Model-Na (WACCM-Na) global
- sodium model (Dunker et al., 2015), the meteoric input function was modeled by placing a flux curve on
- 86 each radiant meteor source with a definite ratio (more details can be found in Marsh et al., 2013). The
- 87 flux curve model is based on observations carried out exclusively by the Arecibo Observatory. Although
- 88 the model can reproduce some of the flux characteristics of the meteors observed at Arecibo, it is a
- 89 relatively simple model and therefore has several limitations (Li et al., 2022). One of the limitations is
- 90 that the model cannot reproduce the velocity distribution of the meteors in observations.
- 91 This study introduces a new numerical model for sodium chemistry that utilizes the continuity equations
- 92 for all Na-related reactions without steady-state approximations. The main objective is to investigate the
- 93 relationship between the apparent sodium concentration and the MIF in the MLT region. We then
- 94 compare the results of the new model with measurements from two lidar instruments, namely the
- 95 Colorado State University (CSU) and the Andes Lidar Observatory (ALO). Furthermore, we compared the
- 96 MIF derived from the new sodium chemistry model and lidar measurements from CSU and ALO, against
- 97 the results of the high-resolution meteor radiant distribution recently deduced from observations
- 98 conducted at AO. Finally, we discuss the implications of these comparisons and suggest possible
- 99 explanations for the discrepancy between the MIF derived from radar and those obtained from lidar 100 observations.
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# 102 **2. The sodium chemistry model (NaChem)**

# 103 2.1 Sodium chemistry

104 Numerical airglow models have been extensively used to investigate atmospheric airglow chemistry and 105 gravity waves (Huang and Hickey, 2008; Huang and Richard, 2014; Huang, 2015). A new numerical 106 sodium chemistry model, hereafter referred to as NaChem, was developed for this study. Table 1 lists 107 the complete reactions and their corresponding rate coefficients used in NaChem, which includes 108 neutral chemistry, ion chemistry, and photochemistry. The dimerization reaction of NaHCO<sub>3</sub> (reaction 25 109 in Table 1) is the outlet that removes Na atoms in the chemistry scheme. The Na atoms can also be 110 removed by the uptake of sodium species onto meteoric smoke particles (Hunten et al., 1980; 111 Kalashnikova et al., 2000; Plane, 2004), a process that can be turned on or off in the model. This study 112 estimates the MIF in the numerical model by matching the amount of sodium atoms removed by the 113 dimerization reaction and uptake, i.e., sodium sink, to maintain the observed sodium presence in the 114 MLT. MIF is a function of time and latitude, representing the mass of meteoroid material entering 115 Earth's atmosphere. Throughout the rest of the paper, the MIF estimated from the sodium chemistry 116 numerical model will be referred to as MIF(s). On the other hand, the MIF derived from meteor radiant 117 distribution, referred to as MIF(m). The MIF(m) is determined through a 3-D meteoroid orbital 118 simulation, a process similar to the seeding process discussed in section 3.1 of Li et al. (2022), based on 119 the meteor radiant distribution. MIF(m) is in arbitrary units. Note that the meteor mass cannot be accurately determined via radar measurements, however, the seasonal variation of meteoroid material 120 121 input can be represented by MIF(m). The estimation of meteor mass is further discussed in Section 5. In 122 contrast, MIF(s) is expressed in units of 1/cm<sup>3</sup>/second.

- 123 The numerical model utilizes the continuity equation to track the time evolution of all 14 Na-related
- species. Table 2 presents a comprehensive list of these species, along with their corresponding
- production and loss rates. The background gas species, including O<sub>3</sub>, O<sub>2</sub>, O, H, H<sub>2</sub>, H<sub>2</sub>O, etc., and the
- temperature are provided by WACCM version 6 (Jiao et al., 2022). Here we use the dynamic version of
- 127 WACCM nudged with NASA's Modern Era Retrospective Analysis for Research and Application MERRA2
- reanalysis data set (Hunziker & Wendt, 1974; Molod et al., 2015; Gettelman et al., 2019). The WACCM
- 129 reference profiles are linearly interpolated to a resolution of one minute and updated every minute
- during the simulation. It is worth noting that the Na-related reactions, which are illustrated in Table 2,
- do not significantly impact the background gas species, as the effect is orders of magnitude smaller than
- 132 the variation of the major gas species themselves. Therefore, the major gas species are simulated
- 133 independently of Na-related reactions.

## 134 2.2 Numerical scheme

As discussed earlier, it is worth noting that the reactions of sodium chemistry in NaChem share

- 136 similarities with those in previous models (e.g., Plane et al., 2015 and references therein); however, the
- 137 implementation of the numerical chemistry scheme differs. NaChem uses continuity equations to treat
- all chemicals involved, including short-lived intermediate species. Treating all species with the continuity
- equation is a straightforward and more accurate approach than using steady-state approximations.
- 140 Moreover, by treating all species in a uniform procedure, the numerical model is more compact and
- easier to interpret. The computational capability of a personal computer nowadays has advanced
- 142 enough to process an ultra-fine time step (microseconds) that is necessary for numerical simulations of
- short-lived species in a reasonable duration. Still, the differential equations for production and loss of
- short-lived species can be numerically unstable unless microsecond or even sub-microsecond time step
- is used (Higham, 2002). The concern of the differential equation instability can be largely mitigated by a
- 146 first-order exponential integrator (Hochbruck and Ostermann, 2010), i.e.,
- 147  $c = x_0 \frac{a_0}{b_0}$  (1a)
- 148

	0	$b_0$	•	1
$x_1 =$	$\frac{a_0}{b_0}$ -	$+ ce^{-b_0\Delta t}$	(1	b)

- 149 Where  $x_0$  is the value of the current step. In the simulation, it is the number density of the species.  $a_0$ 150  $(1/cm^3/s)$  is the production of the species,  $b_0$  (1/s) is the loss rate of the species,  $\Delta t$  is the step size in
- 151 time, and  $x_1$  is the value of the next step. The units for  $x_0$ ,  $x_1$ , and c are  $1/cm^3$ .
- 152 The exponential integrator, as expressed in Eq. 1a and 1b, is the solution to the continuity equation.

153 Note that reaction 25 listed in Table 1 is an exception, which was carried out using explicit Euler

- 154 integrator in the simulation. This reaction's continuity equation is structured differently from the others
- 155 because it represents the only mechanism for removing Na atoms from the chemistry simulation, apart
- 156 from the uptakes of sodium species. Our testing indicates that both the exponential integrator and
- 157 explicit Euler integrator yield nearly identical results. However, for numerical stability, the explicit Euler
- 158 integrator requires a step size of ~1μs, which is orders of magnitude smaller than the exponential
- 159 integrator. The default time step of NaChem is 0.1 seconds with the exponential integrator.
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neutral chemistry1 $Na + O_3 \rightarrow NaO(A) + O_2$ $K_1 = 1.1 \times 10^9 \exp(-116/T)$ 12 $NaO(A) + O \rightarrow Na^{(2}P) + O_2$ $K_2 = 2.2 \times 10^{-10}(T/200)^{0.5}, f_n = 0.1440.4$ 1.33 $NaO(A) + O \rightarrow Na^{(2}S) + O_2$ $K_3 = 2.2 \times 10^{-10}(T/200)^{0.5}, (1-f_n)$ 1.34 $NaO(A) + O_2 \rightarrow Na^{(2}S) + O_2$ $K_n = 1.2 \times 10^{-10}$ 15 $Na + O_2 + M \rightarrow NaO_2 + M$ $K_3 = 5.0 \times 10^{-20}(200/T)^{1.22}$ 16 $NaO_4 + O_2 \rightarrow NaO(X) + O_2$ $K_n = 5 \times 10^{-20}\exp(-940/T)$ 17 $NaO(X) + O \rightarrow Na^{(2}P) + O_2$ $K_7 = 2.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 1,28 $NaO(X) + O \rightarrow Na^{(2}S) + O_2$ $k_8 = 2.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 1,29 $NaO(X) + O_3 \rightarrow NaO_2 + O_2$ $k_8 = 1.1 \times 10^9 \exp(-568/T)$ 110 $NaO(X) + O_4 \rightarrow Na + 2O_2$ $k_{10} = 3.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 111 $NaO(X) + O_4 \rightarrow Na + 2O_2$ $k_{10} = 3.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 112 $NaO(X) + O_4 \rightarrow Na + 2O_2$ $k_{10} = 3.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 113 $NaO(X) + O_4 \rightarrow Na + AO_2$ $k_{10} = 3.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 114 $NaO(X) + O_4 \rightarrow Na + AO_2$ $k_{10} = 3.2 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 115 $NaO(X) + H_2 \rightarrow NaO_4 + M$ $k_{12} = 4.4 \times 10^{-10}\exp(-100/T)$ 116 $NaO(X) + H_2 \rightarrow NaO_4 + M$ $k_{12} = 4.5 \times 10^{-10}(T/200)^{0.5}, f_X = 0.167$ 117 $NaO(X) + H_2 \rightarrow NaO_4 + M$ $K_{12} = 1.0 \times 10^{-9}\exp(-100/T)$ 1<						
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3NaO(A) + O -> Na(25) + O2K3 = 2.2 x 10^{10}(T/20)^{0.5}, (1-f_A)1,34NaO(A) + O2 -> NaO(X) + O2K4 = 1 x 10^{11}15Na + O2 + M -> NaO2 + MK5 = 5.0 x 10^{30}(200/T)^{1.22}16NaO2 + O -> NaO(X) + O2K6 = 5 x 10^{10}exp(-940/T)17NaO(X) + O -> Na(2P) + O2K7 = 2.2 x 10^{10}(T/200)^{0.5}, fx = 0.1671,28NaO(X) + O -> Na(2S) + O2k8 = 2.2 x 10^{10}(T/200)^{0.5}, (1-f_N)1,29NaO(X) + O3 -> Na(2S) + O2k8 = 1.1 x 10^9exp(-568/T)110NaO(X) + O3 -> NaO2 + O2k9 = 1.1 x 10^9exp(-550/T)111NaO(X) + O3 -> NaO3 + MK11 = 5.3 x 10^{30}(200/T)112NaO(X) + H3 -> NaO3 + MK11 = 5.3 x 10^{30}(200/T)113NaO(X) + H3 -> NaO3 + MK11 = 5.3 x 10^{30}(200/T)114NaO(X) + H4 -> NAOH + HK12 = 4.4 x 10^{10}exp(-668/T)115NaO(X) + H2 -> Na + H2OK14 = 1.1 x 10^9 exp(-1100/T)116NaO(X) + H2 -> Na + H2OK15 = 1.3 x 10^{27} (200/T)117NaO(X) + H2 -> Na + H2OK13 = 2.5 x 10^{10}(T/200)^{0.5}118NaO3 + O -> NaO2 + CO2K13 = 2.5 x 10^{10}(T/200)^{0.5}119NaCO3 + H -> NaOH + CO2K20 = 1.0 x 10 <sup>9</sup> exp(-1000/T)120NaCO3 + H -> NaOH + CO2K20 = 1.0 x 10 <sup>9</sup> exp(-1000/T)121NaO4 + H2OK12 = 1.9 x 10^{28}(200/T)122NaO4 + O2 + M2O3K21 = 4.0 x 10^{11} exp(-550/T)123NaHCO3 + H ->						
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6NaO2 + O -> NaO(X) + O2K6 = 5 x 10 <sup>-10</sup> exp(-940/T)17NaO(X) + O -> Na( <sup>2</sup> P) + O2K7 = 2.2 x 10 <sup>-10</sup> (T/200) <sup>0.5</sup> , fx = 0.1671.28NaO(X) + O -> Na( <sup>2</sup> S) + O2k8 = 2.2 x 10 <sup>-10</sup> (T/200) <sup>0.5</sup> , (1.fx)1.29NaO(X) + O3 -> NaO2 + O2k9 = 1.1 x 10 <sup>3</sup> exp(-568/T)110NaO(X) + O3 -> Na + 2O2k10 = 3.2 x 10 <sup>-10</sup> exp(-568/T)111NaO(X) + O2 + M -> NaO3 + Mk11 = 5.3 x 10 <sup>-30</sup> (200/T)112NaO(X) + H2 -> NaO3 + Mk12 = 4.4 x 10 <sup>-10</sup> exp(-668/T)113NaO(X) + H2 -> NaOH + Hk13 = 1.1 x 10 <sup>30</sup> exp(-1100/T)114NaO(X) + H2 -> NaOH + Hk13 = 1.1 x 10 <sup>30</sup> exp(-1400/T)115NaO(X) + H2 -> NaOH + OHk15 = 4.4 x 10 <sup>-10</sup> exp(-507/T)116NaO(X) + CO2 + M -> NaCO3 + MK16 = 1.3 x 10 <sup>-27</sup> (200/T)117NaO2 + H -> Na + HO2K17 = 1.0 x 10 <sup>3</sup> exp(-1000/T)118NaO3 + O -> Na + 2O2k18 = 2.5 x 10 <sup>-10</sup> (T/200) <sup>0.5</sup> 119NaCO3 + O -> NaO2 + CO2k19 = 5.0 x 10 <sup>-10</sup> exp(-1000/T)120NaCO3 + H -> NaOH + CO2k20 = 1.0 x 10 <sup>3</sup> exp(-1400/T)121NaOH + H -> Na + H2Ok21 = 4.0 x 10 <sup>-11</sup> exp(-550/T)122NaOH + H2O + CO2k23 = 1.1 x 10 <sup>-11</sup> exp(-100/T)123NaHCO3 + H -> Na + H2Ok21 = 1.9 x 10 <sup>-28</sup> (200/T) <sup>1</sup> 124NaHCO3 + H -> Na + H2CO3k24 = 1.84 x 10 <sup>-13</sup> T <sup>0.77</sup> exp(-1014/T)1252NaHCO3 + H -> Na + H2CO3k25 = 8.8						
7NaO(X) + O -> Na( <sup>2</sup> P) + O2 $K_7 = 2.2 \times 10^{-10} (T/200)^{0.5}, f.x = 0.167$ 1,28NaO(X) + O -> Na( <sup>2</sup> S) + O2 $k_8 = 2.2 \times 10^{-10} (T/200)^{0.5}, (1-f_X)$ 1,29NaO(X) + O_3 -> Na + 2O2 $k_9 = 1.1 \times 10^{-9} exp(-568/T)$ 110NaO(X) + O_3 -> Na + 2O2 $k_{10} = 3.2 \times 10^{-10} (T/200)^{0.5}, (1-f_X)$ 111NaO(X) + O_2 + M -> NaO_3 + M $k_{11} = 5.3 \times 10^{-10} (200/T)$ 112NaO(X) + H -> Na + OH $k_{12} = 4.4 \times 10^{-10} exp(-568/T)$ 113NaO(X) + H_2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-100/T)$ 114NaO(X) + H_2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1400/T)$ 115NaO(X) + H_2 -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO_2 + M -> NaCO_3 + M $K_{16} = 1.3 \times 10^{-27} (200/T)$ 117NaO_2 + H -> Na + HO2 $K_{17} = 1.0 \times 10^{-9} exp(-1000/T)$ 118NaO_3 + O -> NaO_2 + CO_2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO_3 + O -> NaO_2 + CO_2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO_3 + H -> Na + H_2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 121NaOH + H -> Na + H_2O $k_{21} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO_3 + H -> Na + H_2O3 $k_{24} = 1.84 \times 10^{-13} exp(-1014/T)$ 124NaHCO_3 + H -> Na + H_2CO_3 $k_{24} = 1.84 \times 10^{-13} exp(-1014/T)$ 1252NaHCO_3 + H -> Na + H_2CO_3 $k_{24} = 1.84 \times 10^{-13} exp(-1014/T)$ 126Na( <sup>2</sup> P) -						
NaO(X) + O -> Na(2S) + O2 $k_8 = 2.2 \times 10^{-10} (T/200)^{0.5}, (1-f_x)$ 1,29NaO(X) + O3 -> NaO2 + O2 $k_9 = 1.1 \times 10^{-9} exp(-568/T)$ 110NaO(X) + O3 -> Na + 2O2 $k_1 = 3.2 \times 10^{-10} exp(-550/T)$ 111NaO(X) + O2 + M -> NaO3 + M $k_{11} = 5.3 \times 10^{-30} (200/T)$ 112NaO(X) + H -> Na + OH $k_{12} = 4.4 \times 10^{-10} exp(-668/T)$ 113NaO(X) + H2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1100/T)$ 114NaO(X) + H2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1400/T)$ 115NaO(X) + H2 -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO2 + M -> NaCO3 + M $K_{15} = 1.3 \times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2 $K_{17} = 1.0 \times 10^{-9} exp(-1000/T)$ 118NaO3 + O -> NaO2 + CO2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO3 + H -> Na + H2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 121NaOH + H-> Na + H2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 122NaOH + H-> Na + H2O $k_{22} = 1.9 \times 10^{-28} (200/T)^{-1}$ 123NaHCO3 + H -> Na + H2O + CO2 $k_{22} = 1.9 \times 10^{-28} (200/T)^{-1}$ 124NaHCO3 + H -> Na + H2OA $k_{24} = 1.84 \times 10^{-13} exp(-1010/T)$ 1252NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} exp(-1010/T)$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$						
9 $NaO(X) + O_3 -> NaO_2 + O_2$ $k_9 = 1.1 \times 10^{-9}exp(-568/T)$ 110 $NaO(X) + O_3 -> Na + 2O_2$ $k_{10} = 3.2 \times 10^{-10}exp(-550/T)$ 111 $NaO(X) + O_2 + M -> NaO_3 + M$ $k_{11} = 5.3 \times 10^{-30}(200/T)$ 112 $NaO(X) + H_2 -> NaO + M$ $k_{12} = 4.4 \times 10^{-10}exp(-668/T)$ 113 $NaO(X) + H_2 -> NaO + H$ $k_{12} = 4.4 \times 10^{-10}exp(-668/T)$ 114 $NaO(X) + H_2 -> NaO + H$ $k_{13} = 1.1 \times 10^{-9}exp(-1100/T)$ 115 $NaO(X) + H_2 -> NaO + HQ$ $k_{14} = 1.1 \times 10^{-9}exp(-1400/T)$ 116 $NaO(X) + CO_2 + M -> NaCO_3 + M$ $K_{15} = 4.4 \times 10^{-10}exp(-507/T)$ 117 $NaO_2 + H -> Na + HO_2$ $K_{17} = 1.0 \times 10^{-9}exp(-1000/T)$ 118 $NaO_3 + O -> Na + 2O_2$ $k_{18} = 2.5 \times 10^{-10}(T/200)^{0.5}$ 119 $NaCO_3 + O -> NaO_2 + CO_2$ $k_{19} = 5.0 \times 10^{-10}exp(-1200/T)$ 120 $NaCO_3 + H -> Na + H_2O$ $k_{22} = 1.9 \times 10^{-28}(200/T)^{1}$ 121 $NaOH + CO_2 + M -> NaHCO_3 + M$ $k_{22} = 1.9 \times 10^{-28}(200/T)^{1}$ 123 $NaHCO_3 + H -> Na + H_2O + CO_2$ $k_{23} = 1.1 \times 10^{-11}exp(-910/T)$ 124 $NaHCO_3 + H -> Na + H_2O + CO_2$ $k_{23} = 1.8 \times 10^{-10}exp(-1014/T)$ 125 $2NaHCO_3 + M -> (NaHCO_3)_2 + M$ $k_{25} = 8.8 \times 10^{-10}exp(-1014/T)$ 126 $Na(^2P) -> Na(^2S) + hv(S89.0-589.6 mm)$ $K_26 = 6.26 \times 10^7$ 1						
10NaO(X) + O3 -> Na + 2O2 $k_{10} = 3.2 \times 10^{-10} exp(-550/T)$ 111NaO(X) + O2 + M -> NaO3 + M $k_{11} = 5.3 \times 10^{-30}(200/T)$ 112NaO(X) + H -> Na + OH $k_{12} = 4.4 \times 10^{-10} exp(-668/T)$ 113NaO(X) + H2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1100/T)$ 114NaO(X) + H2 -> Na + H2O $k_{14} = 1.1 \times 10^{-9} exp(-1400/T)$ 115NaO(X) + H2 -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO2 + M -> NaCO3 + MK16 = 1.3 $\times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2K17 = 1.0 $\times 10^{-9} exp(-1000/T)$ 118NaO3 + O -> NaO2 + CO2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO3 + H -> Na + H2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 121NaOH + H -> Na + H2O $k_{21} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO3 + H -> Na + H2O $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 124NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-13} exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
11NaO(X) + O2 + M -> NaO3 + M $k_{11} = 5.3 \times 10^{-30}(200/T)$ 112NaO(X) + H -> Na + OH $k_{12} = 4.4 \times 10^{-10} exp(-668/T)$ 113NaO(X) + H_2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1100/T)$ 114NaO(X) + H_2 -> Na + H_2O $k_{14} = 1.1 \times 10^{-9} exp(-1400/T)$ 115NaO(X) + H_2 O -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO2 + M -> NaCO3 + MK16 = 1.3 $\times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2K17 = 1.0 $\times 10^{-9} exp(-1000/T)$ 118NaO3 + O -> Na + 2O2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO3 + H -> Na + H_2O $k_{21} = 4.0 \times 10^{-19} exp(-550/T)$ 121NaOH + H -> Na + H_2O $k_{21} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO3 + H -> Na + H_2O + CO2 $k_{23} = 1.1 \times 10^{-18} exp(-910/T)$ 124NaHCO3 + H -> Na + H_2CO3 $k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm)K26 = 6.26 \times 10^71						
$ \begin{array}{ c c c c } 11 & NaO(X) + O_2 + M \rightarrow NaO_3 + M & k_{11} = 5.3 \times 10^{-30}(200/T) & 1 \\ 12 & NaO(X) + H \rightarrow Na + OH & k_{12} = 4.4 \times 10^{-10}exp(-668/T) & 1 \\ 13 & NaO(X) + H_2 \rightarrow NaOH + H & k_{13} = 1.1 \times 10^{-9}exp(-1100/T) & 1 \\ 14 & NaO(X) + H_2 \rightarrow Na + H_2O & k_{14} = 1.1 \times 10^{-9}exp(-1400/T) & 1 \\ 15 & NaO(X) + H_2O \rightarrow NaOH + OH & k_{15} = 4.4 \times 10^{-10}exp(-507/T) & 1 \\ 16 & NaO(X) + CO_2 + M \rightarrow NaCO_3 + M & K_{16} = 1.3 \times 10^{-27}(200/T) & 1 \\ 17 & NaO_2 + H \rightarrow Na + HO_2 & K_{17} = 1.0 \times 10^{-9}exp(-1000/T) & 1 \\ 18 & NaO_3 + O \rightarrow Na + 2O_2 & k_{18} = 2.5 \times 10^{-10}(T/200)^{0.5} & 1 \\ 19 & NaCO_3 + O \rightarrow NaO_2 + CO_2 & k_{19} = 5.0 \times 10^{-10}exp(-1200/T) & 1 \\ 20 & NaCO_3 + H \rightarrow NaOH + CO_2 & k_{20} = 1.0 \times 10^{-9}exp(-1400/T) & 1 \\ 21 & NaOH + H \rightarrow Na + H_2O & k_{21} = 4.0 \times 10^{-11}exp(-550/T) & 1 \\ 22 & NaOH + CO_2 + M \rightarrow NaHCO_3 + M & k_{22} = 1.9 \times 10^{-28}(200/T)^1 & 1 \\ 23 & NaHCO_3 + H \rightarrow Na + H_2O + CO_2 & k_{23} = 1.1 \times 10^{-11}exp(-910/T) & 1 \\ 24 & NaHCO_3 + H \rightarrow Na + H_2O + CO_2 & k_{23} = 1.1 \times 10^{-11}exp(-910/T) & 1 \\ 25 & 2NaHCO_3 + M \rightarrow Na + H_2CO_3 & k_{24} = 1.84 \times 10^{-13}T^{0.777}exp(-1014/T) & 1 \\ 25 & 2NaHCO_3 + M \rightarrow (NaHCO_3)_2 + M & k_{25} = 8.8 \times 10^{-10}exp(T/200)^{0.23} & 1 \\ 26 & Na(^{2}P) \rightarrow Na(^{2}S) + hv(589.0-589.6 nm) & K_{26} = 6.26 \times 10^{7} & 1 \\ \end{array}$						
13NaO(X) + H2 -> NaOH + H $k_{13} = 1.1 \times 10^{-9} exp(-1100/T)$ 114NaO(X) + H2 -> Na + H2O $k_{14} = 1.1 \times 10^{-9} exp(-1400/T)$ 115NaO(X) + H2O -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO2 + M -> NaCO3 + M $K_{16} = 1.3 \times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2 $K_{17} = 1.0 \times 10^{-9} exp(-1000/T)$ 118NaO3 + O -> Na + 2O2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO3 + H -> NaOH + CO2 $k_{20} = 1.0 \times 10^{-9} exp(-1400/T)$ 121NaOH + H -> Na + H2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 123NaHCO3 + H -> Na + H2O + CO2 $k_{23} = 1.1 \times 10^{-13} exp(-101/T)$ 124NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} t^{0.777} exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
$ \begin{array}{ c c c c } 13 & NaO(X) + H_2 -> NaOH + H & k_{13} = 1.1 \times 10^{-9} exp(-1100/T) & 1 \\ 14 & NaO(X) + H_2 -> Na + H_2O & k_{14} = 1.1 \times 10^{-9} exp(-1400/T) & 1 \\ 15 & NaO(X) + H_2O -> NaOH + OH & k_{15} = 4.4 \times 10^{-10} exp(-507/T) & 1 \\ 16 & NaO(X) + CO_2 + M -> NaCO_3 + M & K_{16} = 1.3 \times 10^{-27} (200/T) & 1 \\ 17 & NaO_2 + H -> Na + HO_2 & K_{17} = 1.0 \times 10^{-9} exp(-1000/T) & 1 \\ 18 & NaO_3 + O -> Na + 2O_2 & k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5} & 1 \\ 19 & NaCO_3 + O -> NaO_2 + CO_2 & k_{19} = 5.0 \times 10^{-10} exp(-1200/T) & 1 \\ 20 & NaCO_3 + H -> NaOH + CO_2 & k_{20} = 1.0 \times 10^{-9} exp(-1400/T) & 1 \\ 21 & NaOH + H -> Na + H_2O & k_{21} = 4.0 \times 10^{-11} exp(-550/T) & 1 \\ 22 & NaOH + CO_2 + M -> NaHCO_3 + M & k_{22} = 1.9 \times 10^{-28} (200/T)^1 & 1 \\ 23 & NaHCO_3 + H -> Na + H_2O + CO_2 & k_{23} = 1.1 \times 10^{-11} exp(-910/T) & 1 \\ 24 & NaHCO_3 + H -> Na + H_2O + CO_2 & k_{23} = 1.1 \times 10^{-13} exp(-1014/T) & 1 \\ 25 & 2NaHCO_3 + H -> Na + H_2CO_3 & k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T) & 1 \\ 25 & 2NaHCO_3 + M -> (NaHCO_3)_2 + M & k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23} & 1 \\ 26 & Na(^2P) -> Na(^2S) + hv(589.0-589.6 nm) & K_{26} = 6.26 \times 10^7 & 1 \\ \end{array}$						
15NaO(X) + H <sub>2</sub> O -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} exp(-507/T)$ 116NaO(X) + CO <sub>2</sub> + M -> NaCO <sub>3</sub> + M $K_{16} = 1.3 \times 10^{-27} (200/T)$ 117NaO <sub>2</sub> + H -> Na + HO <sub>2</sub> $K_{17} = 1.0 \times 10^{-9} exp(-1000/T)$ 118NaO <sub>3</sub> + O -> Na + 2O <sub>2</sub> $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO <sub>3</sub> + O -> NaO <sub>2</sub> + CO <sub>2</sub> $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO <sub>3</sub> + H -> NaOH + CO <sub>2</sub> $k_{20} = 1.0 \times 10^{-9} exp(-1400/T)$ 121NaOH + H -> Na + H <sub>2</sub> O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 122NaOH + CO <sub>2</sub> + M -> NaHCO <sub>3</sub> + M $k_{22} = 1.9 \times 10^{-28} (200/T)^{1}$ 123NaHCO <sub>3</sub> + H -> Na + H <sub>2</sub> O $k_{24} = 1.84 \times 10^{-13} r^{0.777} exp(-1014/T)$ 124NaHCO <sub>3</sub> + H -> Na + H <sub>2</sub> CO <sub>3</sub> $k_{24} = 1.84 \times 10^{-13} r^{0.777} exp(-1014/T)$ 1252NaHCO <sub>3</sub> + M -> (NaHCO <sub>3</sub> ) <sub>2</sub> + M $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + <i>hv</i> (589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
15NaO(X) + H2O -> NaOH + OH $k_{15} = 4.4 \times 10^{-10} \exp(-507/T)$ 116NaO(X) + CO2 + M -> NaCO3 + M $K_{16} = 1.3 \times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2 $K_{17} = 1.0 \times 10^{-9} \exp(-1000/T)$ 118NaO3 + O -> Na + 2O2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} \exp(-1200/T)$ 120NaCO3 + H -> NaOH + CO2 $k_{20} = 1.0 \times 10^{-9} \exp(-1400/T)$ 121NaOH + H -> Na + H2O $k_{21} = 4.0 \times 10^{-11} \exp(-550/T)$ 122NaOH + CO2 + M -> NaHCO3 + M $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO3 + H -> Na + H2O + CO2 $k_{23} = 1.1 \times 10^{-11} \exp(-910/T)$ 124NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} T^{0.777} \exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-10} \exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
16NaO(X) + CO2 + M -> NaCO3 + M $K_{16} = 1.3 \times 10^{-27} (200/T)$ 117NaO2 + H -> Na + HO2 $K_{17} = 1.0 \times 10^{-9} exp(-1000/T)$ 118NaO3 + O -> Na + 2O2 $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119NaCO3 + O -> NaO2 + CO2 $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120NaCO3 + H -> NaOH + CO2 $k_{20} = 1.0 \times 10^{-9} exp(-1400/T)$ 121NaOH + H -> Na + H2O $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 122NaOH + CO2 + M -> NaHCO3 + M $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO3 + H -> Na + H2O + CO2 $k_{23} = 1.1 \times 10^{-11} exp(-910/T)$ 124NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $						
18 $NaO_3 + O \rightarrow Na + 2O_2$ $k_{18} = 2.5 \times 10^{-10} (T/200)^{0.5}$ 119 $NaCO_3 + O \rightarrow NaO_2 + CO_2$ $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120 $NaCO_3 + H \rightarrow NaOH + CO_2$ $k_{20} = 1.0 \times 10^{-9} exp(-1400/T)$ 121 $NaOH + H \rightarrow Na + H_2O$ $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 122 $NaOH + CO_2 + M \rightarrow NaHCO_3 + M$ $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 123 $NaHCO_3 + H \rightarrow Na + H_2O + CO_2$ $k_{23} = 1.1 \times 10^{-11} exp(-910/T)$ 124 $NaHCO_3 + H \rightarrow Na + H_2CO_3$ $k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T)$ 125 $2NaHCO_3 + M \rightarrow (NaHCO_3)_2 + M$ $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126 $Na(^2P) \rightarrow Na(^2S) + hv(589.0-589.6 nm)$ $K_{26} = 6.26 \times 10^7$ 1						
19 $NaCO_3 + O -> NaO_2 + CO_2$ $k_{19} = 5.0 \times 10^{-10} exp(-1200/T)$ 120 $NaCO_3 + H -> NaOH + CO_2$ $k_{20} = 1.0 \times 10^{-9} exp(-1400/T)$ 121 $NaOH + H -> Na + H_2O$ $k_{21} = 4.0 \times 10^{-11} exp(-550/T)$ 122 $NaOH + CO_2 + M -> NaHCO_3 + M$ $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 123 $NaHCO_3 + H -> Na + H_2O + CO_2$ $k_{23} = 1.1 \times 10^{-11} exp(-910/T)$ 124 $NaHCO_3 + H -> Na + H_2CO_3$ $k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T)$ 125 $2NaHCO_3 + M -> (NaHCO_3)_2 + M$ $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126 $Na(^2P) -> Na(^2S) + hv(589.0-589.6 nm)$ $K_{26} = 6.26 \times 10^7$ 1						
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22NaOH + CO2 + M -> NaHCO3 + M $k_{22} = 1.9 \times 10^{-28} (200/T)^1$ 123NaHCO3 + H -> Na + H2O + CO2 $k_{23} = 1.1 \times 10^{-11} exp(-910/T)$ 124NaHCO3 + H -> Na + H2CO3 $k_{24} = 1.84 \times 10^{-13} T^{0.777} exp(-1014/T)$ 1252NaHCO3 + M -> (NaHCO3)2 + M $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126Na( <sup>2</sup> P) -> Na( <sup>2</sup> S) + hv(589.0-589.6 nm) $K_{26} = 6.26 \times 10^7$ 1						
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25 $2NaHCO_3 + M \rightarrow (NaHCO_3)_2 + M$ $k_{25} = 8.8 \times 10^{-10} exp(T/200)^{-0.23}$ 126 $Na(^2P) \rightarrow Na(^2S) + hv(589.0-589.6 nm)$ $K_{26} = 6.26 \times 10^7$ 1						
26 $Na(^{2}P) \rightarrow Na(^{2}S) + hv(589.0-589.6 \text{ nm})$ $K_{26} = 6.26 \times 10^{7}$ 1						
ion-molecule chemistry						
27 Na + $O_2^+ \rightarrow Na^+ + O_2$ K <sub>27</sub> = 2.7 x 10 <sup>-9</sup> 1						
$\frac{1}{28} \qquad Na + NO^+ -> Na^+ + NO \qquad K_{28} = 8.0 \times 10^{-10} \qquad 1$						
29 $Na^+ + N_2 + M \rightarrow NaN_2^+ + M$ $k_{29} = 4.8 \times 10^{-30} (T/200)^{-2.2}$ 1						
30 Na <sup>+</sup> + CO <sub>2</sub> + M -> NaCO <sub>2</sub> <sup>+</sup> + M $k_{30} = 3.7 \times 10^{-29} (T/200)^{-2.9}$ 1						
$NaN_2^+ + O \implies NaO^+ + N_2 \qquad k_{31} = 4.0 \times 10^{-10} \qquad 1$						
$32  NaO^{+} + N_2 -> NaN_2^{+} + O  k_{32} = 1.0 \times 10^{-12}  1$						
33 NaO <sup>+</sup> + O -> Na <sup>+</sup> + O <sub>2</sub> k <sub>33</sub> = 1.0 x 10 <sup>-11</sup> 1						
$34  NaO^+ + O_2 -> Na^+ + O_3  k_{34} = 5.0 \times 10^{-12}  1$						
35 NaN <sub>2</sub> <sup>+</sup> + X -> NaX <sup>+</sup> + N <sub>2</sub> (X=CO <sub>2</sub> , H <sub>2</sub> O) $k_{35} = 6.0 \times 10^{-10}$ 1						
$36 \qquad NaY^{+} + e \rightarrow Na + Y (Y=N_2, CO_2, H_2O, O) \qquad k_{36} = 1.0 \times 10^{-6} (T/200)^{-0.5} \qquad 1$						
photochemical reactions						
37 NaO(A)/NaO(X) + hv -> Na + O K <sub>37</sub> = 5.5 x 10 <sup>-2</sup> 1						
38     NaO2 + hv -> Na + O2     K <sub>38</sub> = 1.9 x 10 <sup>-2</sup> 1						
39         NaOH + hv -> Na + OH         K <sub>39</sub> = 1.8 x 10 <sup>-2</sup> 1						
40 NaHCO3 + hv -> Na + HCO3 K <sub>40</sub> = 1.3 x 10 <sup>-4</sup> 1						
41 Na + hv -> Na + + e- K <sub>41</sub> = 2 x 10 <sup>-5</sup> 1						

# 162 Table 1. Reactions in NaChem. $f_a$ and $f_x$ are branching ratios.

163

\*1:Plane (2004), 2: Plane (2012), 3: Griffin et al. (2001). Units for rate coefficient: unimolecular, s<sup>-1</sup>;

bimolecular, cm<sup>3</sup> s<sup>-1</sup>, termolecular, cm<sup>6</sup>s<sup>-1</sup>, etc.

## 166 Table 2. The production and loss terms of the sodium-related species.

	Species	Prod	Loss
a1	Na( <sup>2</sup> P)	k <sub>2</sub> [a <sub>3</sub> ][O] + k <sub>7</sub> [a <sub>5</sub> ][O];	1*
a2	Na	$ \begin{array}{l} k_3[a_3][O] + k_8[a_5][O] + k_{10}[a_5][O_3] + k_{12}[a_5][H] + k_{14}[a_5][H_2] + \\ k_{17}[a_4][H] + k_{18}[a_6][O] + k_{21}[a_7][H] + k_{23}[a_9][H] + k_{24}[a_9][H] + \\ k_{36}[a_{11}][e] + k_{36}[a_{13}][e] + k_{36}[a_{12}][e] + k_{36}[a_{14}][e] + [a_1] + k_{37}[a_3][hv] \\ + k_{37}[a_5][hv] + k_{38}[hv][a_4] + k_{39}[hv][a_7] + k_{40}[hv][a_9]; \end{array} $	k1[O3] + k5[O3] + k5[O2][M] + k27[O2+] + k28[NO+] + k41[hv];
a3	NaO(A)	k <sub>1</sub> [a <sub>2</sub> ][O <sub>3</sub> ]	$k_2[O] + k_3[O] + k_4[O_2] + k_{37}[hv]$
a4	NaO <sub>2</sub>	$k_{5}[a_{2}][O_{2}][M] + k_{9}[a_{5}][O_{3}] + k_{19}[a_{8}][O]$	$k_6[O] + k_{17}[H] + k_{38}[hv]$
а5	NaO(X)	$k_{5}[a][O_{3}]+k_{4}[a_{3}][O_{2}]+k_{6}[a_{4}][O]$	$\begin{array}{l} k_7[O] + k_8[O] + k_9[O_3] + k_{10}[O_3] + k_{11}[O_2][M] + \\ k_{12}[H] + k_{13}[H_2] + k_{14}[H_2] + k_{15}[H_2O] + \\ k_{16}[CO_2][M] + k_{37}[hv] \end{array}$
a6	NaO <sub>3</sub>	$k_{11}[a_5][O_2][M]$	k <sub>18</sub> [O]
a7	NaOH	$k_{13}[a_5][H_2] + k_{15}[a_5][H_2O] + k_{20}[a_8][H]$	k <sub>21</sub> [H]+ k <sub>22</sub> [CO <sub>2</sub> ][M] + k <sub>39</sub> [hv]
a8	NaCO <sub>3</sub>	k <sub>16</sub> [a <sub>5</sub> ][CO][M]	k <sub>19</sub> [O] + k <sub>20</sub> [H]
a9	NaHCO <sub>3</sub>	k <sub>22</sub> [a <sub>7</sub> ][CO <sub>2</sub> ][M]	$k_{23}[H] + k_{24}[H] + 2k_{25}[a_9][M] + k_{40}[hv]$
a10	Na <sup>+</sup>	$k_{27}[a_2][O_2^+] + k_{28}[a_2][NO^+] + k_{33}[a_{13}][O] + k_{34}[a_{13}][O_2] + k_{41}[hv][a_2]$	$k_{29}[N_2][M] + k_{30}[CO_2][M]$
a11	$NaN_{2}^{+}$	$k_{29}[a_{10}][N_2][M] + k_{32}[a_{13}][N_2]$	$k_{31}[O]+k_{35}[CO_2] + k_{35}[H_2O] + k_{36}[e]$
a12	$NaCO_{2}^{+}$	$k_{30}[a_{10}][CO_2][M] + k_{35}[a_{11}][CO_2]$	k <sub>36</sub> [e]
a13	NaO+	k <sub>31</sub> [a <sub>11</sub> ][O]	$k_{32}[N2]+k_{33}[O]+k_{34}[O_2]+k_{36}[e]$
a14	$NaH_2O^+$	k <sub>35</sub> [a <sub>11</sub> ][H <sub>2</sub> O]	k <sub>36</sub> [e]

167

168 \*In Species 1, as of the current state of the model, all Na(<sup>2</sup>p) atoms return to their ground state

169 immediately, so the loss term is set to 1. The [hv] is the term that represents loss via photoionization,

170 which is approximately a sinusoidal function based on the solar zenith angle of the respective local solar

171 time.

172

#### 173 3. CSU and ALO Sodium Lidar Observations and data processing

#### 174 3.1 Observations

175 Several aspects of the current research, i.e., the presence of sodium in the MLT, require cross-validation

176 with the measurements. One primary objective of the present model is to match the observed seasonal

variation of the sodium layer. Measurements by the Colorado State University (CSU, 41.4°N, 111.5°W)

178 Lidar, also known as Utah State University (USU) Lidar, and the lidar data acquired by the Andes Lidar

- 179 Observatory (ALO, 30.3°S, 70.7°W), are used to facilitate the research in the current study. We are
- 180 unable to acquire more ALO data after 2019 as the COVID situation disrupted the site operation. The

181 CSU data comprises 27,930 hours of lidar observations between 1990 and 2020, whereas the ALO data

182 consists of 1872 hours between 2014 and 2019.

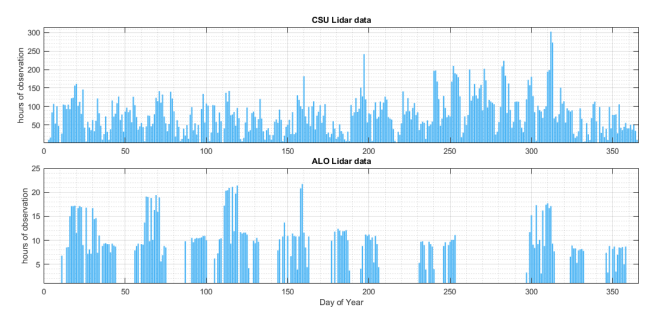


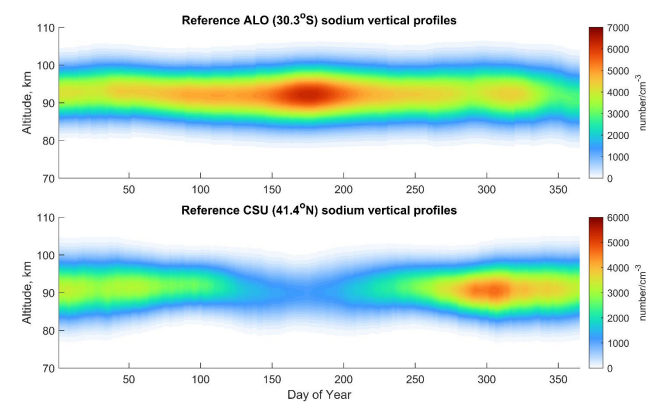
Figure 1. Available hours of lidar observations. CSU lidar (1990-2020, upper plot) and ALO lidar (20142019, lower plot).

186 The statistics of CSU and ALO available data are presented in Figure 1. The Lidar observations of both 187 sites consist of nocturnal observations only, and a typical nocturnal observation lasts between 8 and 11 188 hours. Note that in Figure 1, there could be as many as 300 hours of sodium observations on a single day 189 of year, which means the data of the date comprise observations of many years on that day in different 190 years. The CSU data almost covered every day of the year with only a few exceptions, whereas the ALO 191 data was much more sparse. As a result, due to the significantly larger number of CSU observations, the 192 statistical reliability of the seasonal variation in the sodium layer derived from ALO observations may not 193 be as strong as that of the CSU data. As depicted in Figure 2, the overall seasonal trend of the sodium 194 vertical profile derived from CSU lidar observations closely aligns with the simulation-based estimate by 195 Marsh et al. (2013). In contrast, ALO lidar observations deviate from the findings reported by Marsh et 196 al. (2013). The ALO measurements exhibit a prominent peak around June, while the results in Marsh et 197 al. (2013) show a double peak in March and October.

#### 198 3.2 Data processing

183

199 The sodium layer in atmospheric observations is often affected by perturbations of atmospheric 200 dynamics, which is why sodium is commonly used as a tracer in the study of the MLT dynamics (Plane et 201 al., 2015). However, studying the sodium layer itself can be complicated due to the underlying chemical 202 processes coupled with the dynamics. In order to mitigate the effects of atmospheric dynamics, we 203 process the sodium vertical profiles from observations in three steps. First, we average the profiles by 204 day of the year, meaning we take the average of the data from the same day of the year from different 205 years. Missing data are treated using linear interpolation. Next, we smooth the averaged profiles using a 206 15-day running average. Finally, the height profile for each time step is further smoothed by fitting it 207 with a skew-normal distribution (Azzalini & Valle, 1996), using the least squares error method.



208

**Figure 2**. The reference annual sodium vertical profiles at ALO (top plot) and at CSU (bottom plot). The

reference profiles are the averages throughout all the available data on the same days at the respective

site, then fitted by a skew-normal distribution that mitigates atmospheric dynamics. In essence, the

reference profiles are measurements with small-scale dynamics removed via steps discussed in section

213 *3.2*.

214

Figure 2 displays the processed annual sodium vertical profiles from the lidar measurements, referred to 215 216 as reference profiles hereafter. These profiles serve as references to guide the numerical simulation of 217 the NaChem model. The reference profiles are Na lidar measurements fitted using a skew-normal 218 distribution, smoothed by a 15-day running average, and processed through linear 2-D interpolation 219 across time and altitude. The lidar measurements have an altitude resolution of 500m for ALO and from 220 75m to 140m for CSU. These measurements are interpolated to a 100m resolution as inputs to the 221 NaChem model. The time resolutions of the lidar measurements typically vary between 1 and 10 222 minutes, depending on the experiment, and are linear interpolated to 0.1 seconds. The reference 223 profiles inherently include diffusion and other dynamic effects on the sodium species in the MLT, as 224 these observational data represent snapshots of sodium diffusion at various times. By constantly 225 matching the observed Na profile to the simulated Na profile, the diffusion is included implicitly in the 226 model. The seasonal column densities of both ALO and CSU profiles are similar to a sinusoidal function, 227 with ALO data peaking near June and CSU data peaking in November. The centroid height of the sodium 228 layer is higher in the ALO data than in the CSU data.

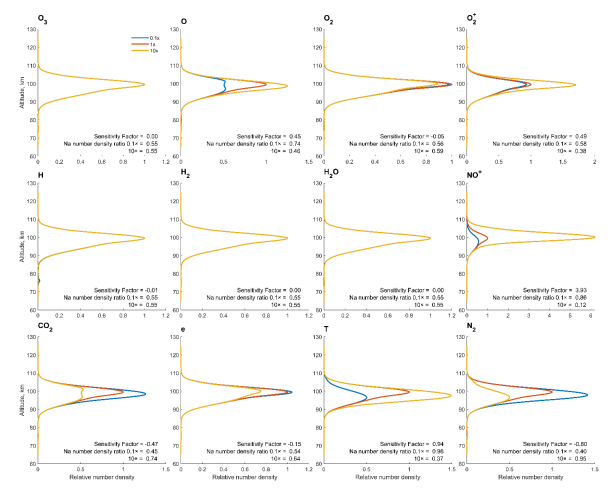
## 230 4. Results

## 231 4.1 Sensitivity test

232 Sodium in the atmosphere could manifest in many forms, i.e., in sodium-bearing neutral chemicals and

233 ionic chemicals. The sodium number densities are typically obtained via lidar measurements. Given the

- complexity of the sodium chemistry, the observed sodium is merely a subset, possibly not even a major
- constituent, of the total number of all the sodium-bearing species in the atmosphere. The total sodium
- content is defined as the total number of sodium atoms in all 14 sodium-bearing species, as listed in
- Table 2. In summary, the sodium that we can detect does not necessarily provide an accurate
- representation of the total sodium content or the overall count of sodium-bearing species, as
- unobservable species such as Na<sup>+</sup> and NaHCO<sub>3</sub> could constitute a substantial portion of the total sodium
   content.
- 241 Understanding the impact of each background species, i.e., species listed in Figure 3., on the total
- sodium content is essential to study the underlying mechanism of the chemical reactions. Therefore, we
- present a sensitivity test by isolating variables. The sensitivity test is done by altering the number
- density of background species in question by two orders of magnitude, i.e., with a factor of 0.1 and 10,
- while keeping the number densities of other background species and the atomic sodium fixed. The
- simulation is kept running until all the numbers are stable. The diurnal variations of the sodium and
- 247 background species are not considered in sensitivity test as they introduce unnecessary complexity. The
- results of the sensitivity test of the 11 background species and temperatures involved in the numerical
- simulation are shown in Figure 3. Each panel contains three lines, where the red curve shows the
- 250 unaltered vertical profile of the total sodium content. The results of the species altered by the factor of
- 251 0.1 and 10 are shown in light blue and yellow, respectively.



252

Figure 3. Sensitivity test of 11 background species and temperature on Na chemistry. The total sodium content vertical profile for the respective background species altered by 10x and 0.1x are shown in yellow and light blue. The reference sodium content vertical profiles are shown in red. Additionally, the sensitivity factor and the Na number density ratio to the concentration of all sodium species are presented on each panel.

In Figure 3, only the yellow curve is visible in some of the panels because the three curves are drawn on
top of each other, indicating that the change of the respective background species bears little to no
effect on the sodium chemistry. A sensitivity factor is defined to better quantify the weight of each
background species in sodium chemistry. The factor is calculated by the following equation:

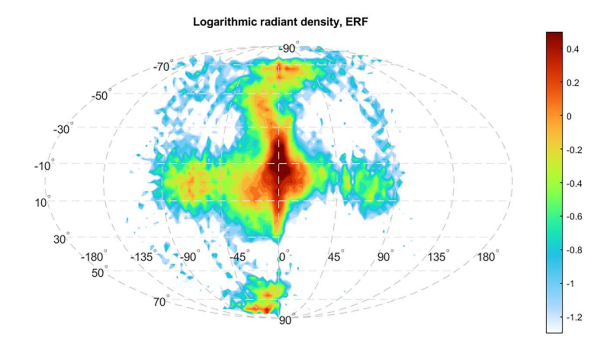
263 
$$Sensitivity Factor = \frac{NaT_c^{10} - NaT_c^{0.1}}{NaT_c}$$
(2)

264 Where  $NaT_c^{10}$  is the column density of the total sodium content with the respective species altered by a 265 factor of 10, and  $NaT_c^{0.1}$  is the same operation as the previous one but altered by a factor of 0.1. The 266 denominator,  $NaT_c$ , is the column density of the reference profile. The sensitivity factor provides a 267 general insight into how variations in the background species correlate with sodium number density. A 268 greater absolute value for the sensitivity factor indicates a stronger correlation. A positive sensitivity 269 factor indicates a positive correlation between the total sodium content and the respective species, and

- vice versa. The reference profile is the total sodium content in steady-state in the background condition
- of the midnight new year of 2002, giving a typical sodium vertical profile similar to the one shown in
- Figure 5 of Plane (2004). In the simulation, a greater total sodium content implies that a smaller
- 273 percentage of the sodium chemicals are present as sodium atoms as the altitude profile of the sodium
- atoms is fixed in the sensitivity test. In reality, instead of the sodium atoms, the total sodium content
- should be more or less conserved. Hence a higher total sodium content in our simulation suggests less
- sodium can be detected by the lidar.
- 277 Although the sensitivity factor could be different upon the change of the reference profile, it still gives
- an insight into the significance of each background species to the sodium chemistry. Apparently, the
- 279 weight of some background species, namely  $O_3$ , H,  $H_2$ , and  $H_2O$ , is negligible in sodium chemistry,
- 280 meaning that removing these species and their associated reactions has no effect on the overall sodium
- 281 chemistry. Nevertheless, these species are still kept in our numerical model for completeness. The
- impact of species that convert Na atom to Na<sup>+</sup>, as listed in reactions 27 and 28 of Table 1, is generally
- strong. The effect of NO<sup>+</sup>, in particular, is the most significant according to the sensitivity factor, greater
- than the combined effect of all the other species. Consequently, the number density of the observable
- [Na] atom by lidar is strongly anti-correlated with the fluctuations of the NO<sup>+</sup>. In a nutshell, more NO<sup>+</sup>
- will directly lead to fewer observable Na atoms. That being said, the interaction between sodium and
- 287 background species is rather complex. The scope of the sensitivity factor in the present paper was
- 288 limited to column density. As a result of such, variations and behaviors of the sodium chemicals by
- altitude are overlooked. The actual impact of the background species may differ at different altitudes.

# 290 4.2 Meteoric input function

- 291 The estimation of meteoric influx is subject to many uncertainties among different techniques (Li et al.,
- 292 2022). Moreover, the meteor flux estimated by the sodium chemistry model also varies (Marsh et al.,
- 2014; Plane et al., 2015). The previous model of Plane (2004) and the following similar models indicate
- that the rate of dimerization, or the speed of removing sodium from the system, is heavily correlated to
- the vertical transport in the MLT. The NaChem model does not explicitly incorporate vertical transport,
- but the vertical transport by diffusion is inherently embedded within the input of the observed sodiumvertical profile.
- 298 Unlike the previous models (Plane 2004; March et al. 2014; and references therein), the present 299 NaChem model took an indirect route to estimate the meteor mass input. During the simulation, the
- 300 NaHCO<sub>3</sub> dimerization and the uptake of the sodium species on meteoric smoke particles, which can be
- NaHCO<sub>3</sub> dimerization and the uptake of the sodium species on meteoric smoke particles, which can be
   turned on or off, create a deficit of sodium atoms. Meanwhile, a meteor input function injects an
- 302 appropriate amount of sodium atoms so that the present sodium vertical profile always matches the
- reference profiles. This is carried out by finding the difference between the current sodium profile (with
- the deficit) and the corresponding reference profile in every iteration and then replacing the former
- 305 with the latter. The study by Plane (2004) found that the diffusion coefficient is highly correlated with
- 306 the sodium sink, primarily because the dimerization reaction occurs predominantly at lower altitudes.
- 307 The simulation circumvents this uncertainty by directly incorporating the observational sodium vertical
- 308 profile, given that diffusion is already inherently in the measurements.

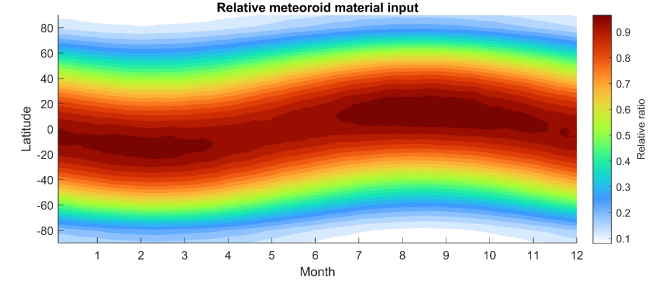


310 **Figure 4.** Logarithmic meteor radiant source distribution derived from the AO observations. The figure is

311 *in a.u. (arbitrary units).* The figure illustrates the relative frequency of meteor occurrence at different

312 radiant directions in the Earth Reference Frame (ERF), equivalent to ground-based observations. The

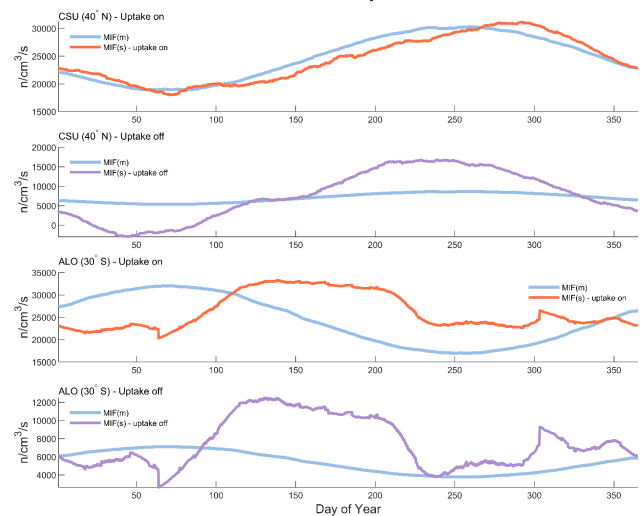
- 313 latitude of the ERF is centered on the ecliptic plane. The longitude of the ERF is centered to the Apex
- direction, the moving direction of the Earth, where the highest number of meteors encounter Earth. The
- radiant distribution is derived from the number of meteor events. Figure reproduced from Li et al. (2022).



316

**Figure 5.** Relative seasonal and latitudinal meteoroid input by meteor occurrence, inferred from the

radiant source distribution shown in Figure 4. The figure is normalized to its max value.



Sodium column injection rate

321 Figure 6. A comparison between two meteor input functions: MIF(m), which is inferred from micro-

322 meteor radiant distribution, and MIF(s), derived from a Na chemistry model with sodium input from lidar 323 observations.

324

325 Figure 4 shows the high-resolution meteor radiant source distribution recently inferred from the AO 326 observations (Li et al., 2022). The typical mass of the Arecibo meteors is estimated to be around 10<sup>-13</sup> kg 327 based on flux rate (Li and Zhou, 2019). Mathews et al. (2001) estimated the limiting meteor mass of 10<sup>-14</sup> 328 kg based on the meteor ballistic parameter. Limiting mass is the smallest mass a meteoroid must have 329 to generate sufficient ionization to be detected by radar. Despite these estimations being based on 330 various simplified assumptions that may lead to inaccurate results, the estimated limiting mass at AO is 331 still at least two orders of magnitude smaller than the estimations of other facilities by similar means. 332 More than 95% of the meteoroid population in the Earth's atmosphere is found to be sporadic meteors 333 by HPLA radar observation (Chau and Galindo, 2007), which typically are low-mass meteors evolved 334 from the outer Solar system due to the Poynting-Robertson drag (Nesvorný et al., 2011; Koschny et al., 335 2019). That being said, the percentage of sporadic meteors, as well as the radiant source distribution,

- are both estimated based on the occurrence. However, the occurrence of sporadic meteors may not be
- able to represent their mass distribution. The relative seasonal and latitudinal meteoroid input by the
- number of occurrence inferred from the new radiant source distribution is depicted in Figure 5. The
- 339 meteoric input generally follows a sinusoidal pattern and differs from the one used in the previous work,
- as shown in Figure 1 of Marsh et al. (2013). Although the interplanetary dust (meteor) background on
- 341 the Earth's orbit could vary in different locations due to a variety of reasons, e.g., Jupiter resonance, it is 342 still safe to assume no change in the interplanetary dust background for our purpose. Taking a stable
- 342 still safe to assume no enange in the interplanetary dust background for our purpose. Taking a stable 343 interplanetary dust background, the MIF(m) 's seasonal sinusoidal pattern should follow the Earth's axis
- 344 rotation relative to the ecliptic plane.
- 345 Figure 6 shows a comparison between two types of meteoric input function: MIF(m), which is inferred
- 346 from the micro-meteor radiant distribution, and MIF(s), derived using the Na chemistry model with
- 347 sodium input from the lidar observations. MIF(m) is in arbitrary units and has been linearly scaled to
- 348 match the amplitude of MIF(s). Same as MIF(s), MIF(m) is also smoothed by a 15-day running average.
- 349 For the MIF(s) model simulations, we did two scenarios, one with and one without uptake by smoke
- particles, for the ALO and CSU data. The MIF(s) with uptake by smoke particles exhibit a good match
- 351 with the MIF(m) on the CSU dataset, while it does not show as good of a match on the ALO dataset. The
- 352 MIF(s) with smoke uptake off is represented by a purple line, while the MIF(s) with smoke uptake turned
- 353 on is depicted by an orange line. The MIF(s) could go negative when the reference sodium vertical
- profile decreases faster than the removal rate by the dimerization, as shown in the purple line in Figure
- 355 6, indicating that the dimerization process alone is not sufficient enough to account for all the sodium
- atom depletion in the MLT region. MIF(m) is derived from a global micro-meteor radiant distribution
- 357 model, as depicted in Figure 4 and Figure 5. The smoke uptake of sodium species in this study is
- 358 implemented using a methodology similar to Plane (2004), but instead of applying smoke uptake solely
- to the three major sodium species, namely Na, NaHCO3, and Na+, it is applied to all 14 sodium-bearing
- 360 species. The optimal uptake factor to obtain the best results was found to be  $2x10^{-2}$ /km/s. The smoke
- uptake and NaHCO<sub>3</sub> dimerization account for approximately 75% and 25% of the Na sink, respectively.
- 362 According to the global meteoroid orbital model outlined in Li et al., (2022), the latitudes spanning 29.5° 363 S to 30.5° S (ALO) account for 0.52% of the total meteor input, while those between 39.5° N and 40.5° N 364 (CSU) represent 0.67%. The CSU site shares more meteor input due to its closer proximity to one of the 365 Apex meteor radiant sources. The global total sodium injection rate inferred from the ALO data-based simulation is (2.01±0.68) x10<sup>23</sup> atoms per second, and the CSU-data-based simulation suggests a global 366 367 sodium injection rate of  $(1.28\pm0.55) \times 10^{23}$  atoms per second. The error is determined by calculating the 368 standard deviation of the detrended, unsmoothed raw MIF(s). Note that both MIF(m) and MIF(s) 369 presented in Fig.6 are smoothed by a 15-day running average. Assuming the relative sodium elemental 370 abundance in meteoroid material is 0.8% (Vondrak et al., 2008), the deduced total meteoroid material
- input of ALO-based simulation is 83±28 t d<sup>-1</sup>. From CSU-based simulation, the rate is 53±23 t d<sup>-1</sup>. Both
- estimations are close to 80-130 t d<sup>-1</sup>, the value reported by the Long Duration Exposure Facility (Love
- and Brownlee, 1993; McBride et al., 1999). It is worth noting that the estimated total daily input of
- 374 meteoroid materials varies among previous studies, ranging from 4.6 t d<sup>-1</sup> (Marsh et al., 2013) to 300 t d<sup>-</sup>
- <sup>1</sup> (Nesvorný et al., 2009), with an intermediate value of 20 t d<sup>-1</sup> reported by Carrillo-Sánchez et al. (2020).
- 376 While these estimates seem quite disparate, the variance is relatively small given that the daily input
- 377 rate is derived from combinations of chemicals that can fluctuate by several orders of magnitude. For

- example, the NO<sup>+</sup>, which exhibits the highest sensitivity factor according to the sensitivity test,
- 379 undergoes diurnal variations of approximately three orders of magnitude.
- 380

## 381 5. Discussion

The sodium concentration in the sodium layer in the MLT region is governed by several factors, including chemistry, dynamics, and the MIF. It's difficult to discern which of these three components is more

important than the others. In this section, we discuss various factors that may contribute to modeling

the sodium concentration in the MLT.

- 386 The mass of the meteoroids has been estimated and measured using various methods. These include
- the ballistic parameter derived from meteor deceleration (Mathews et al., 2001), estimation of meteor
- 388 head echo plasma distribution through a combination of meteor ablation models and radar cross-
- 389 section measurements (Close et al., 2005; Sugar et al., 2021), flux rate determination (Zhou and Kelley,
- 1997), as well as spacecraft in-situ measurements (Leinert and Grun, 1990), among others. The mass
- estimated by the meteor ballistic parameter is commonly referred to as momentum or dynamical mass.
- 392 The mass estimated by the meteor ablation model is usually called the scattering mass. The meteor
- momentum mass from Arecibo Ultra-High-Frequency (UHF) radar observation is estimated to be  $10^{-14} 10^{-7}$  kg, with the typical mass being  $10^{-13}$  kg. On the other hand, the meteor scattering mass is estimated
- to be  $10^{-9} 10^{-5.5}$  kg by data from EISCAT UHF radar (Kero et al., 2008) and  $10^{-7} 10^{-4.5}$  kg by data from
- ALTAIR UHF radar (Close et al., 2005). While the detection sensitivity among different facilities differs,
- 397 these estimations are still off by many orders of magnitude. The assessments of either momentum mass
- 398 or scattering mass are based on a variety of simplified assumptions. They are subject to errors due to
- 399 the complexity of radar beam patterns, background atmosphere conditions, aspect sensitivity, meteor
- 400 radiant sources, and many other possible factors. For example, radar meteor observation is subject to
- 401 bias against low-mass, low-velocity meteors (Close et al., 2007; Janches et al., 2015).
- 402 Another aspect that may contribute to the MIF(m)'s uncertainty is the meteor radiant distribution. The 403 meteor radiant distributions shown in Figure 4 and many others (Chau et al., 2004; Campbell-Brown and 404 Jones, 2006; Kero et al., 2012) are inferred or measured by meteor occurrence instead of mass input. 405 Currently, retrieving a more accurate estimation of the meteor mass input is still a topic under active 406 research, and there is no quantitative study on the disparities between meteor occurrence and meteor 407 mass input. The radiant sources of the meteors are expected to differ by mass as their orbital evolution 408 is highly correlated to their mass. The interplanetary dust interacts with the solar wind while in the Solar 409 System, losing its momentum in the process and evolving into orbits with a smaller semi-major axis and 410 lower eccentricity. The effect is called the Poynting-Robertson effect (Robertson and Russell, 1937), 411 which behaves like a drag force and defines the evolution of interplanetary dust, and it could be the 412 major reason for the existence of sporadic meteors (Li and Zhou, 2019; Koschny et al., 2019). The 413 importance of the Poynting-Robertson effect is highly dependent on the density and mass of the object. 414 By and large, the orbits of the smaller particles evolve exponentially faster. The orbital dynamics of 415 interplanetary particles have been very well summarized in section 2.2 of (Koschny et al., 2019). For the 416 reasons above, the meteor radiant distribution of mass could deviate from the radiant distribution of 417 occurrence. Therefore, the meteor input rates as shown in the blue curves of Figure 6 could be different 418 from those derived from the meteor radiant distribution of mass since they were derived from the 419 meteor radiant distribution by occurrence.

- 420 In the sodium chemistry model presented in this work, the MIF is the sole source of sodium, while the
- 421 sodium sink comprises NaHCO<sub>3</sub> dimerization and smoke uptake. The MIF(s) is determined by matching
- 422 the sink rate of the sodium atoms with the rate of sodium injection. In other words, MIF(s) represents
- the amount of sodium injection needed to keep the sodium concentration equal to the reference
- 424 sodium profiles. If the chemical lifetime of sodium in the MLT is short, then the seasonal variation of
- both the MIF and sodium concentration in the MLT should be similar. After examining Figures 2, 5, and
- 426 6, it can be observed that the averaged seasonal variation of sodium over the years at both sites (ALO
- 427 and CSU) does not correspond to the trend of the MIF(m) at their respective latitudes. This may indicate
- 428 that the chemical lifetime of sodium in the MLT should be relatively long, as there is no immediate effect
- of MIF(m) on the sodium concentration. The MIF(m) displays a sinusoidal pattern which peaks in March
   at the ALO's latitude and in August at the CSU's latitude, whereas the sodium layer shows dual peaks in
- 431 the CSU's lidar observations and one peak in June in the ALO's lidar observations.
- 432 In this study, the MIF(s) derived from the NaChem simulation, based on the CSU lidar measurements
- 433 with uptake turned on, was able to match the amplitude of MIF(m) obtained from the meteor radiant
- 434 distribution. Although the model does not directly incorporate any dynamical processes, the vertical
- 435 transport by diffusion is implicitly included. The model forces the sodium layer to be the same as the
- 436 data, which are derived from the average of many years' measurements, in which the diffusions are
- 437 inherently embedded. The combination of observational data with the numerical chemistry model in
- 438 this paper is a relatively straightforward application of data assimilation (Bouttier & Courtier 2002). The
- lidar data of both sites (CSU and ALO) indicate that the sodium column density consistently increases by
- about 20% from 22:00 to 4:00 LT the next day. This can be attributed to the fact that, during nighttime,
- the large deposits of Na<sup>+</sup> formed by daytime reactions slowly neutralized to Na. As a result, the sodium
- column density consistently increases throughout the night. The same effect can be reproduced in the
- A43 NaChem simulation, albeit with a smaller amplitude. The simulation shows the increase to be about 8%.
- 444 The value is obtained by maintaining a constant total number of sodium-bearing species through the
- deactivation of the sodium sink.
- 446 While meridional transport or atmospheric dynamics both contribute to the seasonal variation of the
- sodium layer in the MLT, the diurnal sodium profile is the mean of observations of thousands of days, of
- 448 which the variation by atmospheric dynamics should be much less prominent. The lack of explicit
- dynamics in the model may be one of the sources of inconsistency when compared to the MIF(m)
- 450 observations. Further, the WACCM 6, which supplied the background species to the NaChem, is an older
- version that does not fully incorporate the dynamics of each ion species. Despite our results showing
- 452 good agreement between the MIF(s) and the MIF(m), there might be several plausible factors that could
- 453 lead to potential errors. For example, the Na sink by NaHCO<sub>3</sub> dimerization varies by the diffusion rate or
- the vertical transport of sodium atoms in the chemistry model (Plane, 2004). Likewise, the MIF(m) may
- also differ if the meteoroid mass input differs from the radiant source distribution by the occurrence of
- 456 meteors, as discussed in the aforementioned paragraph.

#### 457 **5. Conclusion**

- 458 This work introduced a new sodium chemistry model that simulates the time evolution of all sodium-
- 459 bearing species using the continuity equation without making any steady-state assumption. The model
- 460 employs an exponential integrator and runs in high-time resolution to maintain numerical stability. The
- 461 model is simple to maintain in such a configuration and can be scaled up to include additional

462 capabilities more easily. The model is highly optimized for processing efficiency and benefits from the

- use of an exponential integrator. Therefore, within an acceptable total CPU time, the NaChem can afford
- a time resolution of up to milliseconds, several orders of magnitude smaller than those used in other Na
- 465 models. During our testing, the CPU time to simulated real-time ratio is about 1 to 1000 using a 0.1466 second time step.

The model simulation was able to reproduce the seasonal variation of the sodium layer in the MLT by

- simulations of chemical reactions. The simulation results at the CSU's latitude capture the general trend
   of the seasonal variation at the location. The MIF(s) based on the ALO data exhibited less conformity
- of the seasonal variation at the location. The MIF(s) based on the ALO data exhibited less conformity
   with the corresponding MIF(m), which could be attributed to inadequate statistics of the observational
- 471 data. Comparably, the CSU dataset is more reliable as the insufficient lidar hours in the ALO dataset may
- 472 lead to inaccurate statistics. In the simulation, when forcing the sodium layer to be the observation-
- based reference profile, the inferred MIF is estimated to be  $83\pm28$  t d<sup>-1</sup> at ALO and  $53\pm23$  t d<sup>-1</sup> at CSU.
- The numerical simulation by NaChem could reproduce the general trend of diurnal and seasonal
- 475 variation of the sodium layer compared to the observations by the CSU Lidar. There are some
- 476 inconsistencies in MIF(m) and MIF(s) based on data obtained from ALO Lidar. These inconsistencies may
- 477 have originated from poor statistics resulting from insufficient observation hours.
- 478 In summary, a new sodium chemistry model has been developed in this work to investigate the
- relationship between MIF and the sodium layer. We also compared the MIF(m) derived from radar
- 480 meteor observation to the MIF(s) derived from the chemistry model and lidar observations. Our results
- 481 indicate that the uptake of sodium species onto meteoric smoke particles removes approximately three
- times more sodium than the dimerization of NaHCO3. Our future work will focus on incorporating the
- 483 plausible factors that may lead to potential errors discussed above into the chemistry model.
- 484
- 485
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- 493
- 494 Code/Data availability

The CSU lidar data is available through Utah State University data service (Yuan, 2023). The ALO data is

- 496 available through the ALO online database (ALO, 2023). The WACCM data used in this work are available
- 497 through Penn State Scholarsphere (Li, 2023b).
- 498
- 499 Author contribution

- 500 Conceptualization, Yanlin L., Tai-Yin H. and Julio U.; methodology, Yanlin L.; software, Yanlin L.;
- validation, Yanlin L., Tai-Yin H., Fabio V., Julio U. and Wuhu F.; formal analysis, Yanlin L., Tai-Yin H. and
- Julio U.; investigation, Yanlin L., Tai-Yin H., Julio U. and Wuhu F.; resources, Tai-Yin H., Julio U., Fabio V.,
- and Wuhu F.; data curation, Yanlin L.; writing---original draft preparation, Yanlin L.; writing---review and
- editing, Yanlin L., Tai-Yin H., Julio U., Fabio V., and Wuhu F.; visualization, Yanlin L.; supervision, Julio U.
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- 508 Competing interests
- 509 The authors declare no competing interests.
- 510

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